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Physical interventions to interrupt or reduce the spread of respiratory viruses (Review)

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[Intervention Review]

Physical interventions to interrupt or reduce the spread of respiratory viruses

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ABSTRACT

Background

Viral epidemics or pandemics of acute respiratory infections (ARIs) pose a global threat. Examples are influenza (H1N1) caused by the H1N1pdm09 virus in 2009, severe acute respiratory syndrome (SARS) in 2003, and coronavirus disease 2019 (COVID-19) caused by SARS-CoV-2 in 2019. Antiviral drugs and vaccines may be insufficient to prevent their spread. This is an update of a Cochrane Review last published in 2020. We include results from studies from the current COVID-19 pandemic.

Objectives

To assess the effectiveness of physical interventions to interrupt or reduce the spread of acute respiratory viruses.

Search methods

We searched CENTRAL, PubMed, Embase, CINAHL, and two trials registers in October 2022, with backwards and forwards citation analysis on the new studies.

Selection criteria

We included randomised controlled trials (RCTs) and cluster-RCTs investigating physical interventions (screening at entry ports, isolation, quarantine, physical distancing, personal protection, hand hygiene, face masks, glasses, and gargling) to prevent respiratory virus transmission.

Data collection and analysis

We used standard Cochrane methodological procedures.



Main results

We included 11 new RCTs and cluster-RCTs (610,872 participants) in this update, bringing the total number of RCTs to 78. Six of the new trials were conducted during the COVID-19 pandemic; two from Mexico, and one each from Denmark, Bangladesh, England, and Norway. We identified four ongoing studies, of which one is completed, but unreported, evaluating masks concurrent with the COVID-19 pandemic.

Many studies were conducted during non-epidemic influenza periods. Several were conducted during the 2009 H1N1 influenza pandemic, and others in epidemic influenza seasons up to 2016. Therefore, many studies were conducted in the context of lower respiratory viral circulation and transmission compared to COVID-19. The included studies were conducted in heterogeneous settings, ranging from suburban schools to hospital wards in high-income countries; crowded inner city settings in low-income countries; and an immigrant neighbourhood in a high-income country. Adherence with interventions was low in many studies.

The risk of bias for the RCTs and cluster-RCTs was mostly high or unclear.

Medical/surgical masks compared to no masks

We included 12 trials (10 cluster-RCTs) comparing medical/surgical masks versus no masks to prevent the spread of viral respiratory illness (two trials with healthcare workers and 10 in the community). Wearing masks in the community probably makes little or no difference to the outcome of influenza-like illness (ILI)/COVID-19 like illness compared to not wearing masks (risk ratio (RR) 0.95, 95% confidence interval (CI) 0.84 to 1.09; 9 trials, 276,917 participants; moderate-certainty evidence. Wearing masks in the community probably makes little or no difference to the outcome of laboratory-confirmed influenza/SARS-CoV-2 compared to not wearing masks (RR 1.01, 95% CI 0.72 to 1.42; 6 trials, 13,919 participants; moderate-certainty evidence). Harms were rarely measured and poorly reported (very low-certainty evidence).

N95/P2 respirators compared to medical/surgical masks

We pooled trials comparing N95/P2 respirators with medical/surgical masks (four in healthcare settings and one in a household setting). We are very uncertain on the effects of N95/P2 respirators compared with medical/surgical masks on the outcome of clinical respiratory illness (RR 0.70, 95% CI 0.45 to 1.10; 3 trials, 7779 participants; very low-certainty evidence). N95/P2 respirators compared with medical/surgical masks may be effective for ILI (RR 0.82, 95% CI 0.66 to 1.03; 5 trials, 8407 participants; low-certainty evidence). Evidence is limited by imprecision and heterogeneity for these subjective outcomes. The use of a N95/P2 respirators compared to medical/surgical masks probably makes little or no difference for the objective and more precise outcome of laboratory-confirmed influenza infection (RR 1.10, 95% CI 0.90 to 1.34; 5 trials, 8407 participants; moderate-certainty evidence). Restricting pooling to healthcare workers made no difference to the overall findings. Harms were poorly measured and reported, but discomfort wearing medical/surgical masks or N95/P2 respirators was mentioned in several studies (very low-certainty evidence).

One previously reported ongoing RCT has now been published and observed that medical/surgical masks were non-inferior to N95 respirators in a large study of 1009 healthcare workers in four countries providing direct care to COVID-19 patients.

Hand hygiene compared to control

Nineteen trials compared hand hygiene interventions with controls with sufficient data to include in meta-analyses. Settings included schools, childcare centres and homes. Comparing hand hygiene interventions with controls (i.e. no intervention), there was a 14% relative reduction in the number of people with ARIs in the hand hygiene group (RR 0.86, 95% CI 0.81 to 0.90; 9 trials, 52,105 participants; moderate-certainty evidence), suggesting a probable benefit. In absolute terms this benefit would result in a reduction from 380 events per 1000 people to 327 per 1000 people (95% CI 308 to 342). When considering the more strictly defined outcomes of ILI and laboratory-confirmed influenza, the estimates of effect for ILI (RR 0.94, 95% CI 0.81 to 1.09; 11 trials, 34,503 participants; low-certainty evidence), and laboratory-confirmed influenza (RR 0.91, 95% CI 0.63 to 1.30; 8 trials, 8332 participants; low-certainty evidence), suggest the intervention made little or no difference. We pooled 19 trials (71, 210 participants) for the composite outcome of ARI or ILI or influenza, with each study only contributing once and the most comprehensive outcome reported. Pooled data showed that hand hygiene may be beneficial with an 11% relative reduction of respiratory illness (RR 0.89, 95% CI 0.83 to 0.94; low-certainty evidence), but with high heterogeneity. In absolute terms this benefit would result in a reduction from 200 events per 1000 people to 178 per 1000 people (95% CI 166 to 188). Few trials measured and reported harms (very low-certainty evidence).

We found no RCTs on gowns and gloves, face shields, or screening at entry ports.

Authors' conclusions

The high risk of bias in the trials, variation in outcome measurement, and relatively low adherence with the interventions during the studies hampers drawing firm conclusions. There were additional RCTs during the pandemic related to physical interventions but a relative paucity given the importance of the question of masking and its relative effectiveness and the concomitant measures of mask adherence which would be highly relevant to the measurement of effectiveness, especially in the elderly and in young children.

There is uncertainty about the effects of face masks. The low to moderate certainty of evidence means our confidence in the effect estimate is limited, and that the true effect may be different from the observed estimate of the effect. The pooled results of RCTs did not show a clear reduction in respiratory viral infection with the use of medical/surgical masks. There were no clear differences between the use



of medical/surgical masks compared with N95/P2 respirators in healthcare workers when used in routine care to reduce respiratory viral infection. Hand hygiene is likely to modestly reduce the burden of respiratory illness, and although this effect was also present when ILI and laboratory-confirmed influenza were analysed separately, it was not found to be a significant difference for the latter two outcomes. Harms associated with physical interventions were under-investigated.

There is a need for large, well-designed RCTs addressing the effectiveness of many of these interventions in multiple settings and populations, as well as the impact of adherence on effectiveness, especially in those most at risk of ARIs.

PLAIN LANGUAGE SUMMARY

Do physical measures such as hand-washing or wearing masks stop or slow down the spread of respiratory viruses?

Key messages

We are uncertain whether wearing masks or N95/P2 respirators helps to slow the spread of respiratory viruses based on the studies we assessed.

Hand hygiene programmes may help to slow the spread of respiratory viruses.

How do respiratory viruses spread?

Respiratory viruses are viruses that infect the cells in your airways: nose, throat, and lungs. These infections can cause serious problems and affect normal breathing. They can cause flu (influenza), severe acute respiratory syndrome (SARS), and COVID-19.

People infected with a respiratory virus spread virus particles into the air when they cough or sneeze. Other people become infected if they come into contact with these virus particles in the air or on surfaces on which they land. Respiratory viruses can spread quickly through a community, through populations and countries (causing epidemics), and around the world (causing pandemics).

Physical measures to try to prevent respiratory viruses spreading between people include:

- · washing hands often;
- · not touching your eyes, nose, or mouth;
- · sneezing or coughing into your elbow;
- · wiping surfaces with disinfectant;
- · wearing masks, eye protection, gloves, and protective gowns;
- · avoiding contact with other people (isolation or quarantine);
- · keeping a certain distance away from other people (distancing); and
- · examining people entering a country for signs of infection (screening).

What did we want to find out?

We wanted to find out whether physical measures stop or slow the spread of respiratory viruses from well-controlled studies in which one intervention is compared to another, known as randomised controlled trials.

What did we do?

We searched for randomised controlled studies that looked at physical measures to stop people acquiring a respiratory virus infection.

We were interested in how many people in the studies caught a respiratory virus infection, and whether the physical measures had any unwanted effects.

What did we find?

We identified 78 relevant studies. They took place in low-, middle-, and high-income countries worldwide: in hospitals, schools, homes, offices, childcare centres, and communities during non-epidemic influenza periods, the global H1N1 influenza pandemic in 2009, epidemic influenza seasons up to 2016, and during the COVID-19 pandemic. We identified five ongoing, unpublished studies; two of them evaluate masks in COVID-19. Five trials were funded by government and pharmaceutical companies, and nine trials were funded by pharmaceutical companies.

No studies looked at face shields, gowns and gloves, or screening people when they entered a country.

We assessed the effects of:

· medical or surgical masks;



- $\cdot \text{N95/P2} \ respirators \ (close-fitting \ masks \ that \ filter \ the \ air \ breathed \ in, \ more \ commonly \ used \ by \ healthcare \ workers \ than \ the \ general \ public); \ and$
- · hand hygiene (hand-washing and using hand sanitiser).

We obtained the following results:

Medical or surgical masks

Ten studies took place in the community, and two studies in healthcare workers. Compared with wearing no mask in the community studies only, wearing a mask may make little to no difference in how many people caught a flu-like illness/COVID-like illness (9 studies; 276,917 people); and probably makes little or no difference in how many people have flu/COVID confirmed by a laboratory test (6 studies; 13,919 people). Unwanted effects were rarely reported; discomfort was mentioned.

N95/P2 respirators

Four studies were in healthcare workers, and one small study was in the community. Compared with wearing medical or surgical masks, wearing N95/P2 respirators probably makes little to no difference in how many people have confirmed flu (5 studies; 8407 people); and may make little to no difference in how many people catch a flu-like illness (5 studies; 8407 people), or respiratory illness (3 studies; 7799 people). Unwanted effects were not well-reported; discomfort was mentioned.

Hand hygiene

Following a hand hygiene programme may reduce the number of people who catch a respiratory or flu-like illness, or have confirmed flu, compared with people not following such a programme (19 studies; 71,210 people), although this effect was not confirmed as statistically significant reduction when ILI and laboratory-confirmed ILI were analysed separately. Few studies measured unwanted effects; skin irritation in people using hand sanitiser was mentioned.

What are the limitations of the evidence?

Our confidence in these results is generally low to moderate for the subjective outcomes related to respiratory illness, but moderate for the more precisely defined laboratory-confirmed respiratory virus infection, related to masks and N95/P2 respirators. The results might change when further evidence becomes available. Relatively low numbers of people followed the guidance about wearing masks or about hand hygiene, which may have affected the results of the studies.

How up to date is this evidence?

We included evidence published up to October 2022.

SUMMARY OF FINDINGS

Summary of findings 1. Medical/surgical masks compared to no masks for preventing the spread of viral respiratory illness

Randomised studies: medical/surgical masks compared to no masks for preventing the spread of viral respiratory illness

Patient or population: general population

Setting: community and hospitals Intervention: medical/surgical masks

Comparison: no masks

Outcomes	Anticipated absolute effects* (95% CI)		Relative effect (95% CI)	№ of partici- pants	Certainty of the evidence	Comments
	Risk with no masks	Risk with ran- domised studies: masks	,	(studies)	(GRADE)	
Viral respiratory illness - influenza/COVID-like ill- ness	Study population	1	RR 0.95 (0.84 to 1.09)	276,917 (9 RCTs)	⊕⊕⊕⊝ Moderate ^a	
	160 per 1000	152 per 1000 (134 to 174)				
Viral respiratory illness - laboratory-confirmed influenza/SARS-CoV-2	Study population		RR 1.01 (0.72 to 1.42)	13,919 (6 RCTs)	⊕⊕⊕⊝ Moderate ^b	
	40 per 1000	40 per 1000 (29 to 57)	(0.72 to 1.42)		Moderates	
Adverse events	-		-	(3 RCTs)	⊕⊝⊝⊝ Very low ^{a,c}	Adverse events were not reported consistently and could not be meta-analysed.
						Adverse events reported for masks included warmth, discomfort, respiratory difficulties, humidity, pain, and shortness of breath, in up to 45% of participants.

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the median observed risk in the comparison group of included studies and the relative effect of the intervention (and its 95% CI).

CI: confidence interval: RCT: randomised controlled trial: RR: risk ratio

GRADE Working Group grades of evidence

High certainty: we are very confident that the true effect lies close to that of the estimate of the effect.

Moderate certainty: we are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

Low certainty: our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect.

^aDowngraded one level for study limitations (lack of blinding).

bDowngraded one level for imprecision (wide confidence intervals).

^cDowngraded two levels for imprecision (only three studies enumerated adverse events; another study mentioned no adverse events).

Summary of findings 2. N95 respirators compared to medical/surgical masks for preventing the spread of viral respiratory illness

Randomised studies: N95 respirators compared to medical/surgical masks for preventing the spread of viral respiratory illness

Patient or population: general population and healthcare workers

Setting: hospitals and households

Intervention: N95 masks

Comparison: medical/surgical masks

Outcomes	Anticipated absolute effects* (95% CI)		Relative effect (95% CI)	№ of partici- pants (studies)	Certainty of the evidence (GRADE)	Comments	
	Risk with med- ical masks	Risk with ran- domised stud- ies: N95					
Viral respiratory illness - clinical	Study population		RR 0.70 - (0.45 to 1.10)	7799 (3 RCTs)	⊕⊝⊝⊝ Very Low ^{a,b,c}	All studies were conducted in hospital settings with healthcare workers.	
respiratory illness	120 per 1000	84 per 1000 (54 to 132)	(0.13 to 1.10)		very Low-5-5-	neutaleure workers.	
Viral respiratory	illness - influen-		RR 0.82 - (0.66 to 1.03)	8407 (5 RCTs)	⊕⊕⊝⊝ Lowa,b	1 study was conducted in households (MacIntyre 2009).	
za-like illness			(0.00 to 1.00)			2005).	
Viral respiratory illness - laborato-	, , ,		RR 1.10 - (0.90 to 1.34)	8407 (5 RCTs)	⊕⊕⊕⊝ Moderate ^b	1 study was conducted in households (MacIntyre 2009).	
ry-confirmed in- fluenza	70 per 1000	77 per 1000 (63 to 94)	(0.30 to 1.31)		Moderate-	2003).	
Adverse events	-		-	(5 RCTs)	⊕⊝⊝⊝ Very Low ^{a,b,c}	There was insufficient consistent reporting of adverse events to enable meta-analysis.	
						Only 1 study reported detailed adverse events: discomfort was reported in 41.9% of N95 wearers versus 9.8% of medical mask wearers (P < 0.001); headaches	

were more common with N95 (13.4% versus 3.9%; P < 0.001); difficulty breathing was reported more often in the N95 group (19.4% versus 12.5%; P = 0.01); and N95 caused more problems with pressure on the nose (52.2% versus 11.0%; P < 0.001). 4 RCTs either reported no adverse events or only reported on comfort wearing masks.

CI: confidence interval; RCT: randomised controlled trial; RR: risk ratio

GRADE Working Group grades of evidence

High certainty: we are very confident that the true effect lies close to that of the estimate of the effect.

Moderate certainty: we are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

Low certainty: our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect.

Very low certainty: we have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of effect.

^aDowngraded one level for study limitations (lack of blinding).

^bDowngraded one level for imprecision (wide confidence interval or no meta-analysis conducted).

^cDowngraded one level for inconsistency of results (heterogeneity).

Summary of findings 3. Hand hygiene compared to control for preventing the spread of viral respiratory illness

Hand hygiene compared to control for preventing the spread of viral respiratory illness

Patient or population: general population and healthcare workers **Setting:** schools, childcare centres, homes, offices, nursing homes

Intervention: hand hygiene **Comparison:** control

Outcomes	Anticipated absolute effects* (95% CI)			№ of partici- pants	Certainty of the evidence	Comments
	Risk with con- trol	Risk with hand hy- giene	,,,,,,	(studies)	(GRADE)	
Acute respiratory illness	Study population		RR 0.86 52,105 (9 RCTs) - (0.81 to 0.90)	⊕⊕⊕⊝ Moderate ^a		
	380 per 1000	327 per 1000 (308 to 342)	(0.01 to 0.50)		Moderate	

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the median risk in the comparison group and the observed relative effect of the intervention (and its 95% CI).

versus 10.3% in the control group.

Influenza-like illness	Study population		RR 0.94 (0.81 to 1.09)	34,503 (11 RCTs)	⊕⊕⊝⊝ Lowa,b	
	90 per 1000	85 per 1000 (73 to 98)	(0.01 to 1.03)	ite 13)	LOVV	
Laboratory-confirmed in- fluenza	Study population		RR 0.91 (0.63 to 1.30)	8332 (8 RCTs)	⊕⊕⊝⊝ Low ^{b,c}	
	80 per 1000	73 per 1000 (50 to 104)	(0.03 to 1.30)		LOW-	
Composite of acute respiratory illness, influenza-like illness, laboratory-confirmed influenza	Study population			71,210 (19 RCTs)	⊕⊕⊝⊝ Lawa b	
	200 per 1000	178 per 1000 (166 to 188)	(0.83 to 0.94)	RCIS	Lowa,b	
Adverse events	-		-	(2 RCTs)	⊕⊝⊝⊝	Data were insufficient to conduct meta-analysis.
					Very low ^{a,b,c}	1 study reported that no adverse events were observed, and another study reported that skin reaction was recorded for 10.4% of participants in the hand sanitiser group

^{*}The risk in the intervention group (and its 95% confidence interval) is based on the median observed risk in the comparison groups of included studies and the relative effect of the intervention (and its 95% CI).

CI: confidence interval; RCT: randomised controlled trial; RR: risk ratio

GRADE Working Group grades of evidence

High certainty: we are very confident that the true effect lies close to that of the estimate of the effect.

Moderate certainty: we are moderately confident in the effect estimate: the true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different.

Low certainty: our confidence in the effect estimate is limited: the true effect may be substantially different from the estimate of the effect.

Very low certainty: we have very little confidence in the effect estimate: the true effect is likely to be substantially different from the estimate of effect.

^aDowngraded one level for study limitation (majority of studies were unblinded, with participant-assessed outcome).

^bDowngraded one level for inconsistent results across studies.

^cDowngraded one level for imprecision (wide confidence interval or no meta-analysis conducted).



BACKGROUND

Description of the condition

Epidemic and pandemic viral infections pose a serious threat to people worldwide. Epidemics of note include severe acute respiratory syndrome (SARS) in 2003 and the Middle East respiratory syndrome (MERS), which began in 2012, and the current SARS-CoV-2 pandemic. Major pandemics include the H1N1 influenza caused by the H1N1pdm09 virus in 2009 and the coronavirus disease 2019 (COVID-19) caused by SARS-CoV-2.

Even non-epidemic acute respiratory infections (ARIs) place a huge burden on healthcare systems around the world, and are a prominent cause of morbidity (WHO 2017). Furthermore, ARIs are often antecedents to lower respiratory tract infections (RTIs) caused by bacterial pathogens (i.e. pneumonia), which cause millions of deaths worldwide, mostly in low-income countries (Schwartz 2018).

High viral load, high levels of transmissibility, susceptible populations, and symptomatic patients are considered to be the drivers of such epidemics and pandemics (Jefferson 2006a). Preventing the spread of respiratory viruses from person to person may be effective at reducing the spread of outbreaks.

Physical interventions, such as the use of masks and physical distancing measures, might prevent the spread of respiratory viruses which are considered to be transmitted by multiple modes of transmission including by respiratory particles of varying sizes spreading from infected to susceptible people and through direct and indirect contact (Kutter 2018; Leung 2021). It is recognised that there is a continuum of respiratory particle sizes varying between large droplet to fine aerosols, which is an important concept. Particles of a variety of sizes may be expelled from the human airway during coughing, sneezing, singing, talking, and during certain medical procedures (WHO 2021). In addition, transmission of respiratory viruses is likely highly complex, dependent on multiple host, virus and environmental factors, plus the myriad of interactions between these factors, which may influence the predominant modes of transmission in any given setting (Broderick 2008; Hendley 1988; Kutter 2018; Leung 2021). Current evidence suggests that the virus responsible for the current COVID-19 pandemic spreads mainly between people who are in close contact with each other (Onakpoya 2022a).

It is also unknown if all respiratory viruses or different strains of a specific respiratory virus transmit in a similar manner, further adding to the complexity of respiratory virus transmission.

Description of the intervention

Single measures of intervention such as the use of vaccines or antivirals, may be insufficient to contain the spread of influenza, but combinations of interventions may reduce the reproduction number to below 1 (Demicheli 2018a; Demicheli 2018b; Jefferson 2014; Jefferson 2018; Thomas 2010). When the reproduction number (or R0) is below 1, each infection causes less than one new secondary infection and the disease will eventually die out. For some respiratory viruses there are no licensed interventions, and a combination of social and physical interventions may be the only option to reduce the spread of outbreaks, particularly those that may be capable of becoming epidemic or pandemic in nature (Luby 2005). Such interventions were emphasised in the

World Health Organization's latest Global Influenza Strategy 2019 to 2030, and have several possible advantages over other methods of suppressing ARI outbreaks since they may be instituted rapidly and may be independent of any specific type of infective agent, including novel viruses. In addition, the possible effectiveness of public health measures during the Spanish flu pandemic of 1918 to 1919 in US cities supports the impetus to investigate the existing evidence on the effectiveness of such interventions (Bootsma 2007), including quarantine (such as isolation, physical distancing) and the use of disinfectants. We also considered the major societal implications for any community adopting these measures (CDC 2005a; CDC 2005b; WHO 2006b; WHO 2020a; WHO 2020b).

How the intervention might work

Epidemics and pandemics are more likely during antigenic change (changes in the viral composition) in the virus or transmission from animals (domestic or wild) when there is no natural human immunity (Bonn 1997). High viral load, high levels of transmissibility, and symptomatic patients are considered to be the drivers of such epidemics and pandemics (Jefferson 2006b).

Physical interventions, such as the use of masks (Greenhalgh 2020; Howard 2020), physical distancing measures, school closures, and limitations of mass gatherings, might prevent the spread of the virus transmitted by infectious respiratory particles from infected to susceptible individuals. The use of hand hygiene, gloves, and protective gowns can also prevent the spread by limiting the transfer of viral particles onto and from fomites (inanimate objects such as flat surfaces, tabletops, utensils, porous surfaces, or nowadays cell phones, which can transmit the agent if contaminated) (Onakpoya 2022b). Such public health measures were widely adopted during the Spanish flu pandemic and have been the source of considerable debate (Bootsma 2007).

Why it is important to do this review

Although the benefits of physical interventions seem self-evident, given the global importance of interrupting respiratory virus transmission, having up-to-date estimates of their effectiveness is necessary to inform planning, decision-making, and policy. The continuance of outbreaks of COVID-19 and the reporting of several new trials assessing different barrier interventions in preventing the spread of SARS-COV-2 virus, have prompted this update (WHO 2022). Physical methods have several possible advantages over other methods of suppressing ARI outbreaks, including their rapid deployment and ability to be independent of the infective agent, including novel viruses.

The hallmark of the 2020 update was shifting from including all types of studies to a focus on randomised controlled trials (RCTs) only, which had substantially increased in number. This change enabled more robust evidence summaries from high-quality studies, which are much less prone to the risk of the multiple biases associated with observational studies, to help policy and decision makers in making national and global recommendations. The 2020 update identified 67 relevant studies, but none were carried out during the COVID-19 pandemic (Jefferson 2020). The three key messages of that update were: (1) hand hygiene programmes may help to slow the spread of respiratory viruses; (2) uncertainty whether wearing masks or N95/P2 respirators would help in slowing the spread of respiratory viruses; and (3) few studies were identified for other interventions. One study looked



at quarantine, and none looked at eye protection, gowns and gloves, or screening people when they entered a country. However, during the last search of the 2020 update, six ongoing, unpublished studies were identified; three of them evaluate masks in COVID-19. The review authors are aware that several trials have now been published since the publication of the 2020 update, warranting this new update.

This is the fifth update (Jefferson 2009; Jefferson 2010; Jefferson 2011; Jefferson 2020) of a Cochrane Review first published in 2007 (Jefferson 2007).

OBJECTIVES

To assess the effectiveness of physical interventions to interrupt or reduce the spread of acute respiratory viruses.

METHODS

Criteria for considering studies for this review

Types of studies

For this 2022 update we only considered individual-level randomised controlled trials (RCTs), or cluster-RCTs, or quasi-RCTs for inclusion.

In versions of this review prior to 2020 we also included observational studies (cohorts, case-controls, before-after, and time series studies). However, for this update there were sufficient randomised studies to address our study aims, so we excluded observational studies because randomisation is the optimal method to prevent systematic differences between participants in different intervention groups and, further, deciding who receives an intervention and who does not is influenced by many factors, including prognostic factors (Higgins 2011). This point is particularly relevant here because individuals who chose to implement physical interventions are likely to use multiple interventions, thus making it difficult to separate out the effect of single interventions. Further, they are likely to be different from individuals who do not implement physical interventions in ways that are difficult to measure.

Types of participants

People of all ages.

Types of interventions

We included RCTs and cluster-RCTs of trials investigating physical interventions or combinations of interventions to prevent respiratory virus transmission compared with doing nothing or with other interventions. The interventions of interest included: screening at entry ports, isolation, quarantine, physical distancing, personal protection (clothing, gloves, devices), hand hygiene, face masks, gargling, nasal washes, eye protective devices, face shields, disinfecting, and school closure.

Types of outcome measures

For the outcomes listed below we had no predetermined key time points of interest or adverse events of special interest, however, methods of assessment of cases of viral respiratory illness based on laboratory-confirmation needed to be based on an accurate test in combination with critical additional information. For example, a polymerase chain reaction (PCR) test in combination

with symptoms of disease, or a serological test at baseline as well as at the end of follow-up were acceptable methods. Further, we stratified analyses by study-specific definitions for cases of viral respiratory illness which included a broad definition of acute respiratory infection (ARI), a more specific definition of influenza-like-illness (ILI), and the most precise definition of a laboratory-confirmed respiratory infection that identified the actual viral pathogen. For the studies conducted during the COVID-19 pandemic, we assumed that COVID-like illness was interchangeable with ILI. In the case of laboratory-confirmed respiratory infection we separated out SARS-CoV-2/influenza and other viral pathogens. We did not pool these outcomes as it cannot be assumed that the effects of physical interventions will be the same for the different viral pathogens. The one exception was for the comparison of hand-hygiene versus control where the estimated effects for ARI, ILI and laboratory-confirmed infection were highly consistent.

Primary outcomes

- Numbers of cases of viral respiratory illness (including acute respiratory infections (ARI), influenza-like illness (ILI), COVID-like illness and laboratory-confirmed influenza, SARS-CoV-2 or other viral pathogens).
- 2. Adverse events related to the intervention.

Secondary outcomes

- 1. Deaths.
- 2. Severity of viral respiratory illness as reported in the studies.
- 3. Absenteeism.
- 4. Hospital admissions.
- 5. Complications related to the illness, e.g. pneumonia.

Search methods for identification of studies

Electronic searches

For this 2022 update, we refined the original search strategy using a combination of previously included studies and automation tools (Clark 2020). We converted this search using the Polyglot Search Translator (Clark 2020), and ran the searches in the following databases:

- the Cochrane Central Register of Controlled Trials (CENTRAL) (2022, Issue 09), which includes the Acute Respiratory Infections Group's Specialised Register (searched 04 October 2022) (Appendix 1);
- 2. PubMed (01 January 2020 to 04 October 2022) (Appendix 2);
- 3. Embase (01 January 2020 to 04 October 2022) (Appendix 3);
- 4. CINAHL (Cumulative Index to Nursing and Allied Health Literature) (01 January 2020 to 04 October) (Appendix 4);
- 5. US National Institutes of Health Ongoing Trials Register ClinicalTrials.gov (January 2010 to 04 October 2022); and
- 6. World Health Organization International Clinical Trials Registry Platform (January 2010 to 04 October 2022).

We combined the database searches with the Cochrane Highly Sensitive Search Strategy for identifying randomised trials in MEDLINE: sensitivity- and precision-maximising version (2008 revision) (Lefebvre 2011). Details of previous searches are available in Appendix 5.



Searching other resources

We conducted a backwards-and-forwards citation analysis in Scopus on all newly included studies to identify other potentially relevant studies.

Data collection and analysis

Selection of studies

The search and citation analysis results were initially screened via the RobotSearch tool (Marshall 2018) to exclude all studies that were obviously not RCTs. We scanned the titles and abstracts of studies identified by the searches. We obtained the full-text articles of studies that either appeared to meet our eligibility criteria or for which there was insufficient information to exclude it. We then used a standardised form to assess the eligibility of each study based on the full article.

Data extraction and management

Five review authors (LA/GB/EF/EB/TOJ) independently applied the inclusion criteria to all identified and retrieved articles, and extracted data using a standard template that had been developed for and applied to previous versions of the review, but was revised to reflect our focus on RCTs and cluster-RCTs for this update. We resolved any disagreements through discussion with either PG or JMC acting as arbiter. We extracted and reported descriptions of interventions using the Template for Intervention Description and Replication (TIDieR) template (Table 1).

Assessment of risk of bias in included studies

Four review authors (EF/EB/GB/MJ) independently assessed risk of bias for the method of random sequence generation and allocation concealment (selection bias), blinding of participants and personnel (performance bias), blinding of outcome assessment (detection bias), outcome reporting (attrition bias), and selective reporting (reporting bias). In addition, for the cluster trials, we assessed selection bias due to how recruitment of participants was conducted. Participants should be identified before the cluster is randomised or, if not, recruitment should be by someone masked to the cluster allocation. Further, we considered whether there were sufficient numbers of clusters in each treatment group to ensure comparable groups, and excluded one study from the analysis due to insufficient number of clusters. We used the Cochrane risk of bias tool to assess risk of bias, classifying each risk of bias domain as 'low', 'high', or 'unclear'. The following were indications for low risk of bias:

- method of random sequence generation: the method was welldescribed and is likely to produce balanced and truly random groups;
- allocation concealment: the next treatment allocation was not known to participant/cluster or treating staff until after consent to join the study;
- 3. blinding of participants and personnel: the method is likely to maintain blinding throughout the study;
- 4. blinding of outcome assessors: all outcome assessors were unaware of treatment allocation;
- 5. outcome reporting: participant attrition throughout the study is reported, and reasons for loss are appropriately described; and
- 6. selective reporting: all likely planned and collected outcomes have been reported.

Measures of treatment effect

When possible, we performed meta-analysis and summarised effectiveness as risk ratio (RR) using 95% confidence intervals (CIs). For studies that could not be pooled, we used the effect measures reported by the trial authors (such as RR or incidence rate ratio (IRR) with 95% CI or, when these were not available, relevant P values). Where multiple analyses were reported on the same outcome we chose the analysis based on preferences for: (1) an adjusted analysis (over an unadjusted analysis), and (2) an analysis based on a longer follow-up period, or a greater number of outcomes events.

Unit of analysis issues

Many of the included studies were cluster-RCTs. To avoid any unit of analysis issues, we only included treatment effect estimates that were based on methods that were appropriate for the analysis of cluster trials, such as mixed models and generalised estimating equations. Given this restriction, we used the generalised inversevariance method of meta-analysis. Some cluster-RCTs that did not report cluster-adjusted treatment effects provided sufficient data (number of events and participants by treatment group and intraclass correlations) for us to calculate appropriate treatment effect estimates and standard errors using the methods described in the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2021a). For studies with multiple treatment groups but only one control group, where appropriate, we adjusted standard errors upwards to avoid unit of analysis errors in the meta-analyses. We did this by splitting the control group into equal sized groups and adjusting standard errors upwards to account for the reduced sample size of the control subgroups (Higgins 2021b).

Dealing with missing data

Previously, whenever details of studies were unclear, or studies were only known to us by abstracts or communications at meetings, we corresponded with first or corresponding authors. For this 2022 review, we did not contact authors of studies.

Assessment of heterogeneity

Aggregation of data was dependent on types of comparisons, sensitivity and homogeneity of definitions of exposure, populations and outcomes used. We calculated the I²statistic and Chi² test for each pooled estimate to assess the presence of statistical heterogeneity (Higgins 2002; Higgins 2003).

Assessment of reporting biases

Given the widely disparate nature of our evidence base, we limited our assessment of possible reporting biases to funnel plot visual inspection if we had > 10 included studies for any single meta-analysis.

Data synthesis

If possible and appropriate, we combined studies in a metaanalysis. We used the generalised inverse-variance random-effects model where cluster-RCTs were included in the analysis. We chose the random-effects model because we expected clinical heterogeneity due to differences in pooled interventions and outcome definitions, and methodological heterogeneity due to pooling of RCTs and cluster-RCTs.



Subgroup analysis and investigation of heterogeneity

We conducted one post hoc subgroup analyses of adults (18 years +) versus children (0 to 18 years) for the comparison of hand hygiene versus control.

We did not conduct further investigation of heterogeneity due to insufficient numbers of studies included in the comparisons.

Sensitivity analysis

We conducted a sensitivity analysis for hand hygiene versus control where we included the most precise and unequivocal measure of viral respiratory illness reported for each included study.

Summary of findings and assessment of the certainty of the evidence

We created three summary of findings tables using the following outcomes: numbers of cases of viral respiratory illness (including ARIs, ILI, COVID-like illness and laboratory-confirmed influenza/SARS-CoV-2 or other respiratory viruses), and adverse events related to the intervention (Summary of findings 1; Summary of findings 2; Summary of findings 3). We planned to include the secondary outcomes of deaths; severity of viral respiratory illness as reported in the studies; absenteeism; hospital admissions; and complications related to the illness (e.g. pneumonia). However, these data were poorly reported in the included studies. We used the five GRADE considerations (study limitations, consistency of effect, imprecision, indirectness, and publication bias) to assess the certainty of evidence as it related to the studies which contributed

data to the meta-analyses for the prespecified outcomes (Atkins 2004). We used the methods and recommendations described in Section 8.5 and Chapter 12 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011), employing GRADEpro GDT software (GRADEpro GDT). We justified all decisions to down- or upgrade the certainty of the evidence in footnotes, and made comments to aid the reader's understanding of the review where necessary.

RESULTS

Description of studies

See Characteristics of included studies and Characteristics of excluded studies tables. Five trials were funded by government and pharmaceutical companies (Aiello 2010; Aiello 2012; Chard 2019; Yeung 2011; Zomer 2015), and nine trials were funded by pharmaceutical companies (Arbogast 2016; Carabin 1999; Luby 2005; Nicholson 2014; Sandora 2005; Sandora 2008; Turner 2004a; Turner 2012).

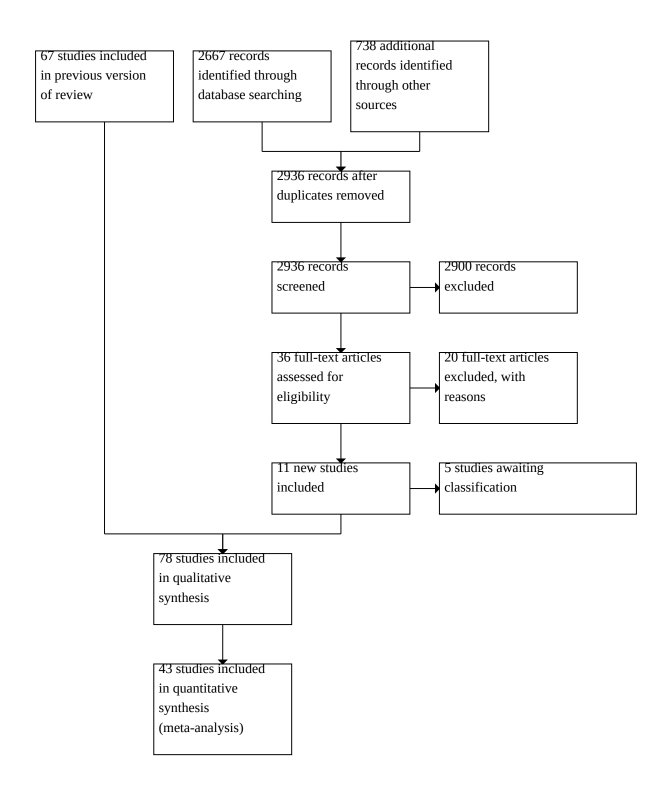
Results of the search

For this 2022 update we found 2667 records through database and trial registry searching, as well as 738 record through citation searching. After removing duplicates we had 2936 records that underwent title and abstract screening.

We identified a total of 202 titles in this 2022 update. We excluded 180 titles and retrieved the full papers of 35 studies, to include 11 new studies. See Figure 1.



Figure 1. Study flow diagram.



Included studies

In this 2022 update we included 11 new studies (610,872 participants); randomised controlled trials (RCTs) (n = 5) or cluster-

RCTs (n = 6) published between 2020 and 2022. In total 78 studies are included in this review update. For detailed descriptions of the interventions of the included studies, see Table 1.



Eighteen trials focused on using masks (Abaluck 2022; Aiello 2010; Aiello 2012; Alfelali 2020; Barasheed 2014; Bundgaard 2021; Canini 2010; Cowling 2008; Ide 2016; Jacobs 2009; Loeb 2009; MacIntyre 2009; MacIntyre 2011; MacIntyre 2013; MacIntyre 2015; MacIntyre 2016; Radonovich 2019; Suess 2012). Thirteen of the 18 trials compared medical/surgical masks to no mask (control) (Abaluck 2022; Aiello 2010; Aiello 2012; Alfelali 2020; Barasheed 2014; Bundgaard 2021; Canini 2010; Cowling 2008; Jacobs 2009; MacIntyre 2009; MacIntyre 2015; MacIntyre 2016; Suess 2012). One study compared catechin-treated masks to no mask (Ide 2016), and one study included cloth masks versus control (third arm in MacIntyre 2015). Three of the 18 trials were in healthcare workers (Ide 2016; Jacobs 2009; MacIntyre 2015), whilst the remaining trials were in non-healthcare workers (students, households, families, or pilgrims). Only one trial was conducted during the H1N1 pandemic season (Suess 2012), and two trials were conducted during the SARS-CoV-2 pandemic (Abaluck 2022; Bundgaard 2021).

Five of the 18 trials compared N95 masks or P2 masks to medical/ surgical masks (Loeb 2009; MacIntyre 2009; MacIntyre 2011; MacIntyre 2013; Radonovich 2019). All of these trials, except for one study that was conducted on household individuals (MacIntyre 2009), included healthcare workers either in a hospital setting, Loeb 2009; MacIntyre 2011; MacIntyre 2013, or an outpatient setting (MacIntyre 2009; Radonovich 2019).

One trial evaluated the effectiveness of quarantining workers of one of two sibling companies in Japan whose family members had developed an influenza-like illness (ILI) during the 2009 to 2010 H1N1 influenza pandemic (Miyaki 2011). Another trial conducted during the SARS-CoV-2 pandemic in Norway investigated fitness centre access with physical distancing compared to no access (Helsingen 2021); and one cluster trial compared daily testing for contacts of individuals with SARS-CoV-2 compared to self-isolation at home in English secondary schools (Young 2021).

Nineteen trials compared hand hygiene interventions with no hand hygiene (control) and provided data suitable for meta-analysis. The populations in these trials included adults, children, and families, in settings such as schools (Biswas 2019; Stebbins 2011), childcare centres (Azor-Martinez 2018; Correa 2012; Roberts 2000; Zomer 2015), homes/households (Cowling 2008; Cowling 2009; Larson 2010; Little 2015; Nicholson 2014; Ram 2015; Sandora 2005; Simmerman 2011), offices (Hubner 2010), military trainees (Millar 2016), villages (Ashraf 2020; Swarthout 2020), and nursing homes (Teesing 2021). None of the trials were conducted during a pandemic, although some of the studies were conducted during peak influenza seasons.

A further 10 trials that compared a variety of hand hygiene modalities to control provided insufficient information to include in meta-analyses. Three trials were in children: one was conducted in daycare centres in Denmark examining a multimodal hygiene programme (Ladegaard 1999), and two trials compared a hand hygiene campaign or workshop in an elementary school environment in Saudi Arabia, Alzaher 2018, and Egypt, Talaat 2011. Three trials tested virucidal hand treatment in an experimental setting, Gwaltney 1980; Turner 2004a, and in a community, Turner 2012, in the USA. Feldman 2016 compared hand-washing with chlorhexidine gluconate amongst Israeli sailors. One trial compared hand sanitiser packaged in a multimodal hygiene programme amongst office employees in the USA (Arbogast 2016). Two trials were conducted in a long-term facility setting: one trial

examined the effect of a bundled hand hygiene programme on infectious risk in nursing home residents in France (Temime 2018), and the other trial compared the effect of using hand sanitisers in healthcare workers on the rate of infections (including respiratory infections) in nursing home residents in Hong Kong (Yeung 2011).

Five trials compared different hand hygiene interventions in a variety of settings such as schools (Morton 2004, in kindergartens and elementary schools in the USA; Priest 2014, in primary schools in New Zealand; and Pandejpong 2012 in kindergartens in Thailand). One study was conducted in low-income neighbourhoods in Karachi, Pakistan (Luby 2005), and one was conducted in a workplace environment in Finland (Savolainen-Kopra 2012). A variety of interventions were used across these trials such as soap and water (Luby 2005; Savolainen-Kopra 2012), hand sanitiser (Morton 2004; Pandejpong 2012; Priest 2014; Savolainen-Kopra 2012), body wash (Luby 2005), and alcohol-based hand wipes (Morton 2004), with or without additional hygiene education. There was considerable variation in interventions, and the information in the trial reports was insufficient to permit meta-analysis.

Seven trials compared a combined intervention of hand hygiene and face masks with control. Four of these trials were carried out in households in Germany (Suess 2012), Thailand (Simmerman 2011), Hispanic immigrant communities in the USA (Larson 2010), and households in Hong Kong (Cowling 2009). Two trials were conducted amongst university student residences (Aiello 2010; Aiello 2012), and two trials in groups of pilgrims at the annual Hajj (Aelami 2015; Alfelali 2020). Moreover, six trials evaluated the incremental benefit of combining surgical masks in addition to hand hygiene with soap (Simmerman 2011), hand sanitiser (Aiello 2010; Aiello 2012; Larson 2010; Suess 2012), or both (Cowling 2009), versus mask or hand hygiene alone on the outcomes of ILI and influenza. Aelami 2015 investigated a hygienic package (alcoholbased hand rub (gel or spray), surgical masks, soap, and paper handkerchiefs) with a control group.

Seven trials compared a multimodal combination of hand hygiene and disinfection of surfaces, toys, linen, or other components of the environment with a control (Ban 2015; Carabin 1999; Ibfelt 2015; Kotch 1994; McConeghy 2017; Sandora 2008; White 2001). Variation in scope and type of interventions and insufficient data in trial reports precluded meta-analysis. All studies except for one were in children (McConeghy 2017), which was in a nursing home population).

Three trials included in two papers investigated the role of virucidal tissues in interrupting transmission of naturally occurring respiratory infections in households (Farr 1988a; Farr 1988b; Longini 1988). Four cluster-RCTs implemented complex, multimodal sanitation, education, cooking, and hygiene interventions (Chard 2019; Hartinger 2016; Huda 2012; Najnin 2019). All four of these trials were conducted in low-income countries in settings with minimal to no access to basic sanitation.

Three trials assessed the effect of gargling on the incidence of upper respiratory tract infections (URTIs) or influenza: gargling with povidone-iodine (Satomura 2005), green tea (Ide 2014), and tap water (Goodall 2014). Two trials investigated the use of mouth/nasal washes on the incidence of SARS-CoV-2 infection in healthcare workers during the COVID-19 pandemic (Almanza-Reyes 2021; Gutiérrez-García 2022). One trial investigated the use of glasses against the transmission of SARS-CoV-2 (Fretheim 2022a).



Ongoing studies

We identified four ongoing studies during the course of the COVID-19 pandemic, of which one is completed, but unreported (NCT04471766). The trials evaluated masks concurrent with the COVID-19 pandemic. Three trials on other interventions are ongoing (Brass 2021; NCT03454009; NCT04267952).

Studies awaiting classification

We identified five studies awaiting classification (Contreras 2022; Croke 2022; Delaguerre 2022; Loeb 2022; Varela 2022).

A previous RCT (NCT04296643) reported as ongoing in the last version has now been recently published but was not able to be included in the summary of findings pooled results (Loeb 2022). In a multicentre, randomised non-inferiority trial of 1009 healthcare workers (HCWs) across four countries randomised to medical mask versus fit-tested N95 respirators for direct care of COVID-19 patients or long-term care residents, laboratory-confirmed SARS-CoV-2 was found in 10.46% (52/497) versus 9.27% (47/507) in the medical/ surgical mask group and fit-tested N95 respirator group (hazard ratio 1.14 (95% CI 0.77 to 1.69), respectively. There was a 1.19% absolute increase in risk of COVID-19 with medical masks versus N95 respirator 95% CI (-2.5% to 4.9%). There were 47 (10.8%) adverse events related to the intervention reported in the medical mask group and 59 (13.6%) in the N95 respirator group. The use of medical masks was found to be non-inferior to N95 respirators in the direct care of COVID-19 patients and the study crossed over into the more transmissible Omicron variant period of the COVID-19 pandemic.

Excluded studies

We excluded a total of 180 studies. We identified 20 new studies for exclusion at the data extraction stage of this 2022 update, all of which appeared to be eligible at screening. Five of the 20

studies were ineligible due to evaluating treatments for patients with disease (Cyril Vitug 2021; Ferrer 2021; Meister 2022; Sanchez Barrueco 2022; Sevinc Gul 2022), two were excluded because they did not assess clinical outcomes (Costa 2021; Seneviratne 2021), four were excluded due to not assessing viral outcomes (Gharebaghi 2020; Giuliano 2021; Karakaya 2021; Kawyannejad 2020), five were excluded as they were experiments that did not measure any of our outcomes of interest (Ahmadian 2022; Dalakoti 2022; Egger 2022; Malaczek 2022; Montero-Vilchez 2022); three were excluded because they were not RCTs (Chen 2022; Lim 2022; Mo 2022), and one was excluded as it was a report of another study (Munoz-Basagoiti 2022).

Risk of bias in included studies

The overall risk of bias is presented graphically in Figure 2 and summarised by included study in Figure 3. Details on the judgements can be found in the descriptions of individual included studies (Characteristics of included studies table). Out of 78 included studies, only two were rated as low risk of bias for all domains. One of those studies compared two different types of masks (Radonovich 2019), and the other compared hand sanitiser to no treatment (Turner 2012). Notably, neither of these two studies was blinded, however, trial procedures were sufficiently robust that the risk of performance bias was low. Overall, approximately only 20% of the studies were rated as low risk of performance bias. This risk of bias domain was particularly problematic because most interventions studied could not be blinded from participants and/ or investigators. The two risks of bias domains that were rated the least problematic were attrition bias and random sequence generation where around 50% of studies were rated as low risk of bias. Allocation concealment, blinded outcome assessment and selective reporting were rated as low risk of bias for around 40% of the included studies. Many of the included studies were cluster-RCTs where the randomisation process was not well reported leading to ratings of unclear risk of bias.

Figure 2. 'Risk of bias' graph: review authors' judgements about each risk of bias item presented as percentages across all included trials.

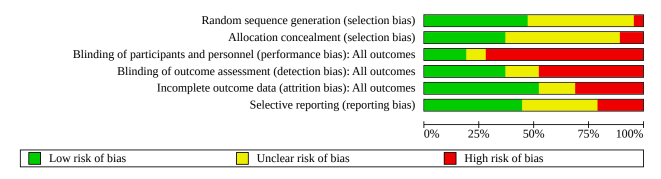




Figure 3. 'Risk of bias' summary: review authors' judgements about each risk of bias item for each included trial.

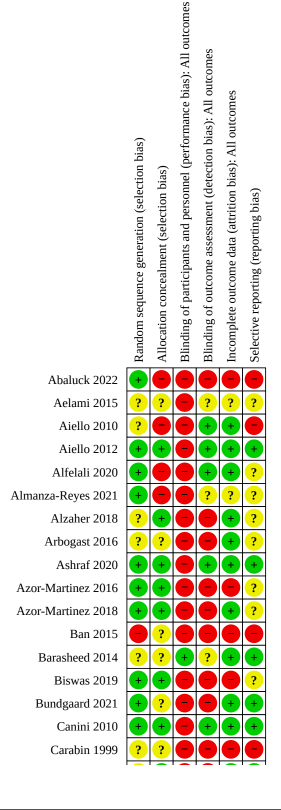




Figure 3. (Continued)

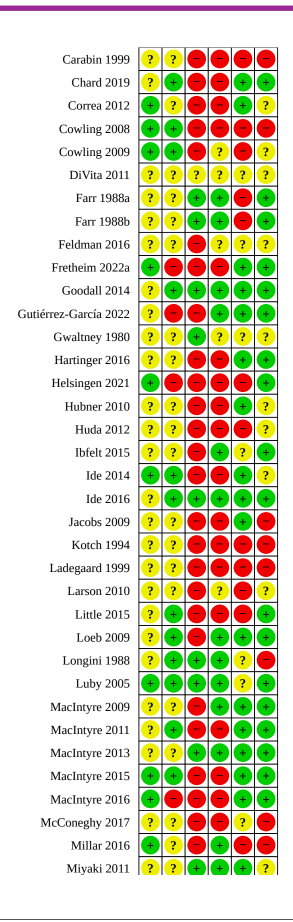
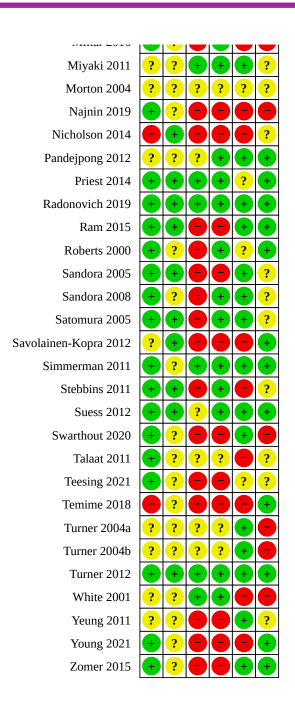




Figure 3. (Continued)



Allocation

For this 2022 review, 10 of the 11 newly included studies provided adequate information on randomisation and were judged to have low risk of bias (Abaluck 2022; Alfelali 2020; Almanza-Reyes 2021; Ashraf 2020; Bundgaard 2021; Fretheim 2022a; Helsingen 2021; Swarthout 2020; Teesing 2021; Young 2021). Six of these studies described the use of a computerised random number generator (Almanza-Reyes 2021; Bundgaard 2021; Helsingen 2021; Swarthout 2020; Teesing 2021; Young 2021). Almanza-Reyes 2021 described the use of computer-generated stratified block scheme, while Bundgaard 2021 reported the use of a computer algorithm stratified by the five regions of Denmark. In Fretheim 2022a, the investigators used a digital platform (Nettskjema)

for recruitment, randomisation and allocation. Three studies mentioned the use of a random number generator, with no additional specifics (Helsingen 2021; Swarthout 2020; Teesing 2021), while Young 2021 mentioned that randomisation was performed in blocks of two and stratified using nine strata to ensure a sample representative of schools and colleges in England. Abaluck 2022 reported pairwise cross randomisation, whilst Ashraf 2020 reported using a block random number generator. Alfelali 2020 described using coin-tossing by an individual who was not a member of the research team (i.e. a fellow pilgrim who was not a participant in the trial, a tour operator, or a medical volunteer). One study provided insufficient information to judge the sequence generation bias (Gutiérrez-García 2022).



The success of randomisation was judged as low risk of bias in one study only that used an off-site investigator to allocate groups (Ashraf 2020). Four new studies provided insufficient information to make a judgment on the adequacy of the process (Bundgaard 2021; Swarthout 2020; Teesing 2021; Young 2021). The remaining six newly included studies were judged as high risk of allocation bias (Abaluck 2022; Alfelali 2020; Almanza-Reyes 2021; Fretheim 2022a; Gutiérrez-García 2022; Helsingen 2021). In Abaluck 2022, there was a significant difference in the numbers of households included in each treatment group, suggestive of a lack of allocation concealment. Alfelali 2020 used coin tossing, which can lead to a large imbalance. In Almanza-Reyes 2021 baseline prognostic factors (vaccination and frequency of handwashing) were unbalanced between the two arms. In Fretheim 2022a, a higher number of participants used face masks in the intervention group. In Gutiérrez-García 2022 there as a significant age difference between the two groups. Helsingen 2021 described assigning the randomised sequence by a member of the research team, with no further description.

For the review published in 2020, information on sequence generation was overall poorly reported in most of the included studies. Nineteen of the included studies provided adequate information on the randomisation scheme and were judged as at low risk of bias (Aiello 2012; Azor-Martinez 2016; Azor-Martinez 2018; Biswas 2019; Canini 2010; Correa 2012; Ide 2014; MacIntyre 2015; MacIntyre 2016; Millar 2016; Najnin 2019; Radonovich 2019; Ram 2015; Simmerman 2011; Stebbins 2011; Suess 2012; Talaat 2011; Turner 2012; Zomer 2015). Nine studies described the use of computerised sequence generation program/software (Aiello 2012; Azor-Martinez 2018; Biswas 2019; Canini 2010; Millar 2016; Najnin 2019; Radonovich 2019; Talaat 2011; Turner 2012). One study used random number tables for sequence generation (Azor-Martinez 2016). Three studies described using the random function in Microsoft Excel (Microsoft Excel 2018) (Correa 2012; MacIntyre 2016; Suess 2012). Two studies used statistical software to generate a randomisation allocation (MacIntyre 2015; Priest 2014). Two studies reported using block randomisation: Ram 2015 used block randomisation, and an independent investigator-generated the list of random assignments, whilst Simmerman 2011 performed block randomisation. Stebbins 2011 used constrained randomisation, and Zomer 2015 reported using stratified randomisation by means of computer generation with a 1:1 ratio in each of the strata.

Fourteen studies reported insufficient information to permit a judgement on the adequacy of the process to minimise selection bias (Aelami 2015; Alzaher 2018; Arbogast 2016; Barasheed 2014; Chard 2019; DiVita 2011; Feldman 2016; Hubner 2010; Ibfelt 2015; McConeghy 2017; Miyaki 2011; Pandejpong 2012; Savolainen-Kopra 2012; Yeung 2011). Six studies provided some description about sequence generation, but it was still unclear (Hartinger 2016; Huda 2012; Ide 2016; Little 2015; MacIntyre 2011; MacIntyre 2013). Huda 2012 mentioned random number tables, but it was unclear if this was for random selection or randomisation. Ide 2016 used computer-generated randomisation, but the method was not stated. Hartinger 2016 used covariate-constrained randomisation, but the method was not described. In Little 2015, participants were automatically randomly assigned by the intervention software, but the sequence generation was not described. Two studies used a secure computerised randomisation program (MacIntyre 2011; MacIntyre 2013), but the sequence generation was not described.

Three of the studies included in the 2020 review, were poorly randomised (Ban 2015; Nicholson 2014; Temime 2018). Ban 2015 included only two clusters, and the randomisation scheme was not reported. Nicholson 2014 used coin tossing, which can lead to a large imbalance. Temime 2018 used "simple randomisation" with no further description.

For the RCTs included in previous versions of the review, three were poorly reported with no description of randomisation sequence or concealment of allocation (Gwaltney 1980; Turner 2004a; Turner 2004b). The quality of the cluster-RCTs varied, with four studies not providing a description of the randomisation procedure (Carabin 1999; Kotch 1994; Morton 2004; White 2001). We rated seven studies as at low risk of bias for sequence generation (Cowling 2008; Cowling 2009; Luby 2005; Roberts 2000; Sandora 2005; Sandora 2008; Satomura 2005), and a further six studies as at unclear risk of bias (Farr 1988a; Farr 1988b; Ladegaard 1999; Loeb 2009; Longini 1988; MacIntyre 2009).

Many of the newly included cluster-RCTs did not report adequately on allocation concealment. Twenty-one of these studies reported adequate allocation and were judged as at low risk of bias (Aiello 2012; Alzaher 2018; Azor-Martinez 2016; Azor-Martinez 2018; Biswas 2019; Canini 2010; Chard 2019; Goodall 2014; Ide 2014; Ide 2016; Little 2015; MacIntyre 2011; MacIntyre 2015; Nicholson 2014; Priest 2014; Radonovich 2019; Ram 2015; Savolainen-Kopra 2012; Stebbins 2011; Suess 2012; Turner 2012). Aiello 2012 randomised all residence houses in each of the residence halls prior to the intervention implementation. Alzaher 2018 allocated schools prior to all schoolgirls attending selected schools being invited to participate. Azor-Martinez 2016 allocated schools/classes prior to children's recruitment. Azor-Martinez 2018 assigned clusters prior to recruitment. Biswas 2019 completed the allocation prior to individuals being recruited. Chard 2019 allocated schools prior to individuals being recruited. Goodall 2014 used opaque, sealed, serially numbered envelopes that were only accessed when two study personnel were present. Ide 2014 also reported using individual drawing of sealed, opaque envelopes to randomly assign participants to the study groups. MacIntyre 2011 randomised hospitals prior to inclusion of participants. In MacIntyre 2015, hospital wards were randomised prior to recruitment of individuals. Nicholson 2014 used coin tossing to assign communities to intervention or control arms. Radonovich 2019 used constrained randomisation to resolve any potential imbalance between covariates between the trial arms. Four studies reported the use of central randomisation: Canini 2010 used central randomisation by employing an interactive voice response system; Ide 2016 used central randomisation services; Little 2015 participants were automatically randomly assigned by the intervention software; and Ram 2015 described a central allocation through data collectors notifying the field research officer, who consulted the block randomisation list to make the assignment of the household compound to intervention or control. Savolainen-Kopra 2012 randomised clusters by matching prior to the onset of the interventions. Four studies reported that allocation was assigned by personnel (investigator, physician, or statistician) unaware of the randomisation sequence (Priest 2014; Stebbins 2011; Suess 2012; Turner 2012). Twenty-two studies reported insufficient information to permit a judgement on the adequacy of the process to minimise selection bias (Aelami 2015; Arbogast 2016; Ban 2015; Barasheed 2014; Correa 2012; DiVita 2011; Feldman 2016; Hartinger 2016; Hubner 2010; Huda 2012; Ibfelt 2015; MacIntyre



2013; McConeghy 2017; Millar 2016; Miyaki 2011; Najnin 2019; Pandejpong 2012; Simmerman 2011; Talaat 2011; Temime 2018; Yeung 2011; Zomer 2015). Two studies provided some information about allocation, but it was not enough to permit a judgement on the risk of bias (Barasheed 2014; Simmerman 2011). Barasheed 2014 randomised pilgrim tents using an independent study coordinator who was not an investigator, but did not describe how this was done. Simmerman 2011 described using a study coordinator to assign households to the study arm (after consent was obtained). Only one of the newly added studies was judged as at high risk of bias, where the random assignment was allocated by doctors enrolling the participants (MacIntyre 2016). Of the previously included RCTs, 14 provided no or an insufficient description of concealment of allocation (Carabin 1999; Farr 1988a; Farr 1988b; Gwaltney 1980; Kotch 1994; Ladegaard 1999; Larson 2010; MacIntyre 2009; Morton 2004; Roberts 2000; Sandora 2008; Turner 2004a; Turner 2004b; White 2001). We assessed all of the remaining studies as at low risk of bias (Canini 2010; Cowling 2008; Cowling 2009; Loeb 2009; Longini 1988; LLuby 2005; Sandora 2005; Satomura 2005). Aiello 2010 used the drawing of a uniform ticket with the name of each hall out of a container and was rated as at high risk of bias.

Blinding

Although blinding is less of a concern in cluster-RCTs, the risk of bias is substantial when the outcomes are subjective and the outcome assessor is not blinded.

In this 2022 review, five RCTs (Almanza-Reyes 2021; Bundgaard 2021; Fretheim 2022a; Gutiérrez-García 2022; Helsingen 2021), and six cluster-RCTs were all judged to have a high risk of detection bias (Abaluck 2022; Alfelali 2020; Ashraf 2020; Swarthout 2020; Teesing 2021; Young 2021).

We judged two of the newly included studies to have a low risk of detection bias as the outcome is laboratory-confirmed (Alfelali 2020; Gutiérrez-García 2022). One study provided insufficient information to enable judgment (Almanza-Reyes 2021). The remaining eight of the 11 new studies have a high risk of detection bias (Abaluck 2022; Ashraf 2020; Bundgaard 2021; Fretheim 2022a; Helsingen 2021; Swarthout 2020; Teesing 2021; Young 2021). In Abaluck 2022, investigators dropped individuals for whom symptom data were missing. In addition, other outcomes were subjective and can be influenced by the unblinded mask promoters, and mask surveillance staff. Moreover, blood testing in the protocol specified baseline testing which was not done, and no further explanation was provided. In Ashraf 2020, although the data collection team was separate from the intervention team, they were not blinded, and the outcome was respiratory illness measured through caregiver-reported symptoms. In Bundgaard 2021, case detection was based on patient-reported symptoms on home tests. In Fretheim 2022a, the outcome was self-reported positive COVID-19 test result, notified to the Norwegian Surveillance System for Communicable Diseases (MSIS). However, the public policy requiring confirmatory PCR-test had changed during the study, which may have affected reporting. In Helsingen 2021, although the outcome was a positive test for COVID-19 based on SARS-CoV-2 ribonucleic acid, the samples were collected and sent by participants, and there was a difference in adherence in testing between the two groups. Swarthout 2020, Teesing 2021, and Young 2021 all had subjective outcomes and assessors were not blinded. As for the detection bias, six of the newly included studies were

considered to have a high risk of detection bias (Bundgaard 2021; Gutiérrez-García 2022; Helsingen 2021; Swarthout 2020; Teesing 2021; Young 2021. In Bundgaard 2021, case detection was based on patient-reported symptoms and results from home point-of-care (POCT) testing. The primary outcome of Gutiérrez-García 2022 was participants' self-reported symptoms. Case detection in Helsingen 2021 was based on a home-test kit. Swarthout 2020, Teesing 2021, and Young 2021 had subjective outcomes.

In the 2020 review, we judged 36 studies to have a high risk of bias (Aiello 2012; Abaluck 2022; Alfelali 2020; Almanza-Reyes 2021; Alzaher 2018; Arbogast 2016; Ashraf 2020; Azor-Martinez 2016; Azor-Martinez 2018; Ban 2015; Biswas 2019; Bundgaard 2021; Carabin 1999; Chard 2019; Correa 2012; Cowling 2008; Gutiérrez-García 2022; Helsingen 2021; Ide 2014; Kotch 1994; Ladegaard 1999; Little 2015; MacIntyre 2011; MacIntyre 2015; MacIntyre 2016; McConeghy 2017; Najnin 2019; Nicholson 2014; Ram 2015; Sandora 2008; Savolainen-Kopra 2012; Swarthout 2020; Teesing 2021; Temime 2018; Young 2021; Zomer 2015). We assessed five cluster-RCTs as at low risk of bias. Farr 1988a and Farr 1988b were double-blinded studies and were judged as at low risk of bias. MacIntyre 2013 and Simmerman 2011 reported laboratoryconfirmed influenza, and blinding would not have affected the result. In Miyaki 2011 the self-reported respiratory symptoms were confirmed by a physician.

We judged four cluster-RCTs to have a low risk of detection bias because the outcome was laboratory-confirmed influenza (Alfelali 2020; Barasheed 2014; Suess 2012), or physician-confirmed ILI, Pandejpong 2012. Another two cluster-RCTs were judged to have a low risk of bias because outcome assessors were blinded (Abaluck 2022; Ashraf 2020). One RCT (Almanza-Reyes 2021) and two cluster-RCTs (Talaat 2011; Yeung 2011) provided insufficient data to judge the effect of non-blinding. Talaat 2011 included outcomes that were both self-reported ILI and laboratoryconfirmed influenza. In Yeung 2011 the detection of cases was based on records for hospitalisation related to infection (including pneumonia). Eleven cluster-RCTs were not blinded, but we judged the primary outcome to be unaffected by non-blinding. Seven trials reported laboratory-confirmed influenza (Aiello 2012; Cowling 2009; Larson 2010; Loeb 2009; MacIntyre 2009; Millar 2016; Stebbins 2011). Four studies reported self-reported outcomes (Canini 2010; Priest 2014; Roberts 2000; Sandora 2008), but outcome assessors were not aware of the intervention assignment. Five RCTs were double-blinded and were judged as at low risk of bias (Goodall 2014; Ide 2016; Longini 1988; Luby 2005; White 2001), whilst two studies were single-blinded where investigators, Radonovich 2019, or laboratory personnel, Turner 2012, were blinded. Four RCTs were not blinded and were judged as at high risk of bias given the subjective nature of the outcome assessed (Hubner 2010; Ibfelt 2015; Jacobs 2009; Satomura 2005). Turner 2004a and Turner 2004b were double-blind studies, but insufficient information was provided to assess the risk of bias.

Incomplete outcome data

In this 2022 review, six of the 11 newly included studies had reasonable attrition and provided sufficient evidence about participant flow throughout the study and reasons of loss to follow-up, and hence were assessed as having a low risk of attrition bias (Alfelali 2020; Ashraf 2020; Bundgaard 2021; Fretheim 2022a; Gutiérrez-García 2022; Swarthout 2020). Two studies provided insufficient information to assess the attrition risk (Almanza-



Reyes 2021; Teesing 2021). The remaining three studies were judged at high risk of attrition bias. In Abaluck 2022, laboratory testing results were only available for 40% of the symptomatic participants. In Helsingen 2021, more people in the control group withdrew from the study and reasons for withdrawal were not provided. In the Young 2021 study there was high attrition at different rates between the two groups.

In the 2020 review, we assessed 26 newly included trials as having a low risk of attrition bias, with sufficient evidence from the participant flow chart, and explanation of loss to follow-up (which was minimal) similar between groups (Aiello 2012; Alzaher 2018; Arbogast 2016; Azor-Martinez 2018; Barasheed 2014; Canini 2010; Chard 2019; Correa 2012; Goodall 2014; Hartinger 2016; Hubner 2010; Ide 2014; Ide 2016; MacIntyre 2011; MacIntyre 2013; MacIntyre 2015; MacIntyre 2016; Miyaki 2011; Pandejpong 2012; Radonovich 2019; Ram 2015; Simmerman 2011; Suess 2012; Turner 2012; Yeung 2011; Zomer 2015). Seven studies did not report sufficient information on incomplete data (attrition bias) (Aelami 2015; DiVita 2011; Feldman 2016; Hartinger 2016; Ibfelt 2015; McConeghy 2017; Priest 2014). Twelve studies had a high risk of attrition bias (Azor-Martinez 2016; Ban 2015; Biswas 2019; Huda 2012; Little 2015; Millar 2016; Najnin 2019; Nicholson 2014; Savolainen-Kopra 2012; Stebbins 2011; Talaat 2011; Temime 2018). In Azor-Martinez 2016, attrition levels were high and differed between the two groups. Ban 2015 did not report on reasons for loss to follow-up. Biswas 2019 did not provide information on missing participants (28 children in the control schools and two children in the intervention schools). Huda 2012 did not provide a flow diagram of study participants. Little 2015 had high attrition that differed between the two groups. Attrition in Millar 2016 differed amongst the three groups. In addition, ARI cases were captured utilising clinic-based medical records for those participants who sought hospital care only. In Najnin 2019, there was high migration movement during the study, which could have distorted the baseline characteristics even more. There was no description of how such migration and changes in the intervention group were dealt with. In Nicholson 2014, households were removed from the study if they provided no data for five consecutive weeks. Although attrition was reported in Savolainen-Kopra 2012, and 76% of volunteers who were recruited at the beginning of the reporting period completed the study, new recruits were added during the study to replace volunteers lost in most clusters. The total number of reporting participants at the end of the trial was 626 (91.7%) compared to the beginning, meaning that 15.7% of participants were replaced during the study. In Stebbins 2011, reasons for episodes of absence in 66% of the study participants were not reported. Talaat 2011 did not provide a flow chart of clusters flow during the study period and provided no information on withdrawal. Temime 2018 was greatly biased due to underreporting of outcomes in the control groups. Furthermore, no study flow chart was provided, and there was no reporting on any exclusions.

Selective reporting

For this 2022 review update, six of the 11 newly included studies reported all specified outcomes and were judged to have a low risk of selective reporting (Ashraf 2020; Bundgaard 2021; Fretheim 2022a; Gutiérrez-García 2022; Helsingen 2021; Young 2021). Three studies had no published protocol and were considered to have an unclear risk of selective reporting (Alfelali 2020; Almanza-Reyes 2021; Teesing 2021). The remaining two new included studies are considered to have a high risk of bias

in this domain. Abaluck 2022 did not report on prespecified seroconversion, while in Swarthout 2020, none of the outcomes reported were prespecified in the trial registry.

In the 2020 review, 22 included studies reported all specified outcomes and were judged as at low risk of reporting bias (Aiello 2012; Barasheed 2014; Canini 2010; Chard 2019; Goodall 2014; Hartinger 2016; Ibfelt 2015; Ide 2016; Little 2015; MacIntyre 2011; MacIntyre 2013; MacIntyre 2015; MacIntyre 2016; Pandejpong 2012; Priest 2014; Radonovich 2019; Savolainen-Kopra 2012; Simmerman 2011; Suess 2012; Temime 2018; Turner 2012; Zomer 2015). For 18 studies, it is unlikely that other outcomes were measured and not reported, although no protocol was available to assess reporting bias (Aelami 2015; Alzaher 2018; Arbogast 2016; Azor-Martinez 2016; Azor-Martinez 2018; Ban 2015; Biswas 2019; Correa 2012; DiVita 2011; Feldman 2016; Hubner 2010; Huda 2012; Ide 2014; Miyaki 2011; Nicholson 2014; Stebbins 2011; Talaat 2011; Yeung 2011). Three studies were at high risk of reporting bias (McConeghy 2017; Millar 2016; Najnin 2019). In McConeghy 2017, URTI was mentioned in the methods (the intervention presumably would have targeted these), but only lower respiratory tract infection (LRTI) and overall infection were reported. Millar 2016 was originally conducted for another purpose; we could not find the respiratory outcomes reported in the study as part of the original study protocol. In Najnin 2019, the published study protocol did not include respiratory illness as an outcome.

Other potential sources of bias

An additional consideration for cluster-RCTs is identification/recruitment bias, where individuals are recruited in the trial after clusters are randomised. Such bias can introduce an imbalance amongst groups.

In this 2022 review, of the six cluster-RCTs included, we judged four to have a low risk of identification/recruitment bias (Abaluck 2022; Ashraf 2020; Swarthout 2020; Teesing 2021). In Abaluck 2022, all of people in the village were assigned to one study arm (control, cloth mask or surgical mask villages). In_Ashraf 2020, participants were unaware of their intervention group assignment until after the baseline survey and randomisation. In Swarthout 2020, village clusters comprised of 12 enrolled households, while in Teesing 2021 randomisation was done per nursing home. Alfelali 2020 recruited individuals after cluster-randomisation and is judged to have a high risk of recruitment bias, while in Young 2021, participation of students and staff contacts were made after random assignment of the school through written consent or electronic completion of a consent form.

Of the cluster-RCTs included in our 2020 review, we judged 13 to have a low risk of identification/recruitment bias (Arbogast 2016; Biswas 2019; Canini 2010; Cowling 2008; Longini 1988; Luby 2005; MacIntyre 2015; MacIntyre 2016; Roberts 2000; Sandora 2005; Suess 2012; Temime 2018; White 2001). In Arbogast 2016, all identified individuals (office workers) were included in the assigned cluster. Schools were identified and then randomised to the clusters; students were then randomly selected from each classroom and school. Nine studies described the identification of participants, consenting/enrolling, and then randomising to the clusters (Canini 2010; Cowling 2008; Longini 1988; Luby 2005; MacIntyre 2015; MacIntyre 2016; Roberts 2000; Sandora 2005; White 2001). Suess 2012 identified and consented patients, then recruitment was performed by physicians unaware of cluster assignment. In Temime



2018, directors of the included nursing homes agreed to participate in the study before randomisation, and written consent was not required from the residents.

Amongst the newly included studies, we judged four cluster-RCTs as at low risk of identification/recruitment bias (Abaluck 2022; Swarthout 2020; Teesing 2021; Young 2021). In Abaluck 2022, the village was the unit of randomisation and all households received one arm of the study (control, surgical mask or cloth mask). In Swarthout 2020, village clusters were each randomised by blocks (group of nine adjacent clusters) into eight groups. In Teesing 2021 nursing homes were computer randomised after baseline hand hygiene measurements to either the intervention arm or the control arm. In Young 2021, schools were randomly assigned (1:1) to either a policy of offering contacts daily testing over seven days to allow continued school attendance (intervention group) or to follow the usual policy of isolation of contacts for 10 days (control group). In two studies there were insufficient details to permit a judgement of the risk of bias (Alfelali 2020; Ashraf 2020).

In the 2020 review, we judged 11 cluster-RCTs as at high risk of identification/recruitment bias (Aiello 2010; Aiello 2012; Azor-Martinez 2018; Chard 2019; Correa 2012; Cowling 2009; Larson 2010; McConeghy 2017; Nicholson 2014; Priest 2014; Savolainen-Kopra 2012). In Aiello 2010 and Aiello 2012, recruitment continued for two weeks after the start of the study, which could have introduced bias. Six trials identified and recruited participants after cluster randomisation (Azor-Martinez 2018; Chard 2019; Cowling 2009; Larson 2010; McConeghy 2017; Nicholson 2014). Three trials recruited new participants after the start of the study to replace those lost to follow-up (Correa 2012; Priest 2014; Savolainen-Kopra 2012). We judged five cluster-RCTs to have probable identification/ recruitment bias (Alzaher 2018; Barasheed 2014; MacIntyre 2011; Najnin 2019; Radonovich 2019), whereas in 19 studies there were insufficient details to permit a judgement of risk of bias (Carabin 1999; DiVita 2011; Feldman 2016; Hartinger 2016; Huda 2012; Ibfelt 2015; Kotch 1994; Ladegaard 1999; MacIntyre 2009; MacIntyre 2013; Millar 2016; Miyaki 2011; Pandejpong 2012; Radonovich 2019; Sandora 2008; Stebbins 2011; Talaat 2011; Yeung 2011; Zomer 2015).

Two of the newly included cluster-RCTs reported intracluster correlation coefficient (ICC) to adjust sample size, taking into consideration clustering effects, and described adjusting outcomes for clustering effect using different statistical methods, or provided justification for not performing adjusted analysis for clustering (Alfelali 2020; Swarthout 2020). For four studies there were insufficient details to permit a judgement of risk of bias (Abaluck 2022; Ashraf 2020; Teesing 2021; Young 2021) since they provided insufficient details on ICC and/or did not perform adjusted analysis or justified the absence of it.

Twenty-six cluster-RCTs identified in the 2020 review reported intracluster correlation coefficient (ICC) to adjust sample size, taking into consideration clustering effects, and described adjusting outcomes for clustering effect using different statistical methods, or provided justification for not performing adjusted analysis for clustering (Aiello 2010; Aiello 2012; Arbogast 2016; Canini 2010; Carabin 1999; Correa 2012; Cowling 2008; Cowling 2009; Hartinger 2016; Huda 2012; Little 2015; Luby 2005; MacIntyre 2009; MacIntyre 2011; MacIntyre 2013; MacIntyre 2015; MacIntyre 2016; McConeghy 2017; Priest 2014; Radonovich 2019; Ram 2015; Roberts 2000; Stebbins 2011; Suess 2012; Talaat 2011; Temime

2018). Five cluster-RCTs did not report the ICC but described adjusting outcomes for clustering effect using different statistical methods, or explained why adjusted analysis for clustering was not performed (Biswas 2019; Chard 2019; McConeghy 2017; Simmerman 2011; Zomer 2015). Thirteen cluster-RCTs provided insufficient details on ICC and/or did not perform adjusted analysis or justified the absence of it (Alzaher 2018; Azor-Martinez 2016; Azor-Martinez 2018; Barasheed 2014; Feldman 2016; Larson 2010; Millar 2016; Miyaki 2011; Najnin 2019; Nicholson 2014; Pandejpong 2012; Savolainen-Kopra 2012; Yeung 2011). Two cluster-RCTs reported the ICC but did not perform adjusted analysis or justified the absence of it (Sandora 2005; Sandora 2008).

Effects of interventions

See: Summary of findings 1 Medical/surgical masks compared to no masks for preventing the spread of viral respiratory illness; Summary of findings 2 N95 respirators compared to medical/surgical masks for preventing the spread of viral respiratory illness; Summary of findings 3 Hand hygiene compared to control for preventing the spread of viral respiratory illness

Comparison 1: Medical/surgical masks compared to no masks

We included 12 trials (10 of which were cluster-RCTs) comparing medical/surgical masks versus no masks (Abaluck 2022; Alfelali 2020; Aiello 2012; Barasheed 2014; Bundgaard 2021; Canini 2010; Cowling 2008; Jacobs 2009; MacIntyre 2009; MacIntyre 2015; MacIntyre 2016; Suess 2012). Two trials were conducted with healthcare workers (HCWs) (Jacobs 2009; MacIntyre 2015), whilst the other 10 studies included people living in the community. In the acute care hospital setting, as opposed to the community setting, variable mask use occurred, according to usual practices in the settings where the studies were undertaken, varying from just under 16% most of the time to 23.6% wearing for > 70% of all working hours (Jacobs 2009; MacIntyre 2015). We therefore excluded the two studies in the acute care hospital setting from the meta-analysis, and report results from these studies narratively. Ten trials were conducted in non-pandemic settings, and two were conducted during the SARS-CoV-2 pandemic (Abaluck 2022; Bundgaard 2021).

Primary outcomes

1. Numbers of cases of viral respiratory illness

Influenza/COVID-like illness

Pooling of nine trials conducted in the community found an estimate of effect for the outcomes of influenza/COVID-like illness cases (risk ratio (RR) 0.95, 95% confidence interval (CI) 0.84 to 1.09; 9 trials; 276,917 participants; moderate-certainty evidence; Analysis 1.1) suggesting that wearing a medical/surgical mask will probably make little or no difference for this outcome. Two studies in healthcare workers provided inconclusive results with very wide confidence intervals: RR 0.88, 95% CI 0.02 to 32; and RR 0.26, 95% CI 0.03 to 2.51, respectively (Jacobs 2009; MacIntyre 2015).

Laboratory-confirmed influenza/SARS-CoV-2 cases

Similarly, the estimate of effect for laboratory-confirmed influenza/ SARS-CoV-2 cases (RR 1.01, 95% CI 0.72 to 1.42; 6 trials, 13,919 participants; moderate-certainty evidence; Analysis 1.1) suggests that wearing a medical/surgical mask probably makes little or no difference compared to not wearing a mask for this outcome.



Laboratory-confirmed other respiratory viruses

One community study reported on laboratory-confirmed other respiratory viruses, showing RR 0.58, 95% CI 0.25 to 1.31; Analysis 1.1, and another study in healthcare workers reported RR 0.79, 95% CI 0.42 to 1.52 (MacIntyre 2015).

Assessing both source control and personal protection

The design of most trials assessed whether masks protected the wearer. Six trials were cluster-RCTs, with all participants in the intervention clusters required to wear masks, thus assessing both source control and personal protection. In two trials the clusters were households with a member with new influenza; neither of these studies found any protective effect (RR 1.03 in 105 households (Canini 2010); RR 1.21 in 145 households (MacIntyre 2009)). In two trials the clusters were college dormitories during the influenza season; neither study found any reduction (RR 1.10 in 37 dormitories (Aiello 2012); RR 0.90 in three dormitories (Aiello 2010)).

Studies conducted during the SARS-CoV-2 pandemic

Two studies were conducted during the SARS-CoV-2 pandemic (Abaluck 2022; Bundgaard 2021), with the former being a very large cluster-RCT of villages in Bangledesh and the latter a large RCT conducted in Denmark.

Exclusion of study due to insufficient number of clusters

We excluded Aiello 2010 from the meta-analysis since we did not consider 'randomisation' of three clusters to three arms to be a proper randomised trial.

2. Adverse events related to the intervention

Canini 2010 reported that 38 (75%) of participants in the intervention arm experienced discomfort with the mask use due to warmth (45%), respiratory difficulties (33%), and humidity (33%). Children reported feeling pain more frequently (3/12) than other participants wearing adult face masks (1/39; P = 0.04). In MacIntyre 2015, adverse events associated with face mask use were reported in 40.4% (227/562) of HCWs in the medical-mask arm. General discomfort (35.1%; 397/1130) and breathing problems (18.3%; 207/1130) were the most frequently reported adverse events. Suess 2012 reported that the majority of participants (107/172; 62%) did not report any problems with mask-wearing. More adults reported no problems (71%) compared to children (36/72; 50%; P = 0.005). The main issues when wearing a face mask for adults as well as for children were "heat/humidity" (18/34; 53% of children; 10/29; 35% of adults; P = 0.1), followed by "pain" and "shortness of breath". Alfelali 2020 reported the most common side effects of wearing a mask in Hajj pilgrims were difficulty in breathing (26%) and discomfort (22%). Although no details were provided, Bundgaard 2021 mentioned that 14% of participants had adverse reactions. Cowling 2008 and Abaluck 2022 mentioned that no adverse events were reported. The other trials did not report measuring adverse outcomes.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Jacobs 2009 reported that participants in the mask group were significantly more likely to experience more days with headache and feeling bad. They found no significant differences between the two groups for symptom severity scores. None of the other trials reported this outcome.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 2: N95/P2 respirators compared to medical/ surgical masks

We included five trials comparing medical/surgical masks with N95/P2 respirators (Loeb 2009; MacIntyre 2009; MacIntyre 2011; MacIntyre 2013; Radonovich 2019). All of these trials except MacIntyre 2009 included HCWs. MacIntyre 2009 included carers and household members of children with a respiratory illness recruited from a paediatric outpatient department and a paediatric primary care practice in Sydney, Australia. None of the trials were conducted during the SARS-CoV-2 pandemic.

Primary outcomes

1. Numbers of cases of viral respiratory illness

Clinical respiratory illness

Pooling of three trials found an estimate of effect suggesting considerable uncertainty as to whether an N95/P2 respirator provides any benefit compared to medical/surgical masks for the outcome of clinical respiratory illness (RR 0.70, 95% CI 0.45 to 1.10; 7799 participants, very low-certainty evidence; Analysis 2.1) (MacIntyre 2011; MacIntyre 2013 (two arms); Radonovich 2019).

Influenza-like-illness

Based on five trials conducted in four healthcare settings and one household, the estimates of effect for the outcome of ILI (RR 0.82, 95% CI 0.66 to 1.03; 8407 participants, low-certainty evidence; Analysis 2.1) suggest that N95/P2 respirators may make little or no difference for this outcome (Loeb 2009; MacIntyre 2009; MacIntyre 2011; MacIntyre 2013; Radonovich 2019).

Laboratory-confirmed influenza

The estimate of the effect for the outcome of laboratory-confirmed influenza infection (RR 1.10, 95% CI 0.90 to 1.34; 8407 participants, moderate-certainty evidence; Analysis 2.1) suggests that the use of a N95/P2 respirator compared to a medical/surgical mask probably makes little or no difference for this more precise and objective outcome.

The outcomes clinical respiratory illness and ILI were reported separately. Considering how these outcomes were defined, it is highly likely that there was considerable overlap between the two, therefore these outcomes were not combined into a single clinical outcome (Analysis 2.1). The laboratory-confirmed viral respiratory infection outcome included influenza primarily but multiple other



common viral respiratory pathogens were also included in several studies. The laboratory-confirmed viral infection outcome was considered more precise and objective in comparison to the clinical outcomes, which were more subjective and considered to be less precise. The findings did not change when we restricted the evidence to HCWs (Analysis 2.2).

2. Adverse events related to the intervention

Harms were poorly reported, but generally discomfort wearing medical/surgical masks and N95/P32 respirators was mentioned in several studies. Radonovich 2019 mentioned that participants wearing the N95 respirator reported skin irritation and worsening of acne. MacIntyre 2011 reported that adverse events were more common with N95 respirators; in particular, discomfort was reported in 41.9% of N95 wearers versus 9.8% of medical-mask wearers (P < 0.01); headaches were more common with N95 (13.4% versus 3.9%; P < 0.01); difficulty breathing was reported more often in the N95 group (19.4% versus 12.5%; P = 0.01); and N95 caused more problems with pressure on the nose (52.2% versus 11.0%; P < 0.01). In MacIntyre 2013, fewer participants using the N95 respirator reported problems (38% (195/512) versus 48% (274/571) of participants in the medical-mask arm; P = 0.001). Loeb 2009 mentioned that no adverse events were reported.

The one trial conducted in the community mentioned that more than 50% of participants reported concerns with both types of masks, mainly that wearing them was uncomfortable, but there were no significant differences between the P2 (N95) and surgical-mask groups (MacIntyre 2009).

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Loeb 2009 reported that 42 participants (19.8%) in the surgical-mask group reported an episode of work-related absenteeism compared with 39 (18.6%) of participants in the N95 respiratory group (absolute risk difference -1.24%, 95% CI -8.75% to 6.27%; P = 0.75).

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Loeb 2009 reported that there were no episodes of LRTIs.

Comparison 3: Hand hygiene compared to control

Nineteen trials compared hand hygiene interventions with control and provided sufficient data to include in meta-analyses (Ashraf

2020; Azor-Martinez 2018; Biswas 2019; Correa 2012; Cowling 2008; Cowling 2009; Hubner 2010; Larson 2010; Little 2015; Millar 2016; Nicholson 2014; Ram 2015; Roberts 2000; Sandora 2005; Simmerman 2011; Stebbins 2011; Swarthout 2020; Teesing 2021; Zomer 2015). The populations of these studies included adults, children, and families, in settings such as schools, childcare centres, homes, and offices. None of the studies was conducted during a pandemic, although a few studies were conducted during peak influenza seasons. A further 16 trials comparing hand hygiene to a control had other outcomes or insufficient information to include in meta-analyses (Alzaher 2018; Arbogast 2016; Azor-Martinez 2016; DiVita 2011; Feldman 2016; Gwaltney 1980; Ladegaard 1999; Luby 2005; Morton 2004; Priest 2014; Savolainen-Kopra 2012; Talaat 2011; Temime 2018; Turner 2012; White 2001; Yeung 2011). The results of these trials were consistent with the findings of our meta-analyses. The results for all outcomes from the 19 trials that were meta-analysed and the 16 trials that were not meta-analysed are shown in Table 2.

Primary outcomes

1. Numbers of cases of viral respiratory illness

Acute respiratory infection (ARI)

Pooling of nine trials for the broad outcome of ARI showed a 14% relative reduction in the numbers of participants with ARI (RR 0.86, 95% CI 0.81 to 0.90; 52,105 participants, moderate-certainty evidence; Analysis 3.1.1) in the hand hygiene group (Analysis 3.1), suggesting a probable benefit (Ashraf 2020; Azor-Martinez 2018; Correa 2012; Larson 2010; Little 2015; Millar 2016; Nicholson 2014; Sandora 2005; Swarthout 2020).

Influenza-like-illness (ILI) and laboratory-confirmed influenza

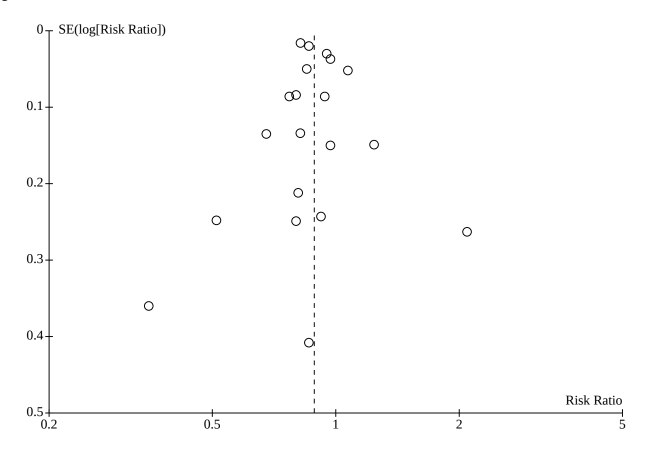
When considering the more strictly defined outcomes of ILI (Biswas 2019; Cowling 2008; Cowling 2009; Hubner 2010; Larson 2010; Little 2015; Ram 2015; Roberts 2000; Simmerman 2011; Teesing 2021; Zomer 2015), and laboratory-confirmed influenza (Biswas 2019; Cowling 2008; Cowling 2009; Hubner 2010; Larson 2010; Ram 2015; Simmerman 2011; Stebbins 2011) the estimates of the effect were heterogeneous, suggesting that hand hygiene may make little or no difference (RR 0.94, 95% CI 0.81 to 1.09 for ILI; 34,503 participants, low-certainty evidence; Analysis 3.1.2); (RR 0.91, 95% CI 0.63 to 1.30 for laboratory-confirmed influenza; 8332 participants; low-certainty evidence; Analysis 3.1.3).

Composite outcome 'ARI or ILI or influenza'

All 19 trials could be pooled for analysis of the composite outcome 'ARI or ILI or influenza', with each study only contributing once with the most comprehensive outcome (in terms of number of events) reported showing an 11% relative reduction in participants with a respiratory illness, suggesting that hand hygiene may offer a benefit (RR 0.89, 95% CI 0.83 to 0.94; low-certainty evidence; Analysis 3.2), but with high heterogeneity. A funnel plot of the 19 trial results did not appear to suggest any small study effects for this outcome (Figure 4).



Figure 4.



Sensitivity analysis

In a sensitivity analysis we used only the most precise and unequivocal (with laboratory confirmed considered the most precise and an undefined ARI considered the least precise) outcome reported in each of 12 studies identified by JMC, an infectious disease physician, and found an estimate of effect in favour of hand hygiene, but with wider CIs (RR 0.88, 95% CI 0.77 to 1.02; Analysis 3.3).

Subgroup analysis by age group

We considered that studies in children might have a different effect than studies in adults, so we conducted subgroup analysis by age group. We found no evidence of a difference in treatment effect by age group (P = 0.18; Analysis 3.4).

2. Adverse events related to the intervention

Correa 2012 reported that no adverse events were observed; in the study by Priest 2014, skin reaction was recorded for 10.4% of participants in the hand sanitiser group versus 10.3% in the control group (RR 1.01, 95% CI 0.78 to 1.30).

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Three trials measured absenteeism from school or work and demonstrated a 36% relative reduction in the numbers of participants with absence in the hand hygiene group (RR 0.64, 95% CI 0.58 to 0.71; Analysis 3.5) (Azor-Martinez 2016; Hubner 2010; Nicholson 2014).

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 4: Hand hygiene + medical/surgical masks compared to control

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza)

Six trials (Aelami 2015; Aiello 2012; Cowling 2009; Larson 2010; Simmerman 2011; Suess 2012) were able to be pooled to compare the use of the combination of hand hygiene and medical/surgical masks with control. Four of these trials were in households, two in university student residences, and one at the annual Hajj pilgrimage. For the outcomes ILI and laboratory-confirmed influenza, pooling demonstrated an estimate of effect suggesting little or no difference between the hand hygiene and medical/surgical mask combination and control. The number of trials and



events was lower than for comparisons of hand hygiene alone, or medical/surgical masks alone, and the confidence interval was wide. For ILI, the RR for intervention compared to control was 1.03 (95% CI 0.77 to 1.37; 4504 participants; Analysis 4.1.1), and for influenza it was 0.97 (95% CI 0.69 to 1.36; 3121 participants; Analysis 4.1.2). Full results of these trials are shown in Table 3

2. Adverse events related to the intervention

Adverse events related to mask wearing in the study by Suess 2012 are reported under Comparison 1 (medical/surgical masks). There was no mention of adverse events related to hand hygiene.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness, e.g. pneumonia

Not reported.

Comparison 5: Hand hygiene + medical/surgical masks compared to hand hygiene

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI and laboratory-confirmed influenza)

Three trials studied the addition of medical/surgical masks to hand hygiene (Cowling 2009; Larson 2010; Simmerman 2011). All three trials had three arms, and are also included in the comparison of hand hygiene plus medical/surgical mask versus control (Comparison 4). All three studies showed no difference between hand hygiene plus medical/surgical mask groups and hand hygiene alone, for all outcomes. The estimates of effect suggested little or no difference when adding masks to hand hygiene compared to hand hygiene alone: for the outcome ILI (RR 1.03, 95% CI 0.69 to 1.53; 3 trials) and the outcome laboratory-confirmed influenza (RR 0.99, 95% CI 0.69 to 1.44), the estimates of effect were not different and the CIs were relatively wide, suggesting little or no difference (Analysis 5.1). However, the CIs around the estimates were wide and do not rule out an important benefit.

2. Adverse events related to the intervention

Not reported.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 6: Medical/surgical masks compared to other (non-N95) masks

One trial compared medical/surgical masks with cloth masks in hospital healthcare workers (MacIntyre 2015), and another trial compared catechin-treated masks versus control masks in healthcare workers and staff of hospitals, rehabilitation centres, and nursing homes in Japan (Ide 2016).

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza)

MacIntyre 2015 found that the rate of ILI was higher in the cloth mask arm compared to the medical/surgical masks arm (RR 13.25, 95% CI 1.74 to 100.97).

Ide 2016 did not find a benefit from the catechin-treated masks over untreated masks on influenza infection rates (adjusted odds ratio (OR) 2.35, 95% CI 0.40 to 13.72; P = 0.34).

2. Adverse events related to the intervention

In MacIntyre 2015 adverse events associated with face mask use were reported in 40.4% (227/562) of HCWs in the medical/surgical mask arm and 42.6% (242/568) in the cloth mask arm (P = 0.45). The most frequently reported adverse events were general discomfort (35.1%; 397/1130) and breathing problems (18.3%; 207/1130). Laboratory tests showed the penetration of particles through the cloth masks to be very high (97%) compared with medical/surgical masks (44%). Ide 2016 reported that there were no serious adverse events associated with the intervention.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.



Comparison 7: Soap + water compared to sanitiser, and comparisons of different types of sanitiser

Two trials compared soap and water with sanitiser (Azor-Martinez 2018; Savolainen-Kopra 2012). Another trial compared different types of hand sanitiser in a virus challenge study (Turner 2004a; Turner 2004b), and one trial studied the frequency of use of hand sanitiser (Pandejpong 2012). The full results of these four trials are shown in Table 4.

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza)

In the trial by Azor-Martinez 2018, ARI incidence was significantly higher in the soap-and-water group compared with the hand sanitiser group (rate ratio 1.21, 95% CI 1.06 to 1.39). In contrast, there was no significant difference between interventions in Savolainen-Kopra 2012. In the rhinovirus challenge study (Turner 2004a; Turner 2004b), all hand sanitisers tested led to a significant lowering of infection rates, but no differences between sanitisers were observed. The study sample size was small.

2. Adverse events related to the intervention

Two trials stated that no adverse events were observed (Pandejpong 2012; Savolainen-Kopra 2012).

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

The authors of Azor-Martinez 2018 also observed a significant benefit for hand sanitiser in reduction in days absent, whereas there was no difference between intervention groups in the Savolainen-Kopra 2012 trial. The study on frequency of use of sanitiser found that use of sanitiser every hour significantly reduced days absent compared with use every two hours or with use only before the lunch break (Pandejpong 2012).

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 8: Surface/object disinfection (with or without hand hygiene) compared to control

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza)

Six trials contributed data to this comparison (Ban 2015; Carabin 1999; Ibfelt 2015; Kotch 1994; McConeghy 2017; Sandora 2008). Full results of these trials are shown in Table 5. Five of the six trials combined disinfection with other interventions such as hand hygiene education, provision of hand hygiene products, and audits. Ban 2015 utilised a combination of provision of hand

hygiene products, and cleaning and disinfection of surfaces, and demonstrated a significant reduction in ARI in the intervention group (OR 0.47, 95% CI 0.48 to 0.65). A similar result was seen in Carabin 1999, with a significant reduction in episodes of ARI. Two studies tested multi component interventions and observed no significant difference in ARI outcomes (Kotch 1994; McConeghy 2017).

One trial compared disinfection alone to usual care (lbfelt 2015). This study demonstrated a significant reduction in some viruses detected on surfaces in the childcare centres (adenovirus, rhinovirus, respiratory syncytial virus (RSV), and metapneumovirus), but not in other viruses, including coronavirus.

2. Adverse events related to the intervention

Not reported.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Only one study measured this outcome (Sandora 2008), observing no significant difference between groups for the outcome of absence due to respiratory illness (rate ratio for intervention to control 1.10, 95% CI 0.97 to 1.24).

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 9: Complex interventions compared to control

Complex interventions are either multifaceted environmental programmes (such as those in low-income countries) or combined interventions including hygiene measures and gloves, gowns, and masks.

Four trials studied complex hygiene and sanitation interventions in low-income country settings (Chard 2019; Hartinger 2016; Huda 2012; Najnin 2019). Full results from these studies are given in Table 6.

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza)

All four trials of complex interventions observed no significant differences between groups in rates of viral respiratory illness.

2. Adverse events related to the intervention

Not reported.

Secondary outcomes

1. Deaths

Not reported.



2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 10: Physical distancing/quarantine

We found three RCTs that assessed physical distancing/quarantine interventions. A quasi-cluster-RCT assessed the effectiveness of quarantining workers of one of two sibling companies in Japan whose family members developed an ILI during the 2009 to 2010 H1N1 influenza pandemic (Miyaki 2011). Workers in the intervention group were asked to stay home on full pay until five days after the household member(s) showed resolution of symptoms or two days after alleviation of fever. A second RCT conducted during the SARS-CoV-2 pandemic investigated whether attending fitness centres with physical distancing was noninferior compared to no access in terms of COVID-19 transmission (Helsingen 2021). The third study was a cluster-RCT conducted during the SARS-CoV-2 pandemic that compared voluntary daily lateral flow device testing for seven days with negative contacts remaining at school to self-isolation of school-based COVID-19 contacts for 10 days in a non-inferiority design (Young 2021).

Primary outcomes

1. Numbers of cases of viral respiratory illness (including laboratory-confirmed influenza and SARS-CoV-2)

Miyaki 2011 reported adherence with the intervention was 100%. In the intervention group 2.75% of workers contracted influenza, compared with 3.18% in the control group (Cox hazard ratio 0.799, 95% CI 0.66 to 0.97; P = 0.02), indicating that the rate of infection was reduced by 20% in the intervention group. However, the risk of a worker being infected was 2.17-fold higher in the intervention group where workers stayed at home with their infected family members. The authors concluded that quarantining workers who have infected household members could be a useful additional measure to control the spread of respiratory viruses in an epidemic setting.

Helsingen 2021 reported 3016 participants were tested for SARS-CoV-2 resulting in one positive case in the fitness centre access arm versus zero in the no access arm at 14 days (risk difference (RD) 0.053%, 95% CI – 0.050 to 0.156%; P = 0.32). In addition, 11 in the fitness centre access arm versus 27 in the no access arm tested positive for SARS-CoV-2 antibodies at one month (RD – 0.87%, 95% CI – 1.52% to – 0.23%; P = 0.001). The authors concluded that access to fitness centres with physical distancing and low population prevalence of SARS-CoV-2 infection did not increase risk of SARS-CoV-2 infection.

Results from Young 2021 suggested no difference between the two treatment arms for SARS-CoV-2 infection (RR 0.96, 95% CI 0.75 to 1.22) leading the study authors to conclude non-inferiority of daily

contact testing of school-based contacts (intervention) compared to self-isolation (control).

2. Adverse events related to the intervention

Not reported.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Young 2021 reported COVID-19 related absences from school were similar in the two treatment groups (RR 0.80, 95% CI 0.54 to 1.19).

4. Hospital admissions

Helsingen 2021 reported no hospital admissions in either treatment arm.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 11: Eye protection compared to control

Primary outcomes

1. Numbers of cases of viral respiratory illness (including laboratory-confirmed influenza and SARS-CoV-2)

We only identified one trial of eye protection which was a preprint only (Fretheim 2022a). This was a pragmatic RCT conducted in Norway from 2 February to 24 April 2022, where 3717 participants were randomised to an intervention group asked to wear glasses (e.g. sunglasses) for two weeks when close to others in public spaces. COVID-19 cases in the national registry were 3.7% in the intervention group (68/1852) and 3.5% (65/1865) in the control group (RR 1.10, 95% CI 0.75 to 1.50). Positive COVID-19 tests based on self-reporting were 9.6% and 11.5% (RR 0.83, 95% CI 0.69 to 1.00). Given the high risk of bias and wide CIs, no policy conclusions can be drawn, but replication studies are clearly warranted. Almost a third of the participants reported respiratory infections. However, a lower proportion of those (215 participants) were in the intervention group compared to the control group (RR 0.90; 95% CI 0.82 to 0.99).

2. Adverse events related to the intervention

A total of 76 participants reported a negative experience from participating in the trial (53 in the intervention group and 23 in the control group). The most common complaint related to the combination of wearing glasses and face masks, and 21 participants in the intervention group cited fogging as an issue. Some participants reported feeling tired or uncomfortable wearing glasses, and a few participants complained of reduced vision when wearing sunglasses or reading glasses. In the control group some participants reported headaches from not being able to wear glasses, and one participant in the intervention group reported a fall due to reduced vision.



Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness, e.g. pneumonia

Not reported.

Comparison 12: Gargling/nose rinsing compared to control

Five trials investigated the effect of gargling/nose rinsing. Satomura 2005 compared throat gargling with povidone-iodine versus tap water in healthy adults. Ide 2014 compared gargling with green tea versus tap water in high school students, and Goodall 2014 compared gargling with tap water with no gargling in university students. Two additional trials were conducted during the SARS-CoV-2 pandemic: Almanza-Reyes 2021 compared silver mouth wash/nose rinse versus conventional mouthwashes and nose rinse in health workers; and Gutiérrez-García 2022 compared neutral electrolysed water mouth and nose rinses versus no rinses in health workers.

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza and SARS-CoV-2)

Satomura 2005 reported that gargling with tap water reduced the incidence of URTIs compared to the control group (usual care) (hazard ratio (HR) 0.60, 95% CI 0.39 to 0.95). Gargling with povidone-iodine did not reduce the incidence of URTIs compared to the control group (HR 0.88, 95% CI 0.58 to 1.34).

Goodall 2014 found no difference in laboratory-confirmed URTIs between the gargling (tap water) and no-gargling groups (RR for gargling versus no gargling 0.82, 95% CI 0.53 to 1.26; P = 0.36).

In a meta-analysis of gargling versus control based on two trials the pooled estimate of effect suggested little or no difference for the outcome of clinical URTI due to gargling (RR 0.91, 95% CI 0.63 to 1.31; 830 participants; Analysis 6.1) (Goodall 2014; Satomura 2005).

There was no difference in the incidence of laboratory-confirmed influenza between high school students gargling with green tea compared with those using tap water (adjusted OR 0.69, 95% CI 0.37 to 1.28; P = 0.24) (Ide 2014). There was also no difference in the incidence of clinically defined influenza (adjusted OR 0.75, 95% CI 0.50 to 1.13; P = 0.17). However, the authors reported that adherence to the interventions amongst students was low.

Almanza-Reyes 2021 reported the incidence of SARS-CoV-2 infection was statistically significantly lower in the silver mouth wash/nose rinse group (two out of 114, 1.8%) compared to the conventional mouthwash group (33 out of 117, 28.2%), and Gutiérrez-García 2022 reported the incidence of COVID-19-

positive cases in the nasal and oral rinses group was 1% compared to 13% in the control group (RR 0.09, 95% CI of 0.01 to 0.72). A metaanalysis of these two studies showed a 93% reduction in risk of SARS-CoV-2 (RR 0.07, 95% CI 0.02 to 0.23; 394 participants; Analysis 6.2).

2. Adverse events related to the intervention

Satomura 2005 reported no adverse events during the 60-day intervention period. Ide 2014 also did not observe any adverse events during the study. Goodall 2014 did not report on adverse effects. There were no adverse reactions in the study by Almanza-Reyes 2021 or side effects in the study by Gutiérrez-García 2022.

Secondary outcomes

1. Deaths

Not reported.

2. Severity of viral respiratory illness as reported in the studies

Satomura 2005 reported that the mean peak score in bronchial symptoms was lower in the water gargling group (0.97) than in the povidone-iodine gargling group (1.41) and the control group (1.40), P=0.055. Other symptoms were not significantly different between groups. Goodall 2014 reported that symptom severity was greater in the gargling group for clinical and laboratory-confirmed URTI, but this was not statistically significant (225.3 versus 191.8, and 210.5 versus 191.8, respectively). Ide 2014 did not report symptom or illness severity.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

Comparison 13: Virucidal tissues compared to control

Two reports (three trials) conducted in the USA studied the effect of virucidal tissues (Farr 1988a; Farr 1988b; Longini 1988). Full results from these studies are given in Table 7.

Primary outcomes

1. Numbers of cases of viral respiratory illness (including ARIs, ILI, and laboratory-confirmed influenza)

The three trials of virucidal tissues reported no differences in infection rates between tissues and placebo, and between tissues and no tissues (Farr 1988a; Farr 1988b; Longini 1988).

2. Adverse events related to the intervention

Farr 1988b reported cough in 4% of participants using virucidal tissues versus 57% in the placebo group, but 24% reported nasal burning in the virucidal tissue group versus 8% in the placebo group. Longini 1988 did not report on adverse effects.

Secondary outcomes

1. Deaths

Not reported.



2. Severity of viral respiratory illness as reported in the studies

Not reported.

3. Absenteeism

Not reported.

4. Hospital admissions

Not reported.

5. Complications related to the illness (e.g. pneumonia)

Not reported.

DISCUSSION

Summary of main results

See Table 8.

1. Medical/surgical masks compared to no masks

The pooled estimates of effect from randomised controlled trials (RCTs) and cluster-RCTs for wearing medical/surgical masks compared to no masks in the community suggests probably little or no difference in interrupting the spread of influenzalike illness (ILI)/COVID-19 like illness (risk ratio (RR) 0.95, 95% confidence interval (CI) 0.84 to 1.09; moderate-certainty evidence), or laboratory-confirmed influenza/SARS-CoV-2 (RR 1.01, 95% CI 0.72 to 1.42; moderate-certainty evidence). Six trials were cluster-RCTs, with all participants in the intervention clusters required to wear masks, thus assessing both source control and personal protection. In two trials the clusters were households with a member with new influenza; neither trial found any protective effect (RR 1.03 in 105 households (Canini 2010); RR 1.21 in 145 households (MacIntyre 2009). In two trials the clusters were college dormitories during the influenza season; neither trial found any reduction (RR 1.10 in 37 dormitories (Aiello 2012); RR 0.90 in three dormitories (Aiello 2010)). Two studies were conducted during the COVID-19 pandemic and their addition had minimal impact on the pooled estimate of effect previously reported from the earlier studies focused on influenza (Abaluck 2022; Bundgaard 2021). We excluded Aiello 2010 from meta-analysis since we did not consider 'randomisation' of three clusters to three arms was a proper randomised trial.

Less than half of the trials comparing masks with no masks addressed harms of mask wearing (Canini 2010; Cowling 2008; MacIntyre 2015; Suess 2012). Warmth, respiratory difficulties, humidity, and general discomfort were the most frequently reported adverse events. Neither of the RCTs conducted during the COVID-19 pandemic directly assessed harms of mask wearing. More adults reported no harms compared to children.

In one trial cloth masks were associated with a significantly higher risk of both ILI and laboratory-confirmed respiratory virus infection in healthcare workers (HCWs) (MacIntyre 2015). In addition, filtration capacity of the two-ply cotton cloth masks was found to be only 3% and markedly less than with medical/surgical masks based on standardised particle testing. The authors suggested moisture retention, poor filtration, and penetration of the virus through the mask as plausible explanations for the increased risk of infection.

We did not find any randomised trials assessing the effectiveness of barrier interventions using a combination of masks, gloves, and gowns.

2. N95 respirators compared to medical/surgical masks

Comparisons between N95 respirators and medical/surgical masks, used as needed for exposure to at-risk patients, for the outcomes of clinical respiratory illness and the outcome of laboratoryconfirmed influenza showed estimates of effect suggesting considerable uncertainty for any benefit of N95 respirators for the former outcome and probably little or no difference for the latter outcome. Five trials (four in healthcare settings and one in a household setting) compared N95/P2 respirators with medical/surgical masks. Pooling of three of these trials showed an estimate of effect suggesting considerable uncertainty as to whether there was any benefit comparing N95 respirators and medical/surgical face masks for the outcome of clinical respiratory illness (RR 0.70, 95% CI 0.45 to 1.10; very low-certainty evidence), and that N95 respirators may make little or no difference for the outcome of ILI (RR 0.82, 95% CI 0.66 to 1.03; low-certainty evidence), and probably little or no difference for the outcome of laboratory-confirmed influenza (RR 1.10, 95% CI 0.90 to 1.34; moderate-certainty evidence). The presence of imprecision (wide CIs) and heterogeneity, particularly for the more subjective and less precise outcomes of clinical respiratory illness and ILI compared to laboratory-confirmed influenza infection, makes it difficult to assess whether there may be a benefit of either medical/surgical masks or N95/P2 respirators. Restricting the pooling to HCWs made no difference to the overall findings. The two trials with the largest event rates were quite consistent in their findings of no significant differences between N95 and medical/surgical masks for the outcomes of laboratory-confirmed influenza and all laboratoryconfirmed viral infections (Loeb 2009; Radonovich 2019). Three of the trials contributing to this analysis were carried out by members of the same group (MacIntyre 2009; MacIntyre 2011; MacIntyre

In general, harms were poorly reported or not reported at all in trials comparing N95 respirators with surgical masks. General discomfort resulting in reduced wear adherence was the most frequently reported harm.

3. Hand hygiene compared to control

We found that the estimate of effect may offer a benefit for hand hygiene for the composite outcome 'acute respiratory infections (ARI) or ILI or influenza' (RR 0.89, 95% CI 0.83 to 0.94; low-certainty evidence), and probably offers a benefit for the outcomes ARI alone (RR 0.86, 95% CI 0.81 to 0.90; moderate-certainty evidence), and absenteeism (RR 0.64, 95% CI 0.58 to 0.71). An observed estimate of effect in favour of hand hygiene for laboratory-confirmed influenza, but with wider CIs may be a consequence of smaller sample sizes in conjunction with a more rigorous outcome measure.

4. Hand hygiene + medical/surgical masks compared to control

The estimate of effect of combined hand hygiene and medical/surgical mask interventions compared to control in six (mostly small) trials suggested that the intervention may make little or no difference for the outcomes ILI (RR 1.03, 95% CI 0.77 to 1.37), and laboratory-confirmed influenza (four trials) (RR 0.97, 95% CI 0.69 to 1.36).



5. Hand hygiene + medical/surgical masks compared to hand hygiene

We also found an estimate of effect suggesting that adding medical/surgical masks to hand hygiene compared to hand hygiene alone may make little or no difference for the outcomes ILI (RR 1.03, 95% CI 0.69 to 1.53; 3 trials), and laboratory-confirmed influenza (RR 0.99, 95% CI 0.69 to 1.44).

6. Medical/surgical masks compared to other (non-N95) masks

One trial found that medical/surgical masks were more effective than cloth masks at reducing the rate of ILI (RR 13.25, 95% CI 1.74 to 100.97) (MacIntyre 2015), but the extremely wide CIs make this finding difficult to interpret. One trial did not find a benefit from catechin-treated masks over untreated masks on influenza infection rates (adjusted odds ratio (OR) 2.35, 95% CI 0.40 to 13.72; P = 0.34) (Ide 2016).

Harms of wearing masks were reported in 40.4% of HCWs using medical/surgical masks, and in 42.6% of those wearing cloth masks (P = 0.45) (MacIntyre 2015). The penetration of particles was higher in cloth masks (97%) compared to medical/surgical masks (44%).

7. Soap + water compared to sanitiser, and comparisons of different types of sanitiser

There were too few trials comparing different types of hand hygiene interventions to be certain of any true differences between soap and water, alcohol-based hand sanitisers, or other types of interventions. Also, it is uncertain whether the incremental effect of adding virucidals or antiseptics to hand-washing actually decreased the respiratory disease burden outside the confines of the rather atypical studies. The extra benefit may have been, at least in part, accrued by confounding additional routines.

8. Surface/object disinfection (with or without hand hygiene) compared to control

We identified six trials on surface/object disinfection (with or without hand hygiene), and although they were heterogeneous (and therefore could not be pooled), three of them showed a clear benefit compared to controls (Ban 2015; Carabin 1999; Ibfelt 2015).

We found no RCTs of nose disinfection, or disinfection of living quarters, as described in observational studies reported in Jefferson 2011.

9. Complex interventions compared to control

Four trials studied complex hygiene and sanitation interventions, all in low-income country settings (Chard 2019; Hartinger 2016; Huda 2012; Najnin 2019). These trials could not be pooled due to the heterogeneity of the interventions and settings. All four trials found no significant differences between groups in the rates of viral respiratory illness.

10. Physical distancing/quarantine compared to control

We identified one trial that evaluated the effect of quarantine and found a reduction in influenza transmission to co-workers when those with infected household members stayed home from work (Miyaki 2011). However, staying home increased their risk of being infected two-fold. Two studies conducted during the COVID-19 pandemic on SARS-cov-2 transmission showed (1) non-inferiority of daily contact testing of school-based contacts (intervention)

compared to self-isolation (control) (Young 2021); and (2) access to fitness centres with physical distancing and low population prevalence of SARS-CoV-2 infection did not increase risk of SARS-cov-2 infection (Helsingen 2021).

11. Eye protection compared to control

We only identified one trial of eye protection which was a preprint only (Fretheim 2022a).

12. Gargling compared to control

Three trials addressed the use of gargling in preventing respiratory infections (Goodall 2014; Ide 2014; Satomura 2005). Although the trials used a variety of liquids and different outcomes, pooling the results of the two trials that compared gargling with tap water versus control did not show a favourable effect in reducing URTIs (RR 0.91, 95% CI 0.63 to 1.31) (Goodall 2014; Satomura 2005). Two trials of mouthwash/nose rinse were conducted during the SARS-cov-2 pandemic in HCWs: Almanza-Reyes 2021 compared silver mouth wash/nose rinse versus conventional mouthwashes and nose rinse; and Gutiérrez-García 2022 compared neutral electrolysed water mouth and nose rinses versus no rinses. Both studies reported large protective effects of the intervention on SARS-CoV-2 infection with reported outcomes of SARS-COV-2 infection in 28.2% and 12.7% in the HCWs not using the interventions versus 1.8% and 1.2% in those using the intervention, despite the use of full personal protective equipment (PPE) and the high outcome rates raise questions about risk of bias, and no data were provided about baseline rates in other settings with full use of

13. Virucidal tissues compared to control

Two reports (three trials) identified in Jefferson 2011 studied the effect of virucidal tissues compared to placebo or no tissues (Farr 1988a; Farr 1988b; Longini 1988). These trials found no differences in infection rates and could not be pooled.

Overall completeness and applicability of evidence

Several features need consideration before making generalisations based on the included studies.

The settings of the included studies, which were conducted over five decades, were heterogeneous and ranged from suburban schools, Carabin 1999, to emergency departments, intensive care units, and paediatric wards, Loeb 2009, in high-income countries; slums in low-income countries (Luby 2005); and an upper Manhattan immigrant Latino neighbourhood (Larson 2010). Few attempts were made to obtain socio-economic diversity by (for example) involving more schools in the evaluations of the same programme. We identified only a few studies from low-income countries, where the vast majority of the burden of ARIs lies and where inexpensive interventions are so critical. Additionally, limited availability of over-the-counter medications and national universal comprehensive health insurance provided with consequent physician prescription of symptomatic treatment may further limit the generalisability of findings.

The included trials generally reported few events and were conducted mostly during non-epidemic periods with the exception of the trials carried out during the influenza H1N1 and SARS-CoV-2 pandemics. The large study by Radonovich 2019 is an exception as it crossed over two of the highest reporting years for influenza in



the USA between 2010 and 2017 (Elflein 2019). None of the trials were conducted during pandemics of SARS-CoV-1or in outbreaks of Middle East respiratory syndrome (MERS).

Of the trials assessing the effect of masks, six were carried out in those at greater exposure (i.e. HCWs) (Jacobs 2009; Loeb 2009; MacIntyre 2011; MacIntyre 2013; MacIntyre 2015; Radonovich 2019). None of these studies included HCWs undertaking aerosolgenerating procedures, for which the World Health Organization (WHO) currently recommends the N95 or equivalent mask. Three trials on hand hygiene interventions were carried out in nursing homes, and included HCWs (McConeghy 2017; Temime 2018; Yeung 2011). The scarcity of RCTs on HCWs limits the generalisability of such results.

The variable quality of the methods of some studies is striking. Incomplete or no reporting of randomisation (Turner 2004a), blinding (Farr 1988a; Farr 1988b), numerators and denominators (Carabin 1999; Kotch 1994), interventions, and cluster coefficients in the relevant trials (Carabin 1999), led to a considerable loss of information. Potential biases were often not discussed.

Inappropriate placebos caused design problems. In some studies the placebo probably carried sufficient effect to dilute the intervention effects (Longini 1988). Two valiant attempts with virucidal tissues probably failed because placebo handkerchiefs were impregnated with a dummy compound that stung the users' nostrils (Farr 1988a; Farr 1988b).

Some studies used impractical interventions. Volunteers subjected to the intervention hand cleaner (organic acids) were not allowed to use their hands between cleaning and virus challenge, so the effect of normal use of the hands on the intervention remains unknown (Turner 2004a; Turner 2004b). Two per cent aqueous iodine painted on the hands, although a successful antiviral intervention, causes unacceptable cosmetic staining, which is impractical for all but those at the highest risk of epidemic contagion (Gwaltney 1980).

Adherence with interventions, especially educational programmes, was a problem for many studies despite the importance of many such low-cost interventions. Adherence with mask wearing varied; it was generally around 60% to 80%, but was reported to be as low as 40% (see Table 1). Overall, the logistics of carrying out trials that involve sustained behaviour change are demanding, particularly in challenging settings such as immigrant neighbourhoods or students' halls of residence.

The identified trials provided sparse and unsystematic data on adverse effects of the intervention, and few of the RCTs measured or reported adherence with the intervention, which is especially important for the use of medical/surgical masks or N95 respirators. No studies investigated how the level of adherence may have influenced the effect size.

We identified one study assessing the effects of eye protection (Fretheim 2022a), and we identified three studies on physical distancing/quarantine (Helsingen 2021; Miyaki 2011; Young 2021). The dearth of evidence and predominant setting of seasonal viral circulation limits generalisability of our findings to other contexts and any future epidemics due to other respiratory viruses such as the COVID-19 pandemic although there have been increasing numbers of RCTs and cluster-RCTs in the latter setting which are adding to the evidence base.

The two recent small trials from Mexico assessing local mouth/ nose rinses airways prophylactic as interventions treatments report large but uncertain reductions in transmission to healthcare workers which warrant further study and replication by other investigator (Almanza-Reyes 2021; Gutiérrez-García 2022).

Certainty of the evidence

We found the available evidence base identified through our search processes to be of variable quality. Reporting of sequence generation and allocation concealment were poor in 30% to 50% of studies across the categories of intervention comparisons. Given the nature of the intervention comparison, blinding of treatment allocation after randomisation was rarely achieved. Although blinding of outcome assessment is highly feasible and desirable, most outcomes were assessed by self-reports. Outcomes in some studies were poorly defined, with a lack of clarity as to the possible aetiological agents (bacterial versus viral). Some studies used laboratory-confirmed outcomes, both adding precision and avoiding indirectness by having an accurate outcome measure and lowering the risk of bias (see Table 9 for heterogeneity of trial outcome definitions). We found no evidence of selective reporting of outcomes within the included studies. We believe publication bias is unlikely, as the included studies demonstrated a range of effects, both positive and negative, over all study sizes. The variable quality of the studies hampers drawing any firm conclusions.

Potential biases in the review process

The non-drug (and often locally manufactured) nature of most of the interventions in this review, the lack of effective regulation in some settings, and the possible endless number of manufacturers make it difficult to gauge the existence of unpublished data. Non-drug interventions typically have no or very loose regulation.

In this 2022 update, we again focused on RCTs and cluster-RCTs, providing a higher level of evidence compared with the previous version of the review, which also meta-analysed observational studies when appropriate (Jefferson 2011). However, many of the trials were small and hence underpowered, and at high or unclear risk of bias due to poor reporting of methods and lack of blinding. The populations, outcomes, comparators, and interventions tested were heterogeneous.

Due to the urgency of this update in the context of the COVID-19 pandemic, we did not contact trial authors to request missing data. This means that we have not considered studies that included other non-respiratory infections, and did not provide stratified data by type of infection.

Agreements and disagreements with other studies or reviews

Several reviews of RCTs have found broadly similar results to this review for face masks. In a meta-analysis comparing surgical masks with N95 respirators, Smith 2016 pooled three trials and found an estimate of effect suggesting no difference for laboratory-confirmed respiratory infections (OR 0.89, 95% CI 0.64 to 1.24) or ILI (OR 0.51, 95% CI 0.19 to 1.41) (Loeb 2009; MacIntyre 2011; MacIntyre 2013). A similar meta-analysis, Offeddu 2017, based on two trials concluded that masks (either N95/P2 respirators or medical/surgical masks) were effective against clinical respiratory infections (RR 0.59, 95% CI 0.46 to 0.77) and ILI (RR 0.34, 95% CI 0.14



to 0.82) (MacIntyre 2011; MacIntyre 2015). Pooling of two studies (MacIntyre 2011; MacIntyre 2013) also found an estimate of effect that favoured N95 respirators to medical/surgical masks for clinical respiratory infections (RR 0.47, 95% CI 0.36 to 0.62), but not for ILI, (RR 0.59, 95% CI 0.27 to 1.28) based on three studies (Loeb 2009: MacIntyre 2011; MacIntyre 2013). The outcome of clinical respiratory infection is considered to be the most subjective and least precise outcome.

A recent meta-analysis included five trials comparing N95/P2 respirators with medical/surgical masks and found no difference between groups for either influenza (RR 1.09, 95% CI 0.92 to 1.28), or respiratory viral infections (RR 0.89, 95% CI 0.70 to 1.11) (Long 2020). By excluding Loeb 2009 (an open, non-inferiority RCT that compared medical/surgical masks with N95 respirators in protecting HCWs against influenza), the authors reported a significant protective effect against viral infections (RR 0.61, 95% CI 0.39 to 0.98). The authors do not report a rationale for the exclusion in the sensitivity analysis, and do not report on exclusion of the studies with low weighting, which arguably would be more relevant in a sensitivity analysis. The two trials that make up 96% of the weighting demonstrated no significant differences in the outcome events (Loeb 2009; Radonovich 2019). A recent metaanalysis of four RCTs adjusting for clustering, which compared N95 respirators with the use of medical/surgical masks, found pooled estimates of effect that did not demonstrate any difference in any laboratory-confirmed viral respiratory infection (OR 1.06, 95% CI 0.90 to 1.25), laboratory-confirmed influenza (OR 0.94, 95% CI 0.73 to 1.20), or clinical respiratory illness (OR 1.49, 95% CI 0.98 to 2.28), with the evidence profile suggesting that there was greater imprecision and inconsistency in the outcome of clinical respiratory illness (Bartoszko 2020). Moreover, in another recent systematic review that assessed the effectiveness of personal protective and environmental measures in non-healthcare settings (funded by the WHO), 10 RCTs reporting estimates of the effectiveness of face masks in reducing laboratory-confirmed influenza virus infections in the community were identified (Xiao 2020). The evidence from these RCTs suggested that the use of face masks either by infected persons or by uninfected persons does not have a substantial effect on influenza transmission.

The findings from several systematic reviews and meta-analyses over the last decade have not demonstrated any difference in the clinical effectiveness of N95 respirators or equivalent compared to the use of surgical masks when used by HCWs in multiple healthcare settings for the prevention of respiratory virus infections, including influenza.

Reviews based on observational studies have usually found a stronger protective effect for face masks, but have important biases. The review by Chu 2020 did not consider RCTs of influenza transmission, but only the observational studies examining impact on SARS, MERS, or SARS-CoV-2. For N95 masks versus no mask in HCWs, there was a large protective effective with an OR of 0.04 (95% CI 0.004 to 0.30); for surgical masks versus no masks, there was an OR of 0.33 (0.17 to 0.61) overall, but four of these studies were in healthcare settings. Chu 2020 has been criticised for several reasons: use of an outdated 'Risk of bias' tool; inaccuracy of distance measures; and not adequately addressing multiple sources of bias, including recall and classification bias and in particular confounding. Confounding is very likely, as preventive behaviours such as mask use, social distancing, and hand hygiene

are correlated behaviours, and hence any effect estimates are likely to be overly optimistic.

The two RCTs of medical/surgical masks during the SARS-CoV-2 pandemic found uncertain evidence of a small or no effect (Abaluck 2022; Bundgaard 2021). The study by Abaluck 2022 found a statistically significant benefit of masks versus no masks for COVID-like-illness, however, this study was rated at high risk of bias for five of the six domains due to issues including baseline imbalance, subjective outcome assessment and incomplete follow-up across the groups. Despite this study contributing 45% of the weight towards the meta-analysis of influenza/COVID-like-illness for masks versus no masks, the updated conclusions from the analysis strengthened around little or no effect of mask use.

Also based on observational studies, Jefferson 2011 found a protective effect of wearing surgical masks with hygienic measures compared to not wearing masks in the SARS 2003 outbreak (OR 0.32, 95% CI 0.26 to 0.39). However, the evidence was based on case-control studies carried out during the outbreak. There was some additional but very limited supportive evidence from the cohort studies in Jefferson 2011.

Although the use of eye protection and physical distancing measures are widely believed to be effective in reducing transmission of respiratory viruses and mitigating the impact of an influenza pandemic, we found only one trial investigating the role of self-quarantine in reducing the incidence of H1N1 influenza events in the workplace, and no trials examining the effect of eye protection. The evidence for these measures was derived largely from observational studies and simulation studies, and the overall certainty of supporting evidence is relatively low. The finding of limited evidence evaluating these interventions was also consistent with a recent review funded by the WHO for the preparation of its guidelines on the use of non-pharmaceutical interventions for pandemic influenza in non-medical settings (Fong 2020).

There are several previous systematic reviews on hand hygiene and respiratory infections. Five of them reviewed the evidence in a community setting (Moncion 2019; Rabie 2006; Saunders-Hastings 2017; Warren-Gash 2013: Wong 2014), and three focused on children (Mbakaya 2017; Willmott 2016; Zivich 2018). The earliest review in 2006 included eight studies, three of which were RCTs (Rabie 2006). The pooled estimate of seven studies was described as "indicative" of the effect of hand hygiene, but the studies were of poor quality. The Warren-Gash 2013 review included 16 studies (10 of which were RCTs) and reported mixed and inconclusive results. A 2014 review identified 10 RCTs and reported that the combination of hand hygiene with face masks in high-income countries (five trials) significantly reduced the incidence of laboratory-confirmed influenza and ILI, whilst hand hygiene alone did not (Wong 2014). This significant reduction in laboratory-confirmed influenza and ILI for hand hygiene and face masks may have been based on the raw numbers without adjusting for any clustering effects in the included cluster trials, which produced inappropriately narrow CIs, and possibly biased treatment effect estimates. Moreover, trials from the low-income countries were not included in the review, and this significant effect was not demonstrated when all the trials identified in the review were combined. The Saunders-Hastings 2017 review of studies evaluating the effectiveness of personal protective measures in interrupting pandemic influenza transmission only



identified two RCTs (Azor-Martinez 2014; Suess 2012), which reported a significant effect of hand hygiene. The Moncion 2019 review identified seven RCTs of hand hygiene compared to control, with mixed results for preventing the transmission of laboratory-confirmed or possible influenza. Systematic reviews of RCTs of hand hygiene interventions amongst children, Mbakaya 2017 and Willmott 2016, or at a non-clinical workplace, Zivich 2018, identified heterogeneous trials with quality problems including small numbers of clusters and participants, inadequate randomisation, and self-reported outcomes. Evidence of impact on respiratory infections was equivocal.

A rapid search for other systematic reviews of RCTs was conducted in September 2022, and none of high quality were found.

AUTHORS' CONCLUSIONS

Implications for practice

The evidence summarised in this review on the use of masks is largely based on studies conducted during traditional peak respiratory virus infection seasons up until 2016. Two relevant randomised trials conducted during the COVID-19 pandemic have been published, but their addition had minimal impact on the overall pooled estimate of effect. The observed lack of effect of mask wearing in interrupting the spread of influenza-like illness (ILI) or influenza/COVID-19 in our review has many potential reasons, including: poor study design; insufficiently powered studies arising from low viral circulation in some studies; lower adherence with mask wearing, especially amongst children; quality of the masks used; self-contamination of the mask by hands; lack of protection from eye exposure from respiratory droplets (allowing a route of entry of respiratory viruses into the nose via the lacrimal duct); saturation of masks with saliva from extended use (promoting virus survival in proteinaceous material); and possible risk compensation behaviour leading to an exaggerated sense of security (Ammann 2022; Brosseau 2020; Byambasuren 2021; Canini 2010; Cassell 2006; Coroiu 2021; MacIntyre 2015; Rengasamy 2010; Zamora 2006).

Our findings show that hand hygiene has a modest effect as a physical intervention to interrupt the spread of respiratory viruses, but several questions remain. First, the high heterogeneity between studies may suggest that there are differences in the effect of different interventions. The poor reporting limited our ability to extract the information needed to assess any 'dose response' relationship, and there are few head-to-head trials comparing hand hygiene materials (such as alcohol-based sanitiser or soap and water). Second, the sustainability of hand hygiene is unclear where participants in some studies achieved 5 to 10 handwashings per day, but adherence may have diminished with time as motivation decreased, or due to adverse effects from frequent hand-washing. Third, there is little evidence about the effectiveness of combinations of hand hygiene with other interventions, and how those are best introduced and sustained. Finally, some interventions were intensively implemented within small organisations, and involved education or training as a component, and the ability to scale these up to broader interventions is unclear.

Our findings with respect to hand hygiene should be considered generally relevant to all viral respiratory infections, given the diverse populations where transmission of viral respiratory infections occurs. The participants were adults, children and

families, and multiple congregation settings including schools, childcare centres, homes, and offices. Most respiratory viruses, including the pandemic SARS-CoV-2, are considered to be predominantly spread via respiratory particles of varying size or contact routes, or both (WHO 2020c). Data from studies of SARS-CoV-2 contamination of the environment based on the presence of viral ribonucleic acid and infectious virus suggest significant fomite contamination (Lin 2022; Onakpoya 2022b; Ong 2020; Wu 2020). Hand hygiene would be expected to be beneficial in reducing the spread of SARS-CoV-2 similar to other beta coronaviruses (SARS-CoV-1, Middle East respiratory syndrome (MERS), and human coronaviruses), which are very susceptible to the concentrations of alcohol commonly found in most hand-sanitiser preparations (Rabenau 2005; WHO 2020c). Support for this effect is the finding that poor hand hygiene, despite the use of full personal protective equipment (PPE), was independently associated with an increased risk of SARS-CoV-2 transmission to healthcare workers in a retrospective cohort study in Wuhan, China in both a high-risk and low-risk clinical unit for patients infected with COVID-19 (Ran 2020). The practice of hand hygiene appears to have a consistent effect in all settings, and should be an essential component of other interventions.

The highest-quality cluster-RCTs indicate that the most effect on preventing respiratory virus spread from hygienic measures occurs in younger children. This may be because younger children are least capable of hygienic behaviour themselves (Roberts 2000), and have longer-lived infections and greater social contact, thereby acting as portals of infection into the household (Monto 1969). Additional benefit from reduced transmission from them to other members of the household is broadly supported by the results of other study designs where the potential for confounding is greater.

Routine long-term implementation of some of the interventions covered in this review may be problematic, particularly maintaining strict hygiene and barrier routines for long periods of time. This would probably only be feasible in highly motivated environments, such as hospitals. Many of the trial authors commented on the major logistical burdens that barrier routines imposed at the community level. However, the threat of a looming epidemic may provide stimulus for their inception.

Implications for research

Public health measures and physical interventions can be highly effective to interrupt the spread of respiratory viral infections, especially when they are part of a structured and co-ordinated programme that includes instruction and education, and when they are delivered together and with high adherence. Our review has provided important insights into research gaps that need to be addressed with respect to these physical interventions and their implementation and have been brought into a sharper focus as a result of the COVID-19 pandemic. The 2014 WHO document 'Infection prevention and control of epidemic - and pandemic-prone acute respiratory infections in health care' identified several research gaps as part of their GRADE assessment of their infection prevention and control recommendations, which remain very relevant (WHO 2014). Research gaps identified during the course of our review and the WHO 2014 document may be considered from the perspective of both general and specific themes.



A general theme identified was the need to provide outcomes with explicitly defined clinical criteria for acute respiratory infections (ARIs) and discrete laboratory-confirmed outcomes of viral ARIs using molecular diagnostic tools which are now widely available. Our review found large disparities between studies with respect to the clinical outcome events, which were imprecisely defined in several studies, and there were differences in the extent to which laboratory-confirmed viruses were included in the studies that assessed them. Another general theme identified was the lack of consideration of sociocultural factors that might affect adherence with the interventions, especially those employed in the community setting. A prime example of this latter point was illustrated by the observations of the use of masks versus mask mandates during the COVID-19 pandemic. In addition, the cost and resource implications of the physical interventions employed in different settings would have important relevance for low- to middle-income countries. Resources have been a major issue with the COVID-19 pandemic, with global shortages of several components of PPE. Several specific research gaps related to physical interventions were identified within the WHO 2014 document and are congruent with many of the findings of this 2022 update, including the following: transmission dynamics of respiratory viruses from patients to healthcare workers during aerosol-generating procedures; a continued lack of precision with regards to defining aerosol-generating procedures; the safety of cohorting of patients with the same suspected but unconfirmed diagnosis in a common unit or ward with patients infected with the same known pathogen in healthcare settings; the optimal duration of the use of physical interruptions to prevent spread of ARI viruses; use of spatial separation or physical distancing (in healthcare and community settings, respectively) alone versus spatial separation or physical distancing with the use of other added physical interventions coupled with examining discrete distance parameters (e.g. one metre, two metres, or > two metres); the effectiveness of respiratory etiquette (i.e. coughing/ sneezing into tissues or a sleeved bent elbow); the effectiveness of triage and early identification of infected individuals with an ARI in both hospital and community settings; the utility of entrance screening to healthcare facilities; use of frequent disinfection techniques appropriate to the setting (high-touch surfaces in the environment, gargling with oral disinfectants, and virucidal tissues or clothing) alone or in combination with facial masks and hand hygiene; the use of visors, goggles or other eyewear; the use of ultraviolet light germicidal irradiation for disinfection of air in healthcare and selected community settings; the use of air scrubbers and /or high-efficiency particulate absorbing filters and the use of widespread adherence with effective vaccination

There is a clear requirement to conduct large, pragmatic trials to evaluate the best combinations in the community and in healthcare settings with multiple respiratory viruses and in different sociocultural settings. Randomised controlled trials (RCTs) with a pragmatic design, similar to the Luby 2005 trial or the Bundgaard 2020 trial, should be conducted whenever possible. Similar to what has been observed in pharmaceutical interventions where multiple RCTs were rapidly and successfully completed during the COVID-19 pandemic, proving they can be accomplished, there should be a deliberate emphasis and directed funding opportunities provided to conduct well-designed RCTs to address the effectiveness of many of the physical interventions in multiple settings and populations, especially in those most at risk,

and in very specific well-defined populations with monitoring of the adherence to the interventions.

Several specific research gaps deserve expedited attention and may be highlighted within the context of the COVID-19 pandemic. The use of face masks in the community setting represents one of the most pressing needs to address, given the polarised opinions around the world, and the increasing concerns over widespread microplastic pollution from the discarding of masks (Shen 2021). Both broad-based ecological studies, adjusting for confounding and high quality RCTs, may be necessary to determine if there is an independent contribution to their use as a physical intervention, and how they may best be deployed to optimise their contribution. The type of fabric and weave used in the face mask is an equally pressing concern, given that surgical masks with their cotton-polypropylene fabric appear to be effective in the healthcare setting, but there are questions about the effectiveness of simple cotton masks. In addition, any masking intervention studies should focus on measuring not only benefits but also adherence, harms, and risk compensation if the latter may lead to a lower protective effect. In addition, although the use of medical/ surgical masks versus N95 respirators demonstrates no differences in clinical effectiveness to date, their use needs to be further studied within the context of a well-designed RCT in the setting of COVID-19, and with concomitant measurement of harms, which to date have been poorly studied. The recently published Loeb RCT conducted over a prolonged course in the current pandemic has provided the only evidence to date in this area (Loeb 2022).

Physical distancing represents another major research gap which needs to be addressed expediently, especially within the context of the COVID-19 pandemic setting as well as in future epidemic settings. The use of quarantine and screening at entry ports needs to be investigated in well-designed, high-quality RCTs given the controversies related to airports and travel restrictions which emerged during the COVID-19 pandemic. We found only one RCT investigating quarantine, and no trials of screening at entry ports or physical distancing. Given that these and other physical interventions are some of the primary strategies applied globally in the face of the COVID-19 pandemic, future trials of high quality should be a major global priority to be conducted within the context of this pandemic, as well as in future epidemics with other respiratory viruses of less virulence.

The variable quality and small scale of some studies is known from descriptive studies (Aiello 2002; Fung 2006; WHO 2006b), and systematic reviews of selected interventions (Meadows 2004). In summary, more high-quality RCTs are needed to evaluate the most effective strategies to implement successful physical interventions in practice, both on a small scale and at a population level. It is very unfortunate that more rigorous planning, effort and funding was not provided during the current COVID-19 pandemic towards high-quality RCTs of the basic public health measures. Finally, we emphasise that more attention should be paid to describing and quantifying the harms of the interventions assessed in this review, and their relationship with adherence.

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The following people conducted the editorial process for this 2022 update:

- Sign-off Editor (final editorial decision): Michael Brown (Michigan State University College of Human Medicine, USA).
- Managing Editor (selected peer reviewers, collated peer reviewer comments, provided editorial guidance to authors): Fiona Russell (Bond University, Australia).

- Contact Editor (assessed peer review comments and recommended an editorial decision): Allen Cheng (Monash University, Australia).
- Statistical Editor (provided comments): Teresa Neeman (Biological Data Science Institute, Australian National University, Australia).
- Copy Editor (copy-editing and production): Heather Maxwell.

Peer reviewers (provided comments and recommended an editorial decision):

- Clinical/content review: Roderick P. Venekamp.
- Consumer review: Janet Wale (Independent consumer representative).
- Methods review: Leslie Choi (Evidence Synthesis Development Editor, Cochrane Central Executive Team).



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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Abaluck 2022

Study characteristics	
Methods	Cluster-RCT
	Randomisation unit: villages (N = 600)
	Intervention duration: 8 weeks "Our intervention was designed to last 8 weeks in each village"
Participants	Inclusion criteria: community level participants
	Intervention = 178,322 individuals, control = 163,861 individuals (Total N = 342,183 adults)
Interventions	2 types of mask used: surgical and cloth masks PLUS a brief video of notable public figures discussing why, how, and when to wear a mask, PLUS a brochure based on WHO materials depicting proper mask-wearing.
	Control villages: the control group did not receive any interventions See Table 1 for details.
Outcomes	Effectiveness: primary outcome: symptomatic seroprevalence (symptomatic and seropositive)
	Laboratory: seropositivity was defined by having detectable IgG antibodies against SARS-CoV-2
	Symptoms defined as per WHO-defined COVID-19 symptoms: (a) fever and cough; (b) 3 or more of the following symptoms (fever, cough, general weakness/fatigue, headache, myalgia, sore throat, coryza, dyspnoea, anorexia/nausea/vomiting, diarrhoea, altered mental status); or (c) loss of taste or smell.
	Secondary outcomes: prevalence of proper mask-wearing as wearing either a project mask or an alternative face-covering over the mouth and nose and improper mask-wearing as wearing a mask in any way that did not fully cover the mouth and nose; prevalence of physical distancing per WHO guideline that defines physical distancing as one meter of separation; prevalence of symptoms consistent with COVID-19: definition (see above)
	Safety not assessed. However, study mentioned that there was no adverse events reported during the study period

^{*} Indicates the major publication for the study



Abaluck 2022 (Continued)

Notes

The authors conclude that: a randomised trial of community-level mask promotion in rural Bangladesh during the COVID-19 pandemic shows that the intervention increased mask usage and reduced symptomatic SARS-CoV-2 infections, demonstrating that promoting community mask-wearing can improve public health (a scalable and effective method to promote mask adoption and reduce symptomatic SARS-CoV-2 infections.)

Funding: this trial was financially supported by a grant from GiveWell.org to Innovations for Poverty Action

The trial authors declare no competing interests.

Risk of bias

Bias	Authors' judgement	Support for judgement
DIAS	Authors judgement	Support for Judgement
Random sequence generation (selection bias)	Low risk	Random number generator used
Allocation concealment (selection bias)	High risk	Significant differences in the numbers of households included in each treatment group suggestive of a lack of allocation concealment
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Participants, mask promoters, and mask surveillance staff were not blinded as intervention materials were clearly visible
Blinding of outcome assessment (detection bias) All outcomes	High risk	Although the pre-specified analyses and sample exclusions were made by analysts blinded to the treatment assignment, investigators dropped individuals who were missing symptom data or who did not consent to blood spot collection from the primary outcome. One of the outcomes is COVID-19 symptoms reported by participants. Mask promoters, and mask surveillance staff were not blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Laboratory testing results were only available for around 40% of the symptomatic participants
Selective reporting (reporting bias)	High risk	Primary outcome of seroconversion was not reported

Aelami 2015

Study characteristics

Mothodo	A

Methods

A prospective cross-sectional study conducted during the Hajj season 2012. Pilgrims were randomised into 2 groups. The intervention group received education on personal hygiene including a hygienic package containing alcohol-based hand rub (gel or spray), surgical masks, soap, paper handkerchiefs, and user instructions; the control group did not receive any intervention. ILI was defined as the presence of at least 2 of the following during their stay: fever, cough, and sore throat. Questionnaires including demographic and clinical information were distributed amongst trained physicians before departure from Iran.

Participants

Total enrolled: 664 Iranian pilgrims (306 in the intervention group and 358 in the control group)

Inclusion criteria: not reported Exclusion criteria: not reported



Aelami 2015 (Continued)			
Interventions	Hygiene education and package. See Table 1 for details.		
Outcomes	ILI defined as the presence of at least 2 of the following during their stay: fever, cough, and sore throat		
	No safety outcomes we	ere reported.	
Notes	This is an abstract, therefore few details were reported. Funding not mentioned. Disclosure of interest: none declared.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Insufficient details provided	
Allocation concealment (selection bias)	Unclear risk	Insufficient details provided	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study	
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Insufficient details provided	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Insufficient details provided	
Selective reporting (reporting bias)	Unclear risk	Insufficient details provided	

Aiello 2010

Alello 2010	
Study characteristics	
Methods	Cluster-RCT assessing the effects of hand sanitiser and masks versus masks or no intervention on ILI symptoms. The trial was conducted in university halls of residence with more than 100 student residents in a US university during the 2006 to 2007 influenza "season". The study lasted 6 weeks.
	The units of randomisation were 7 of the 15 halls. 1 hall was very large (1240 residents), and the 6 remaining ones, which had between 110 and 830 residents, were combined into 2 clusters roughly equivalent in size. The 3 clusters were then randomised by random extraction of the clustered halls' names out of a container. The largest hall (single-cluster) was randomised to the mask and hand sanitiser arm; the 4-halls cluster received masks; and the remaining 2 halls were assigned as controls.
Participants	A total of 1297 with completed baseline survey and at least 1 weekly survey result were analysed (face mask and hand hygiene group = 367; face mask-only group = 378; control group = 552).
	Inclusion criteria: aged 18 or more, willing to wear mask and use alcohol-based hand sanitiser, have a throat swab specimen collected when ill, and complete the baseline and weekly surveys over the 6-week study period



Aiello 2010 (Continued)

Exclusion criteria: individuals reporting a skin allergy to alcohol were excluded

Recruitment of students began in 26 November, but the trial did not go "live" with distribution of intervention materials until 22 January 2007 when the first case of influenza was confirmed on campus by laboratory tests. Enrolment continued until 16 February 2007, and the study was completed on 16 March 2007. During the study period there was a 1-week break when the majority of residents left campus. There were 1327 eligible participants, 1297 of which had a complete baseline survey and at least 1-weekly survey result. It is unclear what the ineligibility criteria were for the 30 missing (1327 minus 1297), but the explanation may be in the appendix.

Interventions

Alcohol-based hand sanitiser (62% ethyl alcohol in a gel base) in a squeeze bottle and TECNOL procedure masks with ear loops (KC Ltd) and educational material or masks and educational material or no intervention. Compliance was encouraged within halls and outside. Sleep wearing was optional.

All participants received basic video-linked instruction on cough etiquette and hand sanitation. At baseline and weekly during the study, participants were asked to fill in a web-based survey collecting demographic and ILI symptom data. This was supplemented by direct observation of compliance by staff.

Compliance with "optimal handwashing" (at least 20 seconds 5 or more times a day) was significantly higher in the sanitiser-and-mask arm. See Table 1 for details.

Outcomes

Laboratory details are described in appendix.

Effectiveness: ILI, defined as cough and at least 1 constitutional symptom (fever/feverishness, chills, headache, myalgia). ILI cases were given contact nurses' phone numbers to record the illness and paid USD 25 to provide a throat swab. 368 participants had ILI, and 94 of these had a throat swab analysed by PCR. 10 of these were positive for influenza (7 for A and 3 for B).

Safety: N/A

Notes

The authors conclude that "These findings suggest that face masks and hand hygiene may reduce respiratory illnesses in shared living settings and mitigate the impact of the influenza A (H1N1) pandemic". This conclusion is based on a significantly lower level of ILI incidence in the mask and hand sanitiser arm compared to the other 2 arms after adjustment for covariates (30% to 50% less in arm 1 compared to controls in the last 2 weeks of the study).

Comparison with the ILI rate of the control arm may not be a reflection of the underlying rate of ILI because the intervention arm received instruction on hand sanitation and hand etiquette.

The play of adjustments is unclear. The intracluster correlation coefficient is reported in the footer of Table 4. Its very small size suggests lack of clustering within halls.

The role of spring break is mentioned in the Discussion, as are the results of this study compared to other studies included in our review (Cowling 2008 and MacIntyre 2009).

The authors report that 147 of 1297 participants (11.3%) had ILI symptoms "at baseline" and were excluded from analysis. During the 6 weeks of the study, 368 of 1150 participants (32%) had ILI. This averages out at about 5% per week. It is unclear what the term "at baseline" means; presumably this means during the 2 to 3 weeks of participant enrolment. If this is so, the reason for the triggering of the interventions (tied to influenza isolation) are obscure, as the trial is supposedly about ILI, and an ILI outbreak was already under way "at baseline".

This study has the same trial registration number as the Aiello 2012 study; the study was funded by government and pharmaceutical industry, i.e. this work was supported by funding from the Centers for Disease Control (CDC) and Prevention Grant U01 C1000441 (www.cdc.gov).

Disclosure of interest: none declared.



Aiello 2010 (Continued)

Bias	Authors' judgement	ment Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Described as randomised, but sequence generation not reported	
Allocation concealment (selection bias)	High risk	The residence hall units were randomised by blindly selecting a uniform ticket with the name of each hall out of a container (A.S.M. and A.A.) for randomisation assignment to each study arm.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study	
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessors blinded.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition is reported as follows: 9, 11, and 19 ineligible and 26, 52, and 21 lost to follow-up (respectively by arm), for a total of 39 and 99 for each reason for attrition. In total, 1297 (97%) of 1331 participants completed a baseline and at least 1-weekly survey.	
		The text reports an ITT analysis with only 1 ILI episode included by participant.	
		No reasons for the attrition of participants and swab volunteers are reported (were the swabs taken from a random sample or not?).	
Selective reporting (reporting bias)	High risk	There is no information on the causes of ILI other than the reporting on the 10 influenza PCR-positive swabs of 94 out of 368 students with ILI. This is a very low rate (and the Discussion confirms that the influenza season was mild), but investigation of the other known causes of ILI is not even mentioned in the text. This is especially important because stress, alcohol intake levels, and influenza vaccination were a significant predictor of ILI symptoms (Table 1). The reason for selective testing and/or reporting of influenza viruses tests over the other causes of ILI are unclear, especially as the study objective was focused on ILI. The text is also difficult to follow, weaving the reporting of ILI and influenza without a clear rationale.	

Aiello 2012

Study characteristics		
Methods	During the 2007 to 2008 influenza season, 1111 students residing in university residence halls were cluster-randomised by residence house (N = 37) to either face mask and hand hygiene, face mask only, or control arms. Discrete time survival analysis using generalised models estimated rate ratios according to study arm, each week and cumulatively over the 6-week intervention period, for clinically verified ILI and laboratory-confirmed influenza A or B.	
Participants	A total of 1187 young adults living in 37 residence halls, randomly assigned to 1 of 3 groups for 6 weeks: face mask use (n = 392), face masks with hand hygiene (n = 349), control (n = 370)	
	Inclusion criteria: aged 18 or more, willing to wear mask and use alcohol-based hand sanitiser, have a throat swab specimen collected when ill, and complete the baseline and weekly surveys over the 6-week study period	
	Exclusion criteria: individuals reporting a skin allergy to alcohol were excluded	



Aiello 2012 (Continued)			
Interventions	Participants were assigned to face mask and hand hygiene, face mask only, or control group during the study. See Table 1 for details.		
Outcomes	Clinically verified ILI: case definition (presence of cough and at least 1 or more of fever/feverishness, chills, or body aches)		
	Laboratory-confirmed influenza A or B. Throat swab specimens were tested for influenza A or B using RT-PCR.		
	No safety outcomes reported.		
Notes	This study has the same trial registration number as the Aiello 2010 study; the study was funded by government and pharmaceutical industry, i.e. this work was supported by funding from the Centers for Disease Control (CDC) and Prevention Grant U01 C1000441 (www.cdc.gov).		
	Disclosure of interest: none declared.		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer generation of sequence described.
Allocation concealment (selection bias)	Low risk	All residence houses in each of the residence halls were randomised prior to the intervention implementation.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	No blinding for study participants and personnel
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessors blinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition low and similar in each group
Selective reporting (reporting bias)	Low risk	2 outcomes specified and reported.

Alfelali 2020

Study characteristic	rs ·
Methods	Cluster open-label RCT
	Location: Mina, Greater Makkah, Saudi Arabia
	Follow up for 4 days
Participants	Arabic or English speaking Hajj pilgrims aged > 18 years from participating countries (Australia, Qatar and KSA) staying in allocated tents and able to provide signed informed consent were included.



Alt	elal	1 2020	(Continued)	
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Interventions	Mask wearing. See Table 1 for details.	
Outcomes	Effectiveness:	
	Laboratory: laboratory-confirmed viral respiratory infections (nasal swab on 650 participants only)	
	Secondary outcomes: clinical respiratory infections in participants	
	Safety reported on side effects of mask wearing	
	The most common side effects: difficulty in breathing (26.2%); discomfort (22%); a small minority (3%) reported feeling hot, sweating, a bad smell or blurred vision with eyeglasses	
Notes	The authors conclude that this trial was unable to provide conclusive evidence on facemask efficacy against viral respiratory infections most likely due to poor adherence to protocol. Funding: this report was made possible by a National Priorities Research Program grant (NPRP 6-1505-3-358) from the Qatar National Research Fund, a member of Qatar Foundation.	
	Disclosure of interests: the other authors have no competing interests to declare.	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Coin-tossing by an individual who was not a member of the research team
Allocation concealment (selection bias)	High risk	Used coin tossing which can introduce imbalance
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	No blinding
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Laboratory staff were blinded to the assigned intervention group
Incomplete outcome data (attrition bias) All outcomes	Low risk	Reported both intention-to-treat and per-protocol analysis and participant flow chart
Selective reporting (reporting bias)	Unclear risk	Insufficient information available.

Almanza-Reyes 2021

Study characteristic	s
Methods	RCT randomised using a computer-generated block scheme and stratified according to duty position, work shifts and the area/department of the service
	FU duration: 9 weeks
Participants	Workers (doctors, nurses, administrators) in a hospital for the exclusive recruitment of patients diagnosed with COVID-19 "General Tijuana Hospital"



Almanza-Reyes 2021 (Continued)

			ns

Experimental group: mouthwash and nose rinse

Silver mouth wash: 50 mL spray bottle containing AgNPs solution with 1 wt% concentration (0.6 mg/mLmetallic silver). Mix 4 to 6 spray shots (corresponding to volume ~ 0.5 mL) of this solution with 20 mL of water and to gargle with obtained solution for 15 to 30 seconds at least 3 times a day. Or use as nasal lavages on the inner part of the nasal alae and nasal passage with the same solution using a cotton swab twice a day.

Mouth spray: cover evenly the oral cavity with the direct 1 to 2 spray shots of solution without its previous dilution in water.

Control group: instructed to do mouth wash and nose rinse with a conventional mouthwash the way they normally did before the study

See Table 1 for details.

Outcomes

Effectiveness:

Laboratory: Lab-confirmed infection using RT-PCR; symptoms of respiratory tract infection (RTI) no definition given; clinical Evacuation: CT (Toshiba Aquilion 16, Japan) chest scan (random selection)

Safety: done using self-reported by participants using a questionnaire. "The present study also showed that no harmful side effects were observed in the 114 participants who used AgNPs as a mouthwash and nose rinse solution for 9 weeks"

Notes

Authors conclude that the mouth and nasal rinse with AgNPs helps in the prevention of SARS-CoV-2 infection in health personnel who are exposed to patients diagnosed with COVID-19. Funding: Funded studies A. Pestryakov Development Program "Priority 2030" Tomsk Polytechnic Uni-

Funding: Funded studies A. Pestryakov Development Program "Priority 2030" Tomsk Polytechnic University https://tpu.ru/en.

Conflict of interest statement: the authors have declared that no competing interests exist.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer-generated stratified block scheme
Allocation concealment (selection bias)	High risk	Unbalanced baseline prognostic factors (vaccination and frequency of handwashing)
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Insufficient information provided.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No participant flow chart reported.
Selective reporting (reporting bias)	Unclear risk	No protocol available



Alzaher 2018

Study characteristics			
Methods	A cluster-RCT conducted amongst girls attending 4 primary schools between January and March 2018. The participants attended a hand-hygiene workshop. The schoolgirls' absences were followed up for 5 weeks. Incidence rate, percentage of absence days, and absence rate were calculated for total and upper respiratory infections absences.		
Participants	A total of 496 schoolgirls aged of 6 to 12 years, attending 4 public primary girls' schools in the city of Riyadh, Saudi Arabia between January and March 2018. Students were randomised to education group (n = 234) or control group (n = 262).		
	Exclusion criteria: not reported		
Interventions	Hand hygiene workshop. See Table 1 for details.		
Outcomes	Incidence rate, percentage of absence days, and absence rate were calculated for total and upper respiratory infections absences.		
	The episode of URIs was defined as having 2 of the following symptoms for a day or 1 of the symptoms for 2 or more consecutive days: 1) a runny nose, 2) a stuffy or blocked nose or noisy breathing, 3) sneezing, 4) a cough, 5) a sore throat, and 6) feeling hot, having a fever or a chill.		
	No safety outcomes reported.		
Notes	Source of funding is unclear.		
	Disclosure of interest: none mentioned.		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Insufficient detail provided.
Allocation concealment (selection bias)	Low risk	Schools allocated prior to all schoolgirls attending selected schools were invited to participate.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (reporting bias)	Unclear risk	No protocol available



Arbogast 2016

Study characteristics	- -	
Methods	A 13.5-month prospective cluster-RCT executed with alcohol-based hand sanitiser in strategic work- place locations and personal use (intervention group) and brief hand hygiene education (both groups). Four years of retrospective data were collected for all participants.	
Participants	Data for a total of 1183 participants were analysed (intervention group = 525, control group = 607).	
	Inclusion criteria: all employees at 3 facilities who were 18 years of age or older, were enrolled in the company health insurance coverage, did not transfer between sites, and worked onsite full time (≥ 32 hours) were eligible for the study	
	Exclusion criteria: not reported	
Interventions	Alcohol-based hand sanitiser in strategic workplace locations and personal use (intervention group) and brief hand hygiene education (both groups). See Table 1 for details.	
Outcomes	The number of healthcare insurance claims, for a defined set of preventable illnesses, per participant per year	
	 Absenteeism, defined as the number of sick episodes per participant per year 	
	Claims based on ICD-9 codes	
	No safety outcomes reported.	
Notes	Only 2 clusters (1 per group) included, hence study data not included in meta-analysis.	
	Industry funded.	
	Disclosure of interest: none mentioned.	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No details provided.
Allocation concealment (selection bias)	Unclear risk	No details provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition minimal and similar in 2 groups
Selective reporting (reporting bias)	Unclear risk	No protocol available



Ashraf 2020

Study characteristics		
Methods	Geographically pair-matched community-based cluster-randomised trial	
	Used a random number generator to block	
	Open-label	
	Block randomised: unit of randomisation was a group of compounds visited by a single local promoter	
Participants	1. Infants (target child) will be eligible to participate in the study if:	
	a. they are in utero at the baseline survey.	
	b. their parents/guardians are planning to stay in the study village for the next 12 months (if a mother is planning to give birth at her natal home and then return, she will still be a candidate for enrolment)	
	2. Children < 36 months old at baseline that are living in the compound of a target child will be eligible to participate in diarrhoea measurement if:	
	a. they are < 36 months old at the baseline survey;	
	b. their parents/guardians are planning to stay in the study village for the next 12 months.	
	3. Children 18 to 27 months old at baseline that are living in the compound of a target child will be eligible to participate in intestinal parasite specimen collection if:	
	a. they are 18 to 27 months old at the baseline survey;	
	b. their parents/guardians are planning to stay in the study village for the next 12 months.	
Interventions	6 intervention arms: water quality, sanitation, hand washing, combined WSH, nutrition, nutrition + WSH	
	Intervention was delivered at the household or the compound level See Table 1 for details.	
Outcomes	Effectiveness:	
	Primary outcome: 7-day prevalence of acute respiratory illness (ARI). Defined as: caregiver-reported symptoms of persistent cough or panting, wheezing, or difficulty breathing (1 or 2) in the 7 days before the interview. No clinical data were collected	
	Secondary analyses: alternate combinations of the measured symptoms: 7-day prevalence of only panting, wheezing, or difficulty breathing (2) and ARI plus fever ([1 or 2] and 3)	
	Outcomes were measured approximately 12 and 24 months following intervention roll out.	
	Safety not assessed	
Notes	The authors conclude that: single targeted water, sanitation, and hygiene interventions reduced reported respiratory illness in young children. There was no apparent respiratory health benefit from combining WASH interventions.	
	Financial support: this research was funded by Global Development grant OPPGD759 from the Bill & Melinda Gates Foundation to the University of California, Berkeley, CA. S. P. L., S. A., M. I., B. F. A., and J. M. C. report grants from the Bill & Melinda Gates Foundation during the conduct of the study. P. K. R. reports grants from Leland Stanford University during the conduct of the study for support to the WASH Benefits project. M. R. reports grants and non financial support from the Bill & Melinda Gates Foundation (through a subcontract from UC Berkeley) during the conduct of the study.	
	Disclosure of interest: none mentioned.	



Ashraf 2020 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random number generator
Allocation concealment (selection bias)	Low risk	Random allocation by an offsite investigator
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The research team who implemented the intervention was separate from the data collection team. The analysis was carried out masked to the allocated group.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Provided participants flow diagram showing minimal attrition.
Selective reporting (reporting bias)	Low risk	Reported the pre-specified outcomes.

Azor-Martinez 2016

Study characteristics	
Methods	Randomised, controlled, and open study with an 8-month follow-up. The experimental group washed their hands with soap and water, together with using hand sanitiser, and the control group followed their usual handwashing procedures. Absenteeism rates due to URIs were compared between the 2 groups through a multivariate Poisson regression analysis. The per cent of days missed in both groups were compared with a z test.
Participants	A sample of 1341 (intervention group = 621, control group = 720)
	Inclusion criteria: children 4 to 12 years old, attending 5 state schools in Almeria (Spain) whose parents/guardians had signed an informed consent document
	Exclusion criteria: children who had any of the following chronic illnesses that predisposed them to infection: neoplasia, primary and secondary immunodeficiencies, cystic fibrosis, chronic treatment with high doses of steroids or immunosuppressants
Interventions	Hand-washing workshops of 2-hour duration. The experimental group washed their hands with soap and water together with using hand sanitiser, whilst the control group followed usual hand-washing procedures. See Table 1 for details.
Outcomes	Absenteeism rates due to URIs
	Per cent of days missed
	Respiratory illness was defined by 2 of the following symptoms during 1 day, or 1 of the symptoms for 2 consecutive days: (1) runny nose; (2) stuffy or blocked nose or noisy breathing; (3) cough; (4) feeling hot or feverish or having chills; (5) sore throat; or (6) sneezing.



Azor-Martinez 2016 (Continued)

A school absenteeism case (episode) was defined as when a child failed to attend school due to an URI. Common infectious illnesses, such as conjunctivitis, and skin infections were not included. Other causes for absenteeism, such as doctors' appointments, family vacations, and accident injuries, were also excluded.

No safety outcomes reported.

Notes Government funded

Disclosure of interest: none mentioned.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	A random number table was used.
Allocation concealment (selection bias)	Low risk	Schools/classes allocated prior to children recruited.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	High risk	Attrition levels high and different in the 2 groups
Selective reporting (reporting bias)	Unclear risk	No protocol available

Azor-Martinez 2018

Study characteristics	
Methods	A cluster-RCT, controlled, and open study of 911 children aged 0 to 3 years attending 24 DCCs in Almería, Spain, with an 8-month follow-up. 2 intervention groups of DCC families performed educational and hand hygiene measures, 1 with soap and water (n = 274), another with hand sanitiser (n = 339), and the control group followed usual hand-washing procedures (n = 298). Respiratory infection (RI) episode rates were compared through multilevel Poisson regression models. The percentage of days missed were compared with Poisson exact tests.
Participants	A total of 911 children attending 24 DCCs in Almería (Spain).
	Inclusion criteria: children between 0 and 3 years old enrolled in DCCs and attending for at least 15 hours per week whose parents or guardians had signed an informed consent
	Exclusion criteria: children with chronic illness or medication that could affect their likelihood of contracting an infection



Azor-Martinez 2018 (Contin	Data were analysed for 911 participants: hand sanitiser group (n = 339), soap and water group (n = 274), and control group (n = 298). 2 intervention groups. 1 group used soap and water, another used hand sanitiser, whilst the control group followed usual hand-washing procedures. Groups received 1-hour hand hygiene workshop. See Table 1 for details.		
Interventions			
Outcomes	Primary: RI incidence rate		
	Secondary: (1) the presence or absence of at least 1 antibiotic prescription for each new RI episode during the study period (topical antibiotics were excluded), and (2) the percentage of RI absenteeism days in the 3 groups calculated as the ratio of RI absenteeism days to all possible days of attendance		
	DCC absenteeism episode was defined as when a child failed to attend a DCC because of an RI.		
	Respiratory illness was defined as the presence of 2 of the following symptoms during 1 day or the presence of 1 of the symptoms for 2 consecutive days: (1) runny nose, (2) stuffy or blocked nose or noisy breathing, (3) cough, (4) feeling hot or feverish or having chills, (5) sore throat, or (6) sneezing.		
	No safety outcomes reported.		
Notes	Government funded. This work was supported by a grant from the Andalusia Department of Health.		
	Competing interests: the authors have indicated they have no potential conflicts of interest to disclose.		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer randomisation using statistical software for the sequence
Allocation concealment (selection bias)	Low risk	Clusters assigned prior to recruitment.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition minimal and similar in 3 groups
Selective reporting (reporting bias)	Unclear risk	No protocol available

Ban 2015

Study characteristics	
Methods	Quote: "Group randomised" trial. Only 2 clusters, which were 2 kindergartens in Xiantao City, Hubei Province, China.



Bias	Authors' judgement Support for judgement		
Risk of bias			
Notes	Funding not mentioned. Disclosure of interest: none mentioned.		
Outcomes	Respiratory illness, defined as: 2 or more of the following: fever, cough and expectoration, runny nose and nasal congestion, collected by parental questionnaire. Axillary temperature higher than 37.3 °C or the range of temperature fluctuation is more than 1 °C. 'Cough and expectoration' were defined as 3 or more coughs in a single hour and lasting for 4 or more hours in a single day, with or without expectoration. 'Runny nose and nasal congestion' were defined as a runny nose lasting for 4 or more hours in 1 day, with or without nasal congestion.		
Interventions	Intervention group: hand hygiene and surface-cleaning education and provision of products for kindergarten and home use. Control group: usual practice. See Table 1 for details.		
	5 classes (221 children) randomly selected from 1 kindergarten in the intervention group and 6 classes (244 children) randomly selected from another kindergarten in the control group. Children were aged 5 or under. There were 72 exclusions from the analysis.		
Participants	Data for a total of 393 participants were analysed (intervention group = 194, control group = 199).		
Ban 2015 (Continued)			

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Method not described, and only 2 clusters.
Allocation concealment (selection bias)	Unclear risk	Method not described.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	High risk	Parental report, and parents were aware of treatment allocation
Selective reporting (reporting bias)	High risk	Attrition reported and balanced between groups, but high rate of attrition in a trial with small numbers of participants.

Barasheed 2014

Study characteristics	
Methods	Pilot, non-blinded, parallel, cluster-RCT
Participants	22 tents were randomly selected from the Australian pilgrims camped in Mina, during Hajj in 2011; 12 tents were allocated to the mask group and 10 tents to the control group. A total of 164 Australian pilgrims were recruited: 75 in the mask group (39 'cases' and 36 'contacts') and 89 in the control group (36 'cases' and 53 'contacts').



Barasheed 2014 (Continued)

Inclusion criteria for index case: 1) Australian pilgrims of any gender aged > 15 years who attend the Hajj 2011, and 2) have symptoms of respiratory infection for 3 days. For close tent contact: 1) Australian pilgrims of any gender aged 15 years or more who attend the Hajj 2011, and 2) pilgrims who share the same tent and sleep "immediately close" to the index case.

Exclusion criteria: for index case: 1) pilgrims who do not suffer from symptoms of respiratory infection, 2) pilgrims who present with symptoms of respiratory infection for > 3 days, and 3) children aged less than 15 years. For close tent contact: 1) pilgrims who are symptomatic at presentation, 2) pilgrims who are not close tent contacts of an index case, and 3) children aged less than 15 years. Only 10% to 15% of potential participants took part in the study.

Interventions

"supervised mask use" versus "no supervised mask use". See Table 1 for details.

Outcomes

Laboratory: 2 nasal swabs from all ILI cases and contacts, 1 for influenza POCT using the QuickVue Influenza (A+B) assay (Quidel Corporation, San Diego, USA) and 1 for later nucleic acid testing for influenza and other respiratory viruses. However, there was a problem with getting POCT on time during Hajj.

Effectiveness: to assess the effectiveness of face masks in the prevention of transmission of ILI. ILI was defined as subjective (or proven) fever plus 1 respiratory symptom (e.g. dry or productive cough, runny nose, sore throat, shortness of breath).

Safety: none planned or reported

Notes

The study was conducted from 4 November 2011 to 10 November 2011.

Compliance with face mask use by pilgrims was 56 of 75 (76%) in the mask group and 11 of 89 (12%) in the control group (P < 0.001). The proportion of face mask user in the 'mask' tents was 76% for both males (19/25) and females (38/50). The most often reported reason for not wearing face masks was discomfort (15%).

Government funded: Qatar National Research Fund (QNRF).

The other authors have declared no conflict of interest in relation to this work.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Insufficient information provided.
Allocation concealment (selection bias)	Unclear risk	Quote: "tents were randomised to either intervention group (supervised mask tent) or control group (no supervised mask tent) by an independent study coordinator who was not an investigator", but did not mention how
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Because advice from the Saudi Ministry of Hajj to all pilgrims included recommending the wearing of masks, all pilgrims, both cases and controls, were asked about mask-wearing"
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Self-reported outcomes (nasal swab was performed for those who reported ILI symptoms and was not intended as systematic detection). ILI was defined as subjective (or proven) fever plus 1 respiratory symptom.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up, all numbers were reported from enrolment to analysis
Selective reporting (reporting bias)	Low risk	All planned outcomes were reported.



Biswas 2019

Study characteristics			
Methods	Cluster-RCT in 24 primary schools in Dhaka to assess the effectiveness of hand sanitiser and a respiratory hygiene education intervention in reducing ILI and laboratory-confirmed influenza during June to September 2015. 12 schools were randomly selected to receive hand sanitiser and respiratory hygiene education, and 12 schools received no intervention. Field staff actively followed children daily to monitor for new ILI episodes (cough with fever) through school visits and by phone if a child was absent. When an illness episode was identified, medical technologists collected nasal swabs to test for influenza viruses.		
Participants	A total of 10,855 stude schools = 5778 children	nts were enrolled in the study (intervention schools = 5077 children; control n).	
	Children aged 5 to 10 y	rears educated in 24 randomly selected primary schools in Dhaka, Bangladesh	
		t offered education above grade 5 because of differences in student populations had previously received a hand or respiratory hygiene intervention	
Interventions	Hand sanitiser and res	piratory hygiene education versus no intervention. See Table 1 for details.	
Outcomes	Incidence of ILI		
	Incidence of laboratory-confirmed influenza (RT-PCR)		
	An ILI episode was defined as measured fever ≥ 38 °C or subjective fever and cough. If a child was absent, the field staff followed up by phone to identify the reason for absenteeism and to determine if the child met the ILI case definition. If a child in a participating school had an ILI episode, a trained medical technologist visited the child's household to obtain consent from the child's parent/guardian and collect a nasal swab from the child within 48 hours of symptom onset. If it was outside the 48-hour window, the sample was not collected.		
	No safety outcomes re	ported.	
Notes	Government funded.		
	Disclosure of interest:	none mentioned.	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Sequence generated using a computer-based random number generator.	
Allocation concealment (selection bias)	Low risk	Allocation completed prior to individuals being recruited.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study	
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study	
Incomplete outcome data (attrition bias)	High risk	Information missing for 30 children (28 children in the control schools and 2 children in the intervention schools)	



Biswas 2019	(Continued)
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All outcomes

Selective reporting (reporting bias)

Unclear risk

No protocol available

Bundgaard 2021

Study characteristics	
Methods	Investigator-initiated, nationwide, unblinded, randomised controlled trial stratified by the 5 regions of Denmark
Participants	Inclusion criteria: community-dwelling adults aged 18 years or older without current or prior symptoms or diagnosis of COVID-19 reported being outside the home amongst others for at least 3 hours per day, and who did not wear masks during their daily work.
	Exclusion criteria: previously tested positive for SARS-CoV-2 and wear face masks at work
Interventions	Exposure: mask (N = 2392)
	Control group: no mask (N = 2470)
	Both groups received materials and instructions for antibody testing on receipt and at 1 month; materials and instructions for collecting an oropharyngeal/nasal swab sample for polymerase chain reaction (PCR) testing at 1 month and whenever symptoms compatible with COVID-19 occurred during follow-up. They registered symptoms and results of the antibody test in the online REDCap system. Written instructions and instructional videos guided antibody testing, oropharyngeal/nasal swabbing, and proper use of masks, and a help line was available to participants. See Table 1 for details.
Outcomes	Study duration: 1 month
	Effectiveness: primary outcome (composite) SARS-CoV-2 infection, defined as a positive result on an oropharyngeal/nasal swab test for SARS-CoV-2, development of a positive SARS-CoV-2 antibody test result (IgM or IgG) during the study period, or a hospital-based diagnosis of SARS-CoV-2 infection or COV-ID-19.
	Secondary outcome: PCR evidence of infection with other respiratory viruses
	Safety: adverse reaction: 14% in mask group (no further descriptions)
Notes	The authors conclude that inconclusive results, missing data, variable adherence, patient-reported findings on home tests, no blinding, and no assessment of whether masks could decrease disease transmission from mask wearers to others. Funding: the primary funding source was The Salling Foundations.
	Disclsure can be viewed at www.acponline.org /authors/icmje/ConflictOfInterestForms.do?msNum=M20-6817.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer algorithm stratified by the 5 regions of Denmark
Allocation concealment (selection bias)	Unclear risk	Insufficient information reported



Bundgaard 2021 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded. Patient reported symptoms, POCT testing, patient-reported findings on home tests.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Participant flow chart showed acceptable attrition
Selective reporting (reporting bias)	Low risk	All pre-specified outcomes reported.

Canini 2010

Study characteristics	
Methods	A cluster-RCT conducted in France during the 2008 to 2009 influenza season. Households were recruited during a medical visit of a household member with a positive rapid influenza A test and symptoms lasting less than 48 hours. Households were randomised either to the mask or control group for 7 days In the intervention arm, the index case had to wear a surgical mask from the medical visit and for a period of 5 days. The trial was initially intended to include 372 households, but was prematurely interrupted after the inclusion of 105 households (306 contacts) following the advice of an independent steering committee. Generalised estimating equations were used to test the association between the intervention and the proportion of household contacts who developed an ILI during the 7 days following the inclusion.
Participants	A total of 105 households were randomised, which represented 148 contacts in the intervention arm and 158 in the control arm.
	The study was conducted in 3 French regions (Ile de France, Aquitaine, and Franche-Comté) and included households of size 3 to 8.
	Exclusion criteria: if index patient was treated for asthma or chronic obstructive pulmonary disease or was hospitalised
Interventions	Surgical mask versus no mask. See Table 1 for details.
Outcomes	The primary endpoint was the proportion of household contacts who developed an ILI during the 7 days following inclusion. Exploratory cluster-level efficacy outcome, the proportion of households with 1 or more secondary illness in household contacts.
	A temperature over 37.8 °C with cough or sore throat was used as primary clinical case definition.
	Adverse reactions due to mask-wearing
Notes	Government funded.
	Competing interests: the authors have declared that no competing interests exist.
Risk of bias	
Bias	Authors' judgement Support for judgement



Canini 2010 (Continued)		
Random sequence generation (selection bias)	Low risk	Randomisation lists were generated by a computerised program.
Allocation concealment (selection bias)	Low risk	Randomisation was performed centrally by the GP after written consent on an interactive voice response system dedicated to the study.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessors blinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	All households included in analysis.
Selective reporting (reporting bias)	Low risk	All specified outcomes reported.

Carabin 1999

Study characteristics	•
Methods	Cluster-RCT carried out in DCCs in the Canadian province of Quebec between 1 September 1996 and 30 November 1997 (15 months). The aim was to test the effects of a hygiene programme on the incidence of diarrhoea and fecal contamination (data not extracted) and on colds and URTIs. The design included before and after periods analysed to assess the Hawthorne effect of study participation on control DCCs. The unit of randomisation was DCC, but analysis was also carried out at classroom and single-child level. This is a common mistake in cluster-RCT analysis. DCCs were stratified by URTI incidence preceding the trial and randomised by location. Cluster coefficients are not reported.
Participants	A total of 1729 children aged 18 to 36 months in 47 DCCs (83 toddler classrooms)
	Inclusion criteria: presence of at least 1 sandbox and 1 play area and of at least 12 available toddler places
	For the autumn of 1997 intervention group (24 DCCs, 43 classrooms, and 414 children), control group (23 DCCs, 23 classrooms, and 374 children). It is not clear what is the distribution and data for the autumn of 1996.
Interventions	Training session (1 day) with washing of hands, toy cleaning, window opening, sand pit cleaning, and repeated exhortations to hand wash. See Table 1 for details.
Outcomes	Laboratory: N/A Effectiveness: diarrhoea and coliform contamination (data not extracted) Colds (nasal discharge with at least 1 of the following: fever, sneezing, cough, sore throat, earache, malaise, irritability) URTI (cold of at least 2 days' duration) Surveillance was carried out by educators, annotating absences or illness on calendars. Researchers al so filled in a phone questionnaire with answers by DCC directors. Safety: N/A
Notes	Risk of bias: high (no description of randomisation; partial reporting of outcomes, numerators, and denominators)



Carabin 1999 (Continued)

Notes: the authors conclude that the intervention reduced the incidence of colds (IRR 0.80, 95% CI 0.68 to 0.93). This was a confusingly written study with unclear interweaving of 2 study designs. For unclear reasons analysis was only carried out for the first autumn. Unclear why colds are not reported in the results. Cluster-coefficients and randomisation process were not described.

Support for the study was provided by the Rhone-Poulenc Rorer Canada Ltd.

Disclosure of interest: none mentioned.

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Block randomisation of DCC according to region, but sequence generation no reported
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Blinding not possible (hygiene session plus educational material versus none)
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Originally 52 eligible DCCs with 89 classrooms agreed to take part, but 5 dropped out (2 closed, 1 was sold, 2 either did not provide data or the data were "unreliable", and 6 classrooms had insufficient data). 43 children failing to attend DCC for at least 5 days in the autumn were also excluded. ITT analysis was carried out including an additional DCC whose director refused to let staff attend the training session. No correction made for clustering.
Selective reporting (reporting bias)	High risk	Denominators unclear and not explained

Chard 2019

Cilaid 2019	
Study characteristic	s
Methods	Cluster-RCT conducted amongst 100 randomly selected primary schools lacking functional WASH facilities in Saravane Province, Lao People's Democratic Republic. Schools were randomly assigned to either the intervention (n = 50) or comparison (n = 50) arm. Intervention schools received a school water supply, sanitation facilities, hand-washing facilities, drinking water filters, and behaviour change education and promotion. Comparison schools received the intervention after research activities had ended. At unannounced visits every 6 to 8 weeks, enumerators recorded pupils' roll-call absence, enrolment, attrition, progression to the next grade, and reported illness (diarrhoea, respiratory infection, conjunctivitis), and conducted structured observations to measure intervention fidelity and adherence. Stool samples were collected annually prior to de-worming and analysed for soil-transmitted helminth (STH) infection. In addition to our primary ITT analysis, we conducted secondary analyses to quantify the role of intervention fidelity and adherence on project impacts.
Participants	100 primary schools (50 intervention, 50 comparison) with a total of 3993 pupils were enrolled throughout the study period (intervention schools = 2021 pupils, control schools = 1972 pupils). Up to 40 pupils



Chard 2019 (Continued)	selected from grades 3 to 5 in each school using systematic stratified sampling, with grade and sex as the stratification variables. Pupils selected at baseline were followed throughout the entire study period; pupils who left the school due to abandonment or transfer were replaced at the beginning of the following academic year, maintaining equal grade and sex ratios when possible. Pupils who progressed from fifth to the sixth grade were replaced with pupils from grade 3 the following academic year.
Interventions	Water supply, sanitation facilities, hand-washing facilities, drinking water filters, and behaviour change education and promotion versus control. See Table 1 for details.
Outcomes	Primary impact of interest was pupil absence, measured by school-wide roll-call at each visit. Secondary health impacts included diarrhoea, symptoms of respiratory infection, and conjunctivitis/non-vision-related eye illness collected through pupil interviews. Pupils were considered to have symptoms of respiratory infection if they reported cough, runny nose, stuffy nose, or sore throat. No safety outcomes reported.
Notes	Funded by government and pharmaceutical industry. Competing interests: all authors have completed the ICMJE uniform disclosure form at http://www.icmje.org/coi_disclosure.pdf (available upon request from the corresponding author) and declare no conflicts of interest.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Insufficient details provided.
Allocation concealment (selection bias)	Low risk	Schools allocated prior to recruitment of individuals.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Exclusions were due to participants leaving school, hence unlikely to cause bias.
Selective reporting (reporting bias)	Low risk	All specified outcomes reported.

Correa 2012

Study characteristics	
Methods	Cluster-RCT in childcare facilities in Colombia from 16 April to 18 December 2008 (3 school terms) testing the effects of hand hygiene using an alcohol-based hand rub versus standard practice



Correa 2012 (Continued)			
Participants	42 childcare facilities in 6 towns in Colombia. A total of 1727 were enrolled (intervention group = 794 from 21 centres, control group = 933 from 21 centres).		
	Inclusion criteria: licensed to care for 12 or more children aged 1 to 5 years for 8 hours a day, 5 times per week, and where availability of tap water was limited		
Interventions	Intervention: alcohol-based hand wash as an addition to hand-washing		
	Control: usual hand-washing practice		
	See Table 1 for details.		
Outcomes	ARI defined as: 2 or more of the following symptoms for at least 24 hours, lasting at least 2 days: runny, stuffy, or blocked nose or noisy breathing; cough; fever, hot sensation, or chills; and/or sore throat. Ear pain alone was considered an ARI.		
Notes	This work was supported by a grant from the Global Development Network (New Delhi, India), "Fifth Global Research Project: Promoting Innovative Programs from the Developing World: Towards Realizing the Health MDG's in Africa and Asia," and the Bill and Melinda Gates Foundation (Seattle, Washington, United States).		
	Authors declare to have no conflicts of interest.		

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"using the random function in Microsoft Excel™ (Microsoft Corp., Redmond, Washington, United States), random numbers (1 or 2) were generated and allotted 1:1 within each group. Finally, a researcher flipped a coin to decide which number would correspond to either arm (heads = 1, intervention; tails = 2, control)."
Allocation concealment (selection bias)	Unclear risk	Method not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	Lost to follow-up similar in each group and not substantial
Selective reporting (reporting bias)	Unclear risk	No protocol available

Cowling 2008

Study characteristics



Cowling 2008 (Continued)

Methods

Cluster-RCT carried out in Hong Kong SARS between February and September 2007. The study assessed the effects of non-pharmaceutical interventions on the household transmission of influenza over a 9-day period. ILI cases whose family contacts had been symptom-free for at least 2 weeks rapid-tested for influenza A and B were used and randomised to 3 interventions. Randomisation was carried out in 2 different schedules (2:1:1 for the first 100 households, and subsequently 8:1:1), but it is unclear why and how this was done.

Participants

A total of 350 of 944 originally enrolled participants representing 122 households were analysed (control group = 71 households with 205 household contacts, face mask = 21 households with 61 household contacts, HH = 30 households with 84 household contacts).

Inclusion criteria: residents of Hong Kong aged at least 2 years, reporting at least 2 symptoms of ILI ((such as fever ≥ 38 degrees, cough, headache, coryza, sore throat, muscle aches and pains) and positive influenza A+B rapid test

and living in a household with at least 2 other individuals, none of whom had ILI in the preceding 14 days

Households were excluded because subsequent laboratory testing (culture) was negative.

Attrition was not explained.

Interventions

Households were randomised to either wearing face masks with education (as the control group plus education about face mask use) or hand-washing with special medicated soap (with alcohol sanitiser) with education (as the control group plus education about hand-washing) or education about general healthy lifestyle and diet (control group). The soap was distributed in special containers that were weighed at the start and end of the study. Interventions visits to the households were done on average 1 day after randomisation of index case household. See Table 1 for details.

Outcomes

Laboratory: QuickVue RTI

MDCK culture

Samples were harvested using NTS, but the text refers to a second procedure from June 2007 onwards testing for non-influenza viruses, with no data reported.

Effectiveness: secondary attack ratios (SAR): SAR is the proportion of household contacts of an index case who were subsequently ill with influenza (symptomatic contact individuals with at least 1 NTS positive for influenza by viral culture or PCR)

3 clinical definitions were used for secondary analysis:

- 1. Fever ≥ 38 degrees, or at least 2 of following symptoms: headache, coryza, sore throat, muscle aches and pains
- 2. At least 2 of the following S/S: fever ≥ 37.8 degrees, cough, headache, sore throat, muscle aches and pains
- 3. Fever ≥ 37.8 degrees plus cough or sore throat

Safety: no harms were reported in any of the arms

Notes

The trial authors conclude that "The secondary attack ratios were lower than anticipated, and lower than reported in other countries, perhaps due to differing patterns of susceptibility, lack of significant antigenic drift in circulating influenza virus strains recently, and/or issues related to the symptomatic recruitment design. Lessons learnt from this pilot have informed changes for the main study in 2008". Although billed as a pilot study, the text is highly confusing and at times contradictory. The intervention was delivered at a home visit up to 36 hours after the index case was seen in the outpatients. This is a long time and perhaps the reason for failure of the intervention. Practically, the intervention will have to be organised before even seeking medical care, i.e. people know to do it when the child gets sick at home.

This work has received financial support from the US Centers for Disease Control and Prevention (grant no. 1 U01 Cl000439-01), the Research Fund for the Control of Infectious Disease, Food and Health Bu-



Cowling 2008 (Continued)

reau, Government of the Hong Kong SAR, and the Area of Excellence Scheme of the Hong Kong University Grants Committee (grant no. AoE/M-12/06).

Competing Interests: the authors have declared that no competing interests exist.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Low risk	Randomisation was computer generated by a biostatistician.
tion (selection bias)		Quote:"A pre-specified table of random numbers will be used to assign one of the three interventions to the household of the index case."
Allocation concealment (selection bias)	Low risk	The households of eligible study index patients were allocated to 3 groups in a 1:1:1 ratio under a block randomisation structure with randomly permuted block sizes of 18, 24, and 30 using a random-number generator. Allocation was concealed from treating physicians and clinics and implemented by study nurses at the time of the initial household visit.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Participants and people who administered the interventions were not blinded to the interventions, but participants were not informed of the specific nature of the interventions applied to other participating households.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Dropout was accounted for. Dropout from the randomised population was high: 32% in control group, 37.5% in hand hygiene group, and 39.4% in face mask and hand hygiene group. Reasons for dropout were distributed evenly across the 3 groups.
		Authors report follow-up as proportion of patients remaining in the study after initial dropout.
Selective reporting (reporting bias)	High risk	The choice of season, change in randomisation schedules, and unexplained dropouts amongst contacts; the use of QuickVue, which proved unreliable, reporting bias on non-influenza isolates resulted in a judgement of high risk of bias.

Cowling 2009

Study characteristics

Methods	Cluster-RCT
Participants	A total of 407 index cases and 794 household contacts were analysed.
	Of 407 enrolled households, 322 received the allocated interventions as follows:
	 control group = 112 households with 346 contacts (only 91 households analysed with 279 contacts); hand hygiene = 106 households with 329 contacts (only 85 households analysed with 257 contacts); face mask + hand hygiene = 104 households with 340 contacts (only 83 households analysed with 258 contacts).



Cowl	ing	2009	(Continued)
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Inclusion criteria: households in Hong Kong. Index cases from 45 outpatient clinics in both the private and public sectors across Hong Kong. They enrolled individuals who reported at least 2 symptoms of ARI (temperature 37.8 °C, cough, headache, sore throat, or myalgia); had symptom onset within 48 hours; and lived in a household with at least 2 other people, none of whom had reported ARI in the preceding 14 days. After giving informed consent, participants provided nasal and throat swab specimens.

2750 patients were eligible and tested between 2 January and 30 September 2008.

Interventions

Participants with a positive rapid-test result and their household contacts were randomly assigned to 1 of 3 study groups: control (lifestyle measures - 134 households), control plus enhanced hand hygiene only (136 households), and control plus face masks and enhanced hand hygiene (137 households) for all household members. No detailed description of the instructions was given to participants. See Table 1 for details.

Outcomes

Influenza virus infection in household contacts, as confirmed by RT-PCR or diagnosed clinically after 7 days

"The primary outcome measure was the secondary attack ratio at the individual level: that is, the proportion of household contacts infected with influenza virus. We evaluated the secondary attack ratio using a laboratory definition (a household contact with a nose and throat swab specimen positive for influenza by RT-PCR) as the primary analysis and 2 secondary clinical definitions of influenza based on self-reported data from the symptom diaries as secondary analyses."

Statistical analysis: adjusted for clustering

Results: no statistically significant difference in secondary attack ratio between groups in total population. Statistically significant reduction in RT-PCR confirmed influenza virus infections in the household contacts in 154 households in which the intervention was applied within 36 hours of symptom onset in the index patient. Adherence to hand hygiene was between 44% and 62%. Adherence of index patient to wearing a face mask between 15% and 49%.

Notes

"In an unintentional deviation from that protocol, 49 of the 407 randomly allocated persons had a household contact with influenza symptoms at recruitment (a potential co-index patient). We also randomly assigned 6 of 407 persons who had symptoms for slightly more than 48 hours."

The trial authors conclude that "Hand hygiene and face masks seemed to prevent household transmission of influenza virus when implemented within 36 hours of index patient symptom onset. These findings suggest that non-pharmaceutical interventions are important for mitigation of pandemic and interpandemic influenza".

Primary funding source: Centers for Disease Control and Prevention.

Potential conflicts of interest: none disclosed.

Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Randomisation was computer generated by a biostatistician.	
		Quote:"A pre-specified table of random numbers will be used to assign one of the three interventions to the household of the index case."	
Allocation concealment (selection bias)	Low risk	The households of eligible study index patients were allocated to 3 groups in a 1:1:1 ratio under a block randomisation structure with randomly permuted block sizes of 18, 24, and 30 using a random-number generator. Allocation was concealed from treating physicians and clinics and implemented by study nurses at the time of the initial household visit.	
Blinding of participants and personnel (perfor- mance bias)	High risk	Quote: "Participants and personnel administering the interventions were not blinded to group assignment."	



Cowling 2009 (Continued) All outcomes		
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	It is not stated if the outcome assessor was blinded.
Incomplete outcome data (attrition bias) All outcomes	High risk	Dropout was accounted for. Dropout from the randomised population was high: 32% in control group, 37.5% in hand hygiene group, and 39.4% in face mask and hand hygiene group. Reasons for dropout were distributed evenly across the 3 groups.
		Trial authors report follow-up as proportion of patients remaining in the study after initial dropout.
Selective reporting (reporting bias)	Unclear risk	In general good reporting

DiVita 2011

Study characteristics		
Methods	The impact of hand-washing promotion on the risk of household transmission of fever was tested in rural Bangladesh. ILI was defined as fever in children < 5 years cough or sore throat in individuals > 5 years old. Households were randomised to trol. The intervention group received hand-washing stations with soap and daily tivation at critical times for pathogen transmission, such as after coughing or sne lance was conducted, and household members with fever were tested for influen condary attack ratios (SAR) were calculated for influenza, ILI, and fever in each ar with generalised estimating equations was used to estimate the significance of the whilst controlling for clustering by household.	s old and fever with intervention or con- hand-washing mo- ezing. Daily surveil- iza viruses by PCR. Se- m. Logistic regression
Participants	The study included 233 patient index cases (intervention group = 100, control group 133) with 2540 household contacts (intervention group = 134, control group = 1226). Inclusion criteria: index case patients (individuals who developed ILI within the previous 2 days and were the only symptomatic person in their household) as well as their household contacts	
Interventions	Hand-washing stations with soap and daily hand-washing motivation versus control. See Table 1 for details.	
Outcomes	SAR were calculated for influenza, ILI, and fever.	
	ILI was defined as fever in children < 5 years old and fever with cough or sore throat in individuals > 5 years old.	
	No safety outcomes reported.	
Notes	Funding source unknown.	
	Disclosure of interest: none declared.	
Risk of bias		
Bias	Authors' judgement Support for judgement	
Random sequence generation (selection bias)	Unclear risk Insufficient details provided	



DiVita 2011 (Continued) Allocation concealment (selection bias)	Unclear risk	Insufficient details provided
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Insufficient details provided
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Insufficient details provided
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Insufficient details provided
Selective reporting (reporting bias)	Unclear risk	Insufficient details provided

Farr 1988a

Study characteristics	s ·	
Methods	6-month cluster-RCT, controlled, double-blind of the efficacy of virucidal nasal tissues in the prevention of natural cold, conducted in Charlottesville, Virginia, USA. Many of the families were enrolled because 1 or more family members worked at the State Farm Insurance Company; the remaining families were recruited from the Charlottesville community by advertisement in a local newspaper. Families were randomly assigned by the sponsoring company to receive boxes of treated tissues, placebo tissues, or no tissues. The randomisation was performed by computer. Study participants and investigators were unaware of the type of tissues each family was randomised to receive. Blinding efficacy was tested using a questionnaire: the mothers in each family were asked twice if she believed her family was using virucidal or placebo tissues. Participants in the treated and placebo groups were instructed to use only tissues received through the study, whilst families in the additional control group without tissues were allowed to continue their usual practice of personal hygiene. Each family member kept a daily listing of respiratory symptoms on a record card. A nurse epidemiologist visited each family monthly to encourage recording.	
Participants	186 families, 58 in the active group, 59 in the placebo group, and 69 in the no-tissues group. A total of 302 families were originally recruited; 116 families who did not comply with the study protocol, lost their surveillance cards, could not complete the protocol were excluded from the analysis.	
Interventions	Use of virucidal tissues versus placebo tissues versus no tissues. The treated tissues were impregnated with malic and citric acids and sodium lauryl sulphate, whilst placebo tissues contained saccharin. See Table 1 for details.	
Outcomes	Laboratory: serological evidence: no Effectiveness: respiratory illness Safety: N/A	
Notes	The authors concluded that virucidal tissues have only a small impact on the overall rate of natural acute respiratory illnesses. The total illness rate was lower in families using virucidal tissues than in both of the other study groups, but only the difference between active and placebo groups was statistically significant (3.4 illness per person versus 3.9 for placebo group, $P = 0.04$, and 3.6 for the no-tissue control group, $P = 0.2$, and overall 14% to 5% reduction). The questionnaire results suggest that some bias may have been present since a majority of mothers in the virucide group believed they were receiving the 'active' tissues. Another possible explanation of the low effectiveness of virucidal tissues	



Farr 1988a (Continued)

is poor compliance by children in use of the virucidal tissues. A well-designed and honestly reported study

Funding source not reported.

Potential conflicts of interest: none disclosed.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "The randomisation was performed by computer in each trial." However, method of sequence generation is not stated.
Allocation concealment (selection bias)	Unclear risk	Quote: "In trial I, families were randomly assigned by the sponsoring company to receive boxes of treated tissues, placebo tissues or no tissues."
		Quote: "Families with one or two children were randomised in one stratum, and families with three or more children were randomised in a second stratum in trial I."
		Concealment of allocation not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Study participants and investigators were unaware of the type of tissues which each family was randomised to receive in both trials. In trial I, the mother in each family was asked twice if she believed her family was using active or placebo tissues, first after three months and then at the end of the study."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Study participants and investigators were unaware of the type of tissues which each family was randomised to receive in both trials. In trial I, the mother in each family was asked twice if she believed her family was using active or placebo tissues, first after three months and then at the end of the study."
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote: "A total of 116 of the 302 families were excluded from the analysis. Families were excluded if they lost their surveillance cards or did not conscientiously record data, did not comply with the study protocol, or simply could not complete the protocol for family reasons. It was discovered that families with five or more members had so many colds that it was not possible to distinguish primary and secondary illnesses. These large families were therefore excluded from the analysis in trial I and were excluded from enrolment in trial II."
Selective reporting (reporting bias)	Low risk	All indicated outcomes are reported.

Farr 1988b

Study characteristics

Methods

Six-month randomised, controlled, double-blind trial of the efficacy of virucidal nasal tissues in the prevention of natural cold, conducted in Charlottesville, Virginia, USA. Families were recruited from the Charlottesville community by advertisement in a local newspaper. Families were randomly assigned by the sponsoring company to receive either virucidal tissues or placebo-treated tissues. Stratified randomisation was performed by computer, and the strata were defined by total number in the family. Study participants and investigators were unaware of the type of tissues each family was randomised to receive. Each family member kept a daily listing of respiratory symptoms on a record card. A nurse



Farr 1988b (Continued)		
	epidemiologist visited each family monthly to encourage recording. In addition, a study monitor visited each family bimonthly to further encourage compliance and reporting of symptoms.	
Participants	98 families, 58 in the active group and 40 in the placebo group. 231 families were initially recruited, 222 completed the trial, data of 98 families were analysed. The other families were excluded from the analysis because they complained of side effects (sneezing, etc.) or reported not using the tissues regularly. See Table 1 for details.	
Interventions	Use of virucidal tissues versus placebo tissues. The treated tissues were impregnated with malic and citric acids and sodium lauryl sulphate, whilst the placebo tissues contained succinic acid. Participant in the treated and placebo groups were instructed to only use tissues received through the study.	
Outcomes	Laboratory: serological evidence: no Effectiveness: respiratory illness Safety: N/A	
Notes	The study suggests that virucidal tissues have only a small impact on the overall rate of natural acute respiratory illnesses. The total illness rate was lower in families using virucidal tissues than in the other study group, but the difference between active and placebo groups was not statistically significant. There was a small, non-significant drop in illness rates across families (5%). The tissues appeared to be ineffective as the drop was confined to primary illness unaffected by tissue use. The placebo (succinic acid) was not inert, and was associated with cough and nasal burning. This impacted on allocation concealment. A well-designed and honestly reported study marred by transparent allocation.	
	Funding source not reported.	
	Potential conflicts of interest: none disclosed.	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"The randomisation was performed by computer in each trial." However, method of sequence generation is not stated.
Allocation concealment (selection bias)	Unclear risk	Quote: "In trial II, families were randomly assigned by the sponsor to receive either virucidal tissues or placebo treated tissues."
		Quote:"In trial II, stratified randomisation was again used, but this time the strata were defined by total number in the family (i.e., one stratum for two-member families, another stratum for three-member families, and a final one for four-member families)."
		Concealment of allocation not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Study participants and investigators were unaware of the type of tissues which each family was randomised to receive in both trials."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Study participants and investigators were unaware of the type of tissues which each family was randomised to receive in both trials."
Incomplete outcome data (attrition bias) All outcomes	High risk	Quote:"A total of 222 (of 231) families completed trial II; 9 families were terminated early (table 1). In 124 families, one or more family members reported not using the tissues regularly and/or reported having significant side effects. The data from these families were not analysed, leaving 58 families (177 persons) and 40 families (114 persons) for analysis in the virucide and placebo groups, respectively."



Farr 1988b (Continued)

Selective reporting (reporting bias)

Low risk

All indicated outcomes are reported.

Feldman 2016

Study characteristics			
Methods	Prospective cluster-RCT. Ships from a single, central naval base. Ships were stratified by vessel classes (corvette, fast missile boat, and patrol boat).		
Participants	All people participating in security operations, routine exercises, and patrol at a single, central naval base were eligible.		
	The actual number of participants in the groups is not reported.		
Interventions	Chlorhexidine gluconate (CHG) dispensers in addition to soap-and-water hand-washing versus soap-and-water hand-washing. See Table 1 for details.		
Outcomes	Laboratory: bacterial palm cultures from 30 sailors from each group using a modified bag broth technique with sterile brain-heart broth, at 0 and 4 months (sample participants)		
	Effectiveness: Primary outcome: incidence of infectious diseases reported by the computerised patient records system using ICD-9 diagnoses and grouped into diarrhoeal, respiratory, and skin infections; the number of sick call visits; and the number of sick leave and light-duty days incurred by the sailors		
	Secondary outcome: subclinical morbidity (i.e. symptoms of self-reported infectious diseases)		
	Safety: not reported		
Notes	No report on adherence		
	Study was conducted between May and September 2014 (4 months follow-up).		
	CHG availability onboard the ships did not reduce the transmission of infectious diseases or colonisation.		
	Government funded (Israeli Defense Force Medical Corps).		
	Potential conflicts of interest: none disclosed.		

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No description of randomisation
Allocation concealment (selection bias)	Unclear risk	No description of allocation
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded. Self-reported outcomes
Blinding of outcome assessment (detection bias)	Unclear risk	No information if personnel collecting data for ICD-9 diagnosis were blinded



Fel	dman	2016	(Continued)
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All outcomes

Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No participants flow chart, no attrition data	
Selective reporting (reporting bias)	Unclear risk	No protocol to compare	

Fretheim 2022a

Study characteristics	s	
Methods	Pragmatic RCT	
Participants	3717 participants in Norway (glasses n = 1852; no glasses n = 1865)	
	Inclusion criteria:	
	 were at least 18 years of age; did not regularly wear glasses; owned or could borrow glasses that they could use (e.g. sun-glasses); had not contracted COVID-19 in the 6 weeks prior to participation; did not have COVID-19 symptoms when providing consent; were willing to be randomised to wear, or not wear glasses outside their home when close to others for a 2-week period; provided informed consent; and contact lenses were allowed in the control group for those dependent on this visual aid. Exclusion criteria: does regularly wear glasses (contact lenses are accepted); and contracted COVID-19 after December 15th 2021. 	
Interventions	Intervention group: wearing eyeglasses (any type) when close to other people outside their home (on public transport, in shopping malls etc.), over a 14-day period. The control: encouraged not to wear glasses when close to others outside their home. See TIDieR Table (Table 1) for details.	
Outcomes	Primary outcome	
	 Any positive COVID-19 test result reported to the Norwegian Surveillance System for Communicable Diseases (MSIS), from day 3 to day 17 of the study period. 	
	Secondary outcomes	
	 Any positive COVID-19 test result based on self-report, from day 1 to day 17 of the study period. Episode of respiratory infection based on self-report of symptoms from day 1 to day 17 of the study period. Respiratory infection was defined as having 1 respiratory symptom (stuffed or runny nose, sore throat, cough, sneezing, heavy breathing) and fever, or 1 respiratory symptom and at least 2 more symptoms (body ache, muscular pain, fatigue, reduced appetite, stomach pain, headache, loss of smell). Healthcare use for respiratory symptoms, self-reported, from day 1 to day 17 of the study period. Healthcare use for injuries, self-reported, from day 1 to day 17 of the study period. Healthcare use for respiratory symptoms as registered in Norwegian Patient Registry (NPR), from day 3 to day 28 of the study period. 	



Fretheim 2022a (Continued)

- 7. Healthcare use for injuries (from day 1 to day 21 as registered in NPR and the Norwegian Registry for Primary Health Care (KPR), from day 3 to day 28 of the study period.
- 8. Healthcare use (all causes) as registered in NPR and KPR from day 1 to day 21 of the study period.

Notes

The study did not report on the <u>latter 4 outcomes</u> due to lack of access to this data at the time of publication

Negative experiences of using the eyeglasses were reported: fogging, feeling uncomfortable and tiring, reduced vision, fall, feeling silly when wearing glasses indoor, headache.

Funding: the costs of running the trial were covered by the Centre for Epidemic Interventions Research (CEIR), Norwegian Institute of Public Health.

Competing interests: all authors declare: no competing interests.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Automatically randomised after signing the consent form in the online recruitment platform (Nettskjema).
Allocation concealment (selection bias)	High risk	A digital recruitment platform (Nettskjema) was used to generate allocation. However, more participants in the intervention group wore face masks.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	An open-label study. Participants and investigators were not blinded.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Outcome is self-reported positive COVID-19 test result reported to the Norwegian Surveillance System for Communicable Diseases (MSIS). However, the public policy requiring confirmatory PCR-test had changed during the study conduct which may have affected case detection.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Participants flow chart was provided.
Selective reporting (reporting bias)	Low risk	No deviation from the published protocol.

Goodall 2014

Study characteristic	s
Methods	A 2X2 factorial RCT with 4 treatment arms
	1. Vitamin D ₃ and gargling
	2. Placebo and gargling
	3. Vitamin D ₃ and no gargling
	4. Placebo and no gargling
Participants	600 students from McMaster University, Hamilton, Ontario, Canada, randomised to the following.
	1. Vitamin D and gargling (N = 150, analysed 135)
	2. Vitamin D and no gargling (N = 150, 123 outcomes included in analysis)



Goodall 2014 (Continued)

- 3. Placebo and gargling (N = 150, 121 known outcomes included in analysis)
- 4. Placebo and no gargling (N = 150, 113 known outcomes included in analysis)

Inclusion criteria: aged ≥ 17 years and lived with at least 1 student house mate.

Exclusion criteria: students with contraindicated medical conditions (hypercalcaemia, parathyroid disorder, chronic kidney disease, use of anticonvulsants, malabsorption syndromes, sarcoidosis), who were currently or planning to become pregnant, who were taking ≥ 1000 international units (IU)/day vitamin D, or who were unable to swallow capsules

Interventions

See Table 1 for details.

Outcomes

Laboratory (influenza assessed via weekly self-collected nasal swabs; only swabs for symptomatic participants were assessed). Lab-confirmed influenza was determined by testing the Day 1 nasal swabs using an in-house enterovirus/rhinovirus PCR and, if negative, a commercial multiplex PCR able to detect 16 respiratory viruses and viral subtypes (xTAG RVP FAST, Luminex, Austin TX).

Clinical URTI assessed via weekly online surveys.

Clinical URTI is defined as the participant's perception of cold in conjunction with 1 or more symptoms (runny/stuffy nose, congestion, cough, sneezing, sore throat, muscle aches, or fever). When participants reported symptoms but were uncertain if they were ill, adjudication was applied by 2 clinicians.

Safety:

None assessed/reported by the investigators.

Notes

Study was conducted during 2 periods: September to October in 2010 and 2011.

Partial governmental funding.

Competing interests: the authors declare that they have no competing interests.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No description on how the randomisation sequence was generated
Allocation concealment (selection bias)	Low risk	Study used opaque, sealed, serially numbered envelopes. Envelopes were only accessed when both personnel were present.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Due to the nature of gargling with tap water, this intervention was not blinded. However, all other aspects of the study were blinded. Self-reported symptoms were adjudicated by 2 clinicians.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Except for gargling, all other participants and study personnel were blinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Study flow chart and reasons for lost to follow-up are provided, imputation used for missing outcomes.
Selective reporting (reporting bias)	Low risk	All planned study outcomes were reported and match the published study protocol.



Gutiérrez-García 2022

Study characteristics	
Methods	Single-blind (analyst) randomised controlled trial carried out in a single centre in Mexico City during September to November 2020. Randomisation was through tokens in opaque envelopes but the trial was open to all except the data analysts. There were some imbalances in age groups post-randomisation at baseline in age and comorbidities
Participants	85 front line healthcare workers, unvaccinated and with no history of COVID infection in each arm. 6 and 1 were excluded from the analysis as they tested positive to CUVID within 14 days of recruitment. Follow-up was 2 weeks
Interventions	Neutral electrolysed water (SES) (pH 6.5 to 7.5) nasal and oral rinses 3 times daily and PPE versus PPE only for the prevention of SARS-CoV-2 infection. See Table 1 for details.
Outcomes	Laboratory
	RT-PCR no further described "according to the WHO guidelines", once only presumably with symptoms.
	Effectiveness
	COVID-19 disease confirmed by RT-PCR, between the 14th day since their recruitment and the 28th day of follow-up. The following are listed as COVID-19 signs and symptoms: dry cough, fever > 37.5°C, headache, myalgia, arthralgia, rhinorrhoea, conjunctivitis, pharyngodynia, odynophagia. 1 and 10 participants were positive in the intervention and control arms respectively. All 11 were nurses.
	Safety
	Local harms from SES applications – none reported
Notes	The authors conclude that quote: "the prophylactic protocol was demonstrated as a protective factor, in more than 90%, for developing the disease, and without adverse effects. Nasal and oral rinses with SES maybe an efficient alternative to reinforce the protective measures against COVID-19 disease and should be further investigated."
	Funding: no funding was received.
	Competing interests: the authors RGG, JCA and IDE declare that they have no competing interests. ACL, NMS and BPM state that they are employees at Esteripharma S.A. de C.V. company but did not participate in the decision to publish the results of the study, nor in the selection of the volunteers or in its development.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Insufficient information provided.
Allocation concealment (selection bias)	High risk	Nurse or doctor chose one of two identical tokens that were placed inside an opaque plastic container. One token was labelled 'with SES' (treatment group) and the other 'without SES' (control group).
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias)	Low risk	Primary endpoint was the number of healthcare professionals, nurses, or physicians, with COVID-19 disease confirmed by



Gutiérrez-García 2022 (Continued) All outcomes		RT-PCR. Researchers that performed the statistical analyses were blinded.	
Incomplete outcome data (attrition bias) All outcomes	Low risk	Minimal exclusions from the analysis.	
Selective reporting (reporting bias)	Low risk	Pre-specified outcomes reported.	

Gwaltney 1980

Study characteristics	
Methods	Study assessed the effectiveness of aqueous iodine applied to the fingers in blocking hand transmission of experimental infection with rhinovirus from 1 volunteer to another. Healthy, young adult volunteers were recruited from the general population at the University of Virginia, Charlottesville. Volunteers were not informed about the contents of the hand preparation until after the study. 2 experiments were conducted to evaluate the virucidal activity of aqueous iodine applied to the fingers immediately before viral contamination. Another 2 experiments were conducted to determine whether there was sufficient residual activity of aqueous iodine after 2 hours to interrupt viral spread by the hand route. Volunteers who were donors of virus for the hand exposures were challenged intranasally on 3 consecutive days with the rhinovirus strain HH. Recipients were randomly assigned to receive iodine or placebo. The donors contaminated their hands with nasal secretions by finger to nose contact before the exposure. Hand contact was made between a donor and a recipient by stroking of the fingers for 10 seconds. Donors and recipients wore masks during the exposure period.
Participants	15 and 20 volunteers in 2 experiments
Interventions	Treatment of fingers with iodine versus placebo. The virucidal preparation used was aqueous iodine (2% iodine and 4% potassium iodide). The placebo was an aqueous solution of food colours. See Table 1 for details.
Outcomes	Experimental rhinovirus infection reduced (P = 0.06) Laboratory: serological evidence Effectiveness: rhinovirus infection (based on serology, isolation, and clinical symptoms) with high- score clinical illness. Score was published elsewhere. Safety: N/A
Notes	Risk of bias: high (poor description of randomisation process, concealment, or allocation) Notes: the study suggests that aqueous iodine applied to the fingers was effective in blocking transmission by hand contact of experimental infection with rhinovirus for up to 2 hours after application (1 out 10 volunteers were infected compared to 6 out of 10 in the placebo preparation arm, P = 0.06 with Fisher's exact test). The effectiveness of iodine treatment of the fingers in interrupting viral transmission in volunteers recommends its use for attempting to block transmission of rhinovirus under natural conditions. Although the cosmetic properties of 2% aqueous iodine make it impractical for routine use, it can be used as an epidemiologic tool to study the importance of the hand transmission route and to develop an effective cosmetically acceptable hand preparation. A summarily reported study.
	Funding source not reported.
	Disclosure of interest: none mentioned.
Risk of bias	
Bias	Authors' judgement Support for judgement



Gwaltney 1980 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Insufficient information
Allocation concealment (selection bias)	Unclear risk	insufficient information
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote:Quote: "The viricidal preparation used was aqueous iodine The placebo was an aqueous solution of food colors mixed to resemble the color of iodine. An odor of iodine was given to the placebo Volunteers were not informed about the contents of the hand preparation until after the study."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	It is not stated whether the outcome assessor was blinded or not.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Insufficient information
Selective reporting (reporting bias)	Unclear risk	Insufficient information

Hartinger 2016

Study characteristics	5
Methods	Communities were randomised to a comprehensive intervention was an improved solid-fuel stove, installation of a kitchen sink with running water, solar drinking water disinfection, education on handwashing, and separating animals from the kitchen environment.
Participants	534 children (267 in each group) in 51 communities (25 in intervention, 26 in control group). 250 children/households in the intervention group and 253 children/households in the control group were available for follow-up. Conducted in a rural farming area
Interventions	Environmental home-based intervention package consisting of improved solid-fuel stoves, kitchen sinks, solar disinfection of drinking water, and hygiene promotion. See Table 1 for details.
Outcomes	Laboratory: Escherichia coli (not relevant to this review)
	Effectiveness: weekly collection of daily diary data on illness. ARI was defined as child presenting cough or difficulty breathing, or both. ALRI was defined as child presenting cough or difficulty breathing, with a raised respiratory rate (> 50 per min in children aged 6 to 11 months and > 40 per min in children aged 12 months) on 2 consecutive measurements.
	Safety: none described in methods and none reported
Notes	The authors conclude that "combined home-based environmental interventions slightly reduced child-hood diarrhoea, but the confidence interval included unity. Effects on growth and respiratory outcomes were not observed, despite high user compliance of the interventions. The absent effect on respiratory health might be due to insufficient household air quality improvements of the improved stoves and additional time needed to achieve attitudinal and behaviour change when providing composite interventions".
	Well-reported trial. Age of children not reported.
	Funding: this work was supported by the UBS Optimus Foundation, Freiwillige Akademische Gesellschaft, Basel, Stiftung EmiliaGuggenheim-Schnurr, Basel.



Hartinger 2016 (Continued)

Conflict of interest: the authors have no conflicts of interest to declare.

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Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Covariate-constrained randomisation is mentioned, but method not described.
Allocation concealment (selection bias)	Unclear risk	Method not mentioned
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Data collected by field worker and recorded by parent. All would be aware of allocation.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Low attrition rate, reasons stated, balanced between groups.
Selective reporting (reporting bias)	Low risk	It is unlikely that other outcomes were measured but not reported.

Helsingen 2021

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Study Characteristics	
Methods	Non-inferiority open randomised trial carried out in May 25 to June 15 2020 during the first lockdown in Norway. Eligible individuals were randomised 1:1 stratified by fitness centre by a computerised random number generator to no access to fitness centre or access to fitness centre with "mitigation measures"
Participants	3825 people aged 18 to 65 with no risk factors for Covid 19 (diabetes, cardiovascular disease including hypertension, age > 65). 61 randomised participants (18 and 43, respectively) withdrew consent before start of the intervention with 3764 remaining
Interventions	The intervention consisted in gym access with: avoidance of body contact; 1 m distance between individuals at all times; 2 m distance for high intensity activities; disinfection of all work stations; cleaning of all equipment after use by participant; regular cleaning of facilities and access control by facility employees to ensure distance measures and avoid overcrowding; open changing rooms with showers and saunas remained closed; staff was present during all opening hours; lids on trash cans removed; individuals were instructed to stay home if they had any Covid-19 related symptoms, participants were advised to avoid touching their eyes, nose and mouth. See Table 1 for details.
Outcomes	Laboratory Self-administered (at times facilitated by HCW) NP, saliva or OP swabs in transport medium taken at day 14 to 15 from beginning sent to central lab. RT-RPC performed. Testing of antibodies (IGG) was carried out in late June with a mailed self-administered spot slide which was then mailed and analysed centrally.
	Effectiveness



Helsingen 2021 (Continued)

Primary: PCR positivity in both arms

Co-primary: hospital admission in the two arms at 21 days (via data linkage)

Secondary: proportion of participants with SARS-CoV-2 antibodies in the 2 study arms at 30 days. Testing also carried out for gym staff.

Safety

NR

Notes

The authors conclude that "Provided good hygiene and physical distancing measures and low population prevalence of SARS-CoV-2infection, there was no increased infection risk of SARS-CoV-2 in fitness centres in Oslo, Norway for individuals without Covid-19-relevant comorbidities." There was low and declining incidence on C19 in the Oslo area during the time of the trial as reported by the authors. The authors call the analysis set ITT but consent withdrawal individuals were not part of the analysis. There was marked difference in PCR uptake (88.7% in the training arm; 71.4% in the no-training arm) and no cycle thresholds are reported.

Funding: this study was funded by the Norwegian Research Council, grant no. 312757. The grant paid for necessary equipment, study personnel and researchers.

Competing interests: Dr. Lise M. Helsingen reports grants from Norwegian Research Council (grant no. 312757), during the conduct of the study. All other authors declare no competing interests in relation to this work.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer random-number generator
Allocation concealment (selection bias)	High risk	Allocation performed by one of the study authors
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded.
Incomplete outcome data (attrition bias) All outcomes	High risk	More women were compliant with SARS-CoV2 testing in the training arm as compared to the no-training arm, and compliant individuals were somewhat younger in the training arm compared to the non-training arm.
Selective reporting (reporting bias)	Low risk	All pre-specified outcomes reported

Hubner 2010

Study	char	acte	ristics
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Methods

A prospective, controlled, intervention-control group design to assess the epidemiological and economical impact of alcohol-based hand disinfectants use at workplace. Volunteers in public administra-



Hubner 2010	(Continued)

tions in the municipality of the city of Greifswald were randomised into 2 groups. Participants in the intervention group were provided with alcoholic hand disinfection, the control group was unchanged. In all, 1230 person-months were evaluated.

Participants

Employees (n = 134) from the administration of the Ernst-Moritz-Arndt University Greifswald, the municipality of Greifswald and the state of Mecklenburg-Pomerania, were recruited for the study and randomised to intervention (N = 67) or control (N = 67). Final analysis was performed on 64 from the intervention and 65 from the control group.

Inclusion criteria: all administrative officers, who did not already apply hand disinfection at work, were considered for participation and were invited by email or mail (n = 850). The 134 participants declared their written consent to participate and completed a pre-study survey with demographic, social, health, and work-related questions to provide data for randomisation.

Exclusion criteria: employees that were already using hand disinfectants at work

Interventions	Alcohol-based hand disinfectants use at workplace versus usual hygiene. See Table 1 for details.
Outcomes	Respiratory and gastrointestinal symptoms and days of work were recorded based on a monthly questionnaire over 1 year.
Notes	Funding source not mentioned.
	Competing interests: the authors declare a financial competing interest: GK is employed by Bode Chemie GmbH, Hamburg, Germany. NOH and AK received financial support for research from Bode

Chemie in the past. All other authors declare no conflict of interest.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No details provided.
Allocation concealment (selection bias)	Unclear risk	No details provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Self-reported outcomes
Incomplete outcome data (attrition bias) All outcomes	Low risk	Lost to follow-up minimal and similar in 2 groups
Selective reporting (reporting bias)	Unclear risk	No protocol available

Huda 2012

Study characteristics



Huda 2012 (Continued)		
Methods	Poorly described cluster-RCT. Partial report of the SHEWA-B trial focused on changing 11 targeted behaviours in villages to measure the impact on diarrhoea and respiratory illness amongst children. Unit of randomisation is not clear, but was probably a village. A group of 10 to 17 households within a village were the participants, based on the household having at least 1 child under the age of 5.	
Participants	A total of 1692 participants (intervention = 848, control = 844) at baseline and 1699 participants at 18 months (intervention = 849, control = 850)	
	Households were eligible if they have a child < 5 years of age and a guardian agreed to participate.	
Interventions	SHEWA-B programme targeting improved latrine coverage and usage, access to and use of arsenic-free water, and improved hygiene practices using soaps. See Table 1 for details.	
Outcomes	Laboratory: none described in methods and none reported	
	Effectiveness: ARI and diarrhoea. ARI defined as cough and fever or difficulty breathing and fever within 48 hours prior to interview.	
	Safety: none described in methods and none reported	
Notes	The authors conclude that quote: "The prevalence of childhood diarrhea and respiratory illness was similar in the intervention and control communities".	
	Poorly reported trial.	
	This research activity was funded by the United Kingdom's Department for International Development (DFID).	
	Disclosure of interest: none mentioned.	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Mentions random-number tables, but not clear if this was for random selection or randomisation
Allocation concealment (selection bias)	Unclear risk	Method not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Data on illness were collected by a resident of the village, who was likely to know treatment allocation.
Incomplete outcome data (attrition bias) All outcomes	High risk	Not reported. No flow diagram
Selective reporting (reporting bias)	Unclear risk	Unlikely that other outcomes were measured and not reported



Ibfelt 2015

Study characteristics	
Methods	Cluster-RCT in 12 daycare nurseries in Denmark. Centres in the intervention group had their linen and children's toys commercially cleaned and disinfected every 2 weeks. Control group centres had usual practice. Swabbing for bacteria and respiratory viruses was conducted at baseline and the end of the intervention period.
Participants	12 nurseries in Copenhagen (intervention = 6, control = 6) with a total of 587 children aged 6 months to 3 years
	Not clear how many children were in each group. Data on illness collected at the individual level, and on presence of bacteria and viruses at the cluster level.
Interventions	Washing and disinfection of toys and linen every 2 weeks for 3 months. See Table 1 for details.
Outcomes	Laboratory: counts of bacteria (not relevant to this review) and 11 respiratory viruses at baseline and end of intervention period, taken from swabs of 10 predefined locations in playroom (7 locations) and toilet area (3 locations). Viruses were influenza A and B; coronavirus NL63229E, OC43, and HKU1; parainfluenza virus 1, 2, 3, and 4; rhinovirus; RSV A/B; adenovirus; enterovirus; parechovirus; metapneumovirus; and bocavirus. Testing by PCR
	Effectiveness: illness counts in the children. Absence due to sickness recorded daily with reason categorised, but no definitions of illness provided.
	Safety: none mentioned in methods and none reported
Notes	The authors conclude that "Although cleaning and disinfection of toys every two weeks can decrease the microbial load in nurseries, it does not appear to reduce sickness absence among nursery children".
	The results of the disinfection are reported as follows: "The most prevalent virus was coronavirus (97% positive samples), followed by bocavirus (96%), adenovirus (73%) and rhinovirus (46%). The intervention reduced the presence of adenovirus, rhinovirus and RSV approximately two- to five-fold [odds ratio (OR) 2.4, 95% confidence interval (CI) 1.1-5.0 for adenovirus; OR 5.3, 95% CI 2.3-12.4 for rhinovirus; OR 4.1, 95% CI 1.5-11.2 for RSV] compared with the control group. On the other hand, metapneumovirus was found significantly less often in the control group than in the intervention group. The intervention had no effect on the detection of other viruses. The fomites with the highest presence of respiratory virus were pillows and sofas, followed by toys and playroom tables. When looking at the samples from the toys alone, there was a significant decrease following the intervention in the intervention group compared with the control group for rhinovirus (OR 3.8, 95% CI 1.3-10.5; P = 0.01) and RSV (OR 5.2, 95% CI 1.1-23.8; P = 0.04), but not adenovirus".
	This a poorly reported cluster-RCT. Its importance lies in the surface viral prevalence data (which could have been overestimated by PCR) and the finding that even in the presence of high viral prevalence, sickness was lower in the control (no surface disinfection) arm. This suggests the absence of other factors that could activate surface respirators viruses.

Funding: this work was supported by the Danish Council for Technology and Innovation under the Ministry of Science, Innovation and Higher Education as part of the Sundhed i Børneinstitutioner innovation consortium.

Conflict of interest statement: Ecolab Denmark, Berendsen Denmark and 3M Denmark supplied materials and cleaning free of charge, but had no influence on the analysis of the data or the writing of the manuscript.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method not mentioned

tors that could activate surface respiratory viruses.



Ibfelt 2015 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Method not mentioned
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Objective measure of bacterial and viral counts. However, illness reporting is unclear.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No attrition or denominators given for results.
Selective reporting (reporting bias)	Low risk	Unlikely that other outcomes were measured but not reported

Ide 2014

Study characteristics			
Methods	Randomised, open-label, 2-group parallel study of 757 high school students (15 to 17 years of age) conducted for 90 days during the influenza epidemic season from 1 December 2011 to 28 February 2012 6 high schools in Shizuoka Prefecture, Japan. The green tea gargling group gargled 3 times a day with bottled green tea, and the water gargling group did the same with tap water. The water group was restricted from gargling with green tea.		
Participants	A total of 747 students were enrolled (green tea gargling group = 384, water gargling group = 363)		
	High school students (15 to 17 years of age) who attended 6 high schools in the Kakegawa and Ogasa districts of Shizuoka Prefecture, Japan		
Interventions	See Table 1 for details.		
Outcomes	Incidence of laboratory-confirmed influenza		
	Incidence of clinically defined influenza infection		
	Time for which the participant was free from clinically-defined influenza infection		
	Clinically-defined influenza infection, specified as fever (≥ 37.8 °C) plus any 2 of the following additional symptoms: cough, sore throat, headache, or myalgia. Influenza infection with viral antigen was detected by immunochromatographic assay.		
	No safety data reported.		
Notes	Funding: this work was supported by Grants-in-Aid for Scientific Research (KAKENHI) Grant Number 23590887.		
	Competing Interests: the authors have declared that no competing interests exist.		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Ide 2014 (Continued)		
Random sequence generation (selection bias)	Low risk	Computer-generated permuted block randomised schema
Allocation concealment (selection bias)	Low risk	Randomised at the Data Management Center of Shizuoka General Hospital in Japan
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	Minimal attrition
Selective reporting (reporting bias)	Unclear risk	Protocol not available

Ide 2016

Study characteristics			
Methods	Randomised controlled study in Japan. Participants were randomly allocated into the catechin-treated (epigallocatechin gallate-treated) or non-treated face mask groups for 60 days from January to March 2016. Incidence of laboratory-confirmed influenza infection was measured and compared between groups using Fisher's exact test. Multivariate analysis was performed to calculate adjusted ORs and associated 95% CIs.		
Participants	Participants included workers in a nursing home, a rehabilitation facility, and a hospital.		
	A total of 234 participants were eligible for the study (catechin group, n = 118; control group, n = 116).		
Interventions	Catechin-treated mask versus non-treated face mask. See Table 1 for details.		
Outcomes	Incidence of laboratory-confirmed influenza infection		
	Laboratory-confirmed influenza infection with viral antigen detected by immunochromatographic assay performed when participants reported ILI.		
	No safety outcomes reported.		
Notes	Funding: this work was supported in part by a grant from the Japan Society for the Promotion of Science (JSPS), through the Grant-in-Aid for JSPS Fellows (No. 15J10190 to KI) and Grants-in-Aid for Scientific Research (C) (15K08924 to HY).		
	Conflict of Interest: the authors declare that they have no conflicts of interest.		
Risk of bias			
Bias	Authors' judgement Support for judgement		



Ide 2016 (Continued)		
Random sequence generation (selection bias)	Unclear risk	Computer-generated randomisation, but method not stated
Allocation concealment (selection bias)	Low risk	Central randomisation service at Data Management Centre of Shizouka General Hospital
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Double-blinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Attrition minimal
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition minimal
Selective reporting (reporting bias)	Low risk	Specified outcomes reported.

Jacobs 2009

Study characteristics	s		
Methods	Open-RCT lasting 77 days from January 2008 to test "superiority" of face masks in preventing "URTI". This term appears as an acronym in the introduction and is not explained. It is assumed that it stands for 'upper respiratory infections', but it is preceded in the text by the term 'common cold', which is also lacking a definition. Randomisation was carried out in blocks within each of 3 professional figures (physicians, nurses, and "co-medical" personnel).		
Participants	33 HCWs mainly females aged around 34 to 37 in a tertiary healthcare hospital in Tokyo, Japan. HCW with quote: "predisposing conditions" (undefined) to "URTI" and those taking antibiotics were excluded.		
	A baseline descriptive survey was carried out including "quality of life".		
	1 participant dropped out at end of week 1, but no reason is reported nor the allocation arm.		
	Analysis was performed on 32 participants (mask = 17, no mask = 15).		
Interventions	Surgical mask MA-3 (Osu Sangyo, Japan) during all phases of hospital work ($n = 17$) or no mask ($n = 15$) (except when specifically required by hospital SOPs). See Table 1 for details.		
Outcomes	Laboratory: N/A		
	Effectiveness: URTI is defined on the basis of a symptoms score, with a score > 14 being a URTI according to Jackson's 1958 criteria ("Jackson score"). These are not explained in text, although the symptoms are listed in Table 3 (any, sore throat, runny nose, stuffy nose, sneeze, cough, headache, ear ache feel bad) together with their mean and scores SD by intervention arm.		
	Safety: the text does not mention or report harms. These appear to be indistinguishable from URTI symptoms (e.g. headache which is reported as of significantly longer duration in the intervention arm) Compliance is self-reported as high (84.3% of participants).		



Jacobs 2009 (Continued)

Notes

The authors conclude that quote: "Face mask use in healthcare workers has not been demonstrated to provide benefit in terms of cold symptoms or getting colds. A larger study is needed to definitively establish non-inferiority of no mask use".

This is a small, badly reported trial. The purpose of trials is to test hypotheses not to prove or disprove 'superiority' of interventions. There is no power calculation, and CIs are not reported (although there is a mention in Discussion). No accurate definitions of a series of important variables (e.g. URTI, runny nose, etc.) are reported, and the Jackson scores are not explained, nor their use in Japanese personnel or language validated.

Intervention arm data not extracted due to the uncertainty of its meaning.

Funding source not mentioned. Conflicts of interest: none to report

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Open RCT, but sequence generation not reported
Allocation concealment (selection bias)	Unclear risk	"Mask and no mask groups were formed using block randomisation of participants within their respective job categories: nurses, doctors, and co-medical personnel." Concealment of allocation not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study. Blinding not possible, as 1 group wore face masks
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	1 dropout in each group accounted for.Quote: "Analyses were performed following the principles of intention-to-treat."
Selective reporting (reporting bias)	High risk	NB: influenza vaccine coverage was 100% in mask group and only 81% in the non-mask-wearing group.

Kotch 1994

Study characteristics	
Methods	Pair-matched, cluster-RCT conducted from 19 October 1988 to 23 May 1989 in 24 childcare centres in North Carolina, USA The trial tested the effects of a hand-washing and environment sterilising programme on diarrhoea (data not extracted) and ARIs. Child daycare centres had to care for 30 children or less, at least 5 of whom had to be in nappies, and intending to stay open for at least another 2 years. Randomisation is not described, nor are cluster coefficients reported.
Participants	389 children aged 3 years or less in daycare for at least 20 hours a week. There were some withdrawals, but attrition of participants is not stated, only that in the end data for 31 intervention classrooms and 36 control classrooms were available. 291 children aged up to 24 months and 80 over 24 months took part. The text is very confusing, as 371 seems to be the total of the number of families that took part.



Kotch 1994 (Continued)	No denominator breakdown by arm is reported, and numerators are only reported as new episodes per child-year.
Interventions	Structured hand-washing and environment (including surfaces, sinks, toilets, and toys) disinfecting programme with waterless disinfectant scrub. See Table 1 for details.
Outcomes	Laboratory: N/A Effectiveness: ARI (coughing, runny nose, wheezing, sore throat, or earache) Safety: N/A
Notes	Risk of bias: high (poor reporting of randomisation, outcomes, numerators and denominators) Note: the authors conclude that the fully adjusted RR for prevention of ARIs was 0.94 (–2.43 to 0.66). A poorly reported study.
	This study was supported in part by grant MCJ-373111 from the Maternal and Child Health Program (Title V. Social Security Act), Health Resources and Services Administration, Department of Health and Human Services. Cal Stat TM was contributed by Cal-gon Vestal Laboratories, a subsidiary of Merck and Co, Inc, St Louis, MO.
	Conflicts of interest: none to report.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Pair-matched cluster-randomised, controlled trial", but sequence generation not reported
Allocation concealment (selection bias)	Unclear risk	Centres were matched in pairs and then randomly allocated to either intervention or control programmes. Allocation concealment was not reported.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not possible (intervention was training session)
Blinding of outcome assessment (detection bias) All outcomes	High risk	"The same staff who conducted the training unobtrusively recorded observa- tions at 5-week intervals"
Incomplete outcome data (attrition bias) All outcomes	High risk	18 families were dropped, denominator not clear.
Selective reporting (reporting bias)	High risk	Denominators not clearly reported

Ladegaard 1999

Study characteristics	
Methods	RCT with cluster-randomisation to intervention or control. Of 10 institutions, 2 were excluded because they wanted institutions to be comparable in uptake area (i.e. housing and income). Interventions were administered to children, parents, and teachers at the institutions.
Participants	Children 0 to 6 years old



Ladegaard 1999 (Continued)

Interventions

Multifaceted: information, t-shirts to the children with: "Clean hands - yes, thank you", performance of a fairytale "The princess who did not want to wash her hands", exercise in hand-washing, importance of clean and fresh air. The aims of the intervention were to:

- 1. increase the hygiene education of the daycare teachers;
- 2. motivate the children by practical learning to have better hand hygiene; and
- 3. inform the parents about better hand hygiene.

See Table 1 for details.

Outcomes

Notes

34% decrease in "sickness" (probably mostly gastroenteritis)

Risk of bias: only limited data available

Note: the authors conclude that there was a 34% decrease in sickness in the intervention arm; this is probably overall sickness, as gastroenteritis is part of the outcomes (data not extracted). Only limited data available from translation by Jørgen Lous.

Funding was received from a local part of the Danish Health Authority (Forebyggelserådet för Fyns Amt).

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not described
Allocation concealment (selection bias)	Unclear risk	Randomisation by "lottery", the same as "flip the coin". Concealment not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not possible
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not possible
Incomplete outcome data (attrition bias) All outcomes	High risk	Total numbers of children included in each arm Not reported.
Selective reporting (reporting bias)	High risk	Limited data reported, in particular denominators missing.

Larson 2010

Study characteristics

Methods

Cluster block-randomised, controlled trial carried out between 20 November 2006 and 20 June 2008 in an upper Manhattan immigrant Latino neighbourhood ("19 month data collection period"). The study aimed at assessing the effects of education versus education and hand sanitiser use versus education and hand sanitiser use and common mask use against upper respiratory infections over a period of under 2 years. Follow-up was through an automated telephone system with a small financial incentive (USD 20) for those with 75% or more compliance. Those reporting an ILI received a visit within 48 hours for swabbing.



Larson 2010 (Continued)

An index case was someone who at the "onset day of illness nobody else in the household had been symptomatic within the previous five days".

A secondary case for each episode quote: "was any member of the household who developed symptoms within five days following the index case"; "The secondary attack rate was defined as the number of secondary cases recorded within 5 days of the onset of symptoms in the index case divided by the number of household members minus one".

The text implies that the unit of observation was the episode ("study subjects contributed more than one episode in which they were considered to be the index case").

Participants

617 households were randomised to the education group (n = 211), the hand-sanitiser group (n = 205), and the hand-sanitiser and mask group (n = 201). There were 2708 participants, mostly adult Latino immigrants to the USA.

Recruitment and allocation were carried out by household. There had to be at least 3 people living in the household, with at least 1 being a preschool or elementary school child, speaking English or Spanish, having a telephone, willingness to complete symptom assessments and have bimonthly home visits, and not using alcohol-based hand sanitiser routinely.

Intracluster correlation coefficients are reported on page 179 of the manuscript.

Interventions

Written Spanish or English language educational materials regarding the prevention and treatment of URTIs and influenza or the same educational materials and hand sanitiser (Purell, J&J), in large (8- and 4-ounce) and small (1-ounce) containers to be carried by individual household members to work or school, or the same interventions as well as regular surgical face masks (Procedure Face Masks for adults and children, Kimberly-Clark) with instructions for both the caretaker and the ill person to wear them when an ILI occurred in any household member. Replenishment of intervention stocks was done at the bimonthly home visit.

Caretakers had to wear a mask for 7 days when within 3 feet of a symptomatic case. They were also encouraged to wear masks within 3 feet of any household member. Reinforcing phone calls were made 3 times in 6 days.

The text clearly reports active influenza vaccine promotion during the bimonthly visits. ("The home visit to each household was made every 2 months to minimise study dropout, reinforce adherence to the assigned intervention, replenish product supplies and record use of supplies, answer questions, and correct ongoing misconceptions. At each visit, new educational materials regarding URTI prevention and treatment and influenza vaccination were distributed." (PDF page 3). Also just before the Discussion as follows: "Influenza vaccination rates: There was an increase between the baseline and exit interview in all three groups that reported 50% of more of members receiving influenza vaccine (preversus post-intervention for each group: 21.1% and 40.8% in the Education group, 19.0% and 57.1% in the hand sanitiser group, and 22.4% and 43.5% in the hand sanitiser and face mask group (P = 0.001). Additionally, those in the hand sanitiser group reported a significantly greater increase than the other 2 groups, controlling for baseline rates (P = 0.002)")

Coverage was unequal across groups, no information on the progressive impact of the vaccine, or indeed the nature of the vaccine(s) is reported. Apparently the first season was mild and the vaccine mismatched, compliance with the trial interventions was low in Arm 3, and a local epidemic of *Staphylococcus aureus* meant that the control group started washing hands.

The trial authors report no effect on reporting rates of vaccine coverage by arms, but with so many confounders who knows?

See Table 1 for details.

Outcomes

Laboratory: PCR carried out on samples from deep nasal swabs for influenza and the most common other pathogens (RSV, rhinovirus, enterovirus, parainfluenza viruses, etc.). The text describing the results of the swabbing is confusing, but in general appears to be non-random "Households reported 669 episodes of ILI (0 to 5 per individual)". Of the 234 deep nasal swabs obtained, 33.3% (n = 78) tested positive for influenza: 43.6% (n = 34) were influenza A and 56.4% (n = 44) were influenza B. Amongst the 66.7% who tested negative for influenza, 30.8% (48/156) tested positive for other viruses: 7 for respiratory syncytial virus, 9 for parainfluenza, 11 for enterovirus, 10 for rhinovirus, 6 for adenovirus, and 5 for metapneumovirus. Swabs were not obtained from the remaining 435 reported ILI episodes for the fol-



Larson 2010 (Continued)

lowing reasons: 72.0% (n = 313) did not meet the CDC definition of an ILI and were therefore included in the URTI symptom count; 21.4% of episodes (n = 93) were reported after 48 hours of ILI onset or the participant refused to be swabbed; and the research staff were unable to reach the participant in 6.7% of episodes (n = 29).

As no definition of URTI is given, it is unclear what kind of biases were introduced by the non-swabbing of the 313/435 "not meeting CDC definition".

 $Effectiveness: ILI (CDC \ definition): ``temperature \ of \ 37.8°C \ or \ more \ and \ cough \ and/or \ sore \ throat \ in \ the \ absence \ of \ a \ known \ cause \ other \ than \ influenza"$

URTI only referred to as "Viral upper respiratory infections (URTIs)".

Safety: N/A

Notes

The authors conclude that quote: "the Hand Sanitizer group was significantly more likely to report that no household member had symptoms (P,0.01), but there were no significant differences in rates of infection by intervention group in multivariate analyses. Knowledge improved significantly more in the Hand Sanitizer group (P,0.0001). The proportion of households that reported >50% of members receiving influenza vaccine increased during the study (P.0.001). Despite the fact that compliance with mask wearing was poor, mask wearing as well as increased crowding, lower education levels of caretakers, and index cases 0–5 years of age (compared with adults) were associated with significantly lower secondary transmission rates (all P,0.02). In this population, there was no detectable additional benefit of hand sanitiser or face masks over targeted education on overall rates of URTIs, but mask wearing was associated with reduced secondary transmission and should be encouraged during outbreak situations. During the study period, community concern about methicillin-resistant *Staphylococcus aureus* was occurring, perhaps contributing to the use of hand sanitiser in the Education control group, and diluting the intervention's measurable impact".

The study is at high risk of bias. Randomisation and reasons for dropout are not described. Differentials in cluster characteristics across arms point to randomisation not having worked, and the confounding effects of a post randomisation staphylococcal scare are difficult to judge. Symptom-driven follow-up gives no idea of the effects on asymptomatic ILI/influenza. Poor definitions (URTI?). There are unexplained dropouts, and the analysis plan is unclear. Finally, the very small number of cases of influenza and an unclear swabbing attrition may introduce further elements of confounding.

Funding: this study was funded by grant #1 U01 Cl000442-01, "Stopping URIs and Flu in the Family: The Stuffy Trial."

Conflicts of interest: none reported

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "Cluster block randomised, controlled trial", but sequence generation not reported
Allocation concealment (selection bias)	Unclear risk	Quote: "Households were block randomised into one of three groups" Allocation concealment not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Blinding of participants and personnel was not possible.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Blinding of outcome assessment is not stated.



Larson	2010	(Continued)
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Incomplete outcome data				
(attrition bias)				
All outcomes				

High risk

In control group households (n = 211), 26 dropped out and 37 did not consent.

In hand-sanitiser group households (n = 205), 21 dropped out and 36 did not

consent.

In hand-sanitiser and face mask group households (n = 201), 19 dropped out

and 35 did not consent.

Reasons for dropout were not described.

Selective reporting (reporting bias)

Unclear risk

617 of 772 eligible households were randomised.

Little 2015

Study ch	aracte	ristics
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Methods

Individuals sharing a household by mailed invitation through general practices in England were recruited. After consent, participants were randomised online by an automated computer-generated random-number program to receive either no access or access to a bespoke automated web-based intervention that maximised hand-washing intention, monitored hand-washing behaviour, provided tailored feedback, reinforced helpful attitudes and norms, and addressed negative beliefs. Participants were enrolled into an additional cohort (randomised to receive intervention or no intervention) to assess whether the baseline questionnaire on hand-washing would affect hand-washing behaviour. Participants were not masked to intervention allocation, but statistical analysis commands were constructed masked to group. The primary outcome was number of episodes of RTIs in index participants in a modified intention-to-treat population of randomly assigned participants who completed follow-up at 16 weeks.

Participants

344 physician offices were recruited over a wide area of England, and 20,066 participants were enrolled and randomised to intervention (N = 16,086) and control (N = 10,026).

Modified ITT was performed on 16,908 participants who completed the follow-up questionnaire at 16 weeks (intervention = 8241 and control = 8667).

Inclusion criteria: adult patients (aged 18 years or older) identified from computerised lists in general practitioner (GP) practices in England, for whom there was at least 1 other individual living in the household who was willing to report illness to the index person

Exclusion criteria: patients with severe mental problems (e.g. major uncontrolled depression or schizophrenia, dementia, or severe mental impairment) or who were terminally ill, and those reporting a skin complaint that would restrict hand-washing

Interventions

Automated web-based intervention that maximised hand-washing intention, monitored hand-washing behaviour, provided tailored feedback, reinforced helpful attitudes and norms, and addressed negative beliefs. Control no access to intervention web pages. See Table 1 for details.

Outcomes

The primary outcome was the number of index individuals that reported 1 or more RTIs (including ILI) at 16 weeks.

Secondary: duration of symptoms, transmission of respiratory infections, gastrointestinal infections, attendance at the practice, and use of health service resources

Infections self-reported by participants. RTI defined as 2 symptoms of an RTI for at least 1 day or 1 symptom for 2 consecutive days. Definition of ILI was a high temperature (feeling very hot or very cold; or measured temperature > 37.5 °C), a respiratory symptom (sore throat, cough, or runny nose), and a systemic symptom (headache, severe fatigue, severe muscle aches, or severe malaise).



Litt	le 2015	(Continued)
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No safety outcomes reported.

Notes

Government funded. The study was funded by the Medical Research Council (study number 09/800/22). Declaration of interests: the authors declare no competing interests.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Participants were automatically randomly assigned by the intervention software, but sequence generation not described.
Allocation concealment (selection bias)	Low risk	Participants were automatically randomly assigned by the intervention software.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	High risk	High attrition that was different in the 2 groups
Selective reporting (reporting bias)	Low risk	Specified outcomes reported.

Loeb 2009

Study characteristics

Methods

Open non-inferiority RCT carried out to compare the surgical mask with the N95 respirator in protecting healthcare workers against influenza. The trial was carried out between 2008 (enrolment started in September and follow-up on 12 January 2009) and 23 April 2009 (when all HCWs caring for febrile patients were told to wear an N95 respirator) because of the appearance of novel A/H1N1). The trial trigger was the beginning of the influenza season, defined as isolation of 2 or more viruses in a district in the same week. Following the 2003 SARS outbreak, all Ontario nurses caring for febrile patients (38 °C or more and new onset cough or SOB) had to wear surgical masks. The randomisation (carried out in blocks of 4 by centre) then consisted of either confirmation to same-maker surgical mask wear or N95 respirator wear. Investigators and laboratory staff were blind to allocation status, but for obvious reasons (the visible difference in interventions), participants were unblinded. "The criterion for non-inferiority was met if the lower limit of the 95% confidence interval (CI) for the reduction in incidence (N95 respirator minus surgical group) was greater than -9%". So this is the non-inferiority margin. It is assumed that the "minus surgical group" means minus surgical mask group.

Participants

Consenting nurses (n = 446 randomised) aged a mean of 36.2 years working full time (\geq 37 hours/week) in 23 acute units (a mix of paediatric, A&E, and acute medical units) in 8 hospitals in Ontario, Canada. 225 were randomised to the surgical mask and 221 to the N95 respirator. There were 13 and 11 dropouts, respectively from each arm (all accounted for), plus 21 and 19 lost to follow-up; 11 in each arm gave no reason, the others are accounted for. There were no deaths. The final total of 212 and 210 was included in the analysis. Table 1 reports the demographic data of participants by arm, which appear comparable.



Loeb 2009 (Continued)

Interventions

Surgical masks (as standard wear by the standard distributor) or fit-tested N95 respirator. All nurses wore gloves or gowns in the presence of a febrile patient. See Table 1 for details.

Outcomes

Laboratory RT-PCR paired sera with 4-fold antibody rise from baseline (only for unvaccinated) nurses

Effectiveness: follow-up (lasting a mean of around 97 days for both arms) was carried out twice-weekly on a web-based instrument. Nurses with new symptoms were asked to swab a nostril if any of the following signs or symptoms had developed: fever (temperature ≥ 38 °C), cough, nasal congestion, sore throat, headache, sinus problems, muscle aches, fatigue, earache, ear infection, or chills.

The text defines influenza with laboratory confirmation, and separately reports criteria for swab triggering and a definition of ILI ("Influenza-like illness was defined as the presence of cough and fever: a temperature $\geq 38^{\circ}$ C"). But this is not formally linked to influenza in the text, as it appears that primary focus was the detection of laboratory-confirmed influenza (either by RT-PCR or serology).

Additional outcome data sought were work-related absenteeism and physician visits for respiratory illness.

Secondary outcomes included detection of the following non-influenza viruses by PCR: parainfluenza virus types 1, 2, 3, and 4; respiratory syncytial virus types A and B; adenovirus; metapneumovirus; rhinovirus-enterovirus; and coronaviruses OC43, 229E, SARS, NL63, and HKU1.

Audits to assess nurse compliance with the interventions were carried out in the room of each patient cared for. The text reports that 50 and 48 nurses in the surgical mask and N95 groups, respectively, had laboratory confirmation of influenza infection, indicating non-inferiority. Interestingly, non-inferiority seemed to be applicable both to seasonal viruses and nH1N1 viruses (as 8% and 11.9% were serologically positive to nH1N1). This finding is explained either by seeding or cross reaction with seasonal H1N1. Equivalent conclusions could be drawn for nurses with complete follow-up. Non-inferiority was applicable also to other ILI agents identified. None of the 52 individuals with positive isolates met the criteria for ILI.

All cases of ILI were confirmed as having influenza (9 and 2 respectively). This means that all the 11 cases of ILI had influenza, but that most of those with a laboratory diagnosis of influenza did not have cough and fever. For example, the text reports that "Of the 44 nurses in each group who had influenza diagnosed by serology, 29 (65.9%) in the surgical mask group and 31 (70.5%) in the N95 respirator group had no symptoms". By implication, of the 88 nurses with antibody rises, 28 had symptoms of some kind, i.e. two-thirds were asymptomatic. Absenteeism was 1 versus 39 episodes in the mask versus respirator arms. No episodes of LRTI were recorded. The number of family contacts with ILI were the same for each arm (45 versus 47). Physician visits were similar in both groups.

Safety: no AEs are reported

Notes

The authors conclude that "Among nurses in Ontario tertiary care hospitals, use of a surgical mask compared with a N95 respirator resulted in non-inferior rates of laboratory-confirmed influenza".

This a well-designed and conducted trial with credible conclusions. The only comment is that the focus in the analysis on influenza (symptomatic and asymptomatic) is not well-described, although the rationale is clear (interruption of transmission).

Funding/Support: this study was supported by the Public Health Agency of Canada.

Financial disclosures: none reported.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Randomisation was performed centrally", but method of sequence generation not described.



Loeb 2009 (Continued)		
Allocation concealment (selection bias)	Low risk	"by an independent clinical trials coordinating group such that investigators were blind to the randomisation procedure and group assignment and was stratified by centre in permuted blocks of 4 participants."
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	"It was not possible to conceal the identity of the N95 respirator or the surgical mask since manipulating these devices would interfere with their function"
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessment blinded: "Laboratory personnel conducting hemagglutinin inhibition assays, polymerase chain reaction (PCR), and viral culture for influenza were blinded to allocation."
Incomplete outcome data (attrition bias) All outcomes	Low risk	21 of 225 randomised to mask group and 19 of 221 randomised to N95 group were lost to follow-up, reasons reported.
Altoutcomes		Study stopped early: Quote: We had planned to stop the study at the end of influenza season. However, because of the 2009 influenza A(H1N1) pandemic, the study was stopped on April 23, 2009, when the Ontario Ministry of Health and Long-Term Care recommended N95 respirators for all healthcare workers taking care of patients with febrile respiratory illness."
Selective reporting (reporting bias)	Low risk	All outcomes reported.

Longini 1988

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Study characteristics	S
Methods	Cluster-controlled, double-blind, randomised trial to assess the efficacy of virucidal tissues in interrupting family transmission of rhinovirus and influenza virus. The study was carried out in the community of Tecumseh, Michigan, USA during the period of 25 November 1984 to 28 April 1985. However, the authors only report results for the period of 13 January to 23 March 1985, when a high circulation of influenza A H3N2 and rhinovirus was detected.
Participants	296 households were enrolled, but 5 households were eliminated from the analysis for "technical reasons". The analysis was carried out in households with 3 to 5 members. The authors report data on 143 households randomised to virucidal tissues and 148 to placebo tissue. The average age in households was around 22, and the difference between arms was not significant. Randomisation was carried out by the sponsor, and tissues were pre-packed in coded boxes with no other identifying features and delivered to households at the beginning of the study period.
Interventions	Disposable 3-layered virucidal tissues (citric and malic acids with sodium lauryl sulphate in the middle layer) or placebo (succinic acid in the middle layer) tissues. They were used to blow the nose and for coughing or sneezing into. Households were also stratified by level of tissue use. Tissue use was significantly higher in the intervention arm (82% versus 71%). See Table 1 for details.
Outcomes	Laboratory: yes - viral culture from nasal and throat swabs from symptomatic participants Effectiveness: ARI (with a proportion of laboratory-confirmed diagnosis in non-randomly chosen participants with symptoms lasting 2 days or more) Follow-up and surveillance was carried out using a telephone questionnaire. Safety: N/A
Notes	Risk of bias: high (inappropriate choice of placebo) Note: the authors conclude that virucidal tissues were up to 36.9% effective in preventing transmission of ARIs as measured by secondary attack rates (18.7% versus 11.8%). This finding was not statistical-



Longini 1988 (Continued)

ly significant, but may well have been affected by the lack of do-nothing community controls. This a well-designed, well-written study despite the unexplained attrition of 5 families, the lack of reporting of cluster coefficients, and the differential in tissue use between the 2 arms, which raises questions about the robustness of double-blinding. Particularly notable is the discussion on the low generalisability of results from the study from the placebo arm given that even the inert barrier of the tissues is likely to have limited spread. Also, the lengths to which the authors went to obtain allocation concealment and maintenance of double-blind conditions.

Funding: The Kimberly-Clark Corporation sponsored this research. This research was also partially supported by National Institutes of Health Grant 1-RO1-Al22877-01 and General Clinical Research Center Public Health Research Grant 5-MO1-RR000039.

Declaration of interests: none declared.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "Treated and placebo tissues were randomly assigned"
tion (selection bias)		Sequence generation not reported
Allocation concealment (selection bias)	Low risk	Quote: "Treated and placebo tissues were randomly assigned by the sponsor to 296 participating households stratified by household size, such that roughly half the households would receive treated tissues. Thus, the investigators were unaware of the assignment of treated tissues."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	"Treated and placebo tissues were randomly assigned by the sponsor to the randomly assigned 296 households stratified by household size The type of tissue was identified by code, and the boxes in which tissues were contained were not marked with any specific identifiers. Therefore, the study was double-blinded."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The investigators were unaware of the assignment of the treated tissues"
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	296 households eligible. "The final sample used for analysis consisted of 143 households in the treatment group and 148 households in the placebo group."
Selective reporting (reporting bias)	High risk	Quote: "The analysis of secondary spread was restricted to households of three to five members for technical reasons, which eliminated five households."
		"The two groups were almost identical in composition."

Luby 2005

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Methods

Partly double-blind, cluster-RCT carried out during 15 April 2002 to 5 April 2003 in Karachi, Pakistan. The trial assessed the effects of mother and child hand-washing on the incidence of respiratory infections, impetigo (data not extracted), and diarrhoea (data not extracted).

Randomisation took place by computer-generated random numbers in 3 phases.

- 1. 25 neighbourhoods were assigned to hand-washing and 11 to standard practice.
- 2. 300 households were assigned to using antiseptic soap.
- 3. 300 households were assigned to using plain soap.



Luby 2005 (Continued)

- 4. 306 households were assigned to standard practice.
- 5. 1523 children younger than 15 years were assigned to using antiseptic soap.
- 6. 1640 children younger than 15 years were assigned to using plain soap.
- 7. 1528 children younger than 15 years were assigned to standard practice.

Soaps were of identical weight, colour, and smell and were packed centrally with a coded packing case matched to households containing 96 bars. Neither field workers nor participants were aware of the content. Control arm households were visited with the same frequency as intervention household but were given books and pens. Codes were held centrally by the manufacturer and broken after the end of the trial to allow analysis.

Participants

Householders of slums in Karachi.

Of the 1523 children younger then 15 years assigned to using antiseptic soap, 117 dropped out (1 died, 51 were born in, and 65 aged out) = 1406; 504 were aged less than 5.

Of 1640 children younger then 15 years assigned to using plain soap, 117 dropped out (3 died, 44 were born in, and 70 aged out) = 1523; 517 were aged less than 5.

Of 1528 children younger then 15 years assigned to standard practice, 125 dropped out (3 died, 40 were born in, and 82 aged out) = 1403; 489 were aged less than 5.

Interventions

Instruction programme and antibacterial soap containing 1.2% triclocarban, or ordinary soap to be used throughout the day by householders, or standard procedure. See Table 1 for details.

Outcomes

Laboratory: N/A

Effectiveness:

- 1. Number of new respiratory illness per person per week
- 2. Pneumonia (cough or difficulty in breathing with a respiratory rate of > 60 min in children less than 60 days old, > 50 min in those less than 1 year old, and > 40 min for those aged 1 to 5 years)

Follow-up was weekly with household interview and direct observation. Children aged less than 5 were weighed, and the report presents stratification of results by child weight. Safety: N/A

Notes

Risk of bias: low (cluster coefficients and analysis by unit of randomisation provided)

Note: the authors conclude that "handwashing" neighbourhoods has significantly fewer episodes of

Note: the authors conclude that "handwashing" neighbourhoods has significantly fewer episodes of respiratory disease than controls (e.g. 50% less cough). "Handwashing" children aged less than 5 had 50% fewer episodes of pneumonia than controls (-65% to -35%). However, there was no difference in respiratory illness between types of soap. The report is confusing, with a shifting focus between children age groups. The impression reading is of an often rewritten manuscript. There is some loss of data (e.g. in the results by weight, i.e. risk group) because of lack of clarity on denominators. Despite this, the trial is a landmark.

Funding: most of the funding for this study was provided by Procter and Gamble, manufacturer of Safeguard Bar Soap. The balance of the funding was provided by the Centers for Disease Control and Prevention

Conflict of interest statement: S Luby was supported by the grant from the Procter & Gamble company that funded this study. W Billhimer is an employee of the Procter & Gamble company. The other authors declare that they have no conflict of interest.

Bias	Authors' judgement	Support for judgement			
Random sequence generation (selection bias)	Low risk	Randomisation took place by computer-generated random numbers in 3 phases.			
Allocation concealment (selection bias)	Low risk	Quote:"One of the investigators (SL) who did not participate in recruiting neighbourhoods or households programmed a spreadsheet to randomly gen-			



Luby 2005 (Continued)		
		erate the integers of a 1 or a 2. He applied the random numbers sequentially to the list of neighbourhoods. Neighbourhoods with a 1 were assigned to control, and those with a 2 were assigned to handwashing promotion. Random assignment continued until neighbourhoods consisted of at least 600 handwashing promotion households and 300 control households were assigned."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote:"The antibacterial soap contained 1-2% triclocarban as an antibacterial substance. The plain soap was identical to the antibacterial soap except that it did not contain triclocarban Neither the fieldworkers nor the families knew whether soaps were antibacterial or plain."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote:"Neither the fieldworkers nor the families knew whether soaps were antibacterial or plain."
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	89% of the study population followed up, but no data on the clusters.
Selective reporting (reporting bias)	Low risk	Quote:"At baseline, households in the three intervention groups were similar."

MacIntyre 2009

Study characteristics	5			
Methods	Prospective cluster-RCT carried out in Sydney, Australia, to assess the use of surgical masks, P2 masks, and no masks in preventing ILI in households. The study was carried out during the 2 winter seasons of 2006 and 2007 (August to the end of October 2006 and June to the end of October 2007). "Gaussian random effects were incorporated in the model to account for the natural clustering of persons in households"			
Participants	290 adults from 145 families. 47 households (94 enrolled adults and 180 children) were randomised to the surgical mask group, 46 (92 enrolled adults and 172 children) to the P2 mask group, and 52 (104 enrolled adults and 192 children) to the no-mask (control) group.			
Interventions	Use of surgical masks and P2 mask versus no mask. The P2 mask is described as very cumbersome. See Table 1 for details.			
Outcomes	Laboratory: serological evidence			
	Effectiveness: ILI (described as fever, history of fever or feeling feverish in the past week, myalgia, arthralgia, sore throat, cough, sneezing, runny nose, nasal congestion, headache) However, a positive laboratory finding for influenza converts the ILI definition into one of influenza. Safety: N/A			
Notes	The study authors conclude that adherence to mask use significantly reduced the risk for ILI-associated infection, but < 50% of participants wore masks most of the time. They concluded that household use of face masks is associated with low adherence and is ineffective for controlling seasonal respiratory disease. Compliance was by self-report, therefore likely to be an underestimate. The primary outcome was ILI or lab-positive illness. This showed no effect. Sensitivity analysis by adherence showed that under the assumption that the incubation period is equal to 1 day (the most probable value for the 2 most common viruses isolated, influenza (21) and rhinovirus (26)), adherent use of P2 or surgical masks significantly reduces the risk for ILI infection, with a hazard ratio = 0.26 (95% CI 0.09 to 0.77; P = 0.015). No other covariate was significant. Under the less likely assumption that the incubation period is equal to 2 days, the quantified effect of complying with P2 or surgical mask use remains strong, although borderline significant; hazard ratio was 0.32 (95% CI			



MacIntyre 2009 (Continued)

0.11 to 0.98; P = 0.046). The study was underpowered to determine if there was a difference in efficacy between P2 and surgical masks (Table 5). The study conclusion appears to be a post hoc data exploration. Regardless of this, the study message is that respirator use in a family setting is unlikely to be effective as compliance is difficult unless there is a situation of real impending risk.

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Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	"Participating households were randomised to 1 of 3 arms by a secure computerised randomisation process", but sequence generation not described.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	"Study participants and trial staff were not blinded, as it is not technically possible to blind the mask type to which participants were randomised."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"However, laboratory staff were blinded to the arm of randomisation."
Incomplete outcome data (attrition bias) All outcomes	Low risk	143 of 145 randomised families were analysed; 2 families in the control group were lost to follow-up during the study, for which no reasons were given.
Selective reporting (reporting bias)	Low risk	No differences between groups at baseline

MacIntyre 2011

Macinityle 2011	
Study characteristic	s
Methods	A cluster-RCT of 1441 HCWs in 15 Beijing hospitals was performed during the 2008 to 2009 winter. Participants wore masks or respirators during the entire work shift for 4 weeks. Outcomes included CRI, ILI, laboratory-confirmed respiratory virus infection, and influenza. A convenience no-mask/respirator group of 481 health workers from 9 hospitals was compared.
Participants	Participants (N = 1441) were hospital HCWs aged > 18 years from the emergency departments and respiratory wards of 15 hospitals. These wards were selected as high-risk settings in which repeated and multiple exposures to respiratory infections are expected.



MacIntyre 2011 (Continued)	Participants were randomised to medical mask (N = 492 staff from 5 hospitals), N95 fit-tested masks (N = 461 staff from 5 hospitals), and N95 non-fit-tested mask (N = 488 staff from 5 hospitals).			
Interventions	Fit-tested N95 respirators versus non-fit-tested N95 respirators versus medical masks. See Table 1 for details.			
Outcomes	Clinical respiratory illness, defined as 2 or more respiratory symptoms or 1 respiratory symptom and a systemic symptom			
	Influenza-like illness, defined as fever ≥ 38 °C plus 1 respiratory symptom (i.e. cough, runny nose, etc.)			
	Laboratory-confirmed viral respiratory infection (detection of adenoviruses, human metapneumovirus, coronavirus 229E/NL63, parainfluenza viruses 1, 2, and 3, influenza viruses A and B, respiratory syncytial virus A and B, rhinovirus A or B, and coronavirus OC43/HKU1 by multiplex PCR)			
	Laboratory-confirmed influenza A or B			
	Adherence with mask or respirator use. Reported problems associated with using the masks or respirators			
Notes	Control arm not randomised so has been ignored. Funding source unknown.			
	Conflict of interests: Raina MacIntyre receives funding from influenza vaccine manufacturers GSK and CSL Biotherapies for investigator-driven research. She has also been on advisory boards for Wyeth, GSK and Merck. Dr Simon Cauchemez received consulting fees from MacIntyre et al. 178 a 2011 Blackwell Publishing Ltd, Influenza and Other Respiratory Viruses, 5, 170–179 Sanofi-Pasteur MSD on the modelling of varicella zoster virus. The remaining authors declare that they have no competing interests. The corresponding author had full access to all the data in the study and had final responsibility for the decision to submit for publication. Prior to the start of this study, NMF acted as a consultant for Roche,			

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Randomisation process (using a secure computerised randomisation program), but sequence generation not described
Allocation concealment (selection bias)	Low risk	Hospitals randomised prior to inclusion of participants.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (reporting bias)	Low risk	Specified outcomes reported.

Novartis and GSK Biologicals (ceasing in 2007).



MacIntyre 2013

Study characteristics				
Methods	A cluster-RCT			
Participants	A total of 1669 nurses and doctors from 68 emergency departments and respiratory wards of 19 Beijing hospitals were included. Inclusion criteria: any nurse or doctor aged 18 years or older who worked full time in the emergency or respiratory wards was eligible. Exclusion: HCWs if they (1) were unable or refused to consent; (2) had beards, long moustaches, or long facial hair stubble; (3) had a current respiratory illness, rhinitis, and/or allergy; or (4) worked part time or did not work in the aforementioned wards or departments			
		ormed on 572 staff and 24 wards in medical mask group, 516 staff and 20 wards sk group, and 581 staff and 24 wards in the N95 mask group.		
Interventions	1817; 3M, St. Paul, MN) Participants wore the r ipants were supplied d Participants using N95	Quote: "Masks used in the study were the 3M Standard Tie-On Surgical Mask (catalog number mask 1817; 3M, St. Paul, MN) and the 3M Health Care N95 Particulate Respirator (catalog number 1860; 3M) Participants wore the mask or respirator on every shift after being shown how to fit and wear it. Participants were supplied daily with either three masks for the medical mask arm or two N95 respirators. Participants using N95 respirators underwent a fit testing procedure using a 3M FT-30 Bitrex Fit Test Kit according to the manufacturer's instructions (3M)." See Table 1 for details.		
Outcomes	Laboratory:			
	adenoviruses; huma viruses 1, 2, and 3; ir by nucleic acid testi Seoul, Korea).	ed viral respiratory infection in symptomatic participants, defined as detection of an metapneumovirus; coronaviruses 229E/NL63 and OC43/HKU1; parainfluenza nfluenza viruses A and B; respiratory syncytial viruses A and B; or rhinoviruses A/B ing (NAT) using a commercial multiplex polymerase chain reaction (Seegen, Inc.,		
	3. Laboratory-confirm Streptococcus pneur	ed influenza A or B in symptomatic participants. ed bacterial colonisation in symptomatic participants, defined as detection of moniae, Legionella, Bordetella pertussis, chlamydia, Mycoplasma pneumoniae, or izae type B by multiplex polymerase chain reaction (Seegen, Inc.).		
		ned as 2 or more respiratory symptoms or 1 respiratory symptom and a systemic as fever (38 °C) plus 1 respiratory symptom		
	Safety: adverse effects measured using a semi-structured questionnaire. Investigators stated that there was higher reported adverse effects and discomfort of N95 respirators compared with the other 2 arms. In terms of comfort, 52% (297 of 571) of the medical mask arm reported no problems, compared with 62% (317 of 512) of the targeted arm and 38% (217 of 574) of the N95 arm (P < 0.001).			
Notes	Compliance with the product was highest in the targeted N95 arm (82%; 422 of 516), then the medical mask arm (66%; 380 of 572), and the N95 arm (57%; 333 of 581); these differences were statistically significant (P < 0.001).			
	The period study conducted: 28 December 2009 to 7 February 2010			
	Funding: unclear Declaration of interests: none declared.			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence generation (selection bias)	Unclear risk	"using a secure computerized randomization program", but sequence generation not described		



MacIntyre 2013 (Continued) Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Outcome was objectively assessed with lab confirmation in addition to clinical illness.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Laboratory outcomes are reported for all subjects (with at least one respiratory symptom or fever) tested, and then for the subset meeting the CRI definition"
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up. Flow chart and text match, investigators conducted ITT and PP analysis. All the outcomes were accounted for amongst all participants.
Selective reporting (reporting bias)	Low risk	All outcomes were reported as planned.

Study characteristics	•		
Methods	A cluster-RCT of cloth masks compared with medical masks in healthcare workers in 14 secondary-/ter-tiary-level hospitals in Hanoi, Vietnam. Hospital wards were randomised to: medical masks, cloth masks, or a control group (usual practice, which included mask wearing). Participants used the mask on every shift for 4 consecutive weeks.		
Participants	1607 hospital HCWs aged ≥ 18 years working full time in selected high-risk wards.		
	Medical mask group (n = 580 HCWs), cloth mask group (n = 569 HCWs), control group (n = 458 HCWs)		
Interventions	Medical masks, cloth masks, or a control group. See Table 1 for details.		
Outcomes	Clinical respiratory illness, influenza-like illness, and laboratory-confirmed respiratory virus infection		
	 Clinical respiratory illness, defined as 2 or more respiratory symptoms or 1 respiratory symptom a systemic symptom 		
	 Influenza-like illness, defined as fever ≥ 38 °C plus 1 respiratory symptom Laboratory-confirmed viral respiratory infection. Laboratory confirmation was by nucleic acid detection. 		
	tion using multiplex reverse transcriptase PCR (RT-PCR) for 17 respiratory viruses.		
	Adverse events associated with mask use		
Notes	Government funded. Competing interests: CRM has held an Australian Research Council Linkage Grant with 3M as the indu try partner, for investigator-driven research. 3M has also contributed masks and respirators for investigator-driven clinical trials. CRM has received research grants and laboratory testing as in-kind support from Pfizer, GSK and Bio-CSL for investigator-driven research. HS had a NHMRC Australian-based Public Health Training Fellowship at the time of the study (1012631). She has also received funding from vaccine manufacturers GSK, bio-CSL and Sanofi Pasteur for investigator-driven research and present tions. AAC used filtration testing of masks for his PhD thesis conducted by 3M Australia.		
Risk of bias			
Bias	Authors' judgement Support for judgement		



MacIntyre 2015 (Continued)		
Random sequence generation (selection bias)	Low risk	Epi info V.6 was used to generate a randomisation allocation.
Allocation concealment (selection bias)	Low risk	74 wards randomised prior to recruitment of individuals.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Unblinded study
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (reporting bias)	Low risk	Specified endpoints reported.

MacIntyre 2016

Study characteristics	5			
Methods	Cluster-RCT to examine medical mask use as source control for people with respiratory illness in 6 major hospitals in 2 districts of Beijing, China. Index cases with ILI were randomly allocated to medical mask (n = 123) and control arms (n = 122). Since 43 index cases in the control arm also used a mask during the study period, an as-treated post hoc analysis was performed by comparing outcomes amongst household members of index cases who used a mask (mask group) with household members of index cases who did not use a mask (no mask group).			
Participants	245 index cases with ILI (medical mask = 123, control group = 122) and 597 household contacts (medical mask = 302, control group = 295)			
Interventions	Medical mask versus no mask (control). See Table 1 for details.			
Outcomes	Clinical respiratory illness, ILI, and laboratory-confirmed viral respiratory infection			
	 Clinical respiratory illness, defined as 2 or more respiratory symptoms (cough, nasal congestion, run- ny nose, sore throat, or sneezes) or 1 respiratory symptom and a systemic symptom (chill, lethargy, loss of appetite, abdominal pain, muscle or joint aches). 			
	2. ILI, defined as fever ≥ 38 °C plus 1 respiratory symptom.			
	3. Laboratory-confirmed viral respiratory infection, defined as detection of adenoviruses, human metapneumovirus, coronaviruses 229E/NL63 and OC43/HKU1, parainfluenza viruses 1, 2, and 3, in- fluenza viruses A and B, respiratory syncytial virus A and B, or rhinovirus A/B by nucleic acid testing using a commercial multiplex PCR.			
	No safety outcomes reported.			
Notes	Government funded. Competing interests: all authors have completed the Unified Competing Interests form (available on request from the corresponding author) and declare that: CRM has held an Australian Research Council Linkage Grant with 3M as the industry partner, for investigator driven research. 3M have also contributed supplies of masks and respirators for investigator-driven clinical trials. She has received re-			



MacIntyre 2016 (Continued)

search grants and laboratory testing as in-kind support from Pfizer, GSK and Bio-CSL for investigator-driven research. HS had an NHMRC Australian based Public Health Training Fellowship at the time of the study (1012631). She has also received funding from vaccine manufacturers GSK, bio-CSL and Saniofi Pasteur for investigator-driven research and presentations. AAC had testing of filtration of masks by 3M for PhD.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Random allocation sequence using Microsoft Excel
Allocation concealment (selection bias)	High risk	Doctors enrolled the participants randomly to intervention and control arms.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study
Blinding of outcome assessment (detection bias) All outcomes	High risk	Clinical endpoints assessed unblinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (reporting bias)	Low risk	Specified outcomes reported.

McConeghy 2017

Study characteristics	S .
Methods	Pilot study of comprehensive intervention (education, cleaning of surfaces, audit and feedback) to staf of nursing homes versus usual care. Pair-matched cluster-randomised design with only 5 clusters (nurs ing homes) in each group
Participants	10 nursing homes in Colorado, USA
	Intervention group = 481 long-stay residents and control group = 380
	'Long-stay' defined as resident at least 90 days prior to baseline, or recently readmitted after previous long stay.
Interventions	A multifaceted hand-washing/surface-cleaning intervention comprised of 1) 1-hour online educational module focused on how to prevent infections; 2) provided with an "essential bundle" of 7 products, ranging from hand sanitiser gel and foam to antiviral facial tissues, disinfecting spray, and hand and face wipe and recommendation to use 4 skin cream and wipe products; 3) audit and feedback system. See Table 1 for details.
Outcomes	Laboratory: surface cultures mentioned in Methods, but no results given
	Effectiveness: LRTI, all infections, hospitalisation, use of antibiotics (not relevant to this review)



McCo	onegh	y 2017	(Continued)
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Safety: none mentioned in Methods and no results given

Notes

The authors conclude that Quote: "This multifaceted hand-washing and surface cleaning intervention was designed to reduce infection rates among nursing homes residents. In our 10-facility randomized, matched pair pilot study, we observed program compliance and satisfaction along with reductions in surface bacterial counts, but did not observe a statistically significant reduction in infection rates, antimicrobial use, or hospitalizations".

Very poorly reported study with results not explained, summarised in Table 3 as RDs. Denominators and attrition are unclear.

This work was supported by Kimberly-Clark Corporation (Contract # 14792008). Declaration of interests: none declared.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Method not described
Allocation concealment (selection bias)	Unclear risk	Method not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Illness and absenteeism reported by treating staff.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	No attrition given. Data were collected from e-medical record at baseline, but not clear whether illness data during the study were collected by the same method.
Selective reporting (reporting bias)	High risk	Upper respiratory tract infection was mentioned in the Methods (intervention presumably would target these), but only LRTI and overall infection reported.

Millar 2016

Study characteristics	
Methods	Cluster-RCT, open-label study, factorial design
Participants	Around 30,000 healthy, male army trainees aged 18 to 42 years at Fort Benning, Georgia were included. Inclusion criteria: trainees assigned to 1 of the 6 selected training battalions, trainees who present with an SSTI at the clinic or the hospital, provide informed consent. Exclusion criteria: fails to meet inclusion criteria. No denominator breakdown by arm is reported.
Interventions	Promotion of hand-washing in addition to a once-weekly application of chlorhexidine-based body wash. See Table 1 for details.
Outcomes	This study was nested in a large field-based RCT and utilised clinic-based medical records.
	Laboratory: none



Millar 2016 (Continued)	Effectiveness: incidence of ARI at 20 months. The case definition was any occurrence of the following ICD-9 symptom or disease-specific codes: 460 to 466, 480 to 488, and specifically 465.9, 482.9, 486, and 487.1.
	Safety: adverse effects neither planned nor reported by the investigators
Notes	The period study conducted: May 2010 to January 2012
	Government funded. Declaration of interests: none declared.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	quote: "computer-generated random numbers to 1 of the 3 study groups"
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The study was open-label and self-reporting of ARI. It is planned as secondary objective of an original trial. Data abstractors were blinded to group assignment.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Data abstractors were blinded to group assignment.
Incomplete outcome data (attrition bias) All outcomes	High risk	There is a statistically significant difference between attrition rates in the 3 groups. The reasons for attrition are briefly reported in Table 1 of the original study (Ellis and colleagues 2014), but are unlikely to be related to the outcomes of this study. ARI cases were captured utilising clinic-based medical records, but this outcome is not prespecified in the protocol.
Selective reporting (reporting bias)	High risk	The study was conducted for another purpose. According to the study protocol, the outcomes of interest in the current report were not mentioned as outcomes when the study was planned. ARI is not prespecified as an outcome in the protocol published on ClinicalTrials.gov.

Miyaki 2011

Study characteristics	
Methods	A quasi-cluster-RCT
Participants	A total of 15,134 assigned to intervention (N = 6634 workers) and control (N = 8500 workers)
	Inclusion criteria: all general employees (aged 19 to 72 years in 2009) of 2 sibling companies of a major car industry in Kanagawa Prefecture, Japan. All workers who regularly reported to the workplace were included, regardless of treatment for chronic diseases.
	All employees have the same health insurance plan and were followed up in the same way.
Interventions	Quote: "The intervention involved asking workers whose family members developed an influenza-like illness (ILI) to stay at home. If any co-habiting family members showed signs of influenza-like illness



Miyaki 2011 (Continued)

(ILI), employees ... were asked to stay at home voluntarily until 5 days has passed since the resolution of the ILS symptoms or 2 days after alleviation of fever." See Table 1 for details.

Outcomes

Workroom: influenza A test kit (rapid test)

Effectiveness: assess the effectiveness of household quarantine in reducing the incidence of influenza A H1N1. ILI was defined as a body temperature greater than 38 $^{\circ}$ C or more than 1 $^{\circ}$ C above the normal temperature accompanied with more than 2 of these symptoms: nasal mucus, pharyngeal pain, cough, chills or heat sensation

Safety: the incidence of influenza A H1N1 amongst workers who were told to stay home if a family member developed ILI was higher (relative risk of 2.17; P < 0.001) compared to control group. No other safety measures/harms reported.

Compliance: quote: "our intervention was not compulsory; we only asked the employees to leave the workplace for a while on full pay, and we succeeded in getting all workers' agreement. In our case, explaining that the home waiting policy might be beneficial to the whole workers and help to avoid stopping the manufacturing lines (explaining it is for the benefit of the public) and guaranteeing payment during the leave (financial support) helped them to obey our request."

Notes

Period study conducted: 1 July 2009 to 19 February 2010

Unfunded

There are no conflicts of interest to declare.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No information given.
Allocation concealment (selection bias)	Unclear risk	No information given.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The nature of the intervention (stay at home) was confirmed in the intervention group, where all workers agree as they were financially supported during absences due to ILI.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	"Company doctors diagnosed the disease through a positive result of an influenza A test or clinical symptoms", but not clear if they were blinded to assignment; however, the diagnostic process is meticulous and objectively confirmed.
Incomplete outcome data (attrition bias) All outcomes	Low risk	All cases are included in the analysis, and none were lost to follow-up.
Selective reporting (reporting bias)	Unclear risk	Although all outcomes of interest are clearly specified, described, and followed up, and text and numbers checked out well and based on the outcome stated for the study, there is no published protocol to match the planned vs the reported outcomes.

Morton 2004

Study characteristics



Morton 2004 (Continued)

Methods

Cross-over study to evaluate the effectiveness of an alcohol gel as an adjunct to regular hand-washing for decreasing absenteeism amongst elementary children by reducing specific communicable diseases such cold, flu, and conjunctivitis. The study was conducted in an elementary school in New England, USA. In the cross-over design, classrooms in each grade level were randomised to begin as the experimental group (alcohol gel) or the control group (regular hand-washing). A study protocol for hand hygiene was introduced following the germ unit education. The hand-washing product was a soapand-water alternative that is approximately 60% ethyl alcohol. In phase 1 (46 days) children in 9 classrooms were in the experimental group, and children in 8 classrooms were in the control group. After a 1-week washout period when no children had access to the alcohol gel, phase 2 (47 days) started, and the classroom that had participated before as experimental group passed into the control group and vice versa. Data were collected by the parents, who informed the secretary or the school nurse of the reasons for a child's absence, including symptoms of any illness. Respiratory illnesses were defined by symptoms of URTI.

Participants

253 children, 120 girls and 133 boys, from kindergarten to 3rd grade. Of the eligible 285 students, 32 children dropped out (10 due to skin irritation and 22 because of lack of parental consent). No denominator breakdown by arm is reported because the study used a cross-over design.

Interventions

Use of an alcohol gel as an adjunct to regular hand-washing and educational programme versus regular hand-washing and educational programme. See Table 1 for details.

Outcomes

Laboratory: no Effectiveness: days of absences from school for respiratory illness Safety: N/A

Notes

Risk of bias: high (no description of randomisation; partial reporting of outcomes, numerators and denominators)

Note: the authors conclude that significantly fewer children became ill whilst using the alcohol gel as an adjunct to regular hand-washing than when using regular hand-washing only (decreased school absenteeism of 43% with the use of alcohol gel on top of hand-washing). The authors also described, as a limitation of the study, the fact that the school nurse served as the data collector, which could be perceived as bias in measurement of the outcome variable.

Randomisation and allocation are not described; no cluster coefficients were reported; and attrition was not taken into consideration during the analysis. Unit of randomisation and analysis are different. No reporting by arm. No ORs, no CIs reported.

Funding: Maine Administrative School District #35 in Eliot, Maine, and South Berwick, Maine. Conlicts of interest: none declared.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Insufficient information
Allocation concealment (selection bias)	Unclear risk	Insufficient information
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "A cross-over design was used. In the crossover design, classrooms in each grade level were randomized to begin as the experimental group (regular hand washing)."
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "The school nurse served as the data collector for the duration of the study. This could be perceived as bias in the measurement of the outcome variable, absenteeism related to infectious illness."



Morton 2004 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Insufficient information
Selective reporting (reporting bias)	Unclear risk	Insufficient information

Najnin 2019

Study characteristics	
Methods	Cluster-RCT, parallel assignment
Participants	Residents of the high-risk, cholera-prone study areas. Low-income communities in Mirpur area of urban Dhaka defined by low per capita income, poor sanitation, unsafe water use, sharing of water source, and poor living conditions. 90 geographic clusters were included, with 30-metre buffer zones.
	A total of 7842 households, with 52,237 individuals analysed
	Vaccine-only area: data were analysed for 1965 households consisting of 13,148 individuals
	Vaccine-plus-behaviour-change area: data were analysed for 3886 households consisting of 25,566 individuals
	Control area: data were analysed for 1991 households consisting of 13,523 individuals
	Study criteria from published protocol:
	Inclusion criteria: apparently healthy residents of selected vaccination sites, aged 1 year and above, non-pregnant women, written informed consent
	Exclusion criteria: age less than 1 year and pregnant women
Interventions	Hand-washing and water treatment promotion. See Table 1 for details.
Outcomes	Laboratory: none used
	Effectiveness: prevalence of respiratory illness. People were classified as having respiratory illness if they reported having fever plus either cough or nasal congestion or fever plus breathing difficulty in the past 2 days of unannounced home visits: in each intervention group and amongst those who had soap/ soapy water with water present in the hand-washing station (35% of all groups combined) versus those without this (regardless of the intervention group). Planned secondary outcome: prevalence of reported respiratory illness during 2-year intervention period
	Safety: no adverse effects planned or reported
Notes	The period study conducted: 2011 to 2013
	Funding: government and private Bill & Melinda Gates Foundation Conlicts of interest: none declared.
Risk of bias	
Bias	Authors' judgement Support for judgement

Low risk

Random sequence genera-

tion (selection bias)

Computer-generated randomisation sequence was used to allocate 90 geo-

graphical clusters to 1 of 3 groups. Before randomisation, clusters were strat-



Najnin 2019 (Continued)		ified blocked into 2 categories according to the distance to the hospital. (parent article: Lancet. 2015 Oct 3;386(10001):1362-1371)
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	All trial participants and investigators were aware of group assignment. Several in and out migrations across all groups before, after, and during outcome monitoring, and large number of changes in intervention areas
Blinding of outcome assessment (detection bias) All outcomes	High risk	Several in and out migrations across all groups before, after, and during outcome monitoring, and large number of changes in intervention areas
Incomplete outcome data (attrition bias) All outcomes	High risk	High migration movement. This could have distorted the baseline characteristics even more. Very hard to assess because the numbers in the index paper are different from the parent paper (Qadri 2015). In addition to that, for each intervention, data were analysed for 15% to 30% of those allocated on start date. Each group started with approximately 80,000 people; the number analysed is much lower (237,216 people were in the study area on start date of outcome monitoring, the total number analysed across all groups was 52,237). No info about data on migrated individuals or on those who changed intervention areas was dealt with? Also data for prevalence of ARI adjusted for age and wealth were not shown. The outcome is addressed in the 2 days preceding an unannounced visit. This means that if there was a respiratory illness in the past week it would not have been reported. Moreover, these monthly unannounced visits were done to a different set of participants in each group!
Selective reporting (reporting bias)	High risk	Published protocol does not include respiratory illness as an outcome.

Nicholson 2014

VICTORSOIT 2014	
Study characteristics	
Methods	Cluster-RCT
Participants	70 low-income communities in Mumbai, India (35 communities per arm) were randomised to intervention arm (N = 1025) and control arm (N = 1026).
	Households located in low-income urban communities in west and south Mumbai, India. Each household contains 1 target child in the first year of a municipal school (typically aged 5 years).
Interventions	Combination of hand-washing promotion with provision of free soap aimed at 5-year-olds with provision of free soap. See Table 1 for details.
Outcomes	Laboratory: none reported
	Effectiveness:
	Primary outcomes: episodes of diarrhoea, ARIs, and school absences amongst target children, and episodes of diarrhoea and ARIs among their families
	Secondary outcomes: episodes of eye infections, vomiting, abscesses or boils, headaches, and earache



Nicholson 2014 (Continued)	Operational defiinitions for all the illnesses were taken from <i>Black's Medical Dictionary</i> (MacPherson 1999). ARIs as "pneumonia, cough, fever, chest pain and shortness of breath, cold, inflammation of any or all of the airways, that is, nose, sinuses, throat, larynx, trachea and bronchi"	
	Safety: no safety measures planned or reported by the investigators	
Notes	The period study conducted: 22 October 2007 to 2 August 2008	
	Funding: multinational corporate company (Unilever plc.) Conlicts of interest: none declared.	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	Coin tossing used, which could have led to a large imbalance.
Allocation concealment (selection bias)	Low risk	"a coin toss was used to assign one community in each pair to intervention and one to control"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Participants knew to which arm they had been recruited. Households were removed from the study if they provided no data for 5 consecutive weeks.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Data collectors were independent of the behaviour change intervention. Each was assigned exclusively to either households in the intervention group or to control households. However, communities, where very low literacy levels exist, were replaced after randomisation.
Incomplete outcome data (attrition bias) All outcomes	High risk	Data for non-completers were available and similar across groups. ITT and PP were performed. However, households were removed from the study if they provided no data for 5 consecutive weeks.
Selective reporting (reporting bias)	Unclear risk	No information to judge

Pandejpong 2012

Study characteristic	S
Methods	Cluster-RCT, single study centre
Participants	Children (total number = 1437) were randomised to alcohol hand gel every 60 minutes (N = 452 children), every 120 minutes (N = 447 children), and once before lunch (N = 540 children).
	Inclusion criteria: all children in a large private school in suburban Bangkok, Thailand, all ages, both genders with parental consent to participate.
	Exclusion criteria: an allergy to alcohol hand gel
Interventions	3 disinfection interventions: Alcohol hand gel applied every 60 minutes vs every 120 minutes vs once before lunch (3 groups). The current school standard for hand hygiene (q lunch group). See Table 1 for details.
Outcomes	Laboratory: none



Pandejpong 2012 (Continued)

Effectiveness:

Primary: rates of absenteeism from physician-confirmed ILI

Secondary: rate of absenteeism caused by total reported ILI (with and without a doctor's confirmation)

In case the child was sick but did not see a doctor, the parents were asked to report any of the following symptoms: runny nose or cough, fever or chills, sore throat, headache, diarrhoea, and presence of hand, foot, or mouth ulcers. If 2 or more of these symptoms were reported, then the child's illness was documented as an ILI.

Safety: investigators reported that no adverse reaction to the alcohol hand gel was reported in any participants

Notes

The period study conducted: December 2009 to February 2010

Funding: Royal College of Physicians of Thailand Conflict of interest: none to report

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	No information provided.
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Parents and teachers are aware of the assignment. Teachers were responsible for recording the absenteeism case record forms. Parents would report child sickness. No diagnostic tests, even in the case of physician-confirmed ILI
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome is physician-confirmed ILI.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "No students were lost to follow-up or discontinued the intervention during the study period."
Selective reporting (reporting bias)	Low risk	All outcomes were reported.

Priest 2014

Ctudy	cha	racta	ristics

Methods	A cluster-RCT
Participants	Study included children aged 5 to 11 years at 68 primary schools in New Zealand. Schools were randomised to hand sanitiser + education session arm (34 schools and 8859 children) and education session arm (34 schools and 7386 children).
	Inclusion criteria:



Priest 2014 (Continued)

School-level inclusion: at least 100 children of primary school age (school years 1 to 6; children will generally range in age from 5 years to 11 years) at November 2008. Schools that are not currently using hand-sanitiser products or are willing to not use them for the period of the trial. Schools are within the City boundaries of Christchurch, Dunedin, or Invercargill in New Zealand. The principal of the school consents to the school being included in the trial. Not "special schools" (e.g. schools for children with deafness or disability) and either not currently using hand-sanitiser products or willing to not use them for the period of the trial if they were randomised to the control group were eligible to participate in the trial.

Student-level inclusion (follow-up children): children were eligible to participate in the follow-up group, for whom more detailed information on absences was collected, if they attended a school year 1 to 6 class in 1 of the included schools at the beginning of the second school term in 2009 (the end of April), and their caregivers completed the consent form indicating that they were willing to be telephoned following their child's absences and that they were able to take part in telephone interviews in English

Exclusion criteria:

School-level exclusion: special needs schools

Student-level exclusion (follow-up children): children of the principal investigators and study personnel of the trial. Or, children of families that the principal of the primary school directs us not to approach

Interventions

Hand sanitiser provision (in addition to hand hygiene education session also provided to control group) in schoolchildren. See Table 1 for details.

Outcomes

Laboratory: none

Effectiveness:

Primary outcome: the incidence rate of absence episodes from school (reported by the parents during telephone calls) due to any illness during the study period (winter term)

Secondary outcomes: assessing whether hand sanitiser was effective in reducing the:

- 1. incidence rate of respiratory illness absence episodes,
- 2. incidence rate of gastrointestinal illness absence episodes,
- 3. incidence rate of absence for any reason,
- 4. length of illness episode,
- 5. length of illness absence episode, and
- 6. incidence rate of subsequent illness amongst other children or adults in the household.

Definition of respiratory illness: at least 2 of the following caregiver-reported symptoms for 1 day, or 1 of the following symptoms for 2 days (but not fever alone): runny nose, stuffy or blocked nose or noisy breathing, cough, fever, sore throat, or sneezing

Safety: examined whether the use of hand sanitiser was associated with an increased risk of any skin reactions during the intervention period. Skin reactions: dryness, redness, flakiness, itchiness, eczema, and any other skin reactions

Notes

The period study conducted: 27 April to 25 September 2009

Government funded: Health Research Council of New Zealand

Competing Interests: the authors have declared that no competing interests exist. All authors affirm that they are not involved in any other trials on the same or a related intervention.

Risk of bias

Bias Authors' judgement Support for judgement



Priest 2014 (Continued)		
Random sequence generation (selection bias)	Low risk	Quote: "Stata/MP 10.1 for Windows was used to generate the random numbers"
Allocation concealment (selection bias)	Low risk	Done by trial statistician provided with school codes and district and randomised the schools to either "A" or "B"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Outcome assessors were blinded to the group allocation until the analysis was completed.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Outcome assessors were blinded to the group allocation until the analysis was completed.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	The study flow diagram gives a clear account on follow-up, with numbers of those lost to follow-up and those who discontinued the intervention along with the reasons for doing so. No child was excluded from the analysis. Only PP analysis was reported.
Selective reporting (reporting bias)	Low risk	All outcomes stated in the published protocol were reported in the study. The exception was quote: "1 planned secondary outcome (that is irrelevant to our study) that was not collected and 2 collected secondary outcomes that were not planned in the original protocol".

Radonovich 2019

Study characteristics	s
Methods	Cluster-RCT, multicentre, pragmatic effectiveness trial
Participants	Study included 280 clusters randomly assigned to N95 respirators (189 clusters and 1993 HCPs) and medical masks (191 clusters and 2058 HCPs).
	All participants in a cluster worked in the same outpatient clinic or outpatient setting. All participants were permitted to participate for 1 or more years and gave written consent for each year of participation.
	Inclusion criteria: healthcare workers in outpatient settings serving adult and paediatric patients with a high prevalence of acute respiratory illness. Participants were aged at least 18 years and employed at 1 of the 7 participating health systems, and self-identified as routinely positioned within 6 feet (1.83 m) of patients. Participants were full-time employees (defined as direct patient care for approximately \geq 24 hours weekly) and worked primarily at the study site (defined as \geq 75% of working hours). Exclusion criteria: medical conditions precluding safe participation or anatomic features that could interfere with respirator fit, such as facial hair or third-trimester pregnancy. Participants self-identified race and sex using fixed categories; these variables were collected because facial anthropometrics related to race and sex may influence N95 respirator fit.
Interventions	Fit-tested N95 respirators versus medical masks when near patients with respiratory illness. See Table 1 for details.
Outcomes	Laboratory. Primary outcome: the incidence of laboratory-confirmed influenza, defined as:
	 detection of influenza A or B virus by RT-PCR in an upper respiratory specimen collected within 7 days of symptom onset; detection of influenza from a randomly obtained swab from an asymptomatic participant; and



Radonovich 2019 (Continued)

3. influenza seroconversion (symptomatic or asymptomatic), defined as at least a 4-fold rise in haemagglutination inhibition antibody titres to influenza A or B virus between pre-season and postseason serological samples deemed not attributable to vaccination.

Effectiveness. Secondary outcomes: the incidence of 4 measures of viral respiratory illness or infection as follows:

- 1. acute respiratory illness with or without laboratory confirmation;
- laboratory-detected respiratory infection, defined as detection of a respiratory pathogen by PCR or serological evidence of infection with a respiratory pathogen during the study surveillance period(s), which was added to the protocol prior to data analysis;
- 3. laboratory-confirmed respiratory illness, identified as previously described (defined as self-reported acute respiratory illness plus the presence of at least PCR-confirmed viral pathogen in a specimen collected from the upper respiratory tract within 7 days of the reported symptoms and/or at least a 4-fold rise from pre-intervention to postintervention serum antibody titres to influenza A or B virus; and
- 4. influenza-like illness, defined as temperature of at least 100 °F (37.8 °C) plus cough and/or a sore throat, with or without laboratory confirmation.

Safety: no serious study-related adverse events were reported. 19 participants reported skin irritation or worsening acne during years 3 and 4 at 1 site in the N95 respirator group.

Notes

The study was conducted from September 2011 to May 2015, with final follow-up on 28 June 2016.

Compliance: adherence was reported on daily surveys 22,330 times in the N95 respirator group and 23,315 times in the medical mask group. Quote: "Always" was reported 14,566 (65.2%) times in the N95 respirator group and 15,186 (65.1%) times in the medical mask group; "sometimes" 5407 (24.2%) times in the N95 respirator group and 5853 (25.1%) times in the medical mask group; "never" 2272 (10.2%) times in the N95 respirator group and 2207 (9.5%) times in the medical mask group; and "did not recall" 85 (0.4%) times in the N95 respirator group and 69 (0.3%) times in the medical mask group. Participant-reported adherence could not be assessed in 784 participants (31.2%) in the N95 respirator group and 822 (30.8%) in the medical mask group (P = 0.84) because of lack of response to surveys or lack of adherence opportunities (i.e. participants did not encounter an individual with respiratory signs or symptoms). Analysed post hoc, participant adherence was reported as always or sometimes 89.4% of the time in the N95 respirator group and 90.2% of the time in the medical mask group.

Government funded.

Conflict of interest disclosures: Dr Bessesen reported receiving grants from the Department of Veterans Affairs during the conduct of the study. Dr Brown reported receiving grants from the US Department of Veterans Affairs during the conduct of the study. Dr Cummings reported receiving grants from the Centers for Disease Control and Prevention, the National Institutes of Health, and MedImmune outside the submitted work and the Biomedical Advanced Research and Development Authority during the conduct of the study. Ms Los reported receiving grants from Centers for Disease Control and Prevention, the Veterans Health Administration, and the Biodefense Advanced Research and Development Agency during the conduct of the study. Dr Gibert reported receiving financial support for the conduct of the study, including research personnel, from the Veterans Health Administration during the conduct of the study. Dr Gorse reported receiving grants from the US Department of Veterans Affairs during the conduct of the study. Dr Nyquist reported receiving grants from the Centers for Disease Control and Prevention/Division of Healthcare Quality Promotion, the National Institute for Occupational Safety and Health, and the Veterans Health Administration during the conduct of the study; personal fees and nonfinancial support from Sequirus outside the submitted work; and serving on a policy making committee regarding infectious disease for the American Academy of Pediatrics Committee on Infectious Diseases. Dr Reich reported receiving grants from Veterans Health Administration during the conduct of the study. Dr Rodriguez-Barradas reported receiving grants from Veterans Affairs Central Office during the conduct of the study. Dr Perl reported receiving grants from the Centers for Disease Control and Prevention and Biomedical Advanced Research and Development Authority during the conduct of the study and grants from Medimmune outside the submitted work. No other disclosures were reported.

Risk of bias

Bias

Authors' judgement Support for judgement



Radonovich 2019 (Continued)		
Random sequence generation (selection bias)	Low risk	Computer-generated random sequences by an individual not involved in the study implementation and data analyses. Used stratified randomisation
Allocation concealment (selection bias)	Low risk	Used constrained randomisation
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The participants cannot be blinded, but it seems that all the measures otherwise were the same with meticulous follow-up. Besides, the primary outcome was lab based (an objective outcome), which is unlikely to be affected by of lack of blinding. Investigators were blinded to the randomisation until completion of the study and analysis.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Primary outcome is laboratory-confirmed diagnosis.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Quote: "Missing outcomes were imputed using standard multiple imputation techniques, creating multiple imputed data sets with no missing values for each analysis"
Selective reporting (reporting bias)	Low risk	Reported study outcomes matched the published protocol. Every outcome was accounted for.

Ram 2015

Study characteristic	s
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Methods	RCT
Participants	377 household compounds (index cases) completed the study. Control arm has 184 compounds with 1607 contacts, and intervention group has 193 compounds with 1814 contacts. Final analysis was performed on 193 index cases and 1661 contacts in the intervention group and 184 index cases and 1498 contacts in the control group.

In 2009, index case-patients with symptom onset within 7 days preceding enrolment were eligible. Eligibility criteria changed in 2010 to include index case-patient with symptom onset within 48 hours preceding enrolment.

Inclusion criteria:

- 1. Individuals ≥ 5 years old: ILI, defined as history of fever and either cough or sore throat with fever onset within the previous 24 hours.
- 2. Individuals < 5 years old: any child with acute fever with onset within the previous 24 hours.
- 3. Return to home within 24 hours of presentation to Upazilla Health Complex, Jahurul Islam Medical College Hospital or the local pharmacies, i.e. the index case cannot be admitted for treatment. If admitted, the patient would not be eligible.
- 4. No fever in any bari resident during the 7 days preceding the patient's presentation to hospital (see definition below).
- 5. At least 2 individuals (in addition to the index case-patient) who intend to reside in the bari during the subsequent 20 days.
- 6. Residence within 30 minutes travel time (1-way) from the Upazilla Health Complex or Jahurul Islam Medical College Hospital or the local pharmacy.

Exclusion criteria: compounds were excluded if any compound member(s) was reported to have fever within 3 days before index case-patient enrolment. At another time point, compounds were excluded



Ram 2015 (Continued)	if any primary household member was reported to have fever (fever occurring within 48 hours prior to enrolment recorded).
Interventions	Promoting intensive hand-washing in households to prevent transmission of ILI. See Table 1 for details.
Outcomes	Laboratory: PCR for influenza A and B, with further subtyping of influenza A isolates for all ILI amongst contacts
	Effectiveness: incidence of ILI. An age-based definition of ILI was used as follows.
	 For individuals > 5 years old, ILI was defined as history of fever with cough or sore throat. For children < 5 years old, ILI was defined as fever (the authors used this relatively liberal case definition in order to include influenza cases with atypical presentations in children).
	Safety: no safety data planned or reported by investigators
Notes	Inclusion/exclusion criteria changed 3 times during the study conduct.
	The period study conducted: June 2009 to December 2010
	Government funded Competing interests: the authors have declared that no competing interests exist.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Block randomisation, with a block size of 4, in order to promote random and even allocation of household compounds to the 2 treatment arms. The list of random assignments was generated by an investigator with no contact with the participants.
Allocation concealment (selection bias)	Low risk	Once baseline data collection was complete, the data collector notified the field research officer, who consulted the block randomisation list to make the assignment of the household compound to intervention or control.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Relied on symptom reporting from the head of family. Inclusion/exclusion criteria changed 3 times during the study conduct. Given the provision of a hand-washing station as part of the intervention, it was not possible to ensure blinding of participants, intervention staff, or data collectors.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Relied on symptom reporting from the head of family. Inclusion/exclusion criteria changed 3 times during the conduct of the study. Given the provision of a hand-washing station as part of the intervention, it was not possible to ensure blinding of participants, intervention staff, or data collectors.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Flow chart followed all households an individuals from recruitment to analysis.
Selective reporting (reporting bias)	Low risk	The specified outcomes are clearly accounted for Investigators report all outcomes for each modified enrolment.



Roberts 2000

Study characteristics	
Methods	Open cluster-RCT carried out between March and November 1996 (the Southern Hemisphere winter season) in 23 childcare centres caring for a minimum of 50 children 10 hours a day, 5 days a week in Australia. The study assessed the effects of an Australian national hand-washing programme compared to standard procedure. Randomisation was according to a random-number table, and cluster coefficients are reported.
Participants	Children (299 in the intervention arm and 259 in the control arm) aged 3 or younger attending the centres at least 3 days a week. Attrition was 51 children in the intervention arm and 72 children in the control arm due mainly to staff leaving the centres.
Interventions	Hand-washing programme with training for staff and children. It is unclear whether any extra hand-cleansing agents were used, as GloGerm (?) is mentioned when it was used in a preliminary study. See Table 1 for details.
Outcomes	Laboratory: N/A Effectiveness: ARI (runny nose, cough, and blocked nose) Follow-up was via a parental phone interview every 2 weeks. Safety: N/A
Notes	Risk of bias: low (cluster coefficients and analysis by unit of randomisation) Note: the authors conclude that although there was no overall decrease in respiratory illness (RR 0.95, 95% CI 0.89 to 1.01), in children up to 24 months the decrease was statistically significant (RR 0.90, 95% CI 0.83 to 0.97). The authors speculated that this was because maximum benefits are likely from this age group due to their limited ability to wipe their nose and hands without a structured programme. Analyses by 3 compliance levels are also reported. A so-so reported and well-conducted trial.
	This work was supported by a grant from the Commonwealth Department of Family Services and Health, Research and Development Scheme. Conflict of interest: none to report.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Randomisation was according to a random-number table.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	It was not possible to blind the intervention.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "The observer was not informed of the content of the training sessions or the intervention status of the centres."
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Recruitment rate 88% (23 of 26 CCCs); loss to follow-up not clear, as no denominator given
Selective reporting (reporting bias)	Low risk	Centres were comparable at baseline.



Sandora 2005

Study characteristics	
Methods	Single-blind, cluster-RCT carried around the Boston area, USA, in the period of November 2002 to April 2003. The trial tested the effects of using a hand sanitiser and a programme of instruction on the transmissions of GI infections (data not extracted) and ARIs in families. Units of randomisation were child-care centres and were carried out on enrolment by an investigator using random block size generated by computer. Assignment was single-blind (i.e. investigator blinded to the status of the centre). Cluster correlation was 0.01.
Participants	292 families with 1 or more children aged 6 months to 5 years who were in child care for 10 or more hours a week
	155 children in 14 centres were allocated to the intervention arm and 137 children in 12 centres to the control arm. The mean age was 3 to 2.7 years. Attrition was respectively 15 (3 lost to follow-up and 12 who discontinued the intervention) and 19 (8 lost to follow-up and 11 who discontinued the intervention). ITT analysis was carried out.
Interventions	Alcohol-based hand sanitiser with biweekly hand hygiene educational materials over 5 months versus biweekly educational material on healthy diet. See Table 1 for details.
Outcomes	Effectiveness: ARI (2 of the following symptoms for 1 day or 1 of the following symptoms for 2 days: runny nose, cough, sneezing, stuffy or blocked nose, fever, sore throat). An illness episode had to be separated by 2 symptom-free days from a previous episode. A secondary illness was when it followed a similar illness in another family member by 2 to 7 days. Follow-up was by means of biweekly phone calls to caregivers. Safety: dry skin (71 reports), stinging (11 reports), bad smell (7 reports), dislike (2 reports), allergic reaction (2 reports), slippery feel (1 report), and irritation (20 reports).
Notes	Risk of bias: low Note: the authors conclude that although the rate of GI illnesses was significantly lower in the intervention group, the IRR was not significantly different for ARIs (0.97, 95% CI 0.72 to 1.30). Compliance and droplet route spread may account for this apparent lack of effect. A well-reported trial.
	Study funds and hand sanitiser were provided by GOJO Industries, Inc (Akron, OH). No conflict of interest declared.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Random assignments were generated by computer using a permuted-blocks design with random block sizes."
Allocation concealment (selection bias)	Low risk	Low riskUnclear riskHigh risk
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "Teachers in the intervention classrooms were responsible for encouraging the use of the disinfecting wipes and hand sanitizer according to the study protocol Given that no placebo was provided and sanitizer use was recorded, neither families nor data collectors could be blinded as to the group assignment of the family."
Blinding of outcome assessment (detection bias) All outcomes	High risk	Quote: "Given that no placebo was provided and sanitizer use was recorded, neither families nor data collectors could be blinded as to the group assignment of the family."



Sandora 2005 (Continued)				
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition was 15 in intervention arm (3 lost to follow-up and 12 who discontinued the intervention) and 19 in the control arm (8 lost to follow-up and 11 who discontinued the intervention). ITT analysis was carried out.		
Selective reporting (reporting bias)	Unclear risk	Well-reported		

Sandora 2008

Study characteristics	
Methods	Cluster-RCT carried out in a single elementary school system located in Avon, Ohio, USA to assess the effectiveness of a multifactorial infection-control intervention, including alcohol-based hand sanitiser and surface disinfection, in reducing absenteeism caused by gastrointestinal and respiratory illnesses amongst elementary school students. The study also aimed to describe the viral and bacterial contamination of common surfaces in the school classroom and to assess the impact of an environmental disinfectant on the presence of selected viruses and bacteria on these surfaces. Clustering was described as "teams of 3-4 classes depending on the class year".
Participants	A total of 363 students in 15 different classrooms were eligible to participate and received letters about the study.
	A sample of 285 of these students provided written informed consent and were randomly assigned to the intervention group (146) or to the control group (139) and contributed to final analysis.
	No students were lost to follow-up or discontinued the intervention during the study period.
	Baseline demographic characteristics were similar in the intervention and control groups. Most families were white and non-Hispanic and in excellent or very good health at baseline.
Interventions	Alcohol-based hand sanitiser to use at school and quaternary ammonium wipes to disinfect classroom surfaces daily for 8 weeks versus usual hand-washing and cleaning practices. See Table 1 for details.
Outcomes	Laboratory: Serological evidence: no Swabs for bacteria and viruses from 3 types of classroom surfaces were taken. Effectiveness: Respiratory illness defined as days absent as measured by a (blinded) school worker who routinely recorded reason for absenteeism either for gastrointestinal or respiratory causes.
	Safety: N/A
Notes	The authors conclude that the multifaceted intervention that included alcohol-based hand sanitiser use and disinfection of common classroom surfaces reduced absenteeism from gastrointestinal illness amongst elementary school students. The intervention did not impact on absenteeism from respiratory illness. In addition, norovirus was detected less frequently on classroom surfaces in the group receiving the intervention. The study is of good quality with low risk of bias. The authors checked compliance by counting discarded wipes. Reasons given for the apparent lack of effect against ARIs but good effect on GI illness are that disinfecting the classroom surfaces (daily at lunchtime with alkali) was important, as were the alcohol wipes. The authors measured the norovirus concentration on surfaces and found this to be reduced. Other reasons may be that droplets are not affected by this method, or that contamination of hands by respiratory infections is likely to be continuous (in orofaecal transmission is mostly at the time of defecation).
	Study funds, hand-sanitiser, and disinfecting wipes were provided by The Clorox Company (Oakland, CA).



Sandora 2008 (Continued)

Financial disclosures: Drs Sandora and Goldmann received a consulting fee from The Clorox Company for their efforts in designing and conducting this study; Dr Shihh as indicated she has no financial relationships relevant to this article to disclose.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "The allocation sequence was generated by computer"
Allocation concealment (selection bias)	Unclear risk	Quote: "and teams were assigned to study groups by a study investigator (Dr Shih)."
		Blinding of allocation cannot be guaranteed.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not possible
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: " All of the students absences were recorded in the usual fashion by the school employee who normally answers this dedicated telephone line. This employee was blinded to the group assignment of the child."
Incomplete outcome data (attrition bias) All outcomes	Low risk	No students were lost to follow-up or discontinued the intervention during the study period.
Selective reporting (reporting bias)	Unclear risk	Well-reported

Satomura 2005

Study characteristics

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RCT. Randomisation was achieved by simple computer-generated random digit. Allocation was concealed using sealed, opaque envelopes. Not clear if there was a central randomisation centre. Post hoc exchange of envelopes was prevented by writing both the name of each participant and the number on the envelope he/she drew before breaking the seal. Participants were not blinded to the intervention; however, disease incidence was determined by 1 study physician who was not informed of the results of assignment. Analysis was done based on the intention-to-treat principle. The study targeted community healthcare all over Japan and was conducted between December 2002 and March 2003 for a follow-up period of 60 days.

Participants

387 participants at 18 sites were recruited, 384 were included in the analysis: water gargling (N = 122), povidone-iodine gargling (N = 132), and control (N = 130).

Follow-up was completed on 338 participants. Attrition was fully explained for URTI analysis; however, 2 participants were not accounted for in the ILI analysis. 46 participants did not complete the follow-up due to either discontinuation of diary use (n = 9) or contracting ILI (n = 37).

Of the 37 participants with ILI, 11 were in the povidone-iodine group, 12 in the water group, and 14 in

the control group. Analysis was performed on 35 participants (Kitamura 2007 [Kitamura 2007]).

Interventions

Participants were randomised to 1 of the following: water gargling, n = 122 (20 mL of water for about 15 seconds 3 times consecutively, at least 3 times a day); povidone-iodine gargling, n = 133 (20 mL of 15 to



Satomura 2005 (Continued)

30 times diluted 7% povidone-iodine (as indicated by the manufacturer) in the same way as water gargling); and control, n = 132 (retain their previous gargling habits).

All groups were asked to fill a daily gargling diary (standardised form to record: gargling habits, handwashing, and influenza complaints).

The frequency of gargling in the water group was higher (3.6); the frequency of hand-washing was similar amongst the 3 groups.

URTI symptom was classified according to Jackson methods. Diary recording was continued throughout the follow-up period and for 1 week after the onset of URTI.

ILI was reported separately.

See Table 1 for details.

Outcomes

Laboratory: none Effectiveness:

Primary outcome: incidence of first URTI. Index cases were defined as all of the following conditions:

- 1. both nasal and pharyngeal symptoms,
- 2. severity of at least 1 symptom increased by 2 grades or more, and
- 3. worsening of a symptom of 1 increment or more for > 3 days.

Secondary outcome: severity of URTI of the incident cases was assessed by grading each symptom during the initial 7 days after the onset of URTI in numeric scores: none = 0, mild = 1, moderate = 2, and severe = 3

ILI was defined as both developing a fever of 38 °C or higher and worsening arthralgia in addition to some respiratory symptoms (Kitamura 2007).

Safety: no harm was reported. However, 2 participants in the povidone-iodine group switched to water gargling (analysed in their assignment group).

Notes

The authors concluded that simple water gargling is effective in preventing URTIs amongst healthy people. However, no statistically significant difference was observed against ILIs.

The study was well-conducted; blinding would have added to the validity of the results. In addition, the study was not powered enough to detect a statistically significant preventative effect against ILI. The study demonstrates that in addition to hand-washing, simple gargling even with water can reduce URTI, but not ILI. However, during periods of endemic influenza, multiple inexpensive and simple modalities (hand-washing, masks, gargling) can be utilised together to reduce infection and transmission.

Overall, the reporting of the 2 combined studies together is highly confusing. In the first study (Satomura 2005), the main outcome is URTI defined as fever and arthralgia. The second study (which is a presentation of further data from the 2005 publication in the guise of a short report) introduces the outcome ILI with a definition similar to that of URTI in the first study but referring to the earlier outcome as common cold. Also of note is reporting of significance without confidence intervals. Overall, this potentially important study should be repeated with a larger denominator.

Unclear risk of bias because of confused reporting and absence of double-blinding.

Partial financial support was provided by the Suzuken Memorial Foundation (2002) and Uehara Memorial Foundation (2003) (trial registry, ISRCTN67680497).

No financial conflict of interest was reported by the authors of this paper.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Group assignment was based on simple computer-generated random digits"
Allocation concealment (selection bias)	Low risk	Quote: "By an individual drawing of sealed opaque envelopes, subjects were randomly assigned to the following three groups"
		Quote: "allocation was completely concealed from study administrators"



Satomura 2005 (Continued)		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "To prevent post hoc exchange of the envelopes, local administrators wrote down both the name of each subject and the number on the envelope he/she drew before breaking the seal."
Incomplete outcome data (attrition bias) All outcomes	Low risk	338 of 385 randomised followed up; reasons reported.
Selective reporting (reporting bias)	Unclear risk	Confusing reporting

Savolainen-Kopra 2012

Study characteristics	
Methods	Open cluster-RCT, 3-arm intervention trial
Participants	A total of 21 clusters (683 individuals) were randomised to implement hand hygiene with soap and water (257 individuals), alcohol-based hand rub (202 individuals), or control (224 individuals).
	The study was conducted in distinct office work units in 6 corporations in the Helsinki Region that together employed some 10,000 staff. All employees (age ≥ 18 years, both genders) were contacted by email survey. Inclusion criteria: quote: "Volunteers working in defined units" Exclusion criteria: quote: "Persons with open wounds or chronic eczema in hands" The designated 21 study clusters were identified as operationally distinct working units, each containing at least 50 people.
Interventions	Hand hygiene with soap and water and standardised instructions on how to limit the transmission of infections. Usual hand hygiene (control). See Table 1 for details.
Outcomes	Laboratory:
	Quote: "Between November 2008 and May 2010, the seven occupational health clinics serving the six participating corporations were advised to collect, using standard techniques, two to three respiratory samples per week from typical RTI patients and also faecal samples from a few representative patients with gastrointestinal symptoms when a GIT outbreak was suspected. The samples could originate from the study participants and also from work units not included in the study. In the laboratory, viral nucleic acids were extracted with well-characterized commercial kits and tested by validated real-time PCR methods to detect influenza A and B viruses, respiratory syncytial virus, parainfluenza virus types 1, 2, and 3, adenoviruses, human rhinoviruses and human enteroviruses from respiratory specimens, and norovirus from faecal specimens (detailed descriptions of the test procedures are available from the authors)."
	Effectiveness:
	Predefined primary endpoints:
	 Number of reported infection episodes in a cluster per total reported weeks. Number of reported sick leave episodes in a cluster per total reported weeks.
	Secondary endpoints and outcome measures:



Savolainen-Kopra 2012 (Continued)

- 1. Number of days with reported symptoms of RTI and/or GTI in a cluster within a time frame of 100 reporting weeks.
- 2. Number of days-off due to own RTI or GTI in a cluster within a time frame of 100 reporting weeks.

Safety: reported 0 adverse events

Notes

The period study conducted: January 2009 to May 2010

Government funded.

Competing interests: the authors declare that they have no competing interests.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Insufficient information
Allocation concealment (selection bias)	Low risk	Quote:"clusters were matched and randomized prior to onset of the interventions"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	The interventions were not blinded to any party involved (i.e. the study group, participants, or the occupational health services). Subjective reporting of disease episodes
Blinding of outcome assessment (detection bias) All outcomes	High risk	Subjective reporting of disease episodes
Incomplete outcome data (attrition bias) All outcomes	High risk	24% loss to follow-up. However, new recruiting in most clusters; the total number of reporting participants at the end of the trial was 91.7% compared to that at the beginning. Attrition was reported, and 76% of volunteers who started reporting continued to do so until the end of the study. Because of new recruiting in most clusters, the total number of reporting participants at the end of the trial was 626, or 91.7%, compared to that at the beginning. This means that 15.7% of the participants were replaced during the study!!! Raw data on the effects of the interventions on the occurrence of respiratory infections and vomiting/diarrhoea diseases were not reported. Zero adverse effects were reported.
Selective reporting (reporting bias)	Low risk	All planned outcomes were reported.

Simmerman 2011

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Study characteristic	S
Methods	Randomised controlled study
Participants	Study recruited 348 households and 885 members and randomised them as follows:
	1. Control (index household = 119, with 302 family members)
	2. Hand-washing (index household = 119, with 292 family members)
	3. Hand-washing and face mask (index household = 110, with 291 family members)



Simmerman 2011 (Continued)

The household members of children (index cases) presenting with ILI at the outpatient department of the Queen Sirikit National Institute of Child Health (QSNICH) in Bangkok, the largest public paediatric hospital in Thailand

Inclusion criteria:

For index cases: children aged 1 month through 15 years, residents of the Bangkok metropolitan area, and had an onset of illness < 48 hours before respiratory specimens tested positive for influenza by an RIDT that was later confirmed by qualitative real-time RT-PCR (rRT-PCR)

Eligible index cases' households must have had at least 2 other members aged ≥ 1 month who planned to sleep inside the house for a period of at least 21 days from the time of enrolment.

Exclusion criteria:

For index cases: children at high risk for severe influenza complications (e.g. chronic lung disease, renal disease, and long-term aspirin therapy) and those treated with influenza antiviral medications

Excluded households: those with any member reporting an ILI that preceded the index case by 7 days or less and households where any member had received influenza vaccination during the preceding 12 months

Interventions

Hand-washing, or hand-washing plus paper surgical face mask, or control. See Table 1 for details.

Outcomes

Laboratory:

To identify index cases:

QuickVue Influenza A+B rapid diagnostic kit (Quidel Co., San Diego, CA, USA), followed by rRT-PCR for influenza viral RNA

Index cases and contacts tested with nasal swab and throat swab both processed for rRT-PCR.

2 blood samples for antibody seroconversion collected on Days 1 and 21 (seroconversion defined as a fourfold rise in HI titre between paired sera for any of the antigens assayed).

Effectiveness:

Laboratory-confirmed secondary influenza virus infections amongst household members described as the secondary attack rate (SAR). A secondary influenza virus infection was defined as a positive rRT-PCR result on Days 3 or 7 or a fourfold rise in influenza HI antibody titres with the virus type and subtype matching the index case.

SAR for ILI defined by the WHO as fever plus cough or sore throat, based on self-reported symptoms.

Safety: no safety measures planned or reported by the investigators

Adherence: participants in the control arm reported an average of 3.9 hand-washing episodes/day (on Day 7), whilst participants in the hand-washing arm reported an average of 4.7 hand-washing episodes/day (95% CI 4.3 to 5.0; P = 0.002 compared to controls), and participants in the hand-washing plus face mask arm reported 4.9 episodes/day (95% CI 4.5 to 5.3; P < 0.001 compared to controls). In the intervention arms, parents had the highest reported daily hand-washing frequency (5.7, 95% CI 5.3 to 6.0) followed by others (4.8, 95% CI 4.3 to 5.3), siblings (4.3, 95% CI 3.7 to 4.8), and the index cases (4.1, 95% CI 3.8 to 4.4). There was no difference in the average amount of soap used in a week in the hand-washing arm (54 mL per person) and the hand-washing plus face mask arm (58.1 mL per person) (P = 0.15). 289 participants in the hand-washing plus face mask arm used an average of 12 masks per person per week (median 11, IQR 7 to 16) and reported wearing a face mask a mean of 211 minutes/day (IQR 17 to 317 minutes/day). Parents wore their masks for a median of 153 (IQR 40 to 411) minutes per day, far more than other relations (median 59; IQR 9 to 266), the index patients themselves (median 35; IQR 4 to 197), or their siblings (median 17; IQR 6 to 107). The study authors note that differences in average usage may be an attenuated measure of appropriate use in relation to the actual unmeasured exposure risk such as proximity to the index case.

Notes

The period study conducted: April 2008 and August 2009



Simmerman 2011 (Continued)

Government funded.

 $\hbox{BJC has received research funding from MedImmune Inc. No other declarations are reported.}\\$

Risk	of	bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	"Randomization was achieved using a block randomization method using a list of blocks each with 12 household IDs, four of which were assigned to each of the three study arms."
Allocation concealment (selection bias)	Unclear risk	Quote: "A study coordinator assigned each household to one study arm after consent was obtained"
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Recruiting clinicians were blinded to the allocation of the specific intervention. The participants were not blinded, but it is unlikely that the outcome would have been affected by lack of blinding.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	The primary outcome is a laboratory-confirmed influenza.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Household flow chart provided with reasons for exclusions, all numbers provided. Analysis was done by ITT and PP.
Selective reporting (reporting bias)	Low risk	All outcomes are accounted for in the ITT analysis of the results.

Stebbins 2011

Study Characteristics	•
Methods	Cluster-RCT, open-label
Participants	Study included 3360 students from 10 Pittsburgh elementary schools. Intervention arm (5 schools, 1695 people) and control arm (5 schools, 1665 people)
	No inclusion or exclusion criteria were provided.
Interventions	Training in hand and respiratory (cough) hygiene. Hand-sanitiser was provided and encouraged to be used regularly. See Table 1 for details.
Outcomes	Laboratory:
	Primary outcome: laboratory-confirmed influenza (RT-PCR) amongst children presenting with ILIs leading to their absence from school
	2 nasal swabs were obtained using test manufacturer-approved sterile Dacron swabs. 1 swab was employed for influenza testing using the QuickVue Influenza A+B test (Quidel Corp, San Diego, CA).
	The second nasal swab was delivered on cold pack to the University of Pittsburgh Medical Center Clinical Virology Laboratory, Pittsburgh, PA for RT-PCR testing (performed within 48 hours). The RT-PCR used viral nucleic acid extract (EasyMag; bioMerieux, Durham, NC)
	and primer/probe sequences for influenza A, influenza B, and influenza A H1 and H3



Stebb	ins	2011	(Continued)
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subtypes (CDC, Atlanta GA).

Effectiveness:

 $Secondary\ outcome:\ absence\ episodes\ and\ cumulative\ days\ of\ absence\ due\ to\ ILI,\ any\ illness,\ and\ all\ outcome$

causes

Safety: none mentioned

Notes

The period study conducted: 1 November 2007 through 24 April 2008

Funding: this research was supported by Cooperative Agreement number 5UCl000435-02 from the Cen-

ters for Disease Control and Prevention (CDC).

DC and DB received support from the NIH MIDAS program (1U01-GM070708). DC holds a Career Award

at the Scientific Interface from the Burroughs Welcome Fund. No other conflicts declared.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "constrained randomization algorithm"
Allocation concealment (selection bias)	Low risk	Quote: "Random allocation of schools to two arms was created by Dr. Cummings and concealed until intervention assignment". "At the beginning of the school year parents and guardians were given the opportunity to decline participation"
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded
Blinding of outcome assessment (detection bias) All outcomes	Low risk	In 76% and 78% of illness in intervention and control group were laboratory confirmed. ILI is objectively defined.
Incomplete outcome data (attrition bias) All outcomes	High risk	Only episodes of identified causes were analysed. Causes of absence episodes in 66% of the study participants were not identified (2092 in the intervention group and 2232 in the control group). The parents could be contacted in only 34% cases of absence. About half of them had an illness, and in one-third of these cases the illness met the criteria of ILI (361 cases (33%)). Of these, 279 (77%) were tested for influenza.
Selective reporting (reporting bias)	Unclear risk	Insufficient information to judge

Suess 2012

Study c	haracteristics
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Methods	Cluster-RCT, open-label, parallel design	
Participants	Study sample included 84 households randomised as follows:	
	 30 control (index cases = 30, household contact = 82) 26 mask group (index cases = 26, household contact = 69) 	



Suess 2012 (Continued)

3. 28 mask and hand hygiene group (index cases = 28, household contact = 67)

Inclusion criteria: patients presenting to general practitioners or family physicians at the study sites within 2 days of symptom onset; had a positive rapid antigen test for influenza (later to be confirmed by quantitative RT-PCR (qRT-PCR); and was at least 2 years old. Index cases also had to be the only household member suffering from respiratory disease within 14 days prior to symptom onset. Exclusion criteria were pregnancy, severely reduced health status, and HIV infection. 1-person households were also not eligible or inclusion.

Interventions

Quote: "facemask and practising intensified hand hygiene (MH group), wearing facemask only (M group) and none of the 2 (control group)". See Table 1 for details.

Outcomes

Primary outcomes: SAR of laboratory-confirmed (qRT-PCR) influenza infection amongst household members (secondary infection cases) presenting with ILI within the observation period (8 days from the date of onset). ILI was defined as fever ($> 38.0 \,^{\circ}$ C) + cough or sore throat. Nasal wash specimens (or if these were not possible, nasal swabs) from all participating household members

Effectiveness:

Secondary outcomes: laboratory-confirmed influenza infection in a household contact (secondary infection cases). The study authors defined a symptomatic secondary influenza virus infection as a laboratory-confirmed influenza infection in a household member who developed fever (> 38.0 °C), cough, or sore throat during the observation period. They termed all other secondary cases as subclinical. A secondary outcome measure was the occurrence of ILI as defined by WHO as fever plus cough or sore throat.

Safety: study reported that the majority of participants (107/172, 62%) did not report any problems with mask-wearing. This proportion was significantly higher in the group of adults (71/100, 71%) compared to the group of children (36/72, 50%) (P = 0.005). The main problem reported by participants (adults as well as children) was "heat/humidity" (18/34, 53% of children; 10/29, 35% of adults) (P = 0.1), followed by "pain" and "shortness of breath" when wearing a face mask.

Notes

Period study conducted: November 2009 to April 2011

Adherence: in general, daily adherence was good, reaching a plateau of over 50% in nearly all groups (M and MH groups; 2009/10 and 2010/11) from the third day on (by then the intervention had been implemented in all households). A gradual decline towards lower adherence began around the sixth day of the index patient's illness.

Government funded.

The authors declare that they have no competing interests.

Risk of bias

Bias	Authors' judgement	Support for judgement	
Random sequence genera- Low risk tion (selection bias)		Quote: "prepared lists of random numbers with Microsoft Excel 2003 (Mircosoft™ Cooperation, Seattle, USA) which were divided between the three tervention groups. Each participating physician received a list of random n bers with the interventions represented in a 1:1:1 ratio"	
Allocation concealment (selection bias)	Low risk	Quote: "the participating physician received a list of random numbers with the interventions represented in a 1:1:1 ratio. Eligible index patients were randomly assigned a number, which was then communicated to the study center. The resulting intervention was only communicated to the households with the physicians. Intervention material was given to the study sites in closed boxes marked only with the randomisation number. Recruiting physicians were not aware of the allocation of the numbers to the interventions and the boxes for the three intervention arms looked identical. After randomisation, participants were given their box by the physician's assistants"	



Suess 2012 (Continued)						
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Outcomes are very objective and therefore unlikely to be influenced by lack of blinding. In addition, Quote: "physicians (as well as laboratory personnel) blinded from the randomisation results".				
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: physicians (as well as laboratory personnel) blinded from the randomisation results". Outcomes are very objective and therefore unlikely to be influenced by lack of blinding.				
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up. Daily follow-up home visits over the short period of data collection (8 days)				
Selective reporting (reporting bias)	Low risk	The follow-up period is very short (8 days) with very good coverage, and the criteria for defining the outcome are highly objective. All planned outcomes were reported.				

Swarthout 2020

Study characteristics	
Methods	Cluster randomised open-label controlled trial carried out over 18 months in Kenyan geographically near villages to test the effect of a package of measures on pregnant mothers and then on prevalence of ARIs in their young children
Participants	7246 pregnant women in 702 clusters were enrolled, with 6960 children in year 1 and 7088 in year 2 children with available ARI data. The mean ages of index children and siblings younger than 3 years were 14.2 months (SD: 6.77 months) and 22.9 months (SD: 5.70 months) for years 1 and 2, respectively. The cluster-level intra-cluster correlation coefficient for ARIs was 0.026 for both years. There were 2212 households with 2279 children lost to follow-up by year 2 for unspecified reasons
Interventions	There were 6 intervention groups: chlorinated drinking water (W), improved sanitation (S), handwashing with soap (H), combined WSH, improved nutrition (N) through counselling lipid based nutrient supplementation (LNS) combined WSHN There were 2 control groups passive control (no promotional visits), a double-sized active control (monthly visits to measure mid-upper arm circumference)
	All were done through health promoters with follow up 1 or 2 years after intervention. See Table 1 for details.
Outcomes	Laboratory NR
	Effectiveness
	Prevalence of ARIs in children (defined as cough or difficulty breathing, including panting or wheezing, within 7 days before the interview - in children younger than 3 years).
	Secondary outcomes included difficulty breathing, including panting or wheezing, in the past 7 days (a more specific indicator of respiratory infection than a cough alone); ARI symptoms presenting with fever in the past 7 days (a potentially more severe infection); and facilitator observed runny nose. As this was a rare outcome, caregiver-reported runny nose was analysed post hoc
	Safety NR
Notes	Quote: "The authors conclude that Water, sanitation, and handwashing interventions with behaviour change messaging did not reduce ARIs. Nutrition counselling and LNS modestly reduced ARI symptoms compared with controls in year 1 [prevalence ratio (PR): 0.87, 95% confidence interval (CI): 0.77–0.99], but no effect in the combined WSHN group weakens this finding" Financial support: this work was supported by the Bill & Melinda Gates Foundation (OPPGD759).



Swarthout 2020 (Continued)

The authors declare no further competing interests.

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer random-number generator
Allocation concealment (selection bias)	Unclear risk	Insufficient information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition balanced across groups and < 20%
Selective reporting (reporting bias)	High risk	None of the outcomes reported were prespecified in the trial registry

Talaat 2011

Study characteristics

Stuay cnaracteristics	
Methods	Cluster-RCT
Participants	Children (N = 44,451) in the first 3 primary grades from 60 governmental elementary schools in Cairo, Egypt were included and randomised to 30 schools in the intervention arm (N = 20,882 students) and 30 control schools (N = 23,569 students).
	No exclusion criteria provided.
Interventions	Students were required to wash their hands at least twice during the school days for about 45 seconds, followed by proper rinsing and drying on a clean towel. Campaign material was developed, and posters were placed near sinks in the classroom and playground to encourage hand-washing with soap and water upon arriving at school, before and after meals, using the bathroom, and after coughing and sneezing. See Table 1 for details.
Outcomes	Laboratory: point-of-care influenza A and B viruses using QuickVue (QuickVue; Quidel Corp., San Diego, CA, USA). School nurses collected nasal swabs from children who visited the school clinic with ILI, and only for students who had prior written approval of a parent.
	Effectiveness: rates of absenteeism caused by ILI and laboratory-confirmed influenza. ILI defined as fever > 38 °C and either cough or sore throat.
	Safety: none planned or reported by the investigators
Notes	The period study conducted: 16 February to 12 May 2008



Talaat 2011 (Continued)

Funding: this work was supported by the Centers of Diseases Prevention and Control, Work Unit no. 6000.000.E0016.

No interests declared.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "computer-generated random number table"
Allocation concealment (selection bias)	Unclear risk	No information given.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	The participants and study personnel were not blinded, although lack of blinding is unlikely to have influenced the outcome. Laboratory-confirmed influenza was only conducted only for students who had prior written approval of a parent.
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "Differential interest of study teams may have contributed to the low rate of testing in students who were absent because of ILI in the control schools compared to the intervention schools (12% vs 22%)"
Incomplete outcome data (attrition bias) All outcomes	High risk	No flow chart of clusters flow during the study period. No information on withdrawal. Differential interest of study teams may have contributed to the low rate of testing in students who were absent because of ILI in the control schools compared to the intervention schools (12% vs 22%) incomplete or loss of data. The total number ILI episodes could be an underestimate, as there is no proactive method to look for symptoms of ILI amongst the students; it depends on the student being absent or in class with symptoms that are picked up by the teachers at school.
Selective reporting (reporting bias)	Unclear risk	Insufficient information to judge

Teesing 2021

Ctudy	charac	teristics

M	etl	าด	ds	

Cluster - trial taking place in 66 nursing homes units (33 nursing homes) in the Netherlands during October to December 2016 with 2 follow-up periods (January to April 2017, May to October 2017). Randomisation was carried out by computer and there were some post-randomisation imbalances: the intervention arm had more small and medium-sized nursing homes (< 88 beds, 88 to 118 beds) and the control arm had more large nursing homes (> 118 beds).

Participants

Nursing home staff whose compliance was measured with direct observation according to the WHO-defined HH moments and recorded in a novel app. "The nurses were blinded by giving distinct names to the lessons (The New Way of Working) and the observations (HANDSOME), so that they appeared to be different projects. Nurses were told that the observers were registering the frequency of health care activities (in general)". Staff worked in 66 nursing home units, 36 (976 beds, median 25 per unit) in the intervention arm, and 30 (886 beds, median 28 per unit) in the control arm. During the trial 8 (12%) units left the study during the follow-up for various reasons: 6 intervention units (four during Follow-up 1 and 2 during Follow-up 2) and 2 control units (both during Follow-up 2)



Teesi	ng	2021	(Continued)
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Interventions Hand hygiene (HH) enhancement activities versus no activities. Activities for staff were: an e-learning session, 3 live lessons, posters, and a photo competition. See Table 1 for details.

Outcomes Laboratory NR

Effectiveness

Incidence of gastroenteritis*, influenza-like illness (ILI), assumed pneumonia*, urinary tract infections (UTIs)*, and infections caused MRSA* in residents

*Data not extracted

Safety NR

Notes

The authors conclude that quote: "This study, similarly to comparable studies, could not conclusively demonstrate the effectiveness of an HH intervention in reducing HAIs among residents of nursing homes, despite the use of clearly defined outcome measures, a standardized illness incident reporting instrument, and directly observed HH in a multicenter cluster-RCT. This could be due to an insufficient increase in HH compliance and/or other factors in the nursing home environment that need to be addressed concurrently in order to decrease illness rates"

The trend of ILI incidence reflects that of the outside community at a higher level. This is probably due to ascertainment bias in the nursing homes in the trial. The trend is seasonal and could be accounted for by visitor transmission.

Funding: this study was funded by the Netherlands Organization for Health Research and Development (ZonMw). Non-financial support was received from Essity during the conduct of the study.

Competing interests: the authors declare that they have no competing interests.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Computer random-number generator
Allocation concealment (selection bias)	Unclear risk	Insufficient information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Nurses blinded but participants and other staff members not blinded.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Staff members of nursing homes in the intervention arm were potentially extra alert to infections and more motivated to register them.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Participant flow diagram not reported.
Selective reporting (reporting bias)	Unclear risk	Insufficient information available



en				

Study characteristics			
Methods	2-arm cluster-RCT		
Participants	All residents and staff of 27 privately held chains of nursing homes owned by Korian. 26 nursing home (13 per arm), with an average of 80 residents per nursing home, were included in the study.		
Interventions	Quote: "The intervention was based on a bundle of HH-related measures aimed at NH staff, residents, visitors, and outside care providers. These measures included facilitated access to handrub solution u ing pocket-sized containers and new dispensers, a campaign to promote HH with posters and event or ganization, the formation of local work groups in each NH to work on HH guidelines, and staff education using e-learning on infection control and HH training performed by the same nurse for all NHs." See Table 1 for details.		
Outcomes	Laboratory: none used		
	Effectiveness:		
	Primary outcomes: incidence rate of ARIs and AGE reported in the context of episodes of clustered cases, defined as at least 5 cases within 4 days amongst nursing home residents or staff. ARIs were defined as the combination of at least 1 respiratory symptom with 1 symptom of systemic infection. AGE was defined as the sudden onset of diarrhoea or vomiting in the absence of a non-infectious aetiology.		
	Secondary endpoints were mortality rate, hospitalisation rate, and antibiotic prescription rate (measured in defined daily doses (DDDs) per 100 resident days).		
	Safety: no adverse event surveillance planned or reported by the investigators		
Notes	The period study conducted: 1 April 2014 to 1 April 2015		
	Funding: private (Institute of Ageing Well Korian (Institut du bien vieillir Korian), which runs the nursing homes included in the study)		
	Conflicts of interest: none to report.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	High risk	"simple" randomisation is used	
Allocation concoalment	Undoorriek	No information provided	

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	High risk	"simple" randomisation is used
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Quote: "we suspected that underreporting occurred. The data were verified qualitatively after the end of the intervention through individual phone interviews with each participating NH. Based on these interviews, ARI clustered cases episodes had actually occurred in 12 out of 13 control NHs; however, only 1 had been notified to health authorities. No unreported clustered cases episodes were identified in the intervention NHs"
Blinding of outcome assessment (detection bias) All outcomes	High risk	Data were collected at NH level and reported to centralised by the NH group headquarters in Paris through computerised databases. There was underreporting of ARI and AGE in the control groups. The trial authors suspected that underreporting occurred. Primary outcome: high risk. Secondary outcomes: low risk



Temime 2018	(Continued)
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Incomplete outcome data (attrition bias)
All outcomes

High risk

For the primary outcome, there was underreporting of ARI and AGE in the control groups; no study flow chart was provided; and no reporting on any exclusions. Surveillance is based on voluntary and standardised notifications to health authorities of any AGE or ARI clustered case episode.

Selective reporting (reporting bias)

Low risk

Reported outcomes match planned outcomes published in the protocol.

Turner 2004a

Study characte	ris	tics
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Methods

Double-blind RCT conducted by Hill Top Research, Inc., Winnipeg, Canada, to assess the efficacy of acids with virucidal activity for the inactivation of virus and prevention of experimental rhinovirus colds. Participants in good health, aged 18 to 60, were recruited from Winnipeg and surrounding communities for participation. Qualified participants were randomised to treatment with vehicle (62% ethanol, 1% ammonium lauryl sulphate, and 1% Klucel), vehicle containing 3.5% salicylic acid, or vehicle containing 1% salicylic acid and 3.5% pyroglutamic acid. The volunteers' hands were disinfected, and then test product was applied to both hands of participant. 15 minutes after application, the fingerprints of each hand were contaminated with rhinovirus type 39. The volunteers touched conjunctiva and the nasal mucosa only with the right hand. Viral contamination of the fingers was assessed in the left hands of the volunteers, and viral infection was assessed by culture of nasal lavage specimens and blood samples.

Participants

85 volunteers; 31 control group, 27 used vehicle with 3.5% salicylic acid, 27 used vehicle with 1% salicylic acid and 3.5% pyroglutamic acid

Interventions

Use of salicylic acid versus salicylic acid and pyroglutamic acid versus "placebo" substance. See Table 1 for details.

Outcomes

Laboratory: yes

Effectiveness: rhinovirus type 39 infection

Safety: N/A

Notes

Risk of bias: unclear (no description of randomisation process, concealment or allocation)

Note: the authors concluded that organic acids commonly used in over-the-counter skin care and cosmetic products have substantial virucidal activity against rhinovirus. These preparations provided effective residual antiviral activity on the hands. The virucidal effect of these hand treatments resulted in a reduction in the incidence of rhinovirus infection in the treated volunteers (P = 0.025). The utility of this observation in the natural setting remains to be determined. The volunteers were not allowed to use their hands in the interval between the hand treatment and the virus challenge, so the effect of normal use of the hands on the virucidal activity of these organic acids is not known. Similarly, the virus challenge method used in these experiments may not simulate the natural setting in all aspects. The effect of nasal secretions that would be transferred with the virus in the natural setting on the activity of the acids or on the transmission of virus was not tested in the model.

We are unsure as to the practical significance of this study and the generalisability of its results to the real world. Poorly reported study

Funding for this study was provided by the Procter & Gamble Co., Cincinnati, Ohio.

No interests declared.

Risk of bias

Bias Authors' judgement Support for judgement



Turner 2004a (Continued)		
Random sequence genera-	Unclear risk	Quote: "randomised"
tion (selection bias)		Sequence generation not described.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "double blind", but no description
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "double blind", but no description
Incomplete outcome data (attrition bias) All outcomes	Low risk	All accounted for (short study).
Selective reporting (reporting bias)	High risk	Poorly reported

Turner 2004b

Study characteristics	•
Methods	Double-blind RCT conducted by Hill Top Research, Inc., Winnipeg, Canada, to assess the residual virucidal activity of a skin cleanser wipe and its effectiveness in preventing experimental rhinovirus colds. Participants in good health, aged 18 to 60 years, were recruited from Winnipeg and surrounding communities for participation. The residual activity of a skin cleanser wipe containing 4% pyroglutamic acid formulated with 0.1% benzalkonium chloride was tested. The negative control treatment was 62% ethanol. Benzalkonium chloride had been previously tested and was found to have no virucidal activity. Volunteers were randomly assigned to use the control preparation or the active preparation. The study material was applied to hands with a towelette. 15 minutes later, when the fingers were completely dry, the fingertips of each hand of the control participants and the volunteers in the active treatment group were contaminated with rhinovirus type 39. An additional volunteer in the active group was challenged with virus 1 hour after application, and the final group of volunteers was challenged 3 hours after application. Viral infection was assessed by culture of nasal lavage specimens and blood samples.
Participants	122 volunteers; 30 in control group, 92 in active group (30 tested after 15 minutes, 30 after 1 hour, 32 after 2 hours)
Interventions	Use of a skin cleanser wipe containing 4% pyroglutamic acid formulated with 0.1% benzalkonium chloride versus skin cleanser wipe containing ethanol. See Table 1 for details.
Outcomes	Laboratory: yes Effectiveness: rhinovirus type 39 infection Safety: N/A
Notes	Risk of bias: unclear (no description of randomisation process, concealment or allocation)
	Funding for this study was provided by the Procter & Gamble Co., Cincinnati, Ohio.
	No interests declared.



Turner 2004b (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "randomised"
tion (selection bias)		Sequence generation not described.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Quote: "double blind", but no description given
Blinding of outcome assessment (detection bias) All outcomes	Unclear risk	Quote: "double blind", but no description given
Incomplete outcome data (attrition bias) All outcomes	Low risk	All accounted for (short study).
Selective reporting (reporting bias)	High risk	Poorly reported

Turner 2012

Study characteristics	S
Methods	Randomised controlled clinical trial
Participants	A total of 212 participants were enrolled (116 in the treatment group, 96 in the control group).
	Healthy adult volunteers aged > 18 years from the University of Virginia community Written informed consent was obtained, and volunteers were compensated for participation.
	Exclusion: individuals with skin conditions that would interfere with safety evaluations or medical conditions that could impact the person's well-being or affect study results, and those whose occupations required frequent hand-washing
Interventions	Antiviral hand treatment containing 2% citric acid, 2% malic acid, and 62% ethanol (n = 116) or to a notreatment control group (n = 96). The hand treatment was applied every 3 hours and after hand-washing whilst the participants were awake. See Table 1 for details.
Outcomes	Laboratory: PCR using AmpliTaq Gold DNA Polymerase from Applied Biosystems
	Effectiveness: reduction of rhinovirus-induced common colds; comparison of the number of RV-associated illnesses per 100 participants in the control group with that in the treatment group over 9 weeks. Definitions: a common cold illness was defined as the presence of any of the symptoms of nasal obstruction, rhinorrhoea, sore throat, or cough on at least 3 consecutive days. Illnesses separated by at least 3 symptom-free days were considered to be separate illnesses. Rhinovirus infection was defined as the detection of RV in nasal lavage. All volunteers were seen weekly for nasal lavage, and specimens were assayed by PCR for the presence of RV. PCR-positive specimens separated by at least 8 days and at least 1 negative PCR specimen were considered to be separate infections. RV-associated illnesses were based on detection of RV either at the time of the illness or at the first weekly visit after the illness.



Turner 2012	(Continued)
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Safety: hand irritation occurred in 11 of the 116 volunteers (9%) in the treatment group, which met protocol criteria for removal from the study. An additional 8 participants who did not meet these protocol criteria voluntarily withdrew due to hand irritation. There was no hand irritation in the control group. No other adverse effects of the study treatment were noted.

Notes

The period study conducted: August 2009 to November 2009

Funding: The Dial Corporation - a Henkel Company, Scottsdale, Arizona, USA

Potential conflicts of interest: R. B. T. is a consultant to Henkel and received grant funding to conduct these studies. All other authors are current or former employees of Henkel. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "A randomization code generated using commercially available software was provided by the sponsor"
Allocation concealment (selection bias)	Low risk	Quote: "staff at the study site assigned sequential subject numbers as they enrolled volunteers into the study, and treatment assignment was determined by the subject number."
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	The outcomes are unlikely to be influenced by lack of blinding.
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Personnel who conducted the laboratory assays were blinded to study groups and to whether the specimen was from a routine or illness related visit"
Incomplete outcome data (attrition bias) All outcomes	Low risk	Attrition (and reasons for it) was reported. Study outcomes reported as ITT and PP.
Selective reporting (reporting bias)	Low risk	All planned outcomes in study protocol were reported on.

White 2001

Study characteristics	
Methods	Double-blind, placebo-controlled, cluster-RCT that took place in 3 schools in California during March to April 1999. The study assessed the incremental value of using an alcohol hand rub together with water-and-soap hand-washing. Both arms were administered an educational programme beginning 2 weeks prior to start of the trial. Randomisation was by classroom, and the placebo hand rub was indistinguishable from the active ingredient. Details of randomisation are not given.
Participants	Of the 72 classes originally recruited, lack of compliance (use of supplementary product at least 3 times a day) reduced the classes to 32 (16 in both arms) and a total of 769 participants aged 5 to 12 (381 students who received the sanitiser, and 388 who received the placebo).



White 2001 (Continued)	
Interventions	Pump-activated antiseptic hand rub with benzalkonium chloride (SAB) (Woodward Laboratories) or inert placebo that "virtually" looked the same in batches of 4 colour-coded bottles. School staff, parents, and participants were blinded. See Table 1 for details.
Outcomes	Laboratory: testing of virucidal and bactericidal activity of the active compound Effectiveness: ARI (cough, sneezing, sinus trouble, bronchitis, fever, red eye, headache, mononucleosis, acute exacerbations of asthma) Gastrointestinal and other illnesses (data not extracted) Follow-up and observation was carried out by classroom staff, and illnesses were described by parents. Safety: 7 students dropped out because of mild sensitivity to the rub
Notes	Risk of bias: high (no description of randomisation; partial reporting of outcomes, numerators and denominators) Note: the authors conclude that addition of the rub led to a 30% to 38% decrease of illness and absenteeism (RR for illness absence incidence 0.69, RR for absence duration 0.71). Very high attrition, unclear randomisation procedure, educational programme and use of placebo hand rub make generalisability of the results debatable. No confidence intervals reported. This study was supported by an Orange County School Nurses Organization Health Promotion Grant. No interests declared.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Quote: "randomised trial", but sequence generation not described
Allocation concealment (selection bias)	Unclear risk	Not described
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "To distinguish content, both the active and placebo formulations were distributed in four color-coded groups of 1oz spritz bottles. The content were and distribution patters were only know to the researchers and were indecipherable by the school staff or students."
Blinding of outcome assessment (detection bias) All outcomes	Low risk	Quote: "Teachers were responsible for recording attendance for each day during the study"
Incomplete outcome data (attrition bias) All outcomes	High risk	Partial reporting of outcomes, numerators and denominators
Selective reporting (reporting bias)	High risk	Poor reporting

Yeung 2011

Study characteristi	cs
Methods	Clustered-RCT of a hand hygiene intervention involving pocket-sized containers of alcohol-based hand rub for the control of infections in long-term care facilities. Staff hand hygiene adherence was directly observed, and residents' infections necessitating hospitalisation were recorded. After a 3-month preintervention period, long-term care facilities (LTCFs) were randomised to receive pocket-sized containers of alcohol-based gel, reminder materials, and education for all HCWs (treatment group) or to re-



Yeung 2011 (Continued)		education and workshops for all HCWs (control group). A 2-week intervention 07) was followed by 7 months of postintervention observations.	
Participants	6 out of 7 community-based, private or semiprivate, residential LTCFs in Hong Kong agreed to participate and were randomised to:		
		o (3 LTCFs, 73 nursing staff and 244 residents analysed); or CFs, 115 nursing staff and 379 residents analysed).	
		s serving an elderly population. All LTCFs were situated in different regions of urban and rural areas. The targets of the intervention were all full- and part-time	
	The LTCFs employed 3	types of HCWs: nurses, nursing assistants, and physiotherapists.	
Interventions		rs of alcohol-based gel, reminder materials, and education (intervention group) ducation and workshop (control group). See Table 1 for details.	
Outcomes	Rates of infection (requ	uiring hospitalisation)	
	Outbreaks		
	Death due to infection		
	Diagnoses of infection coded into 6 categories, all of which were common endemic infections in LTCFs:		
	 pneumonia, urinary tract infection, septicaemia, skin or soft-tissue infection (including cellulitis or pressure sores), gastroenteritis, and fever. 		
	Infections recorded in death certificates were also included, regardless of whether the resident had been hospitalised. The causes of death were categorised as due to infection, not due to infection, or unknown. If the primary or the secondary diagnosis on the death certificate belonged to 1 of the 6 endemic infection categories, the death was coded as due to infection.		
	No safety outcomes reported.		
Notes	University and industry funded.		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	No details provided.	
Allocation concealment (selection bias)	Unclear risk	No details provided.	
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Unblinded study	
Blinding of outcome as-	High risk	Unblinded study	

sessment (detection bias)

All outcomes



Yeung 2011 (Continued)		
Incomplete outcome data (attrition bias) All outcomes	Low risk	No loss to follow-up
Selective reporting (reporting bias)	Unclear risk	No protocol available

Young 2021

Study characteristics		
Methods	Cluster-randomised, controlled trial of daily contact testing in students and staff at secondary sch and colleges in England to show whether daily contact testing increases school attendance and to sess the impact of daily contact testing on SARS-CoV-2 transmission within schools.	
Participants	201 schools, of which 99 were randomly assigned to self-isolation of school-based COVID-19 contacts for 10 days (control) and 102 to voluntary daily lateral flow device (LFD) testing for 7 days with LFD-neative contacts remaining at school (intervention)	
Interventions	All schools in the intervention and control groups followed the national policy of offering twice weekly asymptomatic testing with LFDs. Individuals with positive LFD results were required to self-isolate immediately and requested to obtain a confirmatory PCR test within 2 days. Those with indicator symptoms of possible COVID-19 (new cough, fever, loss or change in taste or smell) were required to self-isolate along with their household and obtain an urgent PCR test. If a student or staff member tested positive by LFD or PCR, close contacts (hereafter referred to as contacts) were identified by schools using national guidelines. Those in close contact with a case less than 48 hours before symptom onset (or a positive test if asymptomatic) were required to self-isolate for 10 days. At schools in the intervention group, contacts were offered daily contact testing as an alternative to self-isolation, provided the contact was school-based (i.e. with a staff member or student), the contact did not have indicator symptoms of COVID-19, and contacts were able to attend for on-site testing at school. See Table 1 for details.	
Outcomes	Laboratory PCR confirmed infections	
	Effectiveness COVID-19-related school absence and symptomatic PCR-confirmed COVID-19.	
	Safety NR	
Notes	The authors conclude that quote: "Daily contact testing of school-based contacts was non-inferio self-isolation for control of COVID-19 transmission, with similar rates of symptomatic infections as students and staff with both approaches."	
	Funding: UK Government Department of Health and Social Care.	
	Declaration of interests: DWE reports lecture fees from Gilead outside the submitted work. VB, RO DC are consultants employed by Department of Health and Social Care as part of Deloitte's broad project work supporting the delivery of NHS Test and Trace. TF reports honoraria from Qatar Nation Research Fund outside the submitted work. All other authors declare no competing interests.	er
	Potential conflicts of interest: all authors report no conflicts of interest relevant to this article.	
Risk of bias		
Bias	Authors' judgement Support for judgement	
Random sequence generation (selection bias)	Low risk Computer random-number generator	



Young 2021 (Continued)		
Allocation concealment (selection bias)	Unclear risk	Insufficient information reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not blinded.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Not blinded.
Incomplete outcome data (attrition bias) All outcomes	High risk	Participant flow diagram reported showing high attrition at different rates in the 2 groups
Selective reporting (reporting bias)	Low risk	Prespecified outcomes reported

Zomer 2015

Study characteristics	
Methods	Cluster-RCT
Participants	71 daycare centres (36 intervention DCCs, and 35 control) in Rotterdam-Rijnmond, Gouda and Leiden in the Netherlands
	Study enrolled 545 children (intervention = 278, control = 267).
	Inclusion/exclusion criteria: children who attended the DCC at least 2 days a week; were aged between 6 months and 3.5 years at start of the trial; intended to attend the DCC throughout the study period; and if their parents consented, were Dutch-speaking, and had access to email or regular post. Children were excluded if they had a chronic illness or medication that predisposed them to infection, a sibling taking part in the trial (i.e. 1 child per family could be included), or if they started attending CCC after the beginning of the trial).
Interventions	4 components:
	1. HH products, paper towel dispensers, soap, alcohol-based hand sanitiser, and hand cream were provided for 6 months.
	2. Training and a booklet outlining the training.
	3. 2 team training sessions aimed at specific HH improvement activities.
	4. Posters and stickers for caregivers and children as reminders.
	See Table 1 for details.
Outcomes	Laboratory: none
	Effectiveness: incidence of respiratory infections in children monitored by parents. The common cold was defined as a blocked or runny nose with at least 1 of the following symptoms: coughing, sneezing, fever, sore throat, or earache.
	Safety: none planned or reported by the investigators
Notes	The period study conducted: September 2011 to April 2012



Zomer 2015 (Continued)

Funding: mixed. The Netherlands Organisation for Health Research and Development (ZonMw). Dispensers and refills were sponsored by SCA Hygiene Products, Sweden.

Declaration of interest: none.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Low risk	Quote: "Stratified randomization is performed by assigning each DCC to one of six strata based on size (i.e. small < 46 children per day versus large ≥ 46 children per day) and geographic location (i.e. highly urban versus urban versus slightly/non-urban). DCCs are assigned to either intervention or control group by means of computer generation with a 1:1 ratio in each of the strata"
Allocation concealment (selection bias)	Unclear risk	No information provided.
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Outcome is subjective.
Blinding of outcome assessment (detection bias) All outcomes	High risk	Symptoms were reported by parents, no validation.
Incomplete outcome data (attrition bias) All outcomes	Low risk	Very few children were excluded or lost to follow-up (reasons for exclusions provided).
Selective reporting (reporting bias)	Low risk	All planned outcomes are reported. However, between published protocol and the paper, secondary outcomes became the primary outcome in the published paper!

AEs: adverse events

AFH: Armed Forces Hospital AGE: acute gastroenteritis

AgNPs: ARGOVIT silver nanoparticles ALRI: acute lower respiratory infection ARI: acute respiratory infection ASR: adverse skin reactions A&E: accident and emergency BIPAP: bilevel positive airway pressure

CCC: childcare centre

CDC: Centers for Disease Control and Prevention

CG: control group

CHG: chlorhexidine gluconate CI: confidence interval

CMF: citric acid: malic acid: sodium lauryl sulphate (a virucidal mixture added to tissue paper)

CoV: coronavirus

cluster-RCT: cluster-randomised controlled trial

CRI: clinical respiratory illness

CXR: chest X-ray DCC: daycare centre EG: experimental group FRI: febrile respiratory illness

FU: follow up GI: gastrointestinal



GTI: gastrointestinal infection GP: general practitioner HCW: healthcare worker HFH: Hanoi French Hospital

HH: hand hygiene HR: high risk

HSG: hand sanitiser group

ICD-9: International Classification of Disease, 9th Revision, Clinical Modification

IgG: immunoglobulin G ICU: intensive care unit ILI: influenza-like illness IQR: interquartile range IRR: incident rate ratio ITT: intention-to-treat

KSA: Kingdom of Saudi Arabia LFD: lateral flow device

LNS: lipid based nutrient supplementation LRTI: lower respiratory tract infection

LTCF: long-term care facility

m: metre

MCU: medical convalescent unit

MDCK: Madin Darby canine kidney cell line

M group: face mask group

MH group: face mask and hand hygiene group

MS: monkey-derived cell line

N/A: not applicable NAT: nucleic acid testing NH: nursing home

NICU: neonatal intensive care unit NOS: Newcastle-Ottawa Scales

NP: non-pharmaceutical NR: not reported

NTS: nasal and throat swab

OR: odds ratio

PCR: polymerase chain reaction PCU: physical conditioning unit POCT: point-of-care testing

PP: per protocol

PPE: personal protective equipment QNAF: Qatar National Research Fund RCT: randomised controlled trial RDS: respiratory distress syndrome

RI: respiratory infection

RIDT: rapid influenza diagnostic test

RNA: ribonucleic acid

RR: risk ratio

rRT-PCR: real-time reverse transcription-polymerase chain reaction

RTI: respiratory tract infection

RT-PCR: reverse-transcriptase polymerase chain reaction

RSV: respiratory syncytial virus

RV: rhinovirus

SAB: surfactant, allantoin, and benzalkonium chloride

SAR: secondary attack rate

SARS: severe acute respiratory syndrome

SCBU: special care baby unit SD: standard deviation SES: electrolysed water

SHEWA-B: Sanitation, Hygiene Education and Water Supply in Bangladesh

SOB: shortness of breath

SOPs: standard operating procedures

S/S: signs/symptoms

SSTI: skin and soft-tissue infection



STH: soil-transmitted helminth SWG: soap and water group

TIDieR: Template for Intervention Description and Replication

UHR-I: ultra high-risk infection UHR-S: ultra high-risk SARS URI: upper respiratory infection URTI: upper respiratory tract infection

WBC: white blood cell

WHO: World Health Organization

WSH: water, sanitation, and handwashing (combined)

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion	
Abou El Hassan 2004	Topic completely extraneous	
Ahmadian 2022	Excluded as study is an experiment that did not measure any of our outcomes of interest.	
Amirav 2005	Randomised controlled trial of aerosol treatment	
Anderson 2004	Mathematical model with interesting discussion of interaction between public health measures	
Anonymous 2002	News item	
Anonymous 2004	News item	
Anonymous 2005a	News item	
Anonymous 2005b	News item	
Anonymous 2005c	News item	
Apisarnthanarak 2009	Intervention bundle not broken down.	
Apisarnthanarak 2010	Participants took antivirals.	
Aragon 2005	Descriptive paper (non-comparative). Has no viral outcomes	
Azor-Martinez 2014	Results reported as respiratory and gastrointestinal infections. No extractable respiratory data	
Barros 1999	Correlational study between incidence of URTI and factors such as overcrowding	
Bauer 2009	Historical comparison with RSV gammaglobulin amongst interventions	
Bell 2004	Has unpublished entry exit screening data and extensive references but no comparative data	
Bellissimo-Rodrigues 2009	Intervention is chlorhexidine.	
Ben-Abraham 2002	Exclude - bacterial illness only	
Black 1981	Diarrhoea only outcome	
Borkow 2010	No human beings involved.	
Bouadma 2010	Hospital-based ventilator routine	



Study	Reason for exclusion
Bowen 2007	Outcomes of composite infections. Respiratory infections are not reported separately.
Breugelmans 2004	Description of risk factors in aircraft
Cai 2009	Compliance study
Cantagalli 2010	Outcome outside inclusion criteria
Carbonell-Estrany 2008	Immunoglobulin intervention and descriptive review
Carter 2002	News item
Castillo-Chavez 2003	Editorial
Cava 2005a	Survey of quarantinees' views
Cava 2005b	Personal experiences of quarantine
CDC 2003a	Case reports
CDC 2003b	No data presented.
Chai 2005	Letter - about MRSA
Chami 2012	Outcomes of composite infections. Respiratory infections are not reported separately.
Chaovavanich 2004	Case report
Chau 2003	No original retrievable data. Mathematical model fitting expected to observed cases with quarantine in the SARS of Hong Kong
Chau 2008	Audit of infection control procedures and compliance with guidelines
Chen 2007	An assessment of the impact of different hand-washing teaching methods. No clinical outcomes
Chen 2022	Not a RCT.
Cheng 2010	Confounded by antiviral use for postexposure prophylaxis
Chia 2005	Knowledge survey
Clynes 2010	Letters
Costa 2021	No clinical outcome assessed
Cowling 2007	Epidemiology, non-comparative, non-interventions study
Cyril Vitug 2021	Is a treatment for COVID-19 infection
Dalakoti 2022	Excluded as study is an experiment that did not measure any of our outcomes of interest.
Daniels 2010	Commentary
Daugherty 2008	No free data presented.



Study	Reason for exclusion	
Davies 1994	Antibody titres as outcomes with so many biases that interpretation of study is problematic	
Day 1993	No acute respiratory infection outcome data	
Day 2006	Mathematical model; no new data	
Dell'Omodarme 2005	Probabilistic and Bayesian mathematical model of screening at entry	
Denbak 2018	Outcomes of composite infections. Respiratory infections are not reported separately.	
Desenclos 2004	Description of transmission	
DiGiovanni 2004	Qualitative study of compliance factors in quarantine	
Doebbeling 1992	RCT respiratory data not present. Only 3 viruses isolated in total with no viral typing available.	
Dwosh 2003	Case series	
Edmonds 2010	Lab study	
Egger 2022	Excluded as study is an experiment that did not measure any of our outcomes of interest.	
Fendler 2002	Cohort study badly biased with differential health profiles and healthcare workers dependency in intervention and control semi-cohorts. No attempt to adjust for confounders was made. No denominators available.	
Ferrer 2021	Is a treatment (not something to interrupt transmission)	
Flint 2003	Description of spread in aircraft and non-comparative data	
Fung 2004	Non-comparative	
Garcia 2010	Commentary	
Gaydos 2001	Editorial linked to Ryan 2001. (Ryan 2001 was an included trial in a previous version of this review (2011). Non-RCTs were removed in this 2020 update).	
Gensini 2004	Interesting historical review	
Gharebaghi 2020	Study on the prevention of ventilator associated pneumonia in mechanical ventilatory patients	
Girou 2002	Non-clinical outcomes	
Giuliano 2021	Outcome is hospital aquired pneumonia which is a syndrome with multiple aetiologies, mainly bacterial and mycotic	
Glass 2006	Mathematical model - no original data presented	
Goel 2007	Non-comparative study	
Gomersall 2006	Non-comparative study	
Gore 2001	Summary of Dyer 2000. (Dyer 2000 was a prospective, cluster open-label cross-over cohort study in cluded in the previous version of this review (2011). Non-RCTs were removed in this 2020 update).	



Study	Reason for exclusion	
Gostin 2003	Not an analytical study	
Gralton 2010	Review	
Guinan 2002	It would appear that 9 classes took part and "acted as their own controls", but it is not clear if there was cross-over of classes or not. In addition, the outcome is combined gastrointestinal/respiratory. The clue lies in the presence of a nested economic analysis which shows considerable savings in time for staff and pupils if the soap is used: in other words this is a (covert) publicity study.	
Gupta 2005	Economic model - no new data	
Gwaltney 1982	No breakdown of cases given by arm.	
Han 2003	Non-comparative	
Hayden 1985	This is an RCT with laboratory-induced colds, small numbers, and uncertain numerators, but almost certainly because of the unique laboratory conditions (placebo tissues not being a placebo at all) of impossible generalisation. It was a pilot to the far bigger trial by Farr 1988a; Farr 1988b.	
Hendley 1988	Inappropriate intervention	
Hens 2009	Model	
Heymann 2009	Already included in review as Heymann 2004. (Heymann 2004 was a controlled before and after study included in the previous version of this review (2011). Non-RCTs were removed in this 2020 update).	
Hilburn 2003	No ARI/viral outcomes (e.g. URTIs)	
Hilmarsson 2007	Animal study	
Hirsch 2006	Study tested pharmacological interventions.	
Ho 2003	Descriptive review	
Hsieh 2007	Mathematical model	
Hugonnet 2007	Letter without any data	
Jiang 2003	Two papers that are probably different versions of the same paper: Jiang SP, Huang LW, Wang JF, Wu W, Yin SM, Chen WX, et al. A study of the architectural factors and the infection rates of health-care workers in isolation units for severe acute respiratory syndrome. Chung-Hua Chieh Ho Hu Hsi Tsa Chih [Chinese Journal of Tuberculosis & Respiratory Diseases]. 26(10):594-7, 2003 Oct	
Johnson 2009	Outcomes are non-clinical.	
Jones 2005	Historical account	
Karakaya 2021	Outcome is ventilator associated pneumonia which is a syndrome with multiple aetiologies, mainly bacterial and mycotic	
Kawyannejad 2020	Trial on mouthwash for VAP patients with no viral infection outcomes	
Kaydos-Daniels 2004	Not an analytical study	
Kelso 2009	Model	



Study	Reason for exclusion
Khaw 2008	Assessing the efficacy of O ₂ delivery
Kilabuko 2007	Aetiological study
Kosugi 2004	Non-comparative study
Lam 2004	Outcomes were generic (infection rates). No laboratory data available for viral diagnosis.
Lange 2004	No data presented.
Larson 2004a	Inappropriate outcomes
Larson 2004b	Inappropriate outcomes
Larson 2005	Cluster-RCT comparing the effects of 2 hand hygiene regimens on infection rates and skin condition and microbial counts of nurses' hands in neonatal intensive care units. Outcomes were generic (e.g. pneumonia and microbial counts of participants' skin). No laboratory data available for viral diagnosis.
Lau 2004	Attitude survey
Lau 2005	Herbal remedy effectiveness assessment
Lee 2005	Descriptive study of risk and protective factors of transmission in households. No assignment took place.
Lee 2010	Cohort study; unclear numbers were vaccinated against influenza
Lennell 2008	Measured absenteeism due to non-specific infection
Lim 2022	Not a RCT.
Lipsitch 2003	Mathematical model fit to evidence
Luckingham 1984	Historical report on Tucson experience during Spanish flu pandemic
Ma 2004	Case-control study of risk factors for SARS
MacIntyre 2010	Commentary on Cowling 2009
Malaczek 2022	Excluded as study is an experiment that did not measure any of our outcomes of interest.
Malone 2009	Model
Marin 1991	Viral resistance study
McSweeny 2007	Historical description
Meister 2022	Excluded as this is a treatment trial (all participants had COVID).
Mielke 2009	Review
Mikolajczyk 2008	No intervention
Mo 2022	Not a RCT.



Study	Reason for exclusion
Monsma 1992	Non-comparative study
Montero-Vilchez 2022	Excluded as study is an experiment that did not measure any of our outcomes of interest.
Munoz-Basagoiti 2022	Excluded as this is a report of another study.
Nandrup-Bus 2009	The trial had only 2 clusters.
Nishiura 2009	Model
O'Callaghan 1993	Letter linked to Isaacs 1991. (Isaacs 1991 was a retrospective and prospective cohort study included in a previous version of this review (2011). Non-RCTs were removed in the 2020 update).
Olsen 2003	Description of transmission
Ooi 2005	Descriptive study, but with interesting organisational chart
Orellano 2010	Confounded by antiviral use
Panchabhai 2009	Pharma intervention
Pang 2004	Descriptive study of Beijing outbreak. Some duplicate data in common with Pang 2003. (Pang 2003 was an eclogical study included in a previous version of this review (2011). Non-RCTs were removed in the 2020 update).
Patel 2012	Although within each district the participating schools and households were randomly selected, the allocation of districts to the intervention and comparison arms was not randomly assigned.
Pittet 2000	Analysis of relationship between hand-washing compliance campaign and nosocomial bacterial infections (e.g. MRSA)
Prasad 2004	Letter about retrospective cohort - behavioural
Rabenau 2005	In vitro test of several disinfectants
Reynolds 2008	Describes the psychological effects of quarantine
Richardson 2010	Non-clinical study
Riley 2003	Mathematical model fit to evidence
Rodriguez 2009	A "reasonable attempt at minimizing bias" (see inclusion criteria) does not include absenteeism
Rosen 2006	Non-specific outcome. Measured absenteeism
Rosenthal 2005	Outcomes were generic (e.g. pneumonia, URTIs). No laboratory data available for viral diagnosis.
Safiulin 1972	Non-comparative set of studies with no clinical outcomes
Sanchez Barrueco 2022	Excluded as this is a treatment trial (all participants had COVID)
Sandrock 2008	Review
Sattar 2000	Experiment assessing virucidal activity of fingertip surface - no clinical outcome data



Study	Reason for exclusion
Schull 2007	Describes the impact of SARS in a Toronto study
Seal 2010	Lab study
Seale 2009	Study looking at whether using respirators in A&E department is feasible
Seneviratne 2021	Not an intervention to reduce transmission and they did not look at ARIs or other clinically relevant outcomes
Sevinc Gul 2022	Excluded as this is a treatment trial (all participants had COVID)
Sizun 1996	This is a review; no original data presented.
Slayton 2016	Compares hand-washing plus (antibacterial) towel versus hand-washing without towel
Stebbins 2009	Attitude survey
Stedman-Smith 2015	Composite outcome. No data on separate respiratory illnesses reported.
Stoner 2007	No study data available.
Stukel 2008	Impact of the SARS disruption on care/mortality for other pathologies (e.g. acute myocardial infarction). There are no interventions, and outcomes are unrelated to acute respiratory infections.
Svoboda 2004	Descriptive study with before-and-after data but shifting denominators
Tracht 2010	Model
Ueno 1990	Experimental study. No clinical intervention
Uhari 1999	No respiratory illness data to be extracted
van der Sande 2008	Laboratory study without any clinical outcomes
Vessey 2007	Composite outcome. No data on separate respiratory illnesses reported.
Viscusi 2009a	Lab study
Viscusi 2009b	Lab study
Wang 2003	Descriptive study
Wang 2005	Case-control study of susceptibility factors
Weber 2004	Editorial linked to Larson 2004a
Wen 2010	Lab study
White 2005	Redundant publication of White 2003. (White 2003 was a prospective, open, cohort study included in a previous version of this review (2011). Non-RCTs were removed in the 2020 update).
Wilczynski 1997	Clinical trial of the effects of breastfeeding
Wilder-Smith 2003	Description of risk factors in aircraft



Study	Reason for exclusion
Wilder-Smith 2005	Descriptive review
Wong 2005	Attitude survey
Yen 2010	Model
Yu 2004	Description of transmission
Zamora 2006	Head-to-head comparison of 2 sets of PPEs with no controls and no clinical outcomes
Zhai 2007	Non-comparative study
Zhao 2003	CCT of SARS treatment

A&E: accident and emergency ARI: acute respiratory infection CCT: controlled clinical trial

MRSA: methicillin-resistant *Staphylococcus aureus*

RCT: randomised controlled trial RSV: respiratory syncytial virus PPE: personal protective equipment SARS: severe acute respiratory syndrome URTI: upper respiratory tract infection VAP: ventilator associated pneumonia

Characteristics of studies awaiting classification [ordered by study ID]

Contreras 2022

Methods	Follow-up of the WASH Benefits Bangladesh cluster-randomised controlled trial. Access to and reported use of latrines was high in both arms, and latrine quality was significantly improved by the intervention, while use of child faeces management tools was low. A random subset of households from the sanitation and control arms was enrolled into a longitudinal substudy, which measured child health with quarterly visits between 1 to 3.5 years after implementation.
Participants	9800 observations on children < 5 years through intention-to-treat analysis using generalised linear models with robust standard errors. 720 households (360 per arm) from the parent trial were enrolled and made 9800 child observations between June 2014 and December 2016.
Interventions	Multicomponent sanitation intervention including periods with differing intensity of behavioural promotion: water, sanitation, hygiene, and nutrition interventions. The sanitation intervention included provision of or upgrades to improved latrines, sani-scoops for faeces removal, children's potties, and in-person behavioural promotion. Promotion was intensive up to 2 years after intervention initiation, decreased in intensity between years 2 to 3, and stopped after 3 years. The study period included approximately 1 year of high-intensity promotion, 1 year of low-intensity promotion, and 6 months with no promotion.
Outcomes	Diarrhoea and ARI, at 1 to 2 years after intervention implementation to 3.5 years (follow-up). Outcomes were caregiver-reported and there were limited data collected after promotion ceased.
Notes	Trial registration: ClinicalTrials.gov; NCT01590095; https://clinicaltrials.gov/ct2/show/NCT01590095



Croke 2022	
Methods	Cluster-randomised trial assessing the effect of a national water, sanitation, and hygiene program on adherence with COVID-19 policies in Congo. The trial is a follow-up of the Villages et Ecoles Assainis programme which was running prior to the COVID-19 pandemic.
Participants	332 communities were randomly assigned to the Villages et Ecoles Assainis program or control. (590/1312; 45%) individuals who owned phones were surveyed by phone 3 times between May 2020 to August 2021.
Interventions	Large-scale water and sanitation programme not described in detail.
Outcomes	Primary outcomes were COVID symptoms, non- COVID illness symptoms, child health, psychological well-being, and vaccine acceptance.
	Secondary outcomes included COVID-19 preventive behaviour and knowledge, and perceptions of governmental performance, including COVID response. All outcomes were self-reported.
	COVID symptoms were defined as the number of household members in the past week with fever, dry cough, difficulty breathing/shortness of breath, or fatigue, while non-COVID illness variable was defined as the number of sick household members in the last 7 days (excluding those with COVID symptoms). The child health index was created using the proportion of children under 5 with fever/cough/diarrhoea in the last 2 weeks. The mental health index is a summary index of scores from answers to questions.
Notes	Cannot find NCT and unclear funders although acknowledgments list a potential load of funders. Probably public.

Delaguerre 2022

Methods	Prospective, open-label, non-inferiority randomised (2:1), controlled trial
Participants	Study included healthy individuals aged 18 to 45 years, with negative RADT test 3 days prior to concert event, with no risk factors and not living with someone with risk factors, and residing in Paris.
	Study excluded people with positive RADT test within 3 days before the gathering. People with clinical signs suggestive of an infectious respiratory disease, or with risk factor for severe COVID-19, or living with someone with risk factors for severe COVID-19. Persons not covered by French National Health Insurance or who cannot stand for the duration of the experiment (about 5 hours from entry line to exit) were excluded. Person under legal guardianship, pregnant woman or woman orally declaring non-use of effective contraception and breastfeeding woman were also excluded.
Interventions	Participants were randomly assigned to:
	 medical face mask wearing during an indoor concert event, or not attending.
	Both groups had RADT test 3 days before the event Saliva samples for RT-PCR were collected from both groups on D0 and D7 using self-saliva-collection kits
Outcomes	Primary outcome:
	1. the number of SARS-CoV-2-positive RT-PCR tests on self-collected saliva at day 7.
	Secondary outcomes:
	1. the conversion rate of salivary carriage between the day 0 and day 7 visits;



Delaguerre 2022 (Continued)	the percentages of adequately masked (nose and mouth covered) faces over the total 4-hour period gathering.
Notes	1. French Ministry of Health. 2. ITT and DD analysis were used. Soveral imputation for missing data.
	2. ITT and PP analysis were used. Several imputation for missing data.
	3. It is not clear if participants had COVID-19 in the past (in the table with baseline characteristics it is reported quote: ""declared Covid-19 history": what does it mean?
	4. Surgical masks were worn also by all attendees, regardless of study participation?
	5. What is the intervention? Combined screening test + surgical mask?

Loeb 2022

Methods	Multicentre, randomised, non-inferiority trial
Participants	1009 healthcare workers who provided direct care to patients with suspected or confirmed COV-ID-19.
	Conducted in 29 healthcare facilities in Canada, Israel, Pakistan, and Egypt from 4 May 2020 to 29 March 2022.
Interventions	Use of medical masks versus fit-tested N95 respirators for 10 weeks, plus universal masking, which was the policy implemented at each site.
Outcomes	The primary outcome was confirmed COVID-19 on reverse transcriptase polymerase chain reaction (RT-PCR) test.
Notes	Financial support was given by the Canadian Institutes of Health Research, World Health Organization, and Juravinski Research Institute.
	Disclosures can be viewed at www.acponline.org/authors/icmje/ConflictOfInterestForms.do?msNum=M22-1966

Varela 2022

Methods	Open-label non-inferiority randomised controlled trial
Participants	Study was conducted in Colombia
	Inclusion criteria:
	people aged ≥ 18 years of both genders and who:
	(a) lived in a geographic area with active COVID-19 transmission and in areas with medium, medium-high, and high vulnerability index; and
	(b) worked outside their homes for at least 2 days during the last week.
	Exclusion criteria:
	retirement, unemployment, home-based working, history of laboratory-confirmed COVID-19, working in health care, and daily N95 mask or face shield use. In addition, during follow-up if participants reported an occupation change from work outside the home to home-based work, or became unemployed
Interventions	1. Intervention group (IG): instructed to wear closed face shields with surgical face masks
	Active control group (ACG): instructed to wear only surgical face mask



Varela 2022 (Continued)	
(,	PPE was sent to their home address for each day of participation
	All participants received a follow-up twice a week by phone
	All participants received recorded educational intervention via email or phone that provided recommendations about COVID-19 prevention measures, guidance to ensure adherence, and appropriate handling of the assigned PPE.
	Weekly short questionnaire was performed on days 7, 14, and 21 to evaluate health status SARS-CoV-2 symptoms, PPE use, and adherence.
Outcomes	Primary outcome was the composite result of positive RT-PCR or seroconversion during follow-up
	Secondary outcomes including PPE use and adherence
Notes	 Study was nested within an observational study (CoVIDA project). Funding was provided by donors administered by the philanthropy department at the Universidad de Los Andes, external financing from the United Nations Development Programme (UNDP), and donations of diagnostic material from the Engineering Services Laboratory S.A.S. (LABSERVING S.A.S. Colombia). Funders had no input on the study at any stage. Provided analysis as ITT and PP. Missing data were imputed with negative results.

ARI: acute respiratory infection

h: hours

ITT: intention-to-treat NCT: trial register number

PPE: personal protective equipment

PP: per protocol

RADT: rapid antigen detection test

 $\hbox{RT-PCR: reverse-transcript ase polymerase chain reaction}\\$

Characteristics of ongoing studies [ordered by study ID]

Brass 2021

Study name	Prevention of SARS-CoV-2 (COVID-19) transmission in residential aged care using ultraviolet light (PETRA)
Methods	A multicentre, 2-arm double-cross-over, randomised controlled trial will be conducted to determine the efficacy of GUV devices to reduce respiratory viral transmission in RACF, as an adjunct to existing infection control measures. The study will be conducted in partnership with 3 aged care providers in metropolitan and regional South Australia. RACF will be separated into paired within-site zones, then randomised to intervention order (GUV or control). The initial 6-week period will be followed by a 2-week washout before cross-over to the second 6-week period. After accounting for estimated within-zone and within-facility correlations of infection, and baseline infection rates (10 per 100 person-days), a sample size of n = 8 zones (n = 40 residents/zone) will provide 89% power to detect a 50% reduction in symptomatic infection rate.
Participants	RACF within metropolitan and regional South Australia will be considered for recruitment if they possess the ability to sub-divide communal living areas into discrete areas that enable a concurrent comparison of interventions, with the facility cohorts otherwise subject to the same facility practices (e.g. environmental cleaning, staffing, and social distancing).
Interventions	The intervention will involve the commercially available Laftech GUV appliances: UV-FLOW-C wall-and ceiling-mounted system, UV-FAN-XS wall-mounted air purifier, and UV-FAN M2/95HP air purification device (LAF Technologies, Melbourne, Australia).



Brass 2021 (Continued)

Outcomes

The primary outcome will be the incidence rate ratio of combined symptomatic respiratory infections for intervention versus control. Secondary outcomes include incidence rates of hospitalisation for complications associated with respiratory infection; respiratory virus detection in facility air and fomite samples; rates of laboratory-confirmed respiratory illnesses and genomic characteristics.

Starting date

Contact information

Andrew P. Shoubridge

- The South Australian Health and Medical Research Institute (SAHMRI), Adelaide, SA, Australia
- The Microbiome and Host Health Programme, College of Medicine and Public Health, Flinders University, Bedford Park, SA, Australia

Notes

NCT03454009

Study name

Appropriate time-interval application of alcohol hand gel on reducing influenza-like illness amongst preschool children: a randomised, controlled trial

Methods

This is a comprehensive randomised cluster hand-hygiene improvement intervention to reduce self-reported ARI/ILI and GI illness, absenteeism, presenteeism and related behavioural and attitudinal change over a 90-day trial. The intervention group will receive hand hygiene supplies and a variety of educational materials, including environmental posters in common areas. The control group will perform their usual hygiene activities and will not receive an intervention.

Identical weekly surveys will be administered to the intervention and control groups to measure self-reported illness, absenteeism, presenteeism, along with behaviour and attitudes measured at specified intervals during the study. The intervention and control groups were randomised by work floors before the onset of the enrolment period. It is hypothesised that employees in the intervention group will experience reduced self-reported illness, absenteeism, and presenteeism along with improved protective hygiene behaviours and related attitudes, relative to those in the control group over the 90-day trial.

Participants

Inclusion criteria

- 1. At least 18 years of age or older
- 2. No known allergies to alcohol or surface disinfecting wipes
- 3. Works at least 30% of office hours at the study host site
- 4. Consent to receiving emails from Kent State University

Exclusion criteria

- 1. Under 18 years of age
- 2. Known allergies to alcohol or surface disinfecting wipes
- 3. Works less than 30% of office hours at the study host site
- 4. Does not consent to receiving emails from Kent State University

Interventions

The intervention group will receive hand hygiene supplies and a variety of educational materials, including environmental posters in common areas. The control group will perform their usual hygiene activities and will not receive an intervention.

Outcomes

Self-reported ARI/ILI and GI illness, absenteeism, presenteeism and related behavioural and attitudinal change over a 90-day trial



NCT03454009 (Continued)							
Starting date	5 February 2018						
Contact information	Maggie Stedman-Smith, PhD, Kent State University College of Public Health						
Notes	Recruitment completed. Last update in ClinicalTrials.gov was 1 May 2019. NCT03454009						
NCT04267952							
Study name	Hand hygiene intervention program on primary school students' health outcomes and absenteeism in school						
Methods	Study Type: interventional (clinical trial)						
	Estimated enrolment: 200 participants						
	Allocation: randomised						
	Intervention model: parallel assignment						
	Masking: single (participant)						
	Masking description: participation will not know whether they are in the experimental or control group						
Participants	Inclusion criteria: primary school student (especially third- and fourth-class student)						
	Exclusion criteria: people with chronic disease						
Interventions	Experimental: first group						
	Hand hygiene intervention programme prepared by using planned behaviour theory will be applied to the students in this group.						
	Active comparator: second group						
	Students in this group will be given classic hand hygiene training.						
Outcomes	Primary outcome measure: children with symptoms of infection will be referred to the family physician to have a rapid antigen test and to report the result to the researcher.						
	10 identified upper respiratory tract symptoms (fever, sore throat, runny nose, etc.) will be recorded weekly by family of children. The researcher will receive symptom information from the family via weekly SMS.						
	The number of days the child does not attend school due to illness and the percentage of absenteeism						
	 Group A streptococcal infections in rapid antigen test (time frame: total 20 weeks) Incidence of symptoms of acute upper respiratory tract infection (time frame: total 20 weeks) School absenteeism (time frame: total 20 weeks) 						
	Secondary outcome measures: Glogerm gel applied hands will shine areas containing micro-organisms. Contamination rate will be calculated by taking a photo of the hands and performing brightness analysis in Adobe Photoshop program.						
	 Pollution rate of hands (time frame: from date of randomisation until the date of first documented progression assessed up to 7 months) 						
Starting date	9 September 2019						



NCT04267952 (Continued) Contact information	Contact: Uyanık +905068949969; gulcinyelten@hotmail.com						
Notes	Recruitment is ongoing. Last update in ClinicalTrials.gov was 13 February 2020. NCT04267952						
NCT04471766	5 1 1 1 1 1 1 1 1 1						
Study name	Evaluation of locally produced cloth face mask on COVID-19 and respiratory illnesses prevention at the community level - a cluster-RCT						
Methods	Study type: interventional (clinical trial)						
	Estimated enrolment: 66,000 participants						
	Allocation: randomised						
	Intervention model: parallel assignment						
	Masking: single (outcomes assessor)						
	Primary purpose: prevention						
Participants	Ages eligible for study: 10 years and older (child, adult, older adult)						
	Sexes eligible for study: all						
	Accepts healthy volunteers: no						
	Criteria						
	Inclusion criteria:						
	 Household resident Age 10 years and older 						
	Exclusion criteria:						
	1. Refusal to participate						
Interventions	Experimental: certified cloth face mask plus preventive information						
	Active comparator: information on COVID-19 prevention						
Outcomes	Self-reported main symptoms of COVID-19 (3 or more of fever, cough, fatigue, shortness of breath, loss of smell/taste)						
	Consultation for COVID-19 like illness or reported positive test, or both						
	Self reported COVID-19 like illness plus hospitalisation or death						
	Any death during the follow-up period:						
	 Reported COVID-19 like illness (time frame: 4 months' follow-up) Consultation (time frame: 4 months' follow-up) 						
	3. Severe illness (time frame: 4 months' follow-up)4. Mortality (time frame: 4 months' follow-up)						
Starting date	Estimated study start date: July 2020						
Contact information	Amabelia Rodrigues, PhD, 00245966078659; a.rodrigues@bandim.org						
	Amabena Nourigues, i fib, 00275300010033, anourigues@banamin.org						



NCT04471766 (Continued)

Notes

The number of cases of COVID-19 is still increasing, and transmission of SARS-CoV-2 seems to occur mainly through person-to-person transmission through respiratory droplets, indirect contact with infected people and surfaces. The use of face masks is recommended as a public health measure, but in many settings only domestic cloth made masks are available to the majority of the people. However, masks can be of different quality, and very little is known about the utility of cloth face masks at the community level.

In Bandim Health Project's Health and Demographic Surveillance System we evaluated the effect of providing locally produced cloth face masks on the severity of COVID-19 like illness and mortality in an urban population. The locally produced cloth mask is made according to a laboratory-certified model and was provided to the intervention group alongside information of how the risk of transmission can be reduced. The control group received information alone.

Follow-up will be implemented through telephone calls and post epidemic home visits.

ARI: acute respiratory tract infections

GUV: germicidal ultraviolet ILI: influenza-like illness GI: gastrointestinal n: number

RACF: residential aged care facilities RCT: randomised controlled trial

SARS: severe acute respiratory syndrome

DATA AND ANALYSES

Comparison 1. Randomised trials: medical/surgical masks versus no masks

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1.1 Viral illness	10		Risk Ratio (IV, Random, 95% CI)	Subtotals only
1.1.1 Influenza/COVID-like illness	9	276917	Risk Ratio (IV, Random, 95% CI)	0.95 [0.84, 1.09]
1.1.2 Laboratory-confirmed influen- za or SARS-cov-2	6	13919	Risk Ratio (IV, Random, 95% CI)	1.01 [0.72, 1.42]
1.1.3 Laboratory-confirmed other respiratory viruses	1	4862	Risk Ratio (IV, Random, 95% CI)	0.58 [0.25, 1.31]



(2) SARS-cov-2

Analysis 1.1. Comparison 1: Randomised trials: medical/surgical masks versus no masks, Outcome 1: Viral illness

		N	/ledical/surgical masks	No masks		Risk Ratio	Risk Ratio
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
1.1.1 Influenza/COVII	D-like illness						
Abaluck 2022 (1)	-0.135	0.036	111525	155268	41.4%	0.87 [0.81, 0.94]	•
Aiello 2012	0.095	0.115	392	370	19.8%	1.10 [0.88, 1.38]	-
Alfelali 2020	0.095	0.105	3864	3823	21.9%	1.10 [0.90, 1.35]	-
Barasheed 2014	-0.55	0.3	75	89	4.6%	0.58 [0.32, 1.04]	
Canini 2010	0.025	0.342	148	158	3.6%	1.03 [0.52, 2.00]	
Cowling 2008	-0.128	0.483	61	205	1.9%	0.88 [0.34, 2.27]	
MacIntyre 2009	0.1	0.28	186	100	5.2%	1.11 [0.64, 1.91]	
MacIntyre 2016	-1.139	1.16	302	295	0.3%	0.32 [0.03, 3.11]	•
Suess 2012	-0.494	0.571	26	30	1.4%	0.61 [0.20, 1.87]	
Subtotal (95% CI)			116579	160338	100.0%	0.95 [0.84, 1.09]	
Heterogeneity: Tau ² = 0	0.01; Chi ² = 11.44, df =	= 8 (P = 0.18	I_{1}); $I_{2} = 30\%$				Y
Test for overall effect: 2	Z = 0.71 (P = 0.48)						
1.1.2 Laboratory-conf	irmed influenza or S	ARS-cov-2					
Aiello 2012	-0.083	0.223	392	370	25.9%	0.92 [0.59 , 1.42]	
Alfelali 2020	0.34	0.215	3864	3823	26.7%	1.40 [0.92, 2.14]	
Bundgaard 2021 (2)	-0.2	0.208	2392	2470	27.4%	0.82 [0.54, 1.23]	
Cowling 2008	0.148	0.674	61	205	5.8%	1.16 [0.31 , 4.34]	
MacIntyre 2009	0.92	0.6225	186	100	6.6%	2.51 [0.74, 8.50]	 -
Suess 2012	-0.942	0.57	26	30	7.7%	0.39 [0.13, 1.19]	
Subtotal (95% CI)			6921	6998	100.0%	1.01 [0.72, 1.42]	•
Heterogeneity: Tau ² = 0	0.07; Chi ² = 8.52, df =	5 (P = 0.13)	; I ² = 41%				Ť
Test for overall effect: 2	Z = 0.07 (P = 0.95)						
1.1.3 Laboratory-conf	irmed other respirat	ory viruses					
Bundgaard 2021	-0.55	0.42	2392	2470	100.0%	0.58 [0.25 , 1.31]	
Subtotal (95% CI)			2392	2470	100.0%	0.58 [0.25, 1.31]	
Heterogeneity: Not app	licable						~
Test for overall effect: 2	Z = 1.31 (P = 0.19)						
						C	0.05 0.2 1 5
Footnotes							ll/surgical masks Favours no n
(1) Covid-like-illness							

Comparison 2. Randomised trials: N95 respirators compared to medical/surgical masks

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.1 Viral illness	5		Risk Ratio (IV, Random, 95% CI)	Subtotals only
2.1.1 Clinical respiratory illness	3	7799	Risk Ratio (IV, Random, 95% CI)	0.70 [0.45, 1.10]
2.1.2 Influenza-like illness	5	8407	Risk Ratio (IV, Random, 95% CI)	0.82 [0.66, 1.03]
2.1.3 Laboratory-confirmed in- fluenza	5	8407	Risk Ratio (IV, Random, 95% CI)	1.10 [0.90, 1.34]
2.2 Viral illness in healthcare workers	4		Risk Ratio (IV, Random, 95% CI)	Subtotals only
2.2.1 Clinical respiratory illness	3	7799	Risk Ratio (IV, Random, 95% CI)	0.70 [0.45, 1.10]
2.2.2 Influenza-like illness	4	8221	Risk Ratio (IV, Random, 95% CI)	0.81 [0.59, 1.11]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.2.3 Laboratory-confirmed in- fluenza	4	8221	Risk Ratio (IV, Random, 95% CI)	1.05 [0.79, 1.40]

Analysis 2.1. Comparison 2: Randomised trials: N95 respirators compared to medical/surgical masks, Outcome 1: Viral illness

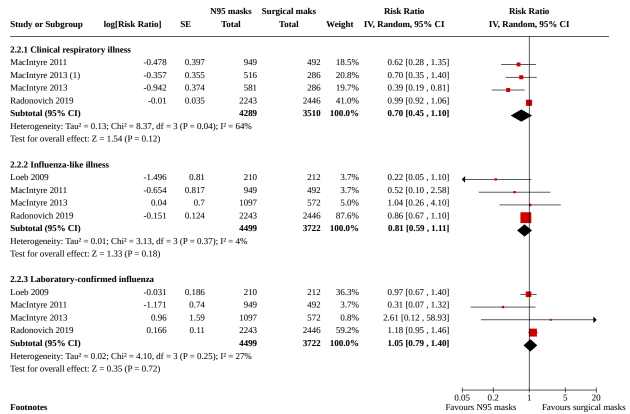
Study or Subgroup	log[Risk Ratio]	SE	N95 respirators Total	Medical/surgical masks Total	Weight	Risk Ratio IV, Random, 95% CI	Risk Ratio IV, Random, 95% CI
2.1.1 Clinical respirate	ory illness						
MacIntyre 2011	-0.478	0.397	949	492	18.5%	0.62 [0.28, 1.35]	
MacIntyre 2013 (1)	-0.357	0.355	516	286	20.8%	0.70 [0.35 , 1.40]	
MacIntyre 2013	-0.942	0.374	581	286	19.7%	0.39 [0.19, 0.81]	
Radonovich 2019	-0.01	0.035	2243	2446	41.0%	0.99 [0.92, 1.06]	•
Subtotal (95% CI)			4289	3510	100.0%	0.70 [0.45, 1.10]	
Heterogeneity: Tau ² = 0	0.13; Chi ² = 8.37, df =	3(P = 0.0	4); I ² = 64%				•
Test for overall effect: 2	Z = 1.54 (P = 0.12)						
2.1.2 Influenza-like ill	ness						
Loeb 2009	-1.496	0.81	210	212	2.0%	0.22 [0.05, 1.10]	—
MacIntyre 2009	-0.306	0.45	92	94	6.6%	0.74 [0.30 , 1.78]	<u> </u>
MacIntyre 2011	-0.654	0.817	949	492	2.0%	0.52 [0.10, 2.58]	
MacIntyre 2013	0.04	0.7	1097	572	2.7%	1.04 [0.26, 4.10]	
Radonovich 2019	-0.151	0.124	2243	2446	86.7%	0.86 [0.67, 1.10]	=
Subtotal (95% CI)			4591	3816	100.0%	0.82 [0.66, 1.03]	⊸
Heterogeneity: Tau ² = 0	0.00; Chi ² = 3.19, df =	4 (P = 0.5)	(3); I ² = 0%				•
Test for overall effect: 2	Z = 1.68 (P = 0.09)						
2.1.3 Laboratory-conf	irmed influenza						
Loeb 2009	-0.031	0.186	210	212	27.7%	0.97 [0.67, 1.40]	<u> </u>
MacIntyre 2009 (2)	0.31	0.94	92	94	1.2%	1.36 [0.22, 8.61]	
MacIntyre 2011	-1.171	0.74	949	492	1.9%	0.31 [0.07, 1.32]	
MacIntyre 2013	0.96	1.59	1097	572	0.4%	2.61 [0.12, 58.93]	-
Radonovich 2019	0.166	0.11	2243	2446	68.8%	1.18 [0.95, 1.46]	=
Subtotal (95% CI)			4591	3816	100.0%	1.10 [0.90, 1.34]	•
Heterogeneity: Tau ² = 0	0.00; Chi ² = 4.15, df =	4 (P = 0.3)	9); I ² = 4%				T
Test for overall effect: 2	Z = 0.89 (P = 0.37)						
							0.1 0.2 0.5 1 2 5 10
Footnotes						Favou	rs N95 respirators Favours medical/surgica

 $^{(1)\} MacIntyre\ 2013\ includes\ 2\ comparisons:\ N95\ vs\ surgical\ masks\ and\ targeted\ N95\ vs\ surgical\ masks$

⁽²⁾ MacIntyre 2009 reported on outcome laboratory confirmed infections



Analysis 2.2. Comparison 2: Randomised trials: N95 respirators compared to medical/surgical masks, Outcome 2: Viral illness in healthcare workers



 $(1)\ MacIntyre\ 2013\ includes\ 2\ comparisons:\ N95\ vs\ surgical\ masks\ and\ targeted\ N95\ vs\ surgical\ masks$

Comparison 3. Randomised trials: hand hygiene compared to control

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3.1 Viral illness	19		Risk Ratio (IV, Random, 95% CI)	Subtotals only
3.1.1 Acute respiratory illness	9	52105	Risk Ratio (IV, Random, 95% CI)	0.86 [0.81, 0.90]
3.1.2 Influenza-like illness	11	34503	Risk Ratio (IV, Random, 95% CI)	0.94 [0.81, 1.09]
3.1.3 Laboratory-confirmed influenza	8	8332	Risk Ratio (IV, Random, 95% CI)	0.91 [0.63, 1.30]
3.2 ARI or ILI or influenza (including outcome with most events from each study)	19	71210	Risk Ratio (IV, Random, 95% CI)	0.89 [0.83, 0.94]
3.3 Influenza or ILI: sensitivity analysis including outcomes with the most precise and unequivocal definitions	12	28205	Risk Ratio (IV, Random, 95% CI)	0.88 [0.77, 1.02]



Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
3.4 ARI or ILI or influenza: subgroup analysis	19	71210	Risk Ratio (IV, Random, 95% CI)	0.89 [0.83, 0.94]
3.4.1 Children	11	29259	Risk Ratio (IV, Random, 95% CI)	0.91 [0.84, 0.98]
3.4.2 Adults	8	41951	Risk Ratio (IV, Random, 95% CI)	0.84 [0.78, 0.91]
3.5 Absenteeism	3	3150	Risk Ratio (IV, Random, 95% CI)	0.64 [0.58, 0.71]



Analysis 3.1. Comparison 3: Randomised trials: hand hygiene compared to control, Outcome 1: Viral illness

			Hand hygiene	Control		Risk Ratio	Risk Ratio
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
3.1.1 Acute respiratory	illness						
Ashraf 2020	-0.39	0.135	588	1123	3.3%	0.68 [0.52, 0.88]	
Azor-Martinez 2018 (1)	-0.261	0.086	339	149	6.7%	0.77 [0.65, 0.91]	
Azor-Martinez 2018	-0.062	0.086	274	149	6.7%	0.94 [0.79, 1.11]	
Correa 2012	-0.223	0.084	794	933	6.9%	0.80 [0.68, 0.94]	
Larson 2010	-0.199	0.134	946	904	3.3%	0.82 [0.63, 1.07]	
Little 2015	-0.151	0.02	8241	8667	20.5%	0.86 [0.83, 0.89]	
Millar 2016	-0.198	0.016	10000	10000	21.4%	0.82 [0.80, 0.85]	
Nicholson 2014	-0.163	0.05	847	833	12.6%	0.85 [0.77, 0.94]	•
Sandora 2005	-0.03	0.15	602	451	2.7%	0.97 [0.72, 1.30]	
Swarthout 2020	-0.03	0.037	1496	4769	15.9%	0.97 [0.90, 1.04]	_
Subtotal (95% CI)			24127	27978	100.0%	0.86 [0.81, 0.90]	A
Heterogeneity: $Tau^2 = 0$.	.00; Chi ² = 24.86, df =	= 9 (P = 0)	.003); I ² = 64%				•
Test for overall effect: Z	L = 5.93 (P < 0.00001)	,	•				
3.1.2 Influenza-like illn	1000						
3.1.2 innuenza-like ilin Biswas 2019	-0.223	0.249	5077	5778	6.2%	0.80 [0.49 , 1.30]	
Cowling 2008	-0.151	0.408	84		2.8%	0.86 [0.39 , 1.91]	
o .							
Cowling 2009 Hubner 2010	-0.083 -1.05	0.243 0.36	257 64	279 65	6.4% 3.5%	0.92 [0.57 , 1.48]	
						0.35 [0.17, 0.71]	
Larson 2010	0.271	0.363	946		3.5%	1.31 [0.64 , 2.67]	
Little 2015	-0.223	0.07	8241	8667	17.0%	0.80 [0.70 , 0.92]	-
Ram 2015	0.215	0.149	193		11.1%	. , ,	 •
Roberts 2000	-0.051	0.03	299		19.4%	0.95 [0.90 , 1.01]	•
Simmerman 2011	0.737	0.263	292		5.7%	2.09 [1.25 , 3.50]	
Teesing 2021	-0.67	0.248	976		6.2%	0.51 [0.31 , 0.83]	
Zomer 2015	0.068	0.052	278	267	18.2%	1.07 [0.97 , 1.19]	
Subtotal (95% CI)		40.00	16707	17796	100.0%	0.94 [0.81, 1.09]	•
Heterogeneity: Tau ² = 0.		= 10 (P <	0.0001); $I^2 = 74\%$				
Test for overall effect: Z	E = 0.83 (P = 0.41)						
3.1.3 Laboratory-confi	rmed influenza						
Biswas 2019	-0.693	0.24	508	689	19.8%	0.50 [0.31, 0.80]	
Cowling 2008	0.07	0.671	84	205	6.0%	1.07 [0.29 , 4.00]	
Cowling 2009	-0.562	0.39	257	279	12.7%	0.57 [0.27 , 1.22]	
Hubner 2010	0.02	0.834	64	65	4.2%	1.02 [0.20 , 5.23]	
Larson 2010	0.648	0.504	946	904	9.2%	1.91 [0.71, 5.13]	
Ram 2015	0.875	0.644	193	184	6.4%	2.40 [0.68 , 8.48]	-
Simmerman 2011	0.182	0.23	292	302	20.4%	1.20 [0.76 , 1.88]	
Stebbins 2011	-0.211	0.212	1695	1665	21.4%	0.81 [0.53 , 1.23]	
Subtotal (95% CI)			4039	4293	100.0%	0.91 [0.63, 1.30]	
Heterogeneity: Tau ² = 0.	.11; Chi ² = 13.58, df =	= 7 (P = 0)	.06); I ² = 48%				\neg
Test for overall effect: Z	L = 0.53 (P = 0.60)						
							0.2 0.5 1 2
							0.2 0.5 1 2

 $(1) Azor\ 2018\ included\ 2\ hand-washing\ groups: one\ using\ soap\ and\ water\ (RR\ 0.94)\ and\ the\ other\ using\ hand\ sanitizer\ (RR\ 0.77)$



Analysis 3.2. Comparison 3: Randomised trials: hand hygiene compared to control, Outcome 2: ARI or ILI or influenza (including outcome with most events from each study)

			Hand hygiene	Control		Risk Ratio	Risk Ratio
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Ashraf 2020	-0.39	0.135	588	1123	3.7%	0.68 [0.52 , 0.88]	
Azor-Martinez 2018 (1)	-0.062	0.086	274	149	6.1%	0.94 [0.79 , 1.11]	_ -
Azor-Martinez 2018	-0.261	0.086	339	149	6.1%	0.77 [0.65, 0.91]	
Biswas 2019	-0.223	0.249	5077	5778	1.4%	0.80 [0.49 , 1.30]	
Correa 2012	-0.223	0.084	794	933	6.3%	0.80 [0.68, 0.94]	
Cowling 2008	-0.151	0.408	84	205	0.6%	0.86 [0.39 , 1.91]	
Cowling 2009	-0.083	0.243	257	279	1.5%	0.92 [0.57 , 1.48]	
Hubner 2010	-1.05	0.36	64	65	0.7%	0.35 [0.17, 0.71]	
Larson 2010	-0.199	0.134	946	904	3.7%	0.82 [0.63, 1.07]	
Little 2015	-0.151	0.02	8241	8667	10.8%	0.86 [0.83, 0.89]	
Millar 2016	-0.198	0.016	10000	10000	11.0%	0.82 [0.80, 0.85]	•
Nicholson 2014	-0.163	0.05	847	833	8.8%	0.85 [0.77, 0.94]	-
Ram 2015	0.215	0.149	193	184	3.2%	1.24 [0.93, 1.66]	 -
Roberts 2000	-0.051	0.03	299	259	10.2%	0.95 [0.90 , 1.01]	-
Sandora 2005	-0.03	0.15	602	451	3.2%	0.97 [0.72 , 1.30]	
Simmerman 2011	0.737	0.263	292	302	1.3%	2.09 [1.25, 3.50]	
Stebbins 2011	-0.211	0.212	1695	1665	1.8%	0.81 [0.53 , 1.23]	
Swarthout 2020	-0.03	0.037	1496	4769	9.8%	0.97 [0.90 , 1.04]	-
Teesing 2021	-0.67	0.248	976	886	1.4%	0.51 [0.31, 0.83]	
Zomer 2015	0.068	0.052	278	267	8.6%	1.07 [0.97 , 1.19]	•
Total (95% CI)			33342	37868	100.0%	0.89 [0.83, 0.94]	•
Heterogeneity: Tau ² = 0.	01; Chi ² = 83.20, df =	= 19 (P <	0.00001); $I^2 = 779$	%			· · · · · · · · · · · · · · · · · · ·
Test for overall effect: Z	= 3.83 (P = 0.0001)						0.2 0.5 1 2 5
Test for subgroup differe	nces: Not applicable					Favo	ours hand hygiene Favours control

Footnotes

(1) Azor 2018 included 2 treatment groups: soap and water (RR 0.94); and hand sanitizer (RR 0.77)

Analysis 3.3. Comparison 3: Randomised trials: hand hygiene compared to control, Outcome 3: Influenza or ILI: sensitivity analysis including outcomes with the most precise and unequivocal definitions

Study or Subgroup	log[Risk Ratio]	SE	Hand hygiene Total	Control Total	Weight	Risk Ratio IV, Random, 95% CI	Risk Ratio IV, Random, 95% CI
Biswas 2019	-0.693	0.24	508	689	6.6%	0.50 [0.31 , 0.80]	
Cowling 2008	0.07	0.671	84	205	1.1%	1.07 [0.29, 4.00]	
Cowling 2009	-0.562	0.39	257	279	3.0%	0.57 [0.27 , 1.22]	
Hubner 2010	0.02	0.834	64	65	0.7%	1.02 [0.20, 5.23]	
Larson 2010	0.648	0.504	946	904	1.9%	1.91 [0.71, 5.13]	
Little 2015	-0.223	0.07	8241	8667	19.7%	0.80 [0.70, 0.92]	-
Ram 2015	0.875	0.644	193	184	1.2%	2.40 [0.68, 8.48]	
Roberts 2000	-0.051	0.03	299	259	23.3%	0.95 [0.90, 1.01]	,
Simmerman 2011	0.182	0.23	292	302	7.0%	1.20 [0.76, 1.88]	
Stebbins 2011	-0.211	0.212	1695	1665	7.8%	0.81 [0.53, 1.23]	
Teesing 2021	-0.67	0.248	976	886	6.3%	0.51 [0.31, 0.83]	
Zomer 2015	0.068	0.052	278	267	21.5%	1.07 [0.97 , 1.19]	•
Total (95% CI)			13833	14372	100.0%	0.88 [0.77 , 1.02]	•
Heterogeneity: Tau ² = 0.	.02; Chi ² = 31.95, df =	= 11 (P =	0.0008); I ² = 66%				•
Test for overall effect: Z	L = 1.75 (P = 0.08)						0.2 0.5 1 2 5
Test for subgroup differe	ences: Not applicable					Favou	ars hand hygiene Favours control



Analysis 3.4. Comparison 3: Randomised trials: hand hygiene compared to control, Outcome 4: ARI or ILI or influenza: subgroup analysis

Study or Subgroup	log[Risk Ratio]	SE	Hand hygiene Total	Control Total	Weight	Risk Ratio IV, Random, 95% CI	Risk Ratio IV, Random, 95% CI
3.4.1 Children							
Ashraf 2020	-0.39	0.135	588	1123	3.7%	0.68 [0.52 , 0.88]	
Azor-Martinez 2018	-0.062	0.086	274	149	6.1%	0.94 [0.79 , 1.11]	
Azor-Martinez 2018 (1)	-0.261	0.086	339	149	6.1%	0.77 [0.65, 0.91]	
Biswas 2019	-0.223	0.249	5077	5778	1.4%	0.80 [0.49 , 1.30]	
Correa 2012	-0.223	0.084	794	933	6.3%	0.80 [0.68, 0.94]	
Nicholson 2014	-0.163	0.05	847	833	8.8%	0.85 [0.77, 0.94]	-
Roberts 2000	-0.051	0.03	299	259	10.2%	0.95 [0.90 , 1.01]	-
Sandora 2005	-0.03	0.15	602	451	3.2%	0.97 [0.72 , 1.30]	
Simmerman 2011	0.737	0.263	292	302	1.3%	2.09 [1.25 , 3.50]	
Stebbins 2011	-0.211	0.212	1695	1665	1.8%	0.81 [0.53 , 1.23]	
Swarthout 2020	-0.03	0.037	1496	4769	9.8%	0.97 [0.90 , 1.04]	4
Zomer 2015	0.068	0.052	278	267	8.6%	1.07 [0.97, 1.19]	-
Subtotal (95% CI)			12581	16678	67.2%	0.91 [0.84, 0.98]	•
Heterogeneity: $Tau^2 = 0$. Test for overall effect: Z		11 (1					
3.4.2 Adults	0.454	0.400	0.4	205	0.60/	0.00.00.00.4.041	
Cowling 2008	-0.151	0.408	84		0.6%	. , ,	-
Cowling 2009	-0.083	0.243	257		1.5%	. , ,	
Hubner 2010	-1.05	0.36	64		0.7%	. , ,	
Larson 2010	-0.199	0.134	946		3.7%	. , ,	
Little 2015	-0.151	0.02	8241		10.8%	. , ,	•
Millar 2016	-0.198	0.016	10000		11.0%	. , ,	•
Ram 2015	0.215	0.149	193		3.2%		+-
Teesing 2021	-0.67	0.248	976		1.4%	. , ,	
Subtotal (95% CI) Heterogeneity: Tau ² = 0. Test for overall effect: Z		= 7 (P = 0.	20761 .005); I ² = 66%	21190	32.8%	0.84 [0.78 , 0.91]	•
Total (95% CI)			33342		100.0%	0.89 [0.83, 0.94]	♦
Heterogeneity: $Tau^2 = 0$.	.01; Chi ² = 83.20, df	= 19 (P < 0	0.00001); $I^2 = 779$	%			1
Test for overall effect: Z Test for subgroup differe	, ,	$\hat{r} = 1 \text{ (P = 0)}$	0.18), I ² = 45.2%			Favo	0.5 0.7 1 1.5 2 ours hand hygiene Favours control

Footnotes

 $(1)\ Azor\ 2018\ includes\ 2\ intervnetion\ groups:\ soap\ and\ water\ (RR\ 0.94)\ and\ hand\ sanitizer\ (RR\ 0.77)$

Analysis 3.5. Comparison 3: Randomised trials: hand hygiene compared to control, Outcome 5: Absenteeism

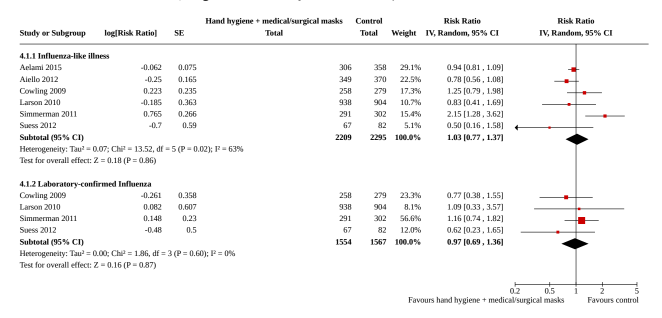
Study or Subgroup	log[Risk Ratio]	SE	Hand Hygiene Total	Control Total	Weight	Risk Ratio IV, Random, 95% CI	Risk I IV, Randon	
Azor-Martinez 2016	-0.478	0.065	621	720	64.8%	0.62 [0.55, 0.70]	_	
Hubner 2010	-0.693	0.435	64	65	1.4%	0.50 [0.21, 1.17]		_
Nicholson 2014	-0.362	0.09	847	833	33.8%	0.70 [0.58 , 0.83]	-	
Total (95% CI)			1532	1618	100.0%	0.64 [0.58, 0.71]	•	
Heterogeneity: Tau ² = 0	0.00; Chi ² = 1.43, df =	2 (P = 0.4)	49); I ² = 0%				•	
Test for overall effect: 2	Z = 8.45 (P < 0.00001)	1					0,2 0,5 1	2 5
Test for subgroup differ	rences: Not applicable					Favo	ours hand hygiene	Favours control



Comparison 4. Randomised trials: hand hygiene + medical/surgical masks compared to control

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
4.1 Viral illness	6		Risk Ratio (IV, Random, 95% CI)	Subtotals only
4.1.1 Influenza-like illness	6	4504	Risk Ratio (IV, Random, 95% CI)	1.03 [0.77, 1.37]
4.1.2 Laboratory-confirmed Influenza	4	3121	Risk Ratio (IV, Random, 95% CI)	0.97 [0.69, 1.36]

Analysis 4.1. Comparison 4: Randomised trials: hand hygiene + medical/surgical masks compared to control, Outcome 1: Viral illness



Comparison 5. Randomised trials: hand hygiene + medical/surgical masks compared to hand hygiene

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
5.1 Viral illness	3		Risk Ratio (IV, Random, 95% CI)	Subtotals only
5.1.1 Influenza-like illness	3	2982	Risk Ratio (IV, Random, 95% CI)	1.03 [0.69, 1.53]
5.1.2 Laboratory-confirmed in- fluenza	3	2982	Risk Ratio (IV, Random, 95% CI)	0.99 [0.69, 1.44]



Analysis 5.1. Comparison 5: Randomised trials: hand hygiene + medical/ surgical masks compared to hand hygiene, Outcome 1: Viral illness

Study or Subgroup	log[Risk Ratio]	SE	Hand hygiene + medical/surgical masks Total	Hand hygiene Total	Weight	Risk Ratio IV, Random, 95% CI	Risk Ratio IV, Random, 95% CI
5.1.1 Influenza-like ill	ness						
Cowling 2009	0.307	0.243	258	257	40.3%	1.36 [0.84, 2.19]	
Larson 2010	-0.456	0.363	938	946	23.6%	0.63 [0.31, 1.29]	
Simmerman 2011	0.028	0.266	291	292	36.2%	1.03 [0.61, 1.73]	
Subtotal (95% CI)			1487	1495	100.0%	1.03 [0.69, 1.53]	—
Heterogeneity: Tau ² = 0	0.04; Chi ² = 3.07, df =	2 (P = 0.2	2); I ² = 35%				T
Test for overall effect:	Z = 0.13 (P = 0.90)						
5.1.2 Laboratory-conf	firmed influenza						
Cowling 2009	0.301	0.39	258	257	23.3%	1.35 [0.63, 2.90]	
Larson 2010	-0.566	0.607	938	946	9.6%	0.57 [0.17, 1.87]	
Simmerman 2011	-0.034	0.23	291	292	67.1%	0.97 [0.62, 1.52]	
Subtotal (95% CI)			1487	1495	100.0%	0.99 [0.69, 1.44]	<u> </u>
Heterogeneity: Tau ² = (0.00; Chi ² = 1.49, df =	2 (P = 0.4	8); I ² = 0%				T
Test for overall effect:	Z = 0.04 (P = 0.97)						
					Fare	ours hand hygiene + medic	0.2 0.5 1 2 5 cal/surgical masks Favours hand hyg

Comparison 6. Randomised trials: gargling compared to control

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
6.1 Viral illness	2	830	Risk Ratio (IV, Random, 95% CI)	0.91 [0.63, 1.31]
6.2 SARS-CoV-2	2	394	Risk Ratio (IV, Random, 95% CI)	0.07 [0.02, 0.23]

Analysis 6.1. Comparison 6: Randomised trials: gargling compared to control, Outcome 1: Viral illness

Canada, au Cash arrang	law[Diala Dagia]	CE.	Gargling	Control	Ta7a: alas	Risk Ratio	Risk Ratio
Study or Subgroup	log[Risk Ratio]	SE	Total	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Goodall 2014	0.18	0.137	256	236	39.5%	1.20 [0.92 , 1.57]	+ = -
Satomura 2005 (1)	-0.44	0.22	104	57	29.5%	0.64 [0.42, 0.99]	-
Satomura 2005	-0.12	0.207	119	58	31.0%	0.89 [0.59 , 1.33]	-
Total (95% CI)			479	351	100.0%	0.91 [0.63, 1.31]	
Heterogeneity: Tau ² = 0	0.07; Chi ² = 6.01, df =	2 (P = 0.0)	05); I ² = 679	6			
Test for overall effect:	Z = 0.52 (P = 0.61)						0.1 0.2 0.5 1 2 5 10
Test for subgroup differ	rences: Not applicable						Favours gargling Favours control

Footnote

(1) Satomura 2005 included 2 intervention groups



Analysis 6.2. Comparison 6: Randomised trials: gargling compared to control, Outcome 2: SARS-CoV-2

	Mouth/no	se rinse	Cont	rol		Risk Ratio	Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI			
Almanza-Reyes 2021	2	114	33	117	67.7%	0.06 [0.02 , 0.25]				
Gutiérrez-García 2022	1	84	10	79	32.3%	0.09 [0.01, 0.72]				
Total (95% CI)		198		196	100.0%	0.07 [0.02 , 0.23]	•			
Total events:	3		43				•			
Heterogeneity: Tau ² = 0.0	0; Chi ² = 0.11,	df = 1 (P =	0.74); I ² =	0%			0.002 0.1	1 10 500		
Test for overall effect: $Z = 4.49$ ($P < 0.00001$)					Favours	Favours control				

Test for subgroup differences: Not applicable

ADDITIONAL TABLES

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist

Au- thor, year	Brief name	Recipi- ent	Why	What (materi- als)	What (procedures)	Who pro- vided	How	Where	When and how much	Tailor- ing	Mod- ifica- tion of inter- ven- tion through- out tri-	Strate- gies to improve or main- tain in- terven- tion fi- delity	Extent of inter- vention fidelity
											al	,	

Masks co	mpared t	o either no	o masks or	different mask ty	pes								
Abaluck 2022 (addi- tional sources: A baluck 2021a, A-	and	Lead- ers and adult house- hold- ers of rural and	In- crease large- scale adop- tion and proper	Masks colour- coded by households, ei- ther: A. cloth masks: an exterior layer of 100%	All villages: 1. household distribution of surgical or cloth masks and showing of maskwearing video;	Local NGO staff and volun- teers (Banglad	Masks and pro- motion deliv- ered e shi ce to face in	House- holds, mar- kets, mosques and streets of 572	8 weeks per vil- lage rolled out over a 6 week period (Novem-	Peri- odic mon- itor- ing and then addi- tional	In the first 5 weeks of the study staff found low en-	Num- bers of masks distrib- uted was noted	Num- bers of masks distrib- uted: A. 370,643
baluck 2021b, K- wong 2021)	bution	peri- urban vil- lages	wear- ing of face masks to slow	non-woven polypropylene (70 grams/m ² [gsm]), 2 interi- or layers of 60%	 distribution and promotion of masks at village markets; mask distribution at mosques; 	Green- Voice) ^[5] and Inno-	house- holds, mar- kets, mosques	vil- lages (in rural Banglade	ber 2020 to Janu- ary 2021)	train- ing of staff provid- ed as	gage- ment in some vil-	Promot- ers peri- odical- ly mon- itored	B. 924,849
	masks or B. Sur- gical masks		the spread of COV- ID-19 and save lives	cotton/40% polyester in- terlocking knit (190 gsm), an elastic loop that goes around the head above	4. mask promotion in public spaces;5. role modelling and advocacy by local leaders, including	vations for Pover- ty Ac- tion (IPA)	and streets of vil- lages both as groups and in-		1 day of training per vil- lage	need- ed Differ-	lages with local mask use, so mask pro-	passers- by and remind- ed peo- ple to put on	Mask- wearing: IGs: 42.3% CG:
	with possi- ble ad- dition- al vil- lage level		in- formed by re- search in pub- lic health,	and below the ears, and a nose bridge; filtra- tion efficiency: 37%[1]	Imams during Friday prayers using a scripted speech. Periodic monitoring	Village Imams and police officers	dividu- ally Text mes-		Once off mask distribu- tion and promo- tion at house-	cations and timing of ob- serva- tion across	motion staff were re- trained by re- searcher	Direct surveil- lance of mask	Increase was largest in mosques (37%
	ele- ments: i) in- centive		psy- chol- ogy, eco-	B. 3 layers of 100% non wo- ven polypropy-	of passers-by and re- minding people to put on masks	No "spe-	sages deliv- ered by		holds (4 days / village)	differ- ent days	part- way through the in-	wearing, correct mask-	points) and 25% to 29% points in

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklis

list (Continued)			. •		•			·	·
ii) sig- nage	nom- ics,	lene ^[2] , elastic ear loops, and a		cial- ized	phone and in-		terven- tion	wearing (wearing	other lo- cations
nage	mar-	nose bridge; fil-	Some villages:	skills"	dividu-	Mask	"to	either a	cations
iii) text	keting,	tration efficien-		need-	ally	distribu-	work	project	
mes-	and	cy: 95%.	village police accom-	ed as	,	tion 3 to	more	mask	_
sage	oth-	-, /	panying mask pro-	inter-		6 days /	close-	or an al-	Proper
re-	er so-	Sticker that	moters, providing	ven-		week	ly with	terna-	mask-
minder	cial sci-	had a logo of a	monetary rewards	tion		at mar-	local	tive face-	wear-
and	ences	mask with an	or certificates to vil- lages if mask-wear-	de-		kets and	leaders	covering	ing in-
and house-	on	outline of the	ing rate improves.	signed		on 3 Fri-	and set	over the	creased by
hold	prod-	Bangladeshi	ing rate improves.	to be		days at	specif-	mouth	Dy
ele-	uct	flag and a		easily		mosques	ic mile-	and	29.0%
ments:	pro-	phrase in Ben-		adopt-		during the first	stones	nose)	
	motion	gali that noted	Some villages:	ed by		4 weeks	for that	and	
i) altru-	and	the mask could be washed and	nublic signalling of	other		4 WEEKS	part-	physi-	Dhysiaal
ism or	dis-	reused ^[3] ; filtra-	public signalling of mask-wearing via	NGOs			ner-	cal dis-	Physical distanc-
self-	semi- nation	tion efficiency	signage, text mes-	or			ship"	tancing (if s/he	ing in-
protec-	strate-	of 76%	sage reminders, mes-	agen- cies		Week-		was at	creased
tion	gies	01 70 70	saging emphasizing	CICS		ly or bi-		least one	from
mes-	gies		either altruistic or			week-	After 5	arm's	24.1%
sages			self-protection mo-			ly mask	weeks,	length	in CG vil-
ii)		Initial 3 masks	tives for mask-wear-	Train-		promo-	mon-	away	lages to
amount		per household	ing, and extracting	ing of		tion	itor-	from the	29.2%
of			verbal commitments	staff			ing of	near-	in IG vil-
house-			from households.	pro- vided			mask-	est per-	lages
holds		Video of no-		by re-		Role-	wear- ing	son) ^[6]	
receiv-		table public		searchers		model-	was		
ing		figures ^[4] dis-	Modelling of safe	for	•	ling and	limit-		No dif-
texts		cussing why,	mask wearing by	mask		leader	ed to	Mone-	ference
:::\		how, and when	study staff	pro-		advo-	those	tary re-	between
iii) com-		to wear a mask	•	motion		cacy at Friday	who	wards or	IGs and
mit-						prayers	ар-	certifi-	CGs in
ment			Detailed procedures			prayers	peared	cates to	number
to		Brochure based	outlined in online				to be	villages	of peo-
mask-		on WHO mate-	protocol supplement				18	if mask-	ple ob-
wear-		rials depicting	osf.io/23mws/			Period-	years	wearing	served
ing		proper mask-				ic moni-	or old-	rate im-	in pub-
		wearing				toring: 1/ week on	er.	proved	lic areas, as an in-
						week on weeks 1,			dication
						2, 4, 6, 8,			of social
		Scripted				and 10;		Addi-	distanc-
		speeches for				ana 20,		tional	ing.
								training	<i>G</i> -

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
checklist (Continued)

ns in included studies, using the items from the T	emplate for Intervention Description
use by role models and lo- cal leaders at Friday prayers	daily schedule provided in Proto- col – 1 hour per
Scripted text messages	site for 9 sites 8am to 5pm
Monetary re- wards (USD 190) or non- monetary re- ward (certifi- cate) for vil- lages	Each village observed on 2 alternating days of the week.
Signage for	Observa-

Sigr household doors declaring they are a mask-wearing household

Smart phone for delivery and receipt of text message reminders

Loudspeaker for announcements in markets by research staff

Observations occurred 7 days of the week (9 am to 7 pm) Detailed schedules provided in

for mask

promotion staff

Record-

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undertaken by interven-

tion staff

includ-

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f.io/23mws/)

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online

protocol

supple-



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

Masks woven by and procured from local Bangladeshi garment factories within 6 weeks after ordering:

\$0.50 per cloth mask and \$0.13 per surgical mask

Masks and hand sanitiser for staff delivering intervention

Costs:

Cloth masks: \$275.10/village

Surgical masks:

\$88.90/village

PPE for staff: \$70/village

Media costs:

\$100/village

Transport and other costs: \$30/village

Handouts and written and some audio scripts for role



(accessed 13 June 2022).



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

models, leaders, surveillance officers and texts etc pro-

				texts etc pro- vided by the research team and in online protocol sup- plement via os- f.io/23mws/	
Alfelali 2020	Face masks	Ha- jj pil- grims aged ≥ 18 years	Prevent and control viral respiratory infections at mass gatherings	50 surgical face masks per participant (3M™ Standard Tie-On surgical mask, Cat No: 1816) Written instructions for mask use (See S1 Appendix)	Proverlinst for redensity of the property of t
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Provide masks and verbal and printed instructions, rules for mask use and demonstration of ap- propriate mask us- age provided (See S1 Appendix)	464 volun- teer trained re- search team mem- bers ap-	Individual ly and face to group of pilgrims in tents
Rules for mask use:	proached pil-	
• "Try to avoid touching the front of the mask.	grims in their tents	
 Change your mask if it is damp, wet or dirty. Always clean your hands before and after changing the masks. 	Train- ing in- clud- ed how to ap- proach pil-	
 Put used masks in a plastic bag and throw it into a rub- bish bin. You will 	grims and ex- plana- tion	

and

stration of

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find bins somewhere close to your tent in

Mina."

al- d to to ps l- s	Tents of pil- grims for Ha- jj in Makkah (Saudi Arabia) 50 to 150 pil- grims per large tent, sleep- ing head- to- head and shar- ing meals and	Mask wear- ing for 24 hours if possi- ble, over days of Hajj sea- son in- side and outside assigned tents 3 con- secu- tive Hajj seasons (5 to 6 days, Oc- tober 2013 to 2015)	Written information provided in preferred language (Arabic or English) Pilgrims who used at least 1 mask each day were considered to have	None de- scribed	4 day diaries of mask use: number of masks used and hours worn each day (see \$1 Appendix)	Mask use: IG: Daily: 24.7% Intermittently: 47.7% None: 20.9% CG: Daily: 14.3% Intermittently: 34.9% None: 43.7%
	rites		used the mask during that day (i.e.			Mask use of at least 4 hours consis- tently greater

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
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hecklist (Continued,	1					-			-		•	•
neckust (conumueu,	,				mask use				could be < 24 hours)			in IG than CG
BarasheedSuper- 2014 vised mask use	Religious pilgrims ≥ 15 years	Pre-vent respi- ratory virus infec- tions at mass gath- erings through mask use	Plain surgical face masks (3M Standard Tie-On Surgical Mask, Cat No: 1816) manufactured by 3M company, USA; 5 masks per day Written instructions on face mask use Special polythene bags for disposal	Masks provided to index case and their contacts with advice on mask use (before prayers, in seminars, and after meals). Written instructions provided on face mask use, need to change them, and disposal.	Not described, presum-ably the medical researchers	Face- to-face provi- sion of masks, in- struc- tions, and re- minders	Tents of pil- grim- age site (Mina Valley, Saudi Arabia)	Advice on mask use given through- out pil- grimage stay (5 days)	None report- ed.	None report- ed.	The medical researchers followed pilgrims each day to remind participants about recording their mask usage in health diary.	Face mask use: mask group: 56/75 (76%), control group: 11/89 (12%) (P < 0.001) 76% of intervention tents wore masks. 10 of 75 (13%) pilgrims in 'mask' tents wore face masks during sleep.
BundgaardFace 2021 masks (surgi- (addi- cal) tional source- Bundgaard 2020)	Com- muni- ty-dwelling adults aged 18 years or old- er with inter-	Re- duce ngvear- ers' risk for SARS- CoV-2 infec- tion out-	Per participant: 50 x 3-layer, disposable, surgical face masks with ear loops (TYPE II EN 14683 (Abena, Denmark); fil- tration rate,	Supply of masks sent to home address by courier Provision of written instructions sent by courier about how and when to wear masks including	Re- searchers provid- ed the masks (fund- ed by Salling Group), in-	Indi- s vidu- ally by mail, email, online and tele- phone	Mask wearing: when outside the home - and in the	Mask wearing: whenever outside the home or when guests in the home,	Chang- ing of mask if worn for more than 8 hours	None de- scribed	Face mask ad- herence: Self-re- port (Yes / Par- tial / No) (Suppl 4)	Face mask ad herence % Adhere: 46% Partial: 47% No: 7%

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)

net ac-	side	98%; made in	links to instructional	struc-	home	up to 8	lf		
cess	the	China)	video for face mask	tions	when	hours for	guests		
	home		use	and	they	1 mask,	in the	Average	Mean
	through			fol-	had	for 1	home,	mask	face
	protec-	1 badge (saying: "I am testing	Instruction to follow	low-up	guests	month	wear mask	use per	masks
	tion			sup-	(in			day	used:
	of the	face masks – for	advice of local health	port	Den-	(April			Week-
	nose	you and me")	authorities (in Den-		mark)	to May			days: 1.7
	and	you and me)	mark)			2020)	Indi-	Self-as-	uays. 1.1
	mouth		iliaik)	Back-			vidu-	sessed	Week-
	from			ground	In-		alised	adher-	ends: 1.3
	droplets	Written instruc-		and	struc-	1 off in-	sup-	ence	
	or	tions and in-	Provision of fol-	train-	tions	struc-	port as	with	
	aerosols	structional	low-up support by	ing	and	tions for	need-	health	
	or con-	videos for prop-	email and a phone	of re-	sup-	mask	ed via	authori-	Health
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		video for prop- er face mask use [in Danish] vimeo.com/40695				M/ I.		hygiene	ported
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79.2% to

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Loeb 2009	2 active interventions A. surgical masks B. N95 respirators	Health-care work-ers (nurs-es)	Re- duce trans- mis- sion of in- fluen- za in health- care set- tings through cough- ing or sneez- ing with pro- tective masks	A. Surgical masks B. N95 respirators	Provision of masks or N95 respirators Instruction in use and proper placement of devices Fit-testing and demonstration of positioning of N95 using standard protocol and procedure (details provided) Qualitative fit-testing using saccharin or Bitrex protocol ^[7]	Provided by research team (not further described) Fittesting by technician for N95	In-per- son face- to-face	Ter- tiary hos- pitals in On- tario, Cana- da	1 in- fluen- za sea- son (12 weeks) Use of mask as re- quired[8] when provid- ing care to or within 1 m of patient with febrile respira- tory ill- ness, ≥ 38 °C, and new or wors- ening cough or short- ness of breath Nurses to wear N95 when caring for pa- tients with "febrile	Fit- test- ing of nurses not al- ready fit-test- ed	Ceased before end of season	Adherence audits during peak of season by trained auditor who stood short distance from patient isolation room	18 episodes: N95: 6/7 partic- ipants (85.7%) wearing assigned device versus 100% for masks

respira-

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

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MacIntyre 2009	2 active interventions in addition to infection control guidelines A. Surgical masks (SM) B. P2 masks (P2)	House-hold- ers with a child with fever and respi- ratory symp- toms	Prevent or reduce respiratory virus transmission in the community through nonpharmaceutical interventions	A. 3M surgical mask, catalogue no. 1820; St Paul, MN, USA for adults B. P2 masks (3M flat-fold P2 mask, catalogue no. 9320; Bracknell, Berkshire, UK) A and B: health guidelines and pamphlets about infection control	Provision of masks and pamphlets and education about infection prevention and mask use Telephone calls and exit interviews to record adherence to mask use All groups: health guidelines, pamphlets about infection control were provided	Not described, presum-ably research team	Face- to-face and by tele- phone	House- holds in Syd- ney, Aus- tralia	2 winter seasons (3 months and 6 months) 2 weeks of follow-up Masks to be worn at all times when in same room as index child, regardless of distance from child	None de- scribed.	None de- scribed.	Daily tele- phone calls to record mask use through- out day Exit in- terviews about adher- ence	Reported mask use: Day 1 SM: 36/94 (38%) P2: 42/92 (46%) stated wearing "most or all" of the time. Other participants were wearing face masks rarely or never. Day 5:
													SM: 29/94 (31%) P2: 23/92 (25%)
MacIn- tyre 2011	3 ac- tive in- terven- tions A. Med- ical masks	Health- care work- ers	Protect HCWs by pre- vent- ing trans- mis-	Daily supply of A. 3 medical masks (3M medical mask, catalogue num- ber 1820, St Paul, MN, USA)	Supply of masks or respirators. Instruction in when to wear it, correct fitting, and storage (in paper bag in personal locker)	Masks provid- ed to hospi- tals. Train- ing of	Masks and train- ing pro- vided face-	Emer- gency de- part- ments and respi-	Entire work shift for 4 weeks	Tak- en off for toi- let and meal breaks and at	None de- scribed.	Mask/ respira- tor use moni- tored by: (i) ob- served	Adher- ence for usage was high for all and not



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Table 1. checklist		ion of int	ervention	s in included stu	idies, using the items	from the	Templat	e for Inte	rvention D	escriptio	n and Rep	olication (1	「IDieR)
	B. N95 respi- rators fit-test- ed C. N95 respi- rators non- fit-test- ed		sion of in- fluen- za and other respi- ratory viruses from pa- tients through mask wear- ing	2 respirators: B. N95 fit-tested mask (3M flat-fold N95 respirator, catalogue number 9132) fit-tested with 3M FT-30 Bitrex Fit Test kit according to manufacturer's instructions (3M, St Paul, MN, USA) C. N95 non-fit-tested mask (3M flat-fold N95 respirator, catalogue number 9132) Diary cards for usage recording	Instruction in importance of hand hygiene before and after removal For fit-tested group: fit-testing procedure	staff provid- ed by 1 mem- ber of re- search team.	to- face, not de- scribed if train- ing was in- divid- ually or in groups.	ratory wards in hos- pitals in Bei- jing, China		end of shift		adherence by head ward nurse recorded daily; (ii) self-report diary cards carried during day recording; (i) no. hours; (ii) usage. Exit interviews	significantly different amongst arms. Medical mask: 76%, 5 hours N95 fit- tested: 74%, 5.2 hours N95 non-fit- tested: 68%, 4.9 hours
MacIntyre 2013	3 active interventions A. N95 respirators at all times B. N95 respirators target ed use C. Medical masks	Health- care work- ers (nurs- es and doc- tors)	Protect HCWs from respi- ratory infec- tions from pa- tients through mask use	Daily supply of: A. and B. 2 respirators (3M Health Care N95 Particulate Respirator; cat- alogue number 1860) 3M FT-30 Bitrex Fit Test Kit C. 3 masks 3 masks (3M Standard Tie-On Surgical Mask cat- alogue number mask 1817; 3M, St Paul, MN, USA) Pocket-sized diary card with	Supply of respirators Instructions in use including times and fit Fit-testing procedure according to the manufacturer's instructions (3M) For targeted N95: checklist of defined high-risk procedures, including common aerosol-generating procedures	3M sup- plied respi- rators and masks. Provider of in- struc- tions not speci- fied.	Masks and training provided faceto-face, not described if training was individually or in groups.	Emergency de- part- ments and respi- ratory wards of ter- tiary hos- pitals in Bei- jing, China	For 4 weeks, A and B worn at all times on shift; B. tar- geted (inter- mittent) use of N95 res- pira- tors on- ly whilst perform- ing high- risk pro- cedures or barri- er.	None de- scribed.	None de- scribed.	Self-re- port- ed daily record of number of hours worked, mask or respira- tor use, number of high- risk pro- cedures under- taken collect- ed by study staff.	Adherence highest for targeted N95 (82%; 422/516) versus N95 (57%; 333/581) versus medical mask (66%; 380/572).

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

tick boxes for	r
mask use	

				tick boxes for mask use									
MacIntyre 2015	2 ac- tive in- terven- tions A. Cloth masks B. Med- ical masks	Hospi- tal health- care work- ers	Prevent respiratory infections in HCWs from patients through mask-wearing	A. 5 cloth masks for study duration (2- layer, cotton) B. 2 medical masks daily for each 8-hour shift for study duration (3 layers, non-woven material) All masks locally manufactured. Written instructions on cleaning cloth masks	Cloth or medical masks to be worn at all times on shift. Cloth masks to be washed with soap and water daily after shifts, and the process of cleaning to be documented. Provision of written instructions for cloth mask cleaning	Re- searchers arranged sup- ply of masks and in- struc- tions and any train- ing of staff assist- ing the deliv- ery.		Hos- pital wards in Viet- nam	4 weeks (25 days) of face mask use	Masks not worn while in the toi- let or during tea or lunch breaks.	None de- scribed.	Monitored adherence with mask use by self-report diary card and exit survey and interviews with a sub- sample (AC- TRN12610	Mask-wearing adher-ence: cloth mask: 56.8% medical mask: 56.6% Report-ed cloth mask washing: 23/25 days (92%)
MacIntyre 2016	Med- ical mask use	Sick house- hold- ers with ILI (index cases) and their well con- tacts of the same house- hold	Protect well people in the com- muni- ty from trans- mis- sion of respi- ratory pathoger by con- tacts with ILI through mask use	21 medical masks (3M 1817 surgical mask) Diary cards for mask use	Supply of masks Instructions for mask wearing and hand- washing protocol Provision of diary cards	Study staff mem- ber pro- vided masks and in- struc- tions in use.	Masks and in- struc- tions pro- vided face- to-face and in- dividu- ally.	Fever clin- ics of major hos- pitals in Bei- jing, China	3 masks/day for 21 days Mask wearing: whenever in the same room as a household member or a visitor to the household Handwashing: before	Al- lowed to re- move their masks during meal- times and whilst asleep and to cease wear- ing once symp- toms	None report- ed.	Self-re- port- ed daily record of mask use us- ing diary card	Mask use: mask group: 4.4 hours; control group: 1.4 hours



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

putting resolved on and after taking off

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Radonov 2019	tic2 active interventions A. N95 respirators (N95) B. Medical masks (MM)	Health-care per-sonnel of out-patient sites within medical centres	Prevent HCP from acquiring work- place viral respi- ratory infections and trans- mitting them to others by effective respi- rato- ry protection by N95 respi- rators which reduce aerosol expo- sure and in-	A. N95 respirators: 3M Corporation 1860, 1860S, and 1870 (St Paul, MN, USA) or Kimberly Clark Technol Fluidshield PFR95-270, PFR95-274 (Dallas, TX, USA) B. Medical mask Precept 15320 (Arden, NC, USA) or Kimberly Clark Technol Fluidshield 47107 (Dallas, TX, USA). Reminder signs posted at each site A portable computer equipped with data recording soft-	Participants instructed to wear assigned protective devices whenever they were positioned within 6 feet (1.83 m) of patients with suspected or confirmed respiratory illness and to don a new N95/MM with each patient interaction. Hand hygiene recommended to all participants in accordance with Centers for Disease Control and Prevention guidelines. Infection prevention policies were followed at each study site. Reminder signs posted at sites and emails cent	Centres provided de- vice sup- plied by study to HCP. Study per- sonnel post- ed re- minder signs and emails and con- ducted adher- ence ob- serva- tions.	Face- to-face indi- vidual provi- sion of de- vices and adher- ence obser- vations Onsite post- ing of signs Oth- er re- minders by email	Outpatient sites within medical centres in USA	As instructed, for each new patient interaction during 12-week period of peak viral respiratory illness each year for 4 years (total of 48 weeks)	Fitting of N95 masks	None de- scribed.	Re-minder signage posted at study sites, and emails sent by study personnel. Self-re-port-ed daily device wearing of "always", "sometimes", "never", or "did not recall" Observation of device-wearing behaviours as participants entered and evit-	Device wearing: N95: 89.4% report- ed "al- ways" or "some- times" versus MM: 90.2% "Never" N95: 10.2% MM: 9.5%
												-	

1	Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
	checklist (Continued)

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onovich Filtration testing performed on the device models in the study. Further details in protocol

(Radonovich 2016).

during unannounced, inconspicuous visits to randomly selected sites documented on portable computer

Posters

strooms as reminders of handwashing hygiene during 5week follow-up period after work-

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Hand hygiene

Alza- her 2018	Hand hy- giene work- shop	Pri- mary school girls	Tar- geted school chil- dren to im- prove hand hy- giene to re- duce school ab- sences due to upper	6-minute video- clip of 2 siblings that attended school-based health educa- tion about hand hygiene Short inter- active lecture about: common infec- tions in schools, methods of	Delivery of workshop and distribution of supporting materials (games and posters) to school and stu- dents	Study inves- tigator deliv- ered work- shop.	Delivered faceto-face in group format for the work-shop	2 pri- mary girls' schools in Sau- di Ara- bia	1-hour once- off work- shop; posters and games provided to school	Not de- scribed	Not de- scribed
			respi-	transmission,							



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

schools Puzzle games related to hand and to hygiene families

> Posters with cartoon princesses' picture promoting

				hand-washing									
Arbo- gast 2016	Multi- modal hand hy- giene inter- ven- tion pro- gramme in ad- dition to con- trol of brief video	Office build-ings and the employ-ees of health insurance company	Reduce hand-to-mouth germ trans-mis-sion from shared work-spaces and work-place facilities and there-by health-care claims and absenteeism through im-proved work-place hand	Alcohol-based hand sanitiser (PURELL Advanced, GOJO Industries Inc, Akron, OH, USA) installed as wall-mounted dispensers, stands, or freestanding bottles One 8-ounce bottle of hand sanitiser (PURELL Advanced) per cubicle One 100-count canister of hand wipes (PURELL Wipes) per cubicle	Hand hygiene supplies installed in offices. Replenishment product was made easily available to individual employees upon request via a simple process. Monitoring of product shipments into sites Physical collection and full replacement of soap, sanitiser, and wipes Intervention and control group: educational video embedded at end of baseline online knowledge survey	Not described, presumably study investigators arranged installations	Hand hy- giene sup- plies pro- vided in office environments and individually at staff cubicles/offices. Video provided in- dividually via email.	High-traffic common areas of 2 US health insurance company offices (e.g. near elevators, at entrances) and appropriate public spaces (e.g. coffee area, break rooms, conference rooms, training	13.5 months overall One-off email video 11 days before study hand hygiene supplies installed. 13 months of provision of supplies 2 times evening collection and	Sani- tis- er in- stalled in high- use ar- eas of the of- fices.	Not de- scribed	Employ- ee sur- vey at 4 months includ- ed ques- tions about hand hy- giene practice adher- ence. Monitor- ing of product ship- ments into the sites and physical collec- tion of the soap, sanitis- er, and wipes products 2 times	Intervention group employ-ees: reported 40% more cleaning of work area regularly; significantly more likely to keep the hand sanitiser with them and use it throughout the day; significant increases in hand sanitiser use for at-risk

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hy- giene	Replenishment products stored in supply room
	(in addition to existing foam hand wash (GO-JO Green Certified Foam Handwash) and an alcohol-based hand sanitiser foam wall-mounted dispenser (PURELL, GO-JO Industries) already provided near the re-

Identical soap in all restrooms

stroom exits prior to intervention)

Intervention and control group:

brief (< 1minute educational video) about proper hand hygiene technique, for both washing and sanitising hands

rooms, lob-bies, reception areas); individual staff cubicles of mostly open plan offices (average 309 square feet).	full re- place- ment of products
Of- fice re-	

strooms

full re- place- ment of products	in the study; collected samples were measured and usage rates were estimated	activities ^[9] Estimated use by average employee from sample collection:
		sanitiser 1.8 to 3.0 times/ day,
		soap
		2.1 to 4.4 times/ day,
		wipes at their desk 1.4 to 1.5

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

> "Wash Your Hands", signage promoting hand hygiene adherence, was already posted next to restroom exits at both the con-

	trol and intervention sites.	
Pre- vent trans- mis- sion of	Brochure about hand-washing awareness and habits	Brochure sent to parents by mail with study information sheet.
upper respi- ratory infec- tions in	Workshop content materials	Workshop provided for pupils and teachers:
schools and to fam- ilies through non- phar- ma- ceuti-	Stories, songs, and classroom posters about hand hygiene and infection transmission	frequent infections in schools, trans- mission and preven- tion, instructions on correct hand-wash- ing (water and soap, soaping > 20 s, dry- ing hands),
cal inter- ven- tion of	Hand sani- tiser (ALCO ALOE GEL hand sanitiser by Americo Gov-	use of hand sanitis- ers and possible side effects
hand- wash- ing pro- gramme in	antes Burguete, S.L. Madrid, Spain con- taining 0.2% chlorhexidine	Classroom activities linked to hand hy- giene and infection transmission

Brochure sent by school admin- istra-	Brochure sent by mail to indi- vidual	Pri- mary school class- es in	8 months overall
Work-shop and verbal and written information presumably provided	work-shops and class-room activities delivered in groups face-to-	Spain (de- tails not provid- ed)	One-off brochure and in- stalla- tion of hand sanitis- er dis- pensers 2-hour work- shop held 1 month
by the study re-	face.		month before study com-
search assis- tant.	Teacher rein- force- ment of		mence- ment
Class- room	hand hy-		Fort- night- ly class-

Not de- scribed	Daily re- inforce- ment by teachers of hand hygiene	Self-re- ported correct hand- washing included in analy- sis but		
	Fort- night- ly sup- port by research assis- tant pro- moting hand- washing	not sep- arately report- ed.		
	Self-re- ported correct hand- wash- ing pro- cedure (wa- ter and soap, soaping			

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

	L (Continuea)			0.1% benzalko- nium chloride, 5% aloe bar- badensis, 70% denat ethyl al- cohol, excipi- ents quantity sufficient for 100 mL alcohol 70%, pH 7.0 to 7.5)	Reinforcement of hand hygiene by teachers Hand sanitiser dispensers fixed to walls with an informational poster about hand-washing	activities provided by research assistant and teachers.	giene provid- ed to class face- to- face. Hand sanitis- er use super-		As required, teacher supervision and administration of hand sanitiser			> than 20 s, drying hands)	
				Informational poster about when and how to wash hands Written and verbal guidance to	Supervision of younger children when using hand sanitiser and administration of sanitiser if needed	Supervision and administration of hand sanitiser for	vision was provid- ed in- divid- ually and face- to-		Daily re- inforce- ment of hand hy- giene by teachers				
				teachers, parents, and students on properties, possible side effects, and precautionary measures for gel use and storage	dren in hand-washing procedures after toilet and when dirty and correct hand sanitiser use ^[10]	younger chil- dren by teach- ers	face.						
Azor- Mar- tinez 2018	Education- al and hand hy- giene pro- gramme	Day care centres and their attending children, their parents,	Prevent transmission of respiratory infections by improved hand	A. Liquid soap (no specific antibacterial components (pH = 5.5)) OR B. Hand sanitiser (70% ethyl alcohol (pH = 7.0 to 7.5)) for home use and	Installation of liquid soap or hand sanitiser dispensers in classrooms Supervision and administration of hand sanitiser if required	Work- shop deliv- ered by re- searcher. Re- search assis- tant	Work-shops deliv-ered face-s. to-face in groups to parents and staff.	Class- room of DCCs (in Spain) for child inter- ven- tions	8 months overall Initial 1-hour work- shop 1 month before study	Administration of hand sanitiser in the case of young children	Not de- scribed	Not described Reported that no monitoring of adherence	Families or DCC staff, or both, used 1660 L of hand sanitiser, estimat- ed use by each child of

dose 6 to 8 times/

day.

(TIDieR) Table 1 checkli

erven- and ions: DCC staff a soap and water and anitis-er	hy- giene of chil- dren, par- ents, and staff through hand- wash- ing prac- tices and use of hand sanitis- er due to its bacte- ricide and viru- cide prop- erties	in dispensers for school class-room Workshop content handout Stories, songs, and posters about hand hygiene and infection transmission	3 hand hygiene workshops for parents and DCC staff: 1. Hand-washing practices, hand sanitiser use, possible side effects and precautionary measures (HSG only) 2. RIs and their treatments 3. Fever Instructions to children, parents, and DCC staff on usual hand-washing practices and protocol ^[11] Classroom activities (stories and songs)	provided hand hygiene materials to DCCs and parents. Parents and staff supervised and administered sanitiser where indicated.	Work-shop content emailed to attendees individually. Individual face-to-face supervision of hand sanitiser use, as indicated	Work- shops provid- ed at DCCs.	commence- ment 3 further identi- cal ses- sions/DCC provid- ed again 1 month apart Fort- night- ly class- rooms and DCC activities One-off instal- lation	DCC staff could attend train- ing at other DCC if unable to at- tend at own DCC.	through continuous observation of hand hygiene behaviours was done, but amount of hand sanitiser was measured
			about hand hygiene and infection trans- mission				As-need-ed su-pervision of hand sanitiser use		

Dose of sanitis-

er: 1 to 2



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

mL/dis-

									infection				
Biswas 2019	Hand sanitiser and respiratory hygiene education	Pri- mary schools and their stu- dents and staff	Re- duce com- muni- ty-wide in- fluenza virus trans- mis- sion by im- prov- ing hand- wash- ing and respi- rato- ry hy- giene and use of sani- tiser in school- child- ren as con- tribu- tors to com- muni- ty-wide	Hand sanitiser (63% ethyl alcohol) in colourless, transparent 1.5-litre local plastic bottles (manufactured by a local pharmaceutical company and was available commercially in Bangladesh (price: USD 5.75/L)) Video clip on respiratory hygiene practices Behavioural change materials – 3 colour posters (see Appendix of paper) Curriculum materials for hy-	Installation of hand sanitiser in wall dispensers in all classrooms and outside all toilets, refilled by field staff as needed Encouragement of use of sanitiser at 5 key times during the day[12] Hand and respiratory hygiene education provided.[13] Integration of hygiene messages into school's hygiene curriculum Delivery of video clip on respiratory hygiene practice Behaviour change	Selected teachers responsible for dissemination of intervention messages throughout were trained over 2 days in these messages, behaviour change communication, sanitiser use, and practices	Hand sanitiser and education materials provided to schools. Education provided in classrooms in groups and faceto-face.	Pri- mary schools (in Banglade Sani- tiser in each class- room and out- side toilets Educa- tion in class- room	10 weeks	Refills provid- ed as need- ed.	Not de- scribed	Structured field observation by 2 field staff of 5 hours/school observing hand-washing and respiratory hygiene behaviours of children at 2 different locations in a class-room or outside Every other day, field staff measured	Hand-washing observed opportunities: IG 604/921 (66%) versus CG 171/802 (21%) Hand sanitiser used in 91% of observed handwashing events in intervention schools. Average consump
			virus trans- mis- sion	terials for hy- giene classes	materials distributed and placed around schools.	for pre- vent- ing spread of res- pira- tory						sured the level of hand sanitis- er in the morn- ing and in the af-	sump- tion of hand sanitis- er/child/ day: 4.3 mL

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Table 1.	Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
checklis	t (Continued)

checklist	is (Continued)				Use of sanitiser by classroom teachers after training Training of selected teachers in consultation with head of school and management committee in key messages Communication of key messages by the selected teachers to other teachers	secretions. Class-room teachers conveyed intervention messages during regular hygiene classes.						ternoon to cal- culate amount of hand sanitis- er used/ day/ school and en- rolled children.	Observation of proper cough or sneeze eti-quette: IG: 33% versus CG: 2%
						Field staff re- placed sup- plies as need- ed.							
Correa 2012	Alco- hol-based hand rubs	Child- I care centres and their staff and chil- dren	Re- duce inci- dence and trans- mis- sion of in- fection in chil-	Dispensers of alcohol-based hand rubs with ethanol 62.0% (PURELL, GO- JO Industries, Akron, OH, USA) Workshop ma- terials ^[14]	ABH and training on proper use to staff and children Pre-trial ABH use workshop to teachers that followed recommended HH teaching tech-	Local representative of GO-JO Industries Inc.	Face- to-face train- ing and provi- sion of mate- rials; group train- ing	Child-care centres in Colombia (centres or community homes)	8 months overall 1 ABH dis-penser per centre with	Re- filled ABH as need- ed	Not de- scribed	Visu- al re- minders and monthly refresher training	Teachers at 7 intervention centres reported almost complete substi-

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dren by im- proved hand hy- giene where wa- ter is scarce includ- ing provi-	Visual re- minders on ABH techniques in bathrooms and next to dis- pensers	niques and instructed teachers to add ABH to routine HH and give preference to hand-washing with soap and water if hands visibly soiled Continuous refilling of ABH	provided dispensers and dispenser installations free of charge.
sion of ABH and train- ing in hand hy- giene teach- ing tech- niques		ABH technique refresher workshops (8/centre) Monitoring of safety, proper use of ABH, amount of ABH used	Field- work team deliv- ered other com- po- nents.

ABH in centres, class-rooms, and common areas depending on size	< 14 children; 1 per class-room in larger centres; 1 per class-room + 1 for common areas in centres	of safe- ty, prop- er use of ABH, amount of ABH used Se- mi-struc- tured survey on com- pletion	tution of HSW with ABH, and HSW decreased from 3 times per day to 1 per day, and ABH rose to 6 per day.
Visu- al re- minders	with > 28 children	of teach- ers' per- ceptions	Teach- ers at re- maining 14 cen-
in bath-rooms and next to dispensers	1 work- shop pre-trial to staff	about changes in HH prac- tices and use of HSW and ABH.	tres reported partial substitution of HSW with ABH.
Work- shops and train- ing	ly 30- minute ABH tech- nique re- fresher training	Mea- sure- ment of con- sump- tion	Controls report- ed HSW 3 times per day.
pre- sum- ably provid- ed in cen- tres.	(8 per centre) Biweekly monitoring	of resources and costs re- lated to ABH use and HSW	Median number of ABH applica- tions per child
	-		

checklist (Continued)

to 4.5 in	
preschools	,
and 3.5	
to 5.5 in	
commu-	
nity cen-	
tres.	

DiVita 2011	House-hold hand-wash-ing pro-motion	House-hold- ers with index patient with ILI	Prevent in- fluenza trans- mis- sion in house- holds in re- source-p set- tings through provi- sion of hand- wash- ing fa- cilities and use of them at crit- ical times for pathoger trans- mis- sion		Provision of handwashing stations Hand-washing motivation to wash at critical times for pathogen transmission (e.g. after coughing or sneezing)	Not specif- ical- ly de- scribed, pre- sum- ably the re- searchers	Face- to-face provi- sion of facili- ties in house- holds "Moti- vation" not de- scribed	House- hold in Banglade	Over 2 influen- esha sea- sons One-off provi- sion of hand- washing facilities Frequen- cy of "moti- vation" not de- scribed	Not de- scribed	Not de- scribed	Not de- scribed	Not de- scribed
Feld- man 2016	2 ac- tive in- terven- tions	Naval ships and	Re- duced infec- tion	Septadine so- lution (Floris, Misgav, Israel) 70% alcohol	Installation of CHG disinfection devices on ships alongside	Provi- sion of CHG pre-	CHG sent to ships	Navy fast missile boats	4 months	CHG replen- ished	Not de- scribed	Total amount of CHG dis-	Mean volume CHG:

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

checklist	(Continued)	their	trans-	and 0.5% CHG;	regular soap and wa-	sum-	direct-	and	Unlimit-	on de-		pensed	8.2 mL
	A. Hand disin- fection with chlorhex- idine glu- conate + hy- giene educa- tion B. Hy- giene educa- tion	sailors	mis- sion and im- proved hand hy- giene in sailors who are at in- creased risk due to closed envi- ron- ments, con- tact with shared sur- faces, and poor HH cul- ture	inactive materials: purified water, glycerin, propylene glycol, and methylene blue	Supply and replenishment of CHG (sent to ships regardless of replenishment demands) Hygiene instruction by a naval physician (to both intervention groups and study control group)	ably by study team and funds Hy- giene in- struc- tion by naval physi- cian	Mode of hygiene instruction not described.	patrol boats of naval base in Israel Dispensers installed in key locations onboard (adjacent to heads (toilets), mess decks (dining rooms), common areas).	ed supply of CHG replenshed on demand for 4 to 5 months. Automatic amount dispensed: 3 mL	mand.		was tal- lied.	per sailor per day (project- ed yearly cost USD 45 per sailor)
Gwalt- ney 1980	A. Viru- cidal hand prepa-	Healthy young adults	Re- duce infec- tion	A. Virucidal hand prepara- tion:	Immersion of each finger and thumb of both hands to proximal interpha-	Re- searchers	Face- to-face and in- dividu-	US uni- versity	Expo- sure to donors on 3	Not de- scribed	Not de- scribed	Re- ported knowl- edge of	Active (n = 24):
	preparation B. Place- bo (no		rates by in- ter-	aqueous iodine (2% iodine and 4% potassium	langeal joint (inter- phalangeal joint of thumb) into desig-		ally		consecu- tive days (days 2,			hand prepa- ration	2 place- bo
			rupting viral spread	iodide)	nated preparation for 5 seconds then air-dried for 5 to 6				3, and 4) after ini- tial ex-			use as active, placebo,	16 don't know
	con- trol)		by hand		min				posure			or don't know	Placebo (n = 22):

6 active

7 place-

9 don't know

bo

lication (TIDieR)

checklist			or self- inocu- lation route	B. Placebo: aqueous solution of food colours (Kroger; Kroger Co., Cincinnati, OH, USA) mixed to resemble the colour of iodine with 0.01% io- dine and 0.02% potassium io- dide to give an odour of iodine	Exposure of recipients to donors either immediately after treatment or after 2-hour delay by hand contact with donor stroking fingers for 10 s Masks worn by donors and recipients during procedure.							
				Masks	Recipients placed in single isolation rooms after second exposure till end of experiment.							
Hubn- er 2010	Alco- holic hand disin- fection	Em- ploy- ees (ad- min- istra- tive of- ficers)	Re- duce absen- teeism and spread of in- fection in ad-	2 alcohol-based hand rubs (500 mL bottles) for desktop use to ensure minimal effort for use: 1. Amphisept E (Bode Chemie,	Provision of hand rub and instruction on use as needed at work only and in accordance with prevailing standard ^[15] : at least 5 times per day, especially after toileting, blow-	Pre- sum- ably provid- ed or arranged by study team	In person to staff	Admin- istra- tion of- fices in Ger- many	12 months overall Hand rub used as much needed	Hand rub use espe- cial- ly af- ter toi- leting, blow- ing	Not de- scribed	S p e h w h g m sı

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Hamburg, Gering nose, before many) ethanol eating, and after (80% w/w) contact with ill colbased formuleagues, customers, la with antibacand archive material

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ing activity.

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needed Hand rubs for comused at plete desk or wetting work of the (not hands (at least outside of 3 mL or work). a palm-

ful)[16]

at least

leagues,

Self-re-Reported mean ported adhand disinfecherence with tion frehand hyquengiene cy times per day: measures > 5: 19% nose, before 3 to 5: eating, 59.8% and after 1 to 2: con-20.5% tact with < 1: 0.7% ill col-

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

wit pap doo me thr im- pro har hy-	pants with skin problems: ork ith Sterillium aper (Bode Chemie, Hamburg, Germany) arough 2-propanol n- (45% w/w), roved 1-propanol and (30% w/w),		pants with skin problems: Sterillium (Bode Chemie, Hamburg, Germany) 2-propanol (45% w/w), 1-propanol (30% w/w), and mecetro- nium etilsul- fate (0.2% w/w), with a refatting effect and has activity against bacteria, fungi and enveloped				5 times per day.	cus- tomers, and archive mater- ial			
	Baki wate emu no n teria (Boo	d cream: colan balm, er-in-oil lsion with on-antibac- il properties de Chemie, nburg, Ger- y)									
Day-Recare duccentres risk and infetheir tionstaff, chi	ce guid of omr ec- for: n in vent	onnel e on rec- nendations hygiene, ilation, out- ay care,	Staff meeting in each DCC and training in microbiological cause of infection spread guided by National Board of	Re- search team pre- sum- ably	Face- to-face with train- ing and activi-	On- site in DCCs	2-month interven- tion peri- od	None de- scribed.	None de- scribed.	None described.	None re- ported.

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1	Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
	checklist (Continued)

scripti	on of int	erventior	ns in included stu	ıdies, using the items
ntinued)	of chil- dren	gien- ic edu- cation of day- care profes- sion-	Fairy tale and poster "The Princess Who Won't Wash Hands"	Education of children in hand-washing (about bacteria and why and when to wash hands)
		als, moti- vation of day- care facili-	Colouring in drawings	Practical hand-washing classes with 4 to 5 children at a time
		ties for regular hand hy- giene,	"Wash hands" song and rhymes	Provision of t-shirt, book, and diploma to children
		and inform- ing par- ents about hand hy- giene	T-shirt for children with the inscription "Clean hands - yes thank you"	Provision of leaflet for parents
		gierie	Diploma for children and book "The Princess Who Won't Wash Hands" to also be used by par- ents with their child	
			Informational	

leaflet for parents in envelope

children

Information sent home to parents via chil-

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)

checklist	(Continued)				, ,		-			-	•		•
Little 2015	Web- based hand- wash- ing in- terven- tion	House-hold- ers (over 18) who were gen- eral prac- tice pa- tients	Prevent trans- mis- sion of respi- ratory tract infec- tions through im- proved hand hy- giene to re- duce spread via close con- tact (via droplets) and hand- to-face con- tact	Website-based programme: provided information about the importance of influenza and role of handwashing; developed a plan to maximise intention formation for hand-washing; reinforced helpful attitudes and norms; addressed negative beliefs (URL provided for demonstration version no longer active; see www.lifeguideonline.org)	Provision of link to website for direct log in Automated emails prompted participants to use sessions and complete monthly questionnaires and maintain hand-washing.	Researchers delivered webbased programme and emails.	Online s indi- vidual- ly	House-holds in Eng-land	4 months overall 4 week-ly web-based sessions Month-ly email ques-tions to maintain hand-washing over 4 months	Tai- lored feed- back pro- vided with- in web pro- gramme	None de- scribed.	Emailed questions monthly to maintain handwashing	None reported.
Luby 2005	Hand- wash- ing pro- mo- tion at neigh- bour- hood level with 2 inter- ven-	Neigh- bour- hoods and their house- holds	Im- prove hand- wash- ing and bathing with soap in set- tings where com- mu-	Slide shows, videotapes, and pamphlets illustrating health problems from contaminated hands and specific handwashing instructions	Hand-washing promotion to neighbourhoods: Neighbourhood meetings of 10 to 15 householders (mothers) from nearby homes and monthly meetings for men Soap to households	Re- search team in col- labo- ration with Health Orient- ed Pre- ventive Edu-	Face- to- face in small groups and in- dividu- ally	Neigh- bour- hoods and homes in Karachi, Pak- istan	1-year weekly house- hold vis- its 30- to 45- minute neigh- bour-	Soap re- placed regu- larly.	None de- scribed.	None de- scribed, though soap use mea- sured.	House-holds' mean use of study soap per week: 3.3 bars Average use per resident

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tions at house- hold level			nica- ble dis- eases are lead-	Soaps: 90- gram white bars without brand names or symbols, same	Fieldworker home visits: discussed im- portance of and cor-	cation (HOPE) ^{[18}	3]		hood meet- ings 2 to 3 times/ week				per day: 4.4 g
		lead- ing caus- es of child- hood mor- bidi- ty and mor-	caus- es of child- hood mor- bidi- ty and mor-	symbols, same smell with identical generic white wrappers with serial numbers matched to households A. Households: 2 to 4 white bars of 90-gram antibacterial soap containing 1.2% triclocarban (Safeguard Bar Soap: Procter & Gamble Company (Cincinnati, OH, USA)	rect hand-washing (wet hands, lather them completely with soap, rub them together for 45 seconds, and rinse off completely) technique and promote regular hand-washing habits ^[17] Encouragement of daily bathing with soap and water	Field- work- ers were trained in in- ter- view- ing and hand- wash- ing pro- mo- tion.			first 2 months then week-ly for months 2 to 9, then monthly Monthly men's meetings first 3 months Weekly house-hold visits				
				B. Households: plain soap (no triclocarban)									
				Soap packets									
Mil- lar 2016 a dition- al de- tails from El- lis 2010	Skin adand soft- tissue infec- tion pre- ven-	Mili- tary trainees	Im- prove per- son- al hy- giene prac- tices	A. Enhanced standard: supplemental materials (a pocket card and posters in the barracks)	Provision of ed- ucation and hy- giene-based mea- sures in addition to standard SSTI pre- vention brief upon entry:	Not de- scribed, pre- sum- ably the re- searchers	Face- to-face and in- dividu- ally for body wash and	US mil- itary train- ing base	One-off educa- tion on entry to training	None de- scribed.	None de- scribed.	None described.	None de- scribed.

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Table 1. checklist		on of int	ervention	s in included stu	idies, using the items	from the	Templat	e for Inte	rvention D	escriptio	n and Re	plication (1	ΓIDieR)
	tion intervention in addition to SSTI brief on entry also provided to control A. Enhanced standard B. Chlorhexidine	-	to prevent infection, especially acute respiratory infection in military trainees who are at increased risk	B. CHG: CHG- based body wash (Hibi- clens, Mölnly- cke Heath Care, Norcross, GA, USA)	Enhanced standard: supplemental materials CHG: as for enhanced standard group, plus a CHG- based body wash and instructions for use		Mode of education not described.		CHG: use of wash 1 per week for entire training period (14 weeks)				
Morton 2004	Healthy hands (alco- hol gel as hand- wash- ing ad- junct)	Ele- men- tary schools and their chil- dren and staff	Prevent infections in elementary schoolage children who are particularly vulnerable through adjunct use of alcohol gel and	Alcohol gel and dispensers: AlcoSCRUB (60% ethyl alcohol) supplied by Erie Scientific Company, Portsmouth, NH, USA "Healthy Hands Rules" protocol [19] (Figure 3 in paper) Healthy Hand Resource Man-	Healthy hands protocol introduced after "Germ unit" education in classes Daily reminders to children on public address system (in first week) then weekly reminders Review of protocol in each classroom after vacation by school nurse	Gel provided by suppliers. Research team provided educational aspects. Classroom teach-	Face- to-face train- ing in class- es and indi- vidual infor- mation giving and moni- toring	Ele-men-tary schools in USA Wall-mount-ed near door en-trance of each class-room at age-appro-priate height	0.5 mL dispensed per application. Use of "special soap" according to "Healthy Hands Protocol" (Fig-	Reinforcement teaching provided if gel usage indicated that it was needed. Germ unit education tailored	1 student was concerned gel was making her sick, so school nurse provided additional classroom visit to allay concerns.	Usage of gel cal- culated.	5 gel applications per day 1 dispenser lasted 1 month.

checklist (Continu	ued)	edu- cation based on Health Belief Model (HBM) (Kirscht 1974)	ual for school nurse, available for parents Monthly newsletters to parents "Healthy Hands" refrigerator magnet for families (see Figure 2 in paper) Informational letter to local primary care providers, paediatricians, family practitioners, and advanced practice nurses "Germ Unit" curriculum and materials including Germ models and Glo Germ	"Healthy Hands" magnet provided to parents and guardians. "Hand Checks on Wednesdays" to identify adverse effects of gel	ers responsible for encouraging use of gel and reinforcing protocol School nurse assisted in monitoring and hand checks for adverse effects.			ure 3 in paper)	for each grade level.			
Nichol- Hand son wash 2014 ing with		Target- ed 5- year- old chil- dren	Initial supply of 5 bars of free soap (90-gram Lifebuoy bars) replenished on submission of	Provision of soap and social marketing programme (Sidibe 2009) (Lifebuoy branding) to edu- cate, motivate, and	Dedi- cated team of "pro- mot-	Face- to- face in groups	"Class- rooms" held in com- munity	41 weeks Weekly "class- rooms"	Moth- ers were asked to pro- vide	Tech- nical diffi- culties with "soap	Regis- ters for "class- rooms" and home	Soar con- sum tion:



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

their	and	empty wrap-	reward children for	ers"	Indi-	build-	after	and	accel-	visits	IG versus
moth-		pers.	HWWS at key times	deliv- ered	vidu-	ings	school	share hand-	eration	where 3-week	CG:
ers	moth- ers as			edu-	ally by moth-		and home	wash-	sen- sors"	gaps in	235 g
	change			cation	er to		visits	ing tips	to	atten-	versus
	agents	Environmental cue reminders	Weeks 1 to 17: hand- washing occasions,	and	child	Home vis-		with	mea-	dance	45 g
	to re- duce	(wall hangers,	germ education,	home visits.		its of		other moth-	sure HWWS	triggered	
	inci-	danglers)	soap's importance in	VISILS.		house-	HWWS	ers,	behav-	supervi- sors to	
	dence		germ removal			holds	encour-	com-	iours	ask par-	
	of res-		Week 18 onward:	Moth-		in Mum-	aged 5 key oc-	peti-	pre-	ticipants	
	pira-	Rewards (e.g.	encouragement of	ers		bai, In-	casions:	tions held	vent- ed suc-	to re- sume or	
	tory infec-	stickers, coins, toy animals)	HWWS on 5 key occa-	provid-		dia	after	for	cessful	be with-	
	tions	toy ammats)	sions supported by environmental cues	ed sup-			defe-	moth-	use.	drawn	
	(and		environmental edes	plied re-			cation, before	ers.			
	diar- rhoeal			wards.			each of				
	dis-		"Classrooms" for				3 meals,			Moni-	
	ease)		children				and during			toring of soap	
	through hand-						bathing.			resale	
	wash-									on open	
	ing us-		Home visits for mothers							market by use of	
	ing be-		mothers				Week 18			unique	
	hav- iour						onward: hand-			iden-	
	change		Parents' evenings to				washing			tifiers	
	prin-		boost morale, build				on 5 oc-			on soap wrap-	
	ciples (Claesser		networks, and run				casions			pers and	
	2008),	ı	competition for ad- herence, assignment				for 10 consecu-			twice	
	includ-		completion, and				tive days			weekly checks	
	ing so-		folder decoration							in local	
	cial norms									shops	
	for						6 weekly				
	child		Establishment of a "Good Mums" club				parents' meet-				
	and mother		for sharing HWWS				ings			Collec-	
	(Perkins		tips				G.			tion of used	
	2003),									soap	
	using									wrap-	
	fear of									pers as	

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ecklist (Continued)		ami- nation and		Rewards provided by mothers.							soap con- sump-	
		disgust (Curtis 2001), peer pres- sure (Sidibe 2003),		Children encouraged to advocate HWWS within families be- fore meals.							tion measure	
		morale boost- ing, and net- work- ing sup- port		Establishment of social norms for child and mother with pledges in front of peers								
ande- 3 ac- cong 2012tive in- terven- tions	Preschoo class- es (stu- dents	geted	1 container of alcohol hand I gel per class- room (active in-	Teachers instructed to: assist each child with	Teach- ers su- per- vised,	Face- to- face to schools,	Kinder- garten school in	12 weeks overall	None de- scribed.	Stu- dents whose fami-	2 re- search assis-	Report- ed that adher- ence wa
(no con- trol) differ-	and teach- ers) and	dren who can have	gredients: eth- yl alcohol, 70%; chlorhexidine gluconate,1%;	dispensing hand gel at required time interval,	stored, and re- filled hand	teach- ers and chil- dren	Bangkok, Thai- land	1 pump of gel per child per dis-		lies de- clined to par- tici-	tants moni- tored hand	ensured for each interver tion
ent time- inter- val ap- plica-	their par- ents	high infec- tion rates in ILI;	Irgasan (tri- closan), 0.3%)	store hand gel prop- erly, and refill gel as needed.	gel. In-	Indi- vidual		infection round at 1 of 3 time in-		pate were not asked	gel use every 60 or 120 minutes for the	group
tions of al- cohol		have close inter-	Cost of hand gel every 60 minutes was	Monitoring of hand gel use at specified	struc- tions to	assis- tance to chil-		tervals of school day:		to use alcohol hand	duration of study.	Cost of hand ge every 60
hand gel		action so at risk of air-	USD 6.39 per child per 12- week period	times	teach- ers pre- sum-	dren with hand gel		A. every 60 min B. every		gel.	Class- room teachers	minute: was US 6.39 pe child pe

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Trusted evidence.
Informed decisions.
Better health.

Table 1.	Description of in	nterventions in inc	cluded studies,	using the items	from the Ten	nplate for Inter	vention Description	on and Replicati	on (TIDieR)

	B. Every 120 min C. Once before lunch		contact trans- mis- sion; and are of in- creas- ingly younger ages through hand gel as a single strat- egy of conve- nient and ef- fective disin- fection	Leaflet describing risk factors for ILI for each family		Leaflets distributed through school. Monitoring of use by 2 research assistants	Leaflets given to each family.		C. once only before lunch, the school standard for hand hygiene		mained in their class-rooms and continued to follow the school standard for hand hy-giene.	sign after each disinfection round.	
Priest 2014	Hand sanitiser provision (in addition to hand hygiene education session also provided to control group)	Pri- mary schools and their stu- dents, teach- ers, and admin- istra- tive staff	Re-duce per-son-to-person community transmis-sion of infectious disease by target-ing improved	"No touch" dispensers (> 60% ethanol) for each classroom that dispensed dose when hands were placed under an infrared sensor Supply of topup sanitiser as needed	Dispensers installed into each classroom. Teachers asked to ensure that the children used sanitiser at particular times and to oversee general use (McKenzie 2010). Weekly classroom visits to top-up of	School liai- son re- search assis- tants topped- up sanitis- er.	Instal- lation of dis- pensers to class- rooms Super- vision of chil- dren by teach- ers de- livered	City schools in New Zealand	20 weeks (2 school terms) Sanitiser to be used by students at least after coughing/sneezing, blowing their nose,	Children were able to use the sanitiser at any time they wished as well as at key times (McKenzie 2010).	Change of sanitiser after week 10 to flavourless type of the same % ethanol in 41 of 396 class-rooms	Week- ly class- room visits by school li- aison re- search assis- tants who record- ed quan- tity of sanitiser used	100% dispensing 45 mL per child Average hand sanitiser dispensed/child for 34 schools: 94 mL



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

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sanitiser and measure quantity used

30-minute in-class hand hygiene education session provided (also to control group) plus instruction in hand sanitiser use.

faceto-face individually and as a

class.

and as they leave for morning break and for

lunch break.

Approximately 0.45 mL of sanitiser dispensed

Weekly top-up of sani-

tiser

Initiation

of inter-

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per

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(10%)Total (in 9 amount of 34 of sanischools) tiser per classdue to room chil-

was dren meatasting sured. it when eat-

ing, afadherfecting ence defined as dispensing a

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equivalent to at least 45 mL

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per child of hand sanitiser solution over the trial period.

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sanitiser usage between first 10 weeks and second 10 weeks amongst classes that switched products was 220 mL.

Median

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Ram	Soap
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House-Rehold duce comhousehold pounds and its transhousemisholdsion of ILI ers (adults and inand fluenza

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that

Hand-washing station in central location of each com-

pound using: large water container with a tap;

plastic case for soap;

Hand-washing station in each compound

Didactic and interactive group-level education and skills training describing influenza symptoms, transmission, and prevention, promot-

Inter-

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staff

provision of handwashing station and pre-

All elements delivered arranged faceto-face

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Soap

present

133 compounds (74%).



d studies, using the items from the Template for Intervention Description and Replication (TIDieR) Table 1. Description checklist (Continued)

on of int	ervention	ns in included st
had a house- holder	wash- ing in house-	bar of soap.
with ILI	holds with house- hold- er with ILI as	Cue cards depicting critical times for handwashing:
	other house-	or sneezing;
	hold- ers who are	after cleaning one's nose or child's nose,
	well are at	after defeca- tion;
	high- est risk of ex- posure	after clearing a child who has defecated;
	due to crowd- ed and poorly	before food preparation or serving;
	venti- lated homes.	before eating.
	Followed constructs of Social Cognitive	

Theory and the Health

Belief Model (Glanz

2008)

•	ares, using the recins	iioiii tiic	Cilipto
	ing health and non- health benefits of hand-washing with soap and identifica- tion of barriers and proposed solutions to hand-washing with soap	sum- ably provid- ed ed- uca- tion.	uca- tion), and indi- vidual levels (rein- force-
	Daily surveillance including weighing of soap and replacing if ≥ 20 g and resupply of water in container if needed	Intervention staff conducted daily surveillance	ment).
	Posting of cue cards	and rein- force- ment visits.	
	Asking household- ers to demonstrate hand-washing with soap technique		

holds with com- mon court- yard, shared latrine,	ing res- olution of index case pa- tient's symp- toms	force- ment and model- ling as need- ed.	haviours includ- ing ob- served hand- washing
water source, and cook- ing fa- cilities	Day 1 set up of hand- washing station		Cue cards in common areas of court- yard

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Ü	er were
	present
	7 or
Cue	more
cards in	of first
common	10 days
areas of	in 99%
court-	of com-
yard	pounds,
	with wa-
	ter and
Presence	soap ob-
or ab-	served
sence	together
of soap	on all 10
during	days in
each of	99 com-
first 10	pounds
days of	(55%)
surveil-	
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from 180	Coon
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	median:
	2.3 g
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and	maxi-
amount	mal: 5 g
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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

Teaching of tech-

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and behaviour change com- muni- cation using social mar- keting con- cepts										
Re- duce trans- mis- sion of	GloGerm (GloGerm, Moab, UT, USA)	Staff training in good health (developed by Kendrick 1994) and practical exer- cise of hand-washing	Train- ing and rein- force- ment	Face- to- face in groups for	Child- care centres in Can- berra,	8 months overall	Train- ing for new staff provid-	None de- scribed.	6-week- ly ad- herence mea- sured by	Adher- ence was report- ed only in rela-
respi- ratory infec- tions in	Newsletters to staff	with GloGerm	activ- ities pro- vided	train- ing and classes and in-	Aus- tralia	3-hour train- ing in	ed as need- ed.		recorded observa- tion of recom-	tion to analysis of out- comes.
child- care centres through	Songs and rhymes on hand-washing	Fortnightly visits and newsletter to reinforce training and to communicate techniques	by 1 of the re- searchers	dividu- ally as aneed- ed to		evening or 1- hour during lunch			mend- ed prac- tice for 3 hours	High ad- herence
im- proved infec- tion control proce- dures	Plastic bags (sandwich bags available at su- permarkets) to	Recommended hand-washing technique as per guidelines of the time ^[21]	Teach- ers de- livered train- ing to	chil- dren or staff		for new staff af- ter study start			in the morning in each centre, graded by quan- tiles of	reported for nose wip- ing and child hand-
	cover hand for nose wiping	and after toileting, before eating, af- ter changing diaper (staff and child), and after wiping nose un- less barrier used	chil- dren based on their train-			Duration of hand- washing: "count to 10" to wash			frequen- cy of recom- mend- ed hand- washing	washing.
		Teaching of tech-	ing.			and "count to 10" to			by chil- dren.	

rinse

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

wash hands for infants

					Turits								
Sando- ra 2005	Healthy Hands Healthy Fami- lies	Families with an index child in out-of-home child-care	Re- duce illness trans- mis- sion in the home through multi- factori- al cam- paign cen- tred on hand hy- giene edu- cation and hand sanitis- er	Alcohol-based hand sanitiser: active ingredient: 62% ethyl alcohol (PURELL Instant Hand Sanitiser; GOJO Industries, Inc, Akron, OH, USA) Hand hygiene educational materials at home (fact sheets, toys, games)	Supply of hand sanitiser and hand hygiene materials Biweekly telephone calls Biweekly educational materials	Study investi- gator	Not stated whether mate- rials mailed or de- livered in per- son	Homes in USA Sani- tiser use in home	5 months overall Biweek- ly edu- cational materi- als Sanitis- er dis- pensed 1 mL each pump.	None de- scribed.	None de- scribed.	Record- ed amount of hand sanitiser used (as reported by the primary caregiv- er)	Median frequency of reported times of hand sanitiser use: 5.2 per day 38% used > 2 ounces of hand sanitiser per fortinight = 4 to 5 uses per day
Savolaine Kopra 2012 further details from Save Kopra 2010	En- hanced hy- giene olainen- 2 ac- tive in- terven- tions IR1. Soap and	Office work- ers of office work units	Prevent transmission of respiratory infections in workplaces through enhanced hand hy-	IR1: Liquid hand soap ("Erisan Non- sid" by Farmos Inc., Turku, Finland) IR2: in addition: Alcohol-based hand rub, 80% ethanol ("LV" by Berner Inc.,	Toilets equipped with liquid hand soap (all groups) or alcohol-based hand rub (IR2). Guidance on other ways to limit transmission of infections, e.g. frequent handwashing in office and at home, coughing, sneezing into disposable handkerchief	In collaboration with occupational health clinics servicing the corporation	In-person provision of soap or hand rub Guidance and written instructions	Office work units in cor- pora- tions in Helsin- ki, Fin- land	15 to 16 months overall Month- ly visits by nurse through- out	Nurses assist- ed with any prac- tical prob- lems with inter- ven- tion as they arose.	None de- scribed.	Adherence assessed by an electronic self-report survey of transmission-limiting habits 3 times (more	Avoiding hand-shaking became more common and remained high in both groups. Recorded use for per-

before,

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KUSI	t (Continued) water wash IR2. Alco- hol-based hand rub	i	giene with behav- ioural recom- men- da- tions to re- duce trans- mis- sion by droplets during cough- ing or sneez- ing	Helsinki, Finland) Bottles of hand hygiene product (free of charge) to be used at home and in the office (IR2). Written instructions on hygiene for further reference	or sleeve, avoiding hand-shaking Visits to work clusters and monitoring of materials availability Monthly electronic "information spot" about viral diseases for motivation to maintain hygiene habits Adherence activities	Specially trained research nurse provided guidance and visited worker clusters throughout intervention period.	given per- sonal- ly. Face- to-face vis- its by study nurse			New employees received guidance on hand hygiene and habits.		details in protocol). Use of soap (IR1) and alcohol-based disinfectant (IR2) for personal use was recorded. Study nurse checked avail-	son-al use small-er than predicted use based on han hygien instructions. Soap of disinfet tant us age per particition pant: IR1: 6.5
eb-	"WHACK		Tar-	Hand sanitiser	Delivery of grade-	Project	Face-	Ele-	Whole	En-	None	ability of soap and alcohol rub.	Teache
5 1	the Flu" (hand sanitis- er and train- ing in hand	men- tary schools and their stu- dents and home-	geted school- aged chil- dren as impor- tant sources of in-	with 62% alco- hol-based hand sanitiser from PURELL (GOJO Industries, Inc, Akron, OH, USA) automatical- ly dispossing 1	specific presenta- tions on "WHACK the Flu" concepts and proper hand-wash- ing technique and sanitiser use: (W)ash or sanitise your hands often;	staff provid- ed ed- uca- tion. Home	to- face at schools, pre- sum- ably as a group in	men- tary schools (Pitts- burgh, USA)	inter- vention over 1 influen- za sea- son	cour- aged to wash hands or use addi- tion- al dos-	report- ed.	teacher surveys of ob- served NPI-re- lated be- haviour in their students	survey of ob- served class- room NPI be haviou indicated ed suc

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Trusted evidence.
Informed decisions.
Better health.

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
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checklis		,	Alexand I			£ 1				la a A I			
neckus	ry hy- giene)		through im- proved cough eti- quette and hand hy- giene in schools including sanitiser as potential in- expensive non- phar- ma-		and mouth; (C) over your coughs and sneezes; and (Keep your distance from sick people (provided URL no longer active) Desired frequency of hand wash use taught to student (see When and how much) Installation of hand sanitiser dispensers	forced mes- sage and moni- tored proper use of sanitis- er.		in each class- room and all major com- mon areas.	one-off 45-minute education presentation and one-off refresher training at onset of influenza season	both, as need- ed		Mea- sure- ment of hand sanitiser use at 2- week in- tervals through- out the interven- tion peri- od	nance of be- haviours through- out in- fluenza season. Average sanitis- er use: 2.4 times per day
			ceuti- cal inter- ven- tions		Reinforcement of message and monitoring of sanitiser				use of 1 dose (0.6 mL) of sanitiser 4 times per day ^[22]				
Talaat 2011	Inten- sive hand hy- giene cam- paign	Schools and their stu- dents, teach- ers, and par- ents	Re- duce or pre- vent trans- mis- sion of in- fluenza viruses amongst chil- dren	Soap supplied as needed. Grade-specific student booklets each including a set of 12 games and fun activities that promoted hand-washing	Establishment of a hand hygiene team in each school Provision of hand hygiene activities: weekly exercises (e.g. games, aerobics, songs, experiments); school activities, (e.g. obliga-	Hand hy- giene team (3 teach- ers from social stud- ies, arts, and	Delivered faceto-face in groups and individually	Ele- men- tary schools (grades 1 to 3) in Cairo, Egypt	12 weeks overall Week- ly hand hygiene cam- paign ac- tivities	Soap and hand- drying ma- terial provid- ed by school admin- istra- tion if chil-	None de- scribed.	Observation by social workers of hand hygiene activities, availability of soap and drying material,	About 93% of the stu- dents had soap and dry- ing ma- terial avail- able.

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

through tory hand-washing sports envi- Week- dren	and stu-
inten- under supervision, and ron- ly visits did not	dents'
sive Hand hygiene morning broadcast, the ment by social bring	hand-
hand activities mate- parent meetings, stu- school and workers their	washing
hy- rials including: dents-parents infor- nurse) class- own	during
giene mation transfer); en- rooms as was	the day
inter- games (e.g. how sured the to escape from specific school inity that all Twice-	
the germs): the germs): that all	
tion pie-de-	Schools
puzzles; hand washing any near torysu-	created
paign active sinks nervised	own ac-
soap activities to seen factory and in hand-	tivities
(e.g. soap draw-	to im-
plant) hv- rooms required	prove
song specially giene and on by stu-	adher-
developed to cam- play- dents for	ence.
promote hand More details in Table paign ground about 45 Schools	
hygiene 1 of paper were followed greate	
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dents' activities Social worker weekly dent towel. ing a	
and methods to visits social weekly	
encourage stu- work- hand dents to prac- ers vis- hy-	
tice these activation ited	
ities. Distribution of flyers the cham-	
to parents schools. pion,	
devel-	
Doctors with oping	
Posters with theatre messages to	
wash hands plays,	
with soan and	
water upon ar-	
riving at school	
before and af-	
ter meals, after tests	
using the bath-	

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

> room, and after coughing or sneezing.

drawings and songs.

Informational flyers for parents reinforcing the messages delivered at the schools.

				30110013.									
Teesing 2021 (addi- tional sources: 2020a an	HANDSO multi- modal nurs- ing Tensine dhu ad-	MNH man- age- ment staff and nurs-	Change hy- giene policy and in- divid- ual HH	Materials for lessons about WHO-defined 5 moments for HH ^[23] using HANDSOME novel method:	See Table 1 of Teesing 2020a and Teesing 2020b for more details	Meet- ing and mate- rials pro- vided by re-	Face to face in groups (man- age- ment and	In residents' rooms or other areas of 2 units	4 months (Jan to Apr 2017)	Per- suasive com- muni- cation used to en-	None de- scribed, except that the process	Unobtru- sive HH direct observa- tion dis- guised as reg-	HH compliance (12 m f/u) IG: 36%
Teesing 2020b)	her- ence inter- ven- tion	es and nurs- ing stu- dents (with or of	behav- iour of nurses through multi- modal inter-	'Room In' (mo- ment 1), 'Room Out' (moments 4 and 5 com- bined), 'Before Clean' (moment	- management meeting (with senior nursing home manager, infection prevention specialist, and facilities are as a serior product.)	Study team mem-	nurs- ing staff)	each of 33 Dutch nurs- ing homes with ≥	Manage- ment meeting (45 to 60 min)	cour- age contin- uing when NH want-	was it- erative in re- sponse to feed- back	istering of fre- quency of health care ac- tivities record-	CG: 21% (OR 2.28, CI 1.67 to 3.11)
		3 or 4- year nurs- ing de- gree) and resi-	ven- tion de- signed specif- ical- ly for	2), and 'After Dirty' (moment 3)[24] Nurse's watch-	ties manager), - personal hygiene rules - HH materials audit	ber de- livered 3 live lessons with in- volve-	groups of maxi- mums of 18/ session	3 nurses providing intense psychogeriatric	Personal hygiene policy presen- tation	ed to stop When < 3	from indi- vidual nurs- ing homes	ed on comput- er tablet (see Fig- ure 2 in- Teesing 2020a and	HH com- pli- ance in- creased more for IG than CG
		dents	nurs- ing homes based on lit- era-	es and certifi- cates earned on completion of e-learning	2. Nursing staff interventions (The New Way of Working)i) 3 live lessons:a. introduction of	ment of se- nior NH man- ager	On- line in- divid- ual e-	and/ or somat- ic care to geri- atric resi-	Live lessons:	nurses work- ing at the unit, either		Table 3 of- Teesing 2020b)	for each WHO- defined moment, except for mo-
			ture, inter- views at nurs-	Paint for washing hands exercise	HANDSOME/WHO HH moments; teaching and discussion re HH when handling medication, food,	Senior NH man-	learn- ing	dents	1 (20 min) 2 (30 min)	the ob- servers con- tinued obser-		Com- pliance regis- tered if	ment 2



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

ing	28 stickers rep-	laundry; when to use	agers	Meet-	3 (40	vations	HH oc-	Estimat-
homes	resenting bar-	hand sanitiser/soap/	in-	ings	min) giv-	at an	curred	ed atten-
and in-	riers to HH in 4	gloves. Team HH	volved	on-site	en mul-	addi-	imme-	dance at
terven-	themes (facili-	goal-setting;	in de-		tiple	tional	diately	lessons:
tion	ties, forgetting,	b. make inventory	livery		times on	ward	before	varied
map-	choosing not to	and solutions for	of as-	Lessons	1 day	(who	(mo-	per unit:
ping	do HH, and the	barriers to HH adher-	pects,	on-site		also re-	ments	23% had
princi-	telephone)	ence; and	includ-	and		ceived the in-	1 and	< 50%
ples, the		crice, and	ing a lesson	online	E-learn-	tne m- terven-	2) or af- ter (mo-	attend-
princi-		c. exercise washing	on NH		ing: 5 to	tion)	ments	ing at
ple of	E-learning ma-	hands with paint to	per-		10 min	or they	3, 4 and	least
repe-	terials including	see where missed;	son-	Posters	each	stopped	5) a HH	1 les-
tition	videos model-	teaching how to dis-	al hy-	through-		ob-	oppor-	son, 18%
and in-	ling knowledge,	infect hands	giene	out NH		serving	tunity	had 50%
formal	guided practice	ii) a laarning, in	poli-	outivii	Adher-	338	without	to 74%
discus-	and promotion	ii) e-learning: in- troduction and 7	cy be-		ence ob-		touching	atten-
sions	of active learn-	lessons showing:	tween		server	1111	anoth-	dance at
with	ing	tessoris snowing.	lessons		training:	HH	er ob-	at least
mem-		 correct/incorrect 	1 and 2		2 to 3	need- ed to	ject (e.g.	1 les-
bers of		HH behaviour			days	hap-	door	son and
over 20	10 posters (mul-					pen	handle)	59% had
nurs-	tiple copies,	- common HH ac-	Nurs-			in the	and only	> 75% atten-
ing	new one each	tions	es and		Adher-	same	if hand	dance at
home	month)	- when to use gloves	doc-		ence	room	sanitiser	least 1
organi- sations			tors in		obser-	as ac-	or soap, water	lesson (n
in an		- food and medica-	train-		vation:	tion	and pa-	= 22).
iter-	Prize for photo	tion preparation	ing		during	OC-	per tow-	,
ative	competition	Ouizzos	pro-		obser-	curred,	el used	
process		Quizzes:	vided 		vation	except		
•		iii) reminder posters	adher-		hours (8	if a		
	NH certificate	hung throughout NH	ence obser-		am to	nurse	Hand-	
See	of good HH	showing large pic-	vation		1.30 pm,	brought a res-	related	
proto-	J	ture of hands and	and as-		week-	ident	person-	
col for		text: "Did you re-	sess-		days)	to an-	al hy-	
more	Small bottle of	member to wash	ment			other	giene ^[28]	
details	hand sanitiser	your hands?" (in				room,	for each	
of in-	for lesson par-	Dutch')				they	nurse ac-	
terven-	ticipants	iv) photo competi-				carried	cording	
tion	panto	tion: prize for best				some-	to Dutch	
map-		photo of hands				thing	guide-	
ping		•				soiled	lines ^[29] 1/	
process						or no	every	
ucinσ						door		

door

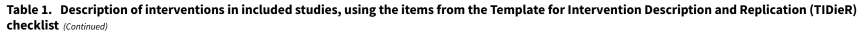
using

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

deter- mi- nants and meth- ods	See website (www.zorgvoor- beter.nl/hy- giene/hand- hygiene-ver- beteren-ver-	3. Arts and craft project for residents involving hands that NH displays
to de- velop strate- gies for inter-	pleeghuis) for materials (in Dutch) used for interven- tion: ^[25]	Adherence recording procedures
ven- tion	- Manual (84p)	Provision of hand sanitiser to lesson
com- po- nents	- E-learning module	participants
	- PowerPoint presentation and script	Provision of good HH certificate to NH if higher than average
	- Assignments	adherence
	- Awareness ac- tivities	Provision of nurse's
	- Audit materi- als	watch on completion of e-learning
	- Policy materi- als	Provision of adher-
	- Posters	ence observers train- ing
	Adherence recording ap- plication and computer table	
	Adherence observer training	

materials using method adapt-

need- ed to be	nurse / day
opened before leav- ing the room; for these in- stances, HH	Attendance at live lessons and elearning was recorded
should take place at the end of action	Participants asked if HH policy information received and if posters seen



ed from a study in Dutch hospital^[26]: videos and case studies and examination using videos from Hand Hygiene Australia^[27]

[1] World Health Organization. (2012). Hand hygiene in outpatient and homebased care and long-term care facilities: a guide to the application of the WHO multimodal hand hygiene improvement strategy and the "My Five Moments For Hand Hygiene" approach. World Health Organization. apps.who.int/iris/ handle/10665/78060 (accessed 15

[2] Moment 1 (before touching a resident) = Room In; Moment 4 (after touching a resident) and Mo-

June 2022)

ident's sur-

roundings) =

Room Out; Mo-

ment 2 (before

a clean/antiseptic procedure)

= Before Clean;

Moment 3 (af-

ter body fluid

exposure risk) –

After Dirty

[3] Hand-

some: hand-

hygiëne in ver-

pleeghuizen.:

Zorg voor

beter; 2019

May 03. URL:

www.zorgvoor-

beter.nl/hand-

some (accessed

7 June 2022)

[4] Veiligheid

en Kwaliteit:

Project Handen

uit de Mouwen.:

Stichting Sa-

menwerk-

ende Rijnmond

Ziekenhuizen

[5] Auditor

training.:

Hand Hy-

giene Australia

URL: www.h-

ha.org.au/audits/audi-

tor-training (ac-

cessed 7 June

2022)

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist

checklist	(Continued)										
Temime 2018	Mul- tifac- eted hand hy- giene pro- gramme (in- clud- ing alco- hol-based hand rub)	Nurs- ing home staff, resi- dents, visi- tors, and out- side care d providers	have an in- creased risk of per- son-to- person trans- mis- sion of	Dispensers and pocket-sized containers of hand rub solution Posters promoting hand hygiene Developed local HH guidelines eLearning module on infection control and HH training with online quizzes srequiring sufficient performance	Facilitated access to hand rub solution Campaign to promote HH with posters and event organisation Formation of local work groups in each NH Development of local HH guidelines Staff education using eLearning Monitoring of quantity of hand rub solution used	Same nurse provided HH training for all NHs. Provision of hand rub by NH Local work group developed guide- line. eLearning mod- ule and posters pre- sum- ably devel- oped by re- search team.	Provision of materials faceto-face Education and quizzes via eLearning	Nurs- ing homes in France	One-off provision of hand rub One-off eLearning repeated if unsatisfactory performance.	If staff did not score sufficiently on online quiz, they were invited to repeat the eLearning.	None de- scribed

ed mean rub soscribed. amount lution of hand used: rub sobaseline lution quantity used per of conresident sumed per day hand rub assessed solution as proxy was 4.5 for HH mL per freresident quency, per day. based on quantity Over the of hand rub solution bought by NH (which was routinely

moni-

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Estimat-

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1 year, mean quantity consumed was significantly higher in intervention NH (7.9 mL per resident per day) than control (5.7 per resident per day).

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Table 1.	Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
checklis	t (Continued)

ing	
homes.	

			nomes.										
Turner 2004a Clinical trial 1	3 active interventions (no control) Product: A. Ethanol B. Salicylic acid C. Salicylic acid with pyroglutamic acid	Healthy volun- teers	Assess the resid- ual viruci- dal ac- tivity of or- ganic acids used in cur- rently avail- able over- the- counter skin prod- ucts for the pre- ven- tion of exper- imen- tal rhi- novirus colds	1.7 mL of hand products: A. 62% ethanol, 1% ammonium lauryl sulphate, and 1% Klucel) B. 3.5% salicylic acid, or vehicle containing C. 1% salicylic acid and 3.5% pyroglutamic acid	Disinfection of hands then application of test product then allowed to dry. 15 min later, fingertips of each hand contaminated with 155 TCID ₅₀ of rhinovirus type 39 in a volume of 100 µL. Hands air-dried for 10 min. Intentional attempted inoculation with virus by contact with fingers, conjunctiva, and nasal mucosa with fingers of right hand. Left hand eluted in 2 mL of virus-collecting broth.	Re- searchers	Face- to-face indi- vidual- ly	Com- muni- ties in Mani- toba, Cana- da	1.7 mL of product applied. See What for timing	Not de- scribed	Not de- scribed	Not de- scribed	Not de- scribed
Turner 2004b Clinical trial 2	2 active interventions (no control)	Healthy volun- teers	Assess the resid- ual viruci- dal ac- tivity of or- ganic acids	Skin cleanser wipe containing: A. 4% pyroglutamic acid formulated with 0.1% benzalkonium chloride B. 62% ethanol	Application of product to hands with towelette then allowed to dry. 15 min later, fingertips of each hand contaminated with 106 TCID ₅₀	Re- searchers	Face- to-face indi- vidual- ly	Com- muni- ties in Mani- toba, Cana- da	Dose not report- ed; see What for timing Addi- tional group	Not described	Not described	Not de- scribed	Not de- scribed

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Table 1. checklist			erventior	ns in included stu	ıdies, using the items	from the	Templat	e for Inte	rvention D	escriptio	n and Re	plication (TIDieR)
CHECKUST	Skin clean- er wipe prod- uct:		used in cur- rently avail- able over- the- counter		of rhinovirus type 39 in a volume of 100 μL. Intentional attempted inoculation with virus by contact with fingers, conjunctiva,	chal- lenged 1 h af- ter appli- cation; final group							
	roglu- tamic acid B. Ethanol		skin prod- ucts for the pre- ven- tion of		and nasal mucosa with fingers of right hand. Left hand eluted in 2 mL of virus-collecting broth.				chal- lenged 3 h after applica- tion (re- mained at study site and				
			exper- imen- tal rhi- novirus colds		S C C C C C C C C C C C C C C C C C C C				not al- lowed to use or wash hands be- tween).				
Turner 2012	An- tiviral hand lotion	Healthy adults	Re- duce rhi- novirus infec- tion and ill- ness through hand disin-	Lotion containing 62% ethanol, 2% citric acid, and 2% malic acid	Provision of lotion and instructions for use Meetings with participants to check compliance	Staff of study site pre- sum- ably sup- plied lotion.	Face- to-face and pre- sum- ably in- divid- ually, but not speci- fied	Study site at uni- versity com- munity in the USA	9 weeks Every 3 hours whilst awake and after hand- wash-	None report- ed.	None report- ed.	Self-re- port- ed dai- ly diary of time of each product applica- tion	"All subjects applied at least 90% of the expected amount of hand treatmen-
			fection with ethanol and or- ganic acid sanitis- er			Study site staff met with partici- pants.			ing for 9 weeks Com- pliance meet- ings			Twice week- ly for 5 weeks then week- ly meet- ings with	t" (p. 1424)

oppor-

tunities

during

30%

	: (Continued)								twice weekly for first 5 weeks then week- ly meet- ings with partici- pants			partici- pants to reinforce com- pliance with treat- ment	
Yeung 2011	Mul- tifac- eted hand	Long- term care fa- cilities	Pro- mote use of alco-	Free supply of pocket-sized containers of alcohol-based	Provision of materials	Study team deliv- ered	Deliv- ered face- to-face	LTCFs in Hong Kong	7 months overall	Re- place- ment of	As ad- her- ence dropped	Direct observa- tion of HCW ad-	90% atten- dance o seminar
	hy- giene pro- gramme (in- clud- ing alco- hol-based hand rub)	and their health- care work- ers	hand rub by staff in LTCFs as an effec- tive, time- ly, and low-ir- ritant	antiseptic hand rub (either WHO formu- lation I (80% ethanol) or II (80% propanol) carried by each HCW (supplier: Vickmans Labo- ratories)	Provision of hand hygiene seminars to HCWs covering: indications, proper method, and importance of antiseptic hand rubbing and washing according to WHO 2006a) guidelines	the materials, seminars, and observer training.	and in- dividu- ally for hand rub and pens; not de- scribed if edu- cation was in-	Posters post- ed in com- mon areas.	Initial 2-week inter- vention period, then 7 months of hand rub pro- vision and re-	hand rub as re- quired	off in the middle months, the feed-back session was delivered.	herence to hand- wash- ing and antisep- tic hand rubbing (record- ed sep- arate- ly and anony-	Hand rubbing with gel increased significantly from 1.5% to 15.9%.
			method of hand hy- giene in a	Replacement hand rub as re- quired	Provision of feed- back session	Admin- istra- tive staff of LTCF	divid- ually or by group, but semi-	ence obser- vations oc- curred in	minders 3 identical sem-			mously) during bedside proce- dures or physical	Hand- wash- ing de-
			high- risk en- viron- ment	Hand hygiene seminar con- tent	Direct, unobtrusive observation of hand hygiene adherence	provid- ed re- place- ment hand	nar im- plies as a group	com- mon rooms and resi-	inars at start of inter- vention; each			contact with res- idents	creased signif- icant- ly from 24.3% t
				Reminder materials (3 to 5	Training of observa- tion staff	rub and com- muni-		dent rooms but not	staff mem- ber to			3300 hand hy- giene	17.4%. Contro

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

CNECKLIST						6 registered nurses conducted direct observation of adherence after 2-hour training (100% interrater reliability).		let areas.	Feed-back session 3 months after start of intervention 2-hour training of observers Adherence observations either 9 am to 12 pm or 3 pm to 6 pm, 1 LTCF at a time			248.5 hours of observa- tion on 92 days	Overall hand-washing adherence increased from 25.8% to 33.3%.
Zomer 2015	Hand hy- giene prod- ucts and train- ing	Day- care centres and their care- givers (staff)	Re- duce infec- tions in chil- dren attend- ing DCCs through im- proved access	HH products: dispensers for paper tow- els, soap, alco- hol-based hand sanitiser, and hand cream, with refills for 6 months	Provision of free HH products spon- sored by SCA Hy- giene Products, Swe- den. Provision of posters and stickers for chil- dren and staff	Study team arranged supply of HH prod- ucts and pre- sum- ably pro- vided	Prod- ucts provid- ed to DCCs in per- son for staff use.	DCCs in re- gions of the Nether- lands	6 months overall Initial one-off supply of products	Re- place- ment hand hy- giene pro- vided as re- quired.	None de- scribed.	6-month follow-up observation of whether intervention dispensers and posters/stickers in use	2 DCCs did not use any HH products. Sanitiser products used in at least 1 of 2 groups



		-		-		-	-
to HH	Reminder	Provision of training	train-	train-	3 train-		
mate-	posters and	about RIVM 2011 for	ing.	ing not	ing ses-		
rials	stickers for chil-	mandatory HH ^[31]	Ü	speci-	sions		Surve
(Zomer	dren and DCC	,		fied.	with 1-		of DC

to HH mate- rials (Zomer 2013a)	posters and stickers for chil- dren and DCC caregivers	about RIVM 2011 for mandatory HH ^[31]	train- ing.
(Zomer 2013a) and compliance of their DCC caregivers to hand hygiene guidelines based on socio-cognitive and environmental determinants of caregivers' HH be-	dren and DCC	Distribution of training booklet Team training sessions aimed at goalsetting and formulating HH improvement activities (Erasmus 2011; Huis 2013)	
hav- iour ^[30] (Zomer 2013b)			

3 training sessions with 1-month interval	Survey of DCC care- givers	in 94%, 89%, 86%, and 45% of inter- vention DCCs.
2 team	НН	

training

sessions

ΗН guide-Posters lines used in compli-86%, ance obstickers served in 74%. at 1, 3, and 6 months' DCC surfolvey relow-up: sults:

no. of 79% at-HH actended tions/no. of opportunities

at least 1 training session; 77% received ΗН guidelines booklet.

HH compliance at 6 months:

IG: 59% vs CG: 44% (Zomer TP, et al, Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

> unpublished data)

> > All intervention DCCs received guidelines training; all but 2 received at least 1 team training.

Hand hygiene and masks

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Aelami 2015	Hy- gien- ic edu- cation and pack- age	Reli- gious pil- grims	Prevent in- fluen- za-like illness by re- duced infec- tion trans- mis- sion through per- son- al hy- giene mea- sures	Hygiene package of: alcohol-based hand rub (gel or spray) surgical masks soap paper handkerchiefs user instructions	Not clearly described, but it appears that packages may have been distributed by trained physicians before departure to or on site of country of pilgrimage	Not specif- ical- ly de- scribed	Not described, but it appears that packages were distributed faceto-face and individually	Not described if before departure (from Iran) or on site (in Saudi Arabia)	One-off during Hajj sea- son	Not de- scribed	Not de- scribed	Not described	None de- scribed
Aiello	2 ac-	Stu-	Re-	7 face masks	Weekly supply of masks through stu-	Not de-	Educa-	Uni-	One-off	Mask	Uni-	Week-	Average
2010	tive in-	dents	duce	(standard med-		scribed,	tion via	versi-	educa-	wear-	versity	ly web-	mask

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Trained staff



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(Continued) terven-	in uni-	inci-	masks with		edu-	and	dence	weeks	during	break	student	hours/
tions:	versi- ty resi- dences	dence of and miti-	ear loops TEC- NOL procedure masks; Kimber-	Provision of basic hand hygiene edu- cation through an	cation provid- ed via	study web- site; provi-	halls in the USA	(ex- cluding spring	sleep option- al and	oc- curred during	survey includ- ed: self-	day: FM + HH 2.99 ver-
A. Face mask (FM) B. Face		gate ILI by use of non- phar- ma-	ly-Clark) 7 re-sealable plastic bags for mask storage	email video link, the study website, and written materials; instruction to wear	study web- site (URL not	web-sion of site masks (URL and not sani-		of face mask and/or	en- cour- aged out- side	weeks 4 and 5 of the study, with	reported average number of times hands	sus FM 3.92
mask and hand hy- giene		ceuti- cal in- terven- tions of per- son-	when not in use (e.g. eating) and for disposal	mask as much as possible; education in correct mask use, change of masks daily, use of provided re-sealable bags for	provid- ed) "Trained	tiser in person to resi- dences		giene mea- sures which com- menced	of residence.	most stu- dents leaving cam- pus	washed/ day and average duration of hand- washing	Average hand- washing times/ day:
(FM + HH)		al pro- tection mea- sures	Alcohol-based hand sanitiser (62% eth- yl alcohol in	mask storage and disposal	staff" for com- pliance moni-			at "the begin- ning of the in-		and trav- elling; they	to obtain compos- ite "op- timal	FM + HI 6.11 ver sus FM 8.18 vs
			a gel base, portable 2- ounce squeeze bottle, 8-ounce pump)	Provision of replace- ment supplies which students signed for upon receipt	toring Study-			fluenza season just af- ter iden- tification		were not re- quired to con- tinue	hand- wash- ing" score (at least 20 s	control group 8.75
			Hand hygiene		affiliat- ed res- idence			of the first case of in-		pro- tective mea-	≥ 5/day); average no. of	Daily wash- ing sec
			education (proper hand hygiene prac-		hall staff provid- ed re-			fluenza on cam- pus" (p.496	6).	sures at that time.	mask hours/ day/	onds/d FM + HI
			tices and cough etiquette) via emailed video, study website, written materi-		place- ment sup- plies.			Replace- ment supplies			week; average hand sanitiser use/day/	20.65 ver- sus FM 23.15 v control
			als detailing ap- propriate hand sanitiser and mask use					provided as need- ed.			week and amount used.	22.35
			mask usc								useu.	Hand sanitis er use

FM + HH:

5.2 versus FM

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

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:	stances
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2.31 vs control 2.02 No. of proper mask wearing participants/hour of observation:

FM + HH 2.26 versus

													FM 1.94
Aiello 2012	2 interventions:	Stu- dents living in uni- versi-	Pre- vent ILI and labo- rato-	Packets of 7 standard med- ical procedure masks with ear loops (TEC-	Intervention materials and educational video provided.	Trained study staff avail- able at	Hy- giene packs deliv- ered	Uni- versi- ty resi- dence halls	One-off educa- tional video at start	Stu- dents en- cour- aged	1-week uni- versity spring break	Weekly student survey includ- ing com-	Self-re- ported mask wearing: no sig-
	A. Face mask (FM) B. Face mask and hand sanitis- er (FM + HH)	ty resi- dences	ry-con- firmed in- fluenza by use of non-	NOL procedure masks, Kim- berly-Clark, Roswell, GA, USA) and plas- tic bags for stor-	Supply of masks and instructions on wearing	tables in each resi- dence hall for surplus	to stu- dent mail- boxes; face- to-face	USA s;	Weekly supply of hygiene	but not oblig- ed to wear masks out-	dur- ing the study when ma- jority	pliance (e.g. masks hours/ day, fre- quen-	nificant differ- ence
				phar- ma- ceuti- cal in- terven- tions of per- son-	terruptions in euti- mask use (e.g. al in- whilst eating, erven- sleeping) and ons for daily dispos- f per- al	Provision of replace- ment masks or sani- tisers as needed on site	masks and sanitis- er and for ob- serving com-	supply also avail- able	-	Masks to be worn at least 6 hours/	side of stu- of resi- dents dence left hall. cam- pus	dents left cam-	cy and amount of sani- tiser use, number of hand wash-
			al pro- tection			pliance			day			es/day, duration of hand-	FM or control groups

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Table 1.	Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
chacklis	Continued)

checklis	t (Continued)												
Hecklis	(Conunued)		mea- sures (e.g. face masks and hand hy- giene)	Hand sanitiser (2-ounce squeeze bottle, 8-ounce pump bottle with 62% ethyl alcohol in a gel base) Replacement face masks and hand sanitiser Educational video: proper hand hygiene and use of standard medical procedure face masks					Study staff available onsite with replacement supplies as needed for duration of intervention (6 weeks, excluding spring break)			washing (seconds) Observed compliance completed by trained study staff who daily and anonymously observed mask wearing in public areas of resi-	More results in S1 of paper. Staff observed an average of 0.0007 participants properly wearing a mask for each hour of observation.
Cowling 2009	2 ac- tive in- terven- tions in ad- dition to con- trol of lifestyle educa-	House-hold-ers with index patient with in-fluenza	Re- duce trans- mis- sion of in- fluen- za in house- holds	A. and B. Liquid soap for each kitchen and bathroom: 221 mL Ivory liquid hand soap (Proctor & Gamble, Cincinnati, OH, USA)	Provision of soap, hand rub, and masks as applicable and when to use them	Trained study nurse provided interventions.	Face- to- face to house- hold- ers	House- holds in Hong Kong	Initial home visit sched-uled within 2 days (ideal-ly 12 h) of in-	Not de- scribed	Not de- scribed	Monitoring of adherence during home visits	Most initial visits completed within 12 h.
	A. En- hanced hand hy-		through per- son- al pro- tective mea- sures	Alcohol hand rub in individ- ual small bot- tles (100 mL) WHO recom-	HH: education about efficacy of hand hygiene Demonstration of proper hand-wash-				dex case identification. Further home			tion of adher- ence on final vis- it by in- terview or self-	groups "reported higher adherence than the

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Table 1.	Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
checklis	st (Continued)

=	erventions in included stu	idies, using the items from the Template for Int
t (Continued)		
giene (HH)	mended formu- lation I, 80% ethanol, 1.45% glycerol, and	ing and antisepsis techniques
B. Face masks and en- hanced hy- giene (FM + HH)	0.125% hydrogen peroxide (Vickmans Laboratories, Hong Kong, China)	+ FM: education about efficacy of sur- gical face masks in reducing disease spread to household contacts if all parties wear masks
	of 50 surgical face masks (Tecnol–The Lite One (Kim- berly-Clark, Roswell, GA, USA) to each	Demonstration of proper wearing and hygienic disposal
	household member or C. Children 3 to 7: box of 75 paedi- atric masks	All groups: provision of education about the importance of a healthy diet and lifestyle, both in terms of illness pre- vention (for house- hold contacts) and symptom alleviation
		(for the index case)

visits day 3 and 6, 7- day fol- low-up		
HH: use of liquid soap after every washroom visit, sneezing or coughing, when their hands were soiled. Use rub when first returning home and immediately after touching any potentially contaminated surfaces		
FM: masks worn as		

often as

reportcontrol ed pracgroup. Self-retices and portcounted daing of amount ta were of soap consisand rub tent with left in meabottles sureand rements of maining amount masks of soap, for FM alcohol hand group rub, and face masks used" (p.443) (see Table 6 in paper). "Adherence to the hand hygiene intervention was slightly higher in the hand hygiene group than the face mask

plus hand hygiene group."

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

CHECKISI	. (Continued)								possible at home (except eating or sleeping) and when the index patient was with the household members outside of the household				Median masks used: Index: 9 Contact: 4 More details in paper and Appendices
Larson 2010	2 active interventions in addition to control of URI education: A. Alcohol-based hand sanitiser (HS) B. Face masks and hand	His- panic house- hold- ers with at least 1 preschoo or ele- men- tary school child	Re- duce inci- dence and sec- ondary I trans- mis- sion of URIs and in- fluenza through non- phar- ma- ceu- tical house- hold level inter-	A. and B. 2-month supply of hand sanitiser in 8-, 4-, and 1-ounce containers: PURELL (Johnson & Johnson, Morris Plains, NJ, USA) B. 2-month supply of masks: Procedure Face Masks for adults and children (Kimberly-Clark,	Provision of materials and instructions for when to use including demonstration of use and observation of return demonstration by householder A. Mask worn when householder had: "temperature of ≥37.8°C and cough and/or sore throat in the absence of a known cause other than influenza" (CDC definition of influenza-like illness at the time).	trained bilingual research assistants (RAs) with minimum baccalaureate degree and experience in community-based research;	Face- to- face to house- hold- ers	House- holds in New York, USA	19- month fol- low-up Initial home visit, then at least every 2 months Sanitiser for use at home, work, and school	Change masks be- tween inter- actions with person with ILL House- hold- ers' ques- tions and mis- con- cep- tions ad- dressed	None de- scribed.	RA home visits for adherence with random accompaniment by project manager, who also made random calls to householders Telephone calls to reinforce	Sanitis- er use (mean ounces/ month) HH: 12.1 FM + HH: 11.6 Mask com- pliance was "poor": 22/44 (50%) used within 48 hours of onset.

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Penlication (TIDIOP)

	_		erventior	ns in included stu	udies, using the items	from the	Templat	e for Inte	rvention D	escriptio	n and Re	olication (ΓIDieR)
checklis	t (Continued) sanitis- er (FM + HS)		ventions	Roswell, GA, USA) Replacement supplies at least once every 2 months Disposable thermometers Educational materials about URI prevention, treatment, and vaccination (written in Spanish or English language)	Home visits to reinforce adherence, replenish supplies and record use, answer questions B. Telephone calls to reinforce mask use All groups received URI educational materials.	procedures were practised with each other until demonstrated proficiency			B. Telephone calls days 1, 3, 6 Masks worn for 7 days when within 3 feet of person with ILL or no symptoms.	on home visits.		mask use Used bottles or face masks, or both, moni- tored for usage.	Mask users re- ported mean mask use of 2.
Simmerman 2011	2 ac- tive in- terven- tions: A. Hand- wash- ing ed- uca- tion and hand- wash- ing kit (HW)	House-holds with a febrile, in-fluen-za-pos-itive child	De- crease in- fluenza virus trans- mis- sion in house- hold with a febrile in- fluen- za-pos- itive child through pro- moted	A. and B. Hand-washing kit per household including graduated dispenser with standard unscented liquid hand soap (Teepol brand. Active ingredients: linear alkyl benzene sulfonate, potassium salt, and sodium lauryl ether sulphate)	A. and B. Provision of intensive hand-washing education on initial home visit to household members with 5 approaches: discussion, individual hand-washing training, self-monitoring diary, provision of soap, and provision of written materials (Kaewchana 2012) Individual handwashing training	Study nurse con- ducted home visits, pro- vided edu- cation and moni- toring activi- ties.	Edu- cation pro- vided face- to-face as a group to house- hold mem- ber and in- dividu- ally for hand- wash- ing	In homes (in Bangkok, Thai- land)	One-off provision of kits at initial home visit conducted within 24 hours of enrolment Subsequent home visits on	B. No face masks whilst eating or sleeping as impractical and could hinder breathing in ill child	None de- scribed.	Self- monitor- ing diary record- ing hand- washing frequen- cy > 20 s and face mask use for that group Rein- force- ment	Report- ed av- erage hand- washing episodes/ day: HW: 4.7 HW + FM: 4.9 Par- ents had highest frequen- cy (5.7), others (4.8),

lings 17

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!	Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
	checklist (Continued)

ו. Description o ist (Continued)	or interventio	ns in included sti	udies, using the items fror	n the Template to	r intervention D	escription and	i Replication (іріек)
B. Hand- wash- ing ed- uca-	use of hand- wash- ing or hand-	Replacement soap as needed	("why to wash", "when to wash", and "how to wash" in 7 hand-washing steps described in Thai-	train- ing.	days 3, 7, and 21	lm- promp- tu edu- cation and	of mes- sages by nurses on sub- sequent	siblings (4.3), index cases (4.1).
tion, hand- wash- ing kit, and face masks (HW + FM)	wash- ing with face mask use	Written materials from education including pamphlets and posters attached near sinks in household.	land Ministry of Public Health guidelines) B. Provision of education of benefits of and appropriate face mask wearing	supply of hand washin supplie 30- minute educa-	of hand- washing supplies 30- minute educa-	train- ing provid- ed by nurs- es as ques- tions arose.	Amount of house-hold liquid soap and	Average soap used/week: HW: 54 mL/person
		B. Box of 50 standard paper surgical face masks and 20	Soap replaced as needed.		tion pro- vided at initial home visit		number of face masks used	HW + FM: 58.1 mL/ person
		paediatric face masks	More details (Kaew- chana 2012)					B. Mask use:
		(Med-con com- pany, Thailand						12/per- son/week
		#14IN-20AM- B-30IN)						Mask wearing medi- an min- utes/day: 211
								Parents 153,
								other re- lations
								59, index patients 35, sib-

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

checklist	(Continued)												
Suess 2012	2 active interventions in addition to written information: A. Mask/hy-giene (MH) B. Mask (M)	House-holds with an in-fluen-za-pos-itive index case in the absence of further respiratory illness within the preceding 14 days	Prevent in- fluenza trans- mis- sion in house- holds through easily applic- able and acces- sible non- phar- ma- ceuti- cal in- terven- tions such as face masks or hand hy- giene mea- sures	A. Alcohol-based hand rub (Sterilium, Bode Chemie, Germany) A. and B. Surgical face masks in 2 different sizes: children < 14 years (Child's Face Mask, Kimberly-Clark, USA) and adults (Aérokyn Masques, LCH Medical Products, France) Written information provided on correct use of intervention and on infection prevention (Suess 2011) (tips and information on the new flu A/H1N1) (URL provided is no longer active) Digital tympanic thermometer	A. Provision of hand rub and masks A. and B. Provision of masks only Provision of thermometer and how to use it Mask fit assessed (at first household visit) Information provided by telephone and written instructions at home visit on proper use of interventions and recommendations to sleep in a different room than the index patient, not to take meals with the index patient, etc. (Suess 2011) In-person demonstration of interventions at first home visit All participating households received general written infor-	Study per- sonnel arranged provi- sion of mate- rials, rang the partici- pants, visit- ed the homes, demon- strated and as- sessed fit of masks.	Provision of materials in person to house-holds Initial telephone delivery of information Face-to-face home visits	House-holds in Berlin, Germany	Over 2 consecutive flu seasons Day 1 house-holds received all necessary material instructions. House-hold visits no later than 2 days after symptom onset of the index case, then days 2, 3, 4, 6, 8 (5 times) or on days 3, 4, 6, 8 (4 times) depending on the day of recruitment	Adult masks worn if masks for under 14-year-olds did not fit properly. If other house-hold members developed fever (> 38.0 °C), cough, or sore throat, they were asked to adopt the same preventive behaviour as the index patient.	In the season 2010/11 participants also recorded number of masks used per day.	Self-re- ported daily ad- herence with face masks, i.e. if they wore masks "al- ways", "most- ly", "some- times", or "nev- er" as in- structed. Partici- pants of the MH house- holds addi- tional- ly not- ed the number of hand disinfec- tions per day. Exit ques- tionnaire about (preven- tive) be- haviour during	Face mask use (median/individual): MH: 12.6 M: 12.9 Daily adherence was good, reaching a plateau of over 50% in nearly all groups from the third day on. MH hand rub use (median): 87 mL (Suess 2011) MH mean frequency of daily hand

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

mation on infection prevention.

General written information on infection prevention

Hand rub use: after direct contact with the index patient

(or other symptomatic household

members), after atrisk activities or contact^[31]

toms

were to-

gether in

1 room

Mask use: at all times when index patient and/ or any other ing the household member housewith respiratory symp-

the past disinfec-8 days, tion: 7.6 (SD 6.4) general attitudes times towards per day NPI, the

actual amount See paof used per and interven-Suess tion ma-2011 for terials,

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able, problems with wearing

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Used intervention material per household member was calculated

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by dividamount used per hold by the numhouse-



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

> Regular change of face masks, not worn during the night or outside the household

See paper and Suess 2011 for more details.

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Hand bygi	iono and	surface/o	hioct disin	faction									
2015	iene and Hand hy- giene and surface clean- ing or disin- fection	surface/ol Kinder- gartens and the fami- lies of their stu- dents	Re- duce trans- mis- sion of infec- tion in young chil- dren from conta- minat- ed sur- faces or hands through hand hy- giene and surface clean- ing or disin- fection	Antibacterial products for hand hygiene and surface cleaning or disinfection: liquid antimicrobial soap for hand-washing (0.2% to 0.3% parachlorometaxy Instant hand sanitiser for hand disinfecting (72% to 75% ethanol), antiseptic germicide (4.5% to 5.5%	Provision of products to kindergartens and families Instruction of parents or guardians and teachers in hand hygiene techniques and use of antibactedial products Daily cleaning of kindergartens with products At least twice/week viewining of homes and weekly cleaning or disinfecting of items such as children's toys, house furnishings, frequently touched ob-	Re-search team pro- vided prod- ucts and in- struc- tions and moni- toring.	Materials provided to kindergartens and families in person and presumably instructions in person to families and staff.	In kinder-gartens (hard sur-faces) and families' homes (Xi-antao, China)	1 year overall Daily hand-washing with soap before eating, after using bathroom, nose blowing, and outdoor activities Hand sanitiser carried daily.	Families and teachers could contact study management at any time as needed. Exchange of empty bottles for new ones at any	Not de- scribed	Close contact with teachers and families for monitoring, e.g. unscheduled parents' meetings, quarterly home visits, phone interviews, and monthly cell phone messages	Consumption of products by person (mL/person/day). Liquid soap: 7.7 Sanitiser: 1.4 Bleach: 25.0 Antiseptic-germicide: 12.5

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lunch



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) che

				use) for surface disinfecting. Produced by Whealth- fields Lohmann (Guangzhou) Company Ltd.	tables or desks), kitchen surfaces (utensils, cutlery, countertops, chop- ping boards, sinks, floors, etc.), bath- room surfaces (toilet, sink, floor, etc.)				Kinder- garten cleaning daily Home cleaning at least twice/ week			Month- ly survey of con- sump- tion of products by vol- ume, to- tal us- age, per- son us- age	
Carabin 1999	Hy- giene pro- gramme	Day- care centres and their staff and chil- dren	Reduce infections in at-risk children (under 3 years old) in DCCs with inexpensive, easily implementable and practical interventions	Hygiene materials and documents, e.g. colouring books, hand-washing posters, hygiene videotapes Materials for training Reimbursement of equivalent of 1 full-time educator's salary Bleach (diluted 1:10) for toy and play area cleaning	Provision of comprehensive hygiene training session to entire DCC staff, especially the educators of participating classrooms Training in recommendations for hygiene practices: i. toy cleaning ii. hand-washing technique and schedule iii. use of creative reminder cues for hand-washing iv. open window for daily period v. sandbox and play area cleaning	Training appears to have been provided by study team.	Ap- pears staff trained as a group, i.e. "entire DCC staff"	Day- care cen- tres in Cana- da Loca- tion of train- ing not de- scribed, except may have been off-site from DCCs since 1 DCC did not "send" staff to train- ing.	15- month trial One-off 1-day training Toy cleaning at least every 2 days Hand- wash- ing at least af- ter DCC arrival, after outside play, af- ter bath- room, before	Teachers to use creative reminder cues for handwashing with children	Not de- scribed	Fol- low-up tele- phone ques- tionnaire for DCC directors about follow- ing train- ing rec- ommen- dations	Use of materials: colouring book: 22/24 poster: 23/24 videotapes: 18/24 staff meetings: 19/24 Increased frequency of toy cleaning: 6/24 Use of rake and shov-

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR	1)
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				Payment of salary of educator for the day to encourage participation DCC meetings to discuss training session with all staff				Open windows at least 30 min/ day				sandpit: 17/24 Frequen- cy of cleaning sandbox 14/24
								cleaning of sand- box/play area				
Kotch Hy- 1994 giene	Care-givers at child day-care centres (CD-CCs)	Develop feasible, multi component hygienic intervention to reduce infections in children at CDCCs who are at increased risk	Hygiene curriculum for caregivers Availability of soap, running water, and disposable towels Waterless disinfectant scrub (Cal Stat) used only if alternative was not washing at all. Handouts posted in CDCC.	Delivery of hygiene curriculum to caregivers through initial training session which required demonstration of participants' handwashing and diapering skills Local procedures: Hand-washing of children and staff Disinfection of toilet and diapering areas Physical separation of diapering areas from food preparation and serving areas Hygienic diaper dis-	Re-search team delivered training. Scrub donated by Calgon Vetal Laboratories.	Face- to-face train- ing and fol- low-up group and in- dividu- ally	Class- rooms of child day- care centres in the USA	8 months overall 3-hour initial training session Cleaning sched- ules as de- scribed in col- umn What (proce- dures) On- site fol-	Follow-up sessions addressed questions and local adaptations to procedures. As-required induction training	During in- terven- tion, re- search team en- cour- aged direc- tors to ad- dress phys- ical barrier to hy- giene prac- tice, such as dis- tance be- tween	Follow-up sessions reinforced training. Meeting with directors 5 weekly unobtrusive recorded observation by training staff	Rate of compliance to barrier modification was better in younger centres, which were more likely to have written guidelines.

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checklist (Continued)		sinks, kitchen and bathroom floors				1 week and 5		ing ar- eas		
		Daily laundering of blankets, sheets, dress-up clothes				weeks later		and sink ac- cess in		
		Hygienic preparation, serving, and clean up of food						rooms.		
		Separate training of food handlers								
		As-required induc- tion training for new staff								
		Onsite follow-up training reinforcing adaptations, demonstrations and discussion of hygiene techniques, responding to questions, and review of handouts								
		Monthly meeting with centre directors to encourage leadership and support								
Coneghy tifac- ing d 2017 eted homes e	Re- Education and luce launch materi- expo- als	Pre-intervention: NH administrators required to:	Study per- sonnel	Face- to-face inter-	Nurs- ing homes	6 months overall:	Sites could use ex-	2 sites re- trained	Cloud- based audit	Online trainir partici

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Table 1. checklist

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ing in-	per-	ule for certified	In Prevention" cham-	and	ning	and at		ucts	pation	login	for 3/5
terven-	son-per-	nursing assis-	pion and team	tools	and	unit/	1-hour	from	rate.	to web	sites,
tion	son	tants about: in-		and	some	team	launch	anoth-		browsers	120/
	trans-	fection preven-	- allow all staff par-	sup-	as-	levels	event	er ven-		on NHs'	13% and
	mis-	tion, product,	ticipation in educa- tion	port.	pects			dor		existing	23% for 2/5
	sion in	and monitoring	tion		and			and fill		comput-	2/3
	high- risk fa-		- iPad use for staff in		deliv-	Online	1 or 2	in any		ers or via iPads in-	
	cility of		each floor or com-	NH	ery of prod-	train-	hygiene	gaps with		cluded	
	close	"Essential bun-	munity	staff	ucts	ing	moni-	study		week-	Admin-
	envi-	dle" of hygiene		(e.g.	ucts		tors/site	prod-		ly prod-	istrators
	ron-	products sup-	- ask staff to incorpo-	cham-				ucts.		uct con-	demon-
	ment	plied at no cost:	rate intervention into workflow	pion,				0.000		sump-	strated
	and	- hand sanitiser	WORKIIOW	hy- ·	Some		1 cham-			tion to	high fi- delity in
	poten-	gel and foam		giene	as-		pion/site	Marri		get mea-	report-
	tially	ger and roam		mon-	pects deliv-			New staff		sure:	ing mea-
	conta-	- antiviral facial	Delivery of 3 compo-	itors, nurs-	ered			provid-			sures of
	minat-	tissues	nents:	ing as-	online		1-hour	ed with		week-	
	ed sur-	l C	- education	sis-	(e.g.		online	educa-		ly count of prod-	hand-
	faces	- disinfecting	- education	tants)	nurs-		module	tion, as		uct units	washing
	through mul-	spray	- cleaning products	deliv-	ing		for se-	need-		con-	(> 80%
	tifac-	- hand and face		ered	mod-		lected	ed and		sumed	of time).
	eted	wipes	- compliance audit	as-	ules,		nursing	came		x no. of	
	inter-		and feedback	pects	com-		assis-	on-		hand hy-	
	ven-	Plus additional:		of in-	pliance		tants	board.		giene oc-	Hand-
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	equip-	and wipe prod-	Education:	tions	ing)						rates in
	ping	ucts		after			iPads	Re-			Figure
	staff to		Launch event for all staff to publicise pro-	spe- cific			for each	train-			1B in pa-
	protect		gramme and explain	train-			commu-	ing of			per re-
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	dents	iPads for com-	10103	6*			floor	with			as rei- ative-
	from	pliance audits	Intensive training of					low			ly con-
	infec- tion		"hygiene monitors"					train-			stant"
	with-		for data collection				Weekly	ing			and "not
	in the	Newsletters for	and compliance au-				telecon-	partici- pation			ideal
	"cul-	support during	dit and feedback tool				ferences	rates			in the
	ture"	intervention	Training of site					iates			first few
	of care		champion				initial-				months",

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Training of select group of certified

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	(Continued)				nursing assistants (online module)				creased in fre- quen- cy over				signif- icant- ly over time.
					Audit and feedback activities				time.				cime.
					Ongoing support during intervention: - newsletter with best practices - teleconferences with each NH - "onboarding" education of new staff				Week- ly mea- sure- ment of prod- uct con- sump- tion				
Sando- ra 2008	Multi- facto- rial in- terven- tion,	Ele- men- tary school and	Re- duce trans- mis- sion of	1 container of disinfecting wipes (Clorox Disinfecting Wipes (The	Sanitiser and wipes provided to class-room/teacher with instructions for use.	Re- search team arranged supply	Prod- ucts provid- ed to schools.	Ele- men- tary schools and	8-week period	Prod- ucts replen- ished as	None de- scribed.	Individ- ually la- belled contain- ers col-	Product usage: average wipes used/
	includ- ing alco- hol-based hand sanitis-	its stu- dents	infec- tions in school- child- ren through	Clorox Company, Oakland, CA, USA); active ingredient, 0.29% quaternary ammoni-	Teachers disinfected desks once daily.	of ma- terials and in- struct- ed teach-	In- struc- tion	their class- rooms in the USA	Desks disin- fected once a day.	need- ed.		lected every 3 weeks from the class- room to	week: 897 (128 wipes/ class- room/w
	er and surface disin- fection		im- proved hand hy- giene	um chloride compound)	Hand sanitiser to be used: before and after lunch, after use of	ers on use.	pro- vided face- to- face to					assess adher- ence.	Average bottles of hand

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cohol-based hand sanitis-

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sanitisers were not

placed there), after

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

> foaming hand sanitiser (DEB SBS Inc, Stanley, NC, USA, for The Clorox Company); active ingredient, 70% ethyl alcohol)

any contact with potentially infectious secretions (e.g. after exposure to other ill children or shared toys that had been mouthed)

lecting empty containers and distributing new product.

Receptacle in classrooms for empty contain-

tion preven-

ers

Quarantine/Physical distancing

Helsin- gen 2021	Rapid-Cycle Re-Im-ple- menta- tion of TRAin- ing Fa- cilities in Nor- way (TRAiN) hy- giene and physi- cal dis- tanc-	Members of health and fitness training facilities aged 18 to 64 years not at increased risk for severe	Enable safe re- open- ing of fitness train- ing fa- cili- ties to main- tain health and fit- ness by reduc- ing the risk of SARS-	Infection mitigation measures described by "Norwegian guidelines for Hygiene and Social Distancing in Training Facilities during the COV-ID-19 Pandemic" (in Norwegian t-i.no/wpcontent/uploads/2020/04/Br sjestandard-forsentre.pdf)	Implementation of the following during regular floor training facilities and group classes: - avoidance of body contact - 1 metre distance between individuals, - 2 metre distance for high intensity activities can-	Facility employees controlled access and enforced implementation of guidelines and procedures	Face- to-face indi- vidual- ly and as a group	5 health and fit- ness train- ing fa- cili- ties in Oslo, Nor- way	3 weeks May 22nd to June 15th, 2020 Hours of access not re- ported; presum- ably the partic- ipants	Masks not required, so were option- al Change rooms avail- able Access	None de- scribed	Staff moni- tored ac- cess and distanc- ing No ap- parent mea- sures of fidelity	None de- scribed
	tanc- ing mea- sures	severe COV- ID-19	SARS- CoV2 trans- mis- sion	See Supple- mentary Appen- dix for "Stan- dard for COV- ID-19 infec-	Provision of disinfectants at all workstations Requirement of HW and cleaning of all	dures at all times Staff present			ipants had un- limited access to train- ing facili- ty within the pro-	con- trolled to avoid over- crowd- ing			

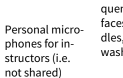
dur-

equipment by mem-

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eplication (TIDieR) Table 1. Description of interventions checklist (Continued)

tion measures	bers before and after	ing all	for dis-	Staff
in fitness cen-	use with utensils pro-	open-	tancing	moni-
ters during the	vided	ing		tored
TRAiN-study"		hours		that
,				dis-
				tance
	No physical contact			mea-
Disinfectant	between participants	Not re-		sures
readily avail-	or participants and	ported		were
able at work-	instructors	if train-		en-
stations and		ing		sured
strategic places		need-		
(reception, booking sta-	Regular cleaning of	ed for facility		
tion, changing	facilities by facility	staff		Num-
rooms, toilets,	employees			ber of
water taps used				people
for drinking or				attend-
refilling bottles)	Create lists of what			ing de-
	should be cleaned			pend-
	and how often			ed on
Rubbish cans				size of
without lids				gym
Without has	Disinfection of in-			and as-
	structor micro-			soci-
	phones			ated
Washbasin with	priories			chang-
soap or hand disinfection				ing rooms,
	Extra cleaning of fre-			show-
	quently touched sur-			ers and
	faces (e.g. door han-			toilets.
Personal micro-	dles, card readers,			Facility



dles, card readers, washbasin batteries)

Frequent refilling at all hygiene stations

Infection preventive measures reminders online and via posters in facilities

Avoid queuing by making sure group classes do not start and stop at same

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

time and keep 15 min minimum between group classes

Access control by facility employees

Closure of showers and sauna but changing rooms open

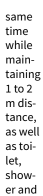
Staff presence during all opening hours

Removal of lids on trash cans

Reminders of infection preventive measures

Communication to members about changes to training for social distancing

Advice to members to stay home if any COVID-19 related symptoms



change

room capacity

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

Advice to members to avoid touching eyes, nose and mouth

					mouth								
					Closure of childcare facilities								
Miyaki 2011	Quar- antine from work (stay- at- home order)	Em- ploy- ees	Prevent spread of influenza in work-places by quarantining workers who had a cohabiting family member with an ILI	Full wages to employee	Non-compulsory asking of workers whose family members developed an ILI to stay at home voluntarily on full wages. Daily measuring of temperature before leaving work. Where symptoms were doubtful, industrial physician made judgement. Company doctors provided input on cancelling of stay-athome orders as required.	Health manage-ment department oversaw the procedures and decisions.	Mode of ad- vice to em- ploy- ees not de- scribed.	Car industries in Japan	Stay-at- home or- der for 5 days af- ter reso- lution of ILI symp- toms or 2 days after al- leviation of fever over 7.5 months	Strict stan- dard for can- celling of stay- at- home orders de- scribed.	None de- scribed.	Recording of compliance with stay-athome request	100% compliance to stay at home reported.
Young 2021 (addi- tional source: Den- ford 2022)	Daily con- tact testing (DCT) with Later- al Flow Device (LFD)	Stu- dents and staff from sec- ondary schools and further	Provide a quicker, more convenient and alternative	SARS-CoV-2 Lateral Flow Device (LFD) (Orient Gene, Huzhou, China) ^[47]	In addition to twice weekly asymptomatic testing with LFD according to national policy: students and staff who were close contacts [48] of students or staff members	A study work- er was funded at each school but role not	Indi- vidual- ly and face to face	172 sec- ondary gov- ern- ment fund- ed, res- iden- tial,	March to May 2021 Daily contact testing was per- formed at arrival	When testing could not start immediately following iden-	None report- ed	Daily participation rates in IG measured per day and per participant	Testing did not occur on 15.8% of per- son-school- days due to school or pub- lic health

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Con

(Continued)		
for	edu-	testing
con-	cation	option
tacts	col-	and
of COV-	leges	poli-
ID-19		cy for
cases		COV-
		ID-19
		close
		con-
		tact
		test-
		ing in
		schools,
		as an
		alter-

native

to self-

isola-

tion

willo mad a pos	SILIVE
LFD or PCR we	ere
identified and	of-
fered daily LFI	D test-
ing on arrival	at
school or colle	ege
each morning	(if
asymptomation	and
no household	mem-
ber isolating o	lue to
testing positiv	e for
COVID-19)	
Participants	
swabbed own	
(anterior nare	
pervised by tr	
staff. Swabs te	
by school staf	fusing
LFC	
Contacts with	
ative LFC atte	
education but	
asked to self-i	
late at home a	
school and on	
ends/holidays	
Contacts with	
ative tests (tes	
done over 7 co	
utive days) ind	
ing one on or	
the 7th day of	
were released	trom
self-isolation	

who had a positive

Contacts with positive test were required to self-isolate for 10 days, along with their contacts. Their school-based contacts were identified and process repeated

specified School staff tested the swabs that were taken by students Study	special and independent day schools and further education colleges in England	at se each more payed the after case identified.
staff trained ac- cord- ing to nation- al NHS Test and Trace stan-		don ove con tive (allo ing no t ing wee
dard process super- vised LFD testing		Sch acti part twe 19 A 202 27 J 202 (cor erec riod low

week- end), cesting could start within	school / week, and participant type, (= sum of all study school
3 days of case den- cifica- cion	days of individuals eligible for DCT returning a test result or already having completed follow
	up each day, divided by the sum of individuals eligible for DCT. Qualitative interviews conducted to understand reasons for participa-
	3 days of case den- ifica-

as cal-	tives
ılated /	IC man
hool /	IG par-
eek,	ticipa-
nd par-	tion rate:
cipant	42.4%
pe, (=	with
ım of	marked
l study	variation
hool	between
ays of	schools
divid-	(range
als eli-	0% to
ble for	100%).
CT re-	
rning	See Fig-
test re-	ure 2
ılt or	for non-
ready	partici-
aving	pation
om-	reasons
eted	break-
llow	down
each	(e.g.
ay, di-	testing
ded by	kit un-
e sum	avail-
indi-	able,
duals	whole
igible	cohort
r DCT.	moved
I DCI.	to isola-
ualita-	tion).
/e in-	•
rviews	Staff
nduct-	more
to un-	likely
erstand	to par-
	ticipate
asons	than stu-
r par-	

dents.

tion and

agency

direc-

ported sepa-

rately in Denford 2022)

ure 2 for participation by school type break-

down

"Al-

though contacts at government-funded schools with students 11-16 years old with a low proportion of free school meals were most likely to participate, other school types were simi-

lar, such that differences in participation related

to fac-

checklist (Continued)

tative analysis of interviews indicated daily testing may be feasible and acceptable but needs improved communication to students and parents about rationale, test interpretation and actions (Denford 2022)

Other (miscellaneous/multimodal) interventions

Ashraf 2020	6 ac- tive in-	Resi- dents	lm- prove	Free technolo- gies and sup-	Provision and de- livery of supplies or	540 CHW	Mostly face to	House- holds	2 years from	CHWs iden-	S: la- trine	Mea- sured by	CHWs visited
	terven-	of	envi-	plies:	installations as de-	or 'pro-	face in	and	May	tified	pits	a sep-	more
(addi-	tions	house-	ron-		scribed in Materials	mot-	groups	com-	2012	and	adapt-	arate	than
tional	of Wa-	holds	mental		column according to	ers'	and in-	pounds		ad-	ed	trained	planned

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) cł

checklist		ion or int	erventioi	is in included sti	idles, using the items	from the	remptat	te for intei	vention D	escriptio	n and Ke	pucation (пріек)
sources:		of vil-	condi-	W: chlorine	intervention type or	who	divid-	(n =		dressed	when	team	(5 to 7 /
2013, Lu-	,	lage	tions	(sodium	combination.	were	ually	5551)		any	insuf-	(uni-	month)
by	tion,	com-	to in-	dichloroisocya-		local	with	of rur-	6 to 8	bar-	ficient	versity	which
2018, Pai		pounds	terrupt	nurate) tablets		women	some	al vil-	house-	riers	space	gradu-	re-
2018, Ra		and for	trans-	(Aquatabs,		and	activi-	lages	holds /	that	(2% of	ates) at	searchers
man	(WASH)	some	mis-	Medentech,	Interventions de-	resi-	ties by	in	CHW	arose	cases)	regular	suggest
2018, Un	,	inter-	sion of	Wexford, Ire-	ployed so that they	dents	phone	Gazipur,		through	ouoco,	intervals	may
comb	nutri-	ven-	respi-	land)	were in place before	of	phone	Kishore-		ongo-		using a	have af-
2018)	tion	tions,	ratory	tarray	index children were	study		ganj,		ing di-		priori	fected
2010)	com-	partic-	nathoge	ns- 10 L insulat-	born	vil-		My-	1:12 su-	alogue	Func-	bench-	uptake
	po-	ularly	and	ed safe stor-		lages		mensingh	pervisor	with	tional	marks:	арсакс
	nents:	preg-	im-	age vessel (Li-		re-		and	to City	care-	water	mants.	
	memes.	nant	prove	on Star Plastics,	In combined inter-	cruited		Tangail	ratio	givers	seals	a) sur-	
	A. Wa-	women	child	Sri Lanka) with	vention arms, the	through		Dis-		giveis	count	veys	Report-
	ter (W)	and	malnu-	a lid and tap for	sanitation measures	trans-		tricts			was	and spot	ed "high
		their	trition	drinking water	were delivered first,	parent		in	CHWs		low (<	checks	adher-
	B. San-	infants	there-	per household	followed by hand-	mer-		Banglade		CHWs	80%	in 30	ence to
	itation	and	by re-	·	washing, then water	it-based		Danglaac	house-	met	bench-	to 35	all in-
	(S)	chil-	ducing		treatment.	selec-			holds	with	mark)	house-	terven-
	_	dren <	child-		treatment.	tion			1 / week	super-	in ini-	holds /	tions"
	C.	5 years	hood	S: Dual-pit pour		meth-		House-	for first 6	visors	tial	IG / per	with
	Hand-	5 years	respi-	flush latrines		ods		holds	months,	month-	months	month,	"marked
	wash-		ratory	with water seals	Household visits and	and		spread	then at	ly to	which	over 20-	differ-
	ing (H)		illness	for all com-	community discus-	consul-		across	least 1 /	adapt	trig-	month	ences in
	D. Wa-		and	pound house-	sions based on be-	tation		0.2 to	fortnight	tech-	gered	period;	promot-
	ter+		im-	holds. Each	haviour change strat-	with		2.2 km		nolo-	a rapid	,	ed be-
	sanita-		prov-	pit had 5 con-	egy by CHWs (paid a	com-		radius		gy and	re-	b) 5-	haviors
	tion +		ing	crete rings 0.3	monthly stipend), in-	munity				behav-	sponse	hours	from the
	hand-		child-	m high;	cluding interactive	leaders			Promot-	iour-char	ngwahich	of struc-	control
	wash-		hood	- Pot-	sessions for develop-				er train-	ар-	im-	tured	group
	ing		mor-	ties ^[34] (RFL,	ing solutions to im-				ing:	proach-	proved	observa-	at both
	(WSH)		bidity	Bangladesh)	prove practice. Key				11411.	es to	uptake	tions in	year 1
	(**311)		based	Dangiauesii)	recommendations	CHWs			Initial:	meet	(Rah-	324 IG	and year
	E. Nu-		on the	- Sani-	per IG:	had			W, S,	evolv-	man	and 108	2," with
	trition		Inte-	scoops ^[35] (lo-		com-			HW: 4	ing	2018);	control	over
			grated	cally devel-		pleted			days;	condi-	house-	house-	75% ad-
	F. Nu-		Behav-	oped hand-tool	W: children drink	mini-			uays,	tions	holds	holds,	herence
	trition		ioural	made for the	treated, safely stored	mum			N, WSH:		were	approx-	in the
	+WSH		Mod-	trial for removal	water from ves-	of 8			5 days;		using	imate-	single
	(WSHN		el for	of faeces from	sel (filled vessel	years			-	CHW	own la-	ly 15	IG and
			Water	compound)	with added 1 33 mg	formal			WSHN: 9		trines	months	com-
			Sani-	for households	tablet, wait 30 min	educa-			days	super- visors	with	after in-	bined
			tation	with index chil-	before drinking)	tion,				avail-	broken	terven-	IGs.
			and	dren		lived				avaii- able	water	tions	
			Ну-	ulell		within				able	seals in		
			,										



rvention	s in included st	uaies,
giene ^[33] and 2 years of iter- ative testing and re- vision.	H: 2 HW stations, 1 water reservoir near kitchen (16 L) and 1 near latrine (40 L), each with basins for rinsing with a	S: fa ble p ty tr how pose clea latri
Intervention specific behavioural objectives:	soapy water bottle (RFL, Bangladesh) and detergent sachets for index house- holds ^[36]	H: fa with cation a ch cate ing co a ch

W:

drink

treat-

ed and

safely

stored

water

S: safe

faeces

dispos-

H: HW

with

soap

at key

times

al

N: supply of lipid-based nutrient supplements (LNS, Nutriset; Malaunay, France) (for 6 to 24 months olds) 2 10g sachets per day per child; (118 kcal, 9.6g fat, 2.6g protein, 12 vitamins and 10 minerals) Cost: USD 0.08/ day

18-month shelf N: agelife appropriate nutri-Stipends for tion CHWs (USD 20/ birth

month for 24

amily use doupit latrines, potrain children and v to safely dise of faeces and an and maintain ines

amily wash hands h soap after defeion, after cleaning nild who has defeed, before eator before feeding nild, and before d preparation

N: recommendations for exclusive breastfeeding up to 180 davs and maternal and infant nutrition to mothers and index children: introduce diverse complementary food at 6 months; feed LNS from 6 to 24 months, mixed into the child's food (not intended as a replacement for breastfeeding or complementary foods). Messages adapted from the Alive & Thrive programme^[37]

solu-

tions

techni-

and

walk-Refreshing diser traintance ing: 1 of IG day each cluster and passed 21 day a writtraining ten of adand herence oral exteam amination. They at-Monthly tended CHW sumulpervisor tiple meettrainings ing sessions and quarterly refreshers. Training covered active listening, strategies for developing collaborative

by cell paralcomphone lel with menced. as trial laneedtrines ed so pre-Meaexistsured: ing latrines W: Pres-Trainwere ence of ing of closed. stored provisdrinking moter its by water varied **CHWs** with dein conwere tectable tent infree and creased chlorine length and (> 0.1)dewamg/L) pendter-seal ing on re-S: a laintermoval trine venor with tion breakfunctiontype age al wawas ter seal, dissani-Potties courscoop aged proaccessivided bility Initial if chilprofes-H: presdren < sional ence of 3 vears trainsoap at er for primary CHW HW statraintions ing did not en-N: regage ported contrainees enough sumption of so re-LNS saplaced with chets

inter-

and N IGs compared with WSH and WSHN S: observed use of latrines: 94% to

Similar

adher-

ence in

single

W, S, H

97%;

child

sani-

tation

practices

(37% to

54%)

H: HW

soap in

IG more

common

after toi-

let use

to 74%)

versus

18% to

40% in

non-IGs

and after

cleaning

child's

(61% to

72%) but

anus

(67%

with

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

to 24 months	months) delivered through mobile phone network to ensure timely payments Promoter's guide for visits for each relevant interven-	On household visits, following a structured plan, CHWs greeted targeted household members, checked presence and functionality of relevant hardware and signs of use, observed recommended practice using a guide.	cal aspects of interventions (see Table 1 of Luby 2018 for more details)
	tion including:	CHWs used discussions, video dramas,	
	- visit objective,	storytelling, games	CHWs
	- target audi- ence	and songs and pro- vided training on hardware mainte-	were trained by 47
	- steps and ma- terials to be used	nance, where applicable	CHW super- visors who
	CHW ID badges	Adherence observed and measured by separate team	re- ceived direct train- ing on
	Cell phones for CHW supervi- sors	Supervision meet- ings of CHWs and pe- riodic internal moni- toring of their perfor- mance	inter- ven- tion deliv- ery
	Training Plan and Manual for CHW supervi-		Hard-
	sors covering:	Intervention Delivery Team managed de-	ware instal-
	i) basic training	livery through regular team phone calls,	lation team
	- introduc- tion of project, CHW roles and responsibili-	field meetings, field reports and liaison with relevant gov-	(n = 18)

ernment and other

stakeholders. It coordinated CHWs to

responsibili-

ties, introduc-

IIal	See K-	tow pe-
train-	ahman	fore food
ing re-	2018 for	handling
source	more de-	
group	tails (Ta-	W: > 65%
8.000	ble 1)	mothers
	Dic 1)	and chil-
		dren ob-
Due to		served
obser-	Contin-	drink-
vation	uous	
of in-	over-	ing chlo-
		rine-treat-
terven-	sight	ed wa-
tion fa-	and pe-	ter from
tigue	riodic	safe con-
report-	moni-	tainer
ed by	toring	
CHWs	of CHWs	N: LNS
and	perfor-	feeding >
sub-	mance	80%
opti-	(CHW re-	
mal	placed	
prac-	within 1	_
tices	month	33 low
ob-	of attri-	per-
served,	tion or	forming
new	critical-	CHWs
behav-	ly low	discon-
iour	perfor-	tinued
	mance	
change	mance	
activ-		
ities		See Luby
were		2018, Parvez
devel-		2018, Arnold
oped		2013, Uni-
(e.g.		comb
further		2018 for
tech-		more de-
nology		tails
use, in-		cuito
creas-		
ing		
self-		
effica-		
cy and		

nal

See R-

low be-

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

tion to behav-
iour-change
principles
based on the
IBM-WASH the-
oretical frame-
work and inter-
personal and
counselling
communication
skills.
ii) Interven-
tion-specific
training
ti dillillig
iii) classroom

practice / role

playing

ensure rapid identification of issues with delivery. Including a dedicated training officer, it also trained the CHW supervisors who then trained the CHWs under their supervision ("train the trainer" approach)

9 field research officers

The Intervention Delivery Team^[38] co-ordinated delivery including CHWs, overseen by Principal In-

vestigators with consultation from Technical Advisory Group (see

Dedicated

Unicomb, 2018)



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

Training Officer and Communication Development officer

ecklist (Continued)		., g	

3-ply tissues

A. 5.1 mg/inch²

mixture (58.8%

(2.54 cm²) of

the virucidal

citric acid,

29.4% malic

acid, 11.8%

sulphate)

sodium lauryl

B. 3 mg/inch²

(2.54 cm²) of

saccharin ap-

with:

Re-

duce

trans-

sion of

viruses

from

hand

conta-

mina-

to-

hand

con-

tact or

tion via hand-

mis-

Fami-

lies

Adher-	
ence	
ob-	
served	
by sep-	
arate	
team	
who	
re-	
ceived	
formal	
21 day	
train-	
ing	

	'''S							
Family visits to dis- tribute tissues	Nurse epi- demi- ologist	Face- to-face visits to fam-	Com- mu- nities in the	6 months overall	Not de- scribed	Not de- scribed	Fami- ly vis- its and week-	Not de- scribed
Weekly contact of mother	visited fami- lies.	ilies and in- divid- uals in fam-	USA	Month- ly family visits			ly con- tact with moth- er to en- courage	
Families instructed to only use supplied tissues.		ilies (espe- cially moth- ers)		Week- ly con- tact with mother			compli- ance	

Farr

1988a

trial 1

2 ac-

tive in-

terven-

tions

in ad-

dition

to control of

no tis-

sues:

A. Viru-

cidal

<u>,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,</u>
Cochran Library

Table 1. checklist			large- par- ticle aerosol through tissues for nose blow- ing and coughs and sneezes	plied uniformly to all 3 plies of the tissue Tissues prepared by Kimberly-Clark Corporation, Neenah, WI, USA.	udies, using the item	s from the	: Templat	e for Int	ervention D	escriptio	n and Re	olication (TIDieR)
Farr 1988b trial 2	2 active interventions (no control): A. Virucidal nasal tissues B. Placebo tissues	Fami- lies	Re- duce trans- mis- sion of viruses from hand conta- mina- tion via hand- to- hand con- tact or large- par- ticle aerosol through tissues for nose blow- ing and coughs and sneezes	2-ply tissues containing: A. 4.0 mg/inch² (2.54 cm²) of antiviral mixture (53.3% citric acid, 26.7% malic acid, 20% sodium lauryl sulphate) B. 3 mg/inch² (2.54 cm²) of succinic acid, malic acid, sodium hydroxide, and polyethylene glycol Tissues prepared by Kimberly-Clark Corporation, Neenah, WI, USA.	Family visits to distribute tissues and encourage compliance Weekly contact of mother Families instructed to only use supplied tissues.	Nurse epi- demi- ologist visited fam- ilies month- ly. Study moni- tor vis- ited bi- month- ly.	Face- to-face visits to fam- ilies and in- divid- uals in fam- ilies (espe- cially moth- ers)	Com- mu- nities in the USA	6 months overall Month- ly family visits Week- ly con- tact with mother Bi- month- ly study monitor visit	None de- scribed.	None de- scribed.	Bi- month- ly study moni- tor vis- its to en- courage compli- ance as well as month- ly and weekly contact by nurse	In 124/222 fami- lies, 1 or more family mem- bers re- ported not us- ing the tissues regular- ly and/or report- ed hav- ing side effects from the tissues.

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
checklist (Continued)

Fretheim	GI ASSV	Adult	Pro-	Instructions via	Request to wear	Re-	Indi-	Out-	14 days	Could	None	No con-	Report-
cadditional source: F 2022b (pi tocol)		Adult members of the public who did not regularly wear glasses and who owned or could borrow glasses to use (e.g. sunglasses)	Provide a simple, readily available, environmentally friendly, safe and sustainable means of personal protection from infection with respiratory viruses including SARS-CoV-2	Regular eye- wear, e.g. sun- glasses owned by participant or that could be borrowed by participant	Request to wear sunglasses or other types of glasses when outside home and close to others in public spaces for 14 days	Re- search team	Individual- ly In- struc- tions provid- ed via email and online portal (Nettskje ma-plat- for- m)ac- cessed via web- page hosted by the Norwe- gian Insti- tute of Public Health	Out- side the home, e.g. on public trans- port, in shop- ping malls (in Nor- way)	14 days when out- side and close to others in public spaces Over 11 to 12 week period (Feb- ruary – April 2022)	Could borrow glass- es if did not own any	None report- ed.	No contact was made with participants between enrolment and data collection.	Report- ed use of glasses often, al- most al- ways, or always: IG: 71% CG: 11% Negative experi- ences (espe- cially fogging with mask use): IG: 21/76
Longi- ni 1988	2 active interventions (no control):	House- holds and their fami- lies	Prevent intrafamilial transmission of viral agents in a com-	Treated tissues of 3-ply mate- rial identified with no specif- ic identifiers (Kimberly-Clark Corporation) with inside lay- er containing:	Tissues delivered to households with specific instructions on use (all purposes, when blowing nose, coughing or sneezing) and to discard after use and to help young children use tissues if develop a cold.	Tissues as- signed by study spon- sor (Kim- ber- ly-Clark	Supply of tis- sues through- out 5- month trial period	House- holds in the USA	5 months' overall supply	Resup- ply of tissues as re- quired	None de- scribed.	Report- ed use of tissues "not at all, some of the time, most of the time, or	Reported use "all of the time": A. versus B. 82% versus 71%



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Deplication (TIDioD)

	A. Viru- cidal nasal tissues		munity setting	A. citric and malic acid plus sodium lauryl sulphate;		Corpo- ration).						all of the time"	
	B. Place- bo tis- sues			B. succinic acid.									
Chard 2019 (addi- tional details from- Chard 2018)	Water, Sani- tation, and Hy- giene for Health and Educa- tion in Laot- ian Pri- mary Schools (WASH HELPS)	Primary schools and their students	Prevent the spread of pathogen within schools through im- proved water sup- ply and hy- giene facil- ities and im- proved WASH habits in chil- dren at home and through- out the life course	For each school: Water supply for school compound: (borehole, protected dug well with pump, or gravity-fed system) Water tank to supply toilet and handwashing station School sanitation facilities (3 toilet compartments) Hand-washing facilities: 2 sinks with tapped water and supply of soap available	Provision of school: Water supply, sanitation facilities, handwashing facilities (individual and group), drinking water filters Behaviour change education and promotion including daily group hygiene activities Daily hand-washing and cleaning schedules	UNICEF paid for materials. School and teachers conducted daily handing activities with children. Students participated in daily group cleaning activities.	Facilities provided within schools. Children participated in group handwashing and cleaning.	Primary schools and their class-rooms (in Laos)	One-off provision of water and hygiene facilities Daily hand-washing activities and cleaning for 1 school year Cleaning schedules posted in at least 1 classroom near toilet.	Water supply tailored to the school requirements/er vironment. Sanitation facilities provided as needed and designated for boys, girls, and students with disabilities.	Rain water tank provi- sion af- fected by rain water n-sup- ply, so changed to tanks with mo- torised hand pumps or gravi- ty-fed water sup- ply sys- tems. Theft and animal con- sump-	Unan- nounced visits every 6 to 8 weeks for struc- tured observa- tions to measure fidelity and ad- herence Fideli- ty Index score (0 to 20): for hard- ware provid- ed see Table 1 in paper and pro- tocol Adher- ence in-	Fidelity: 30.9% across all schools and visits Adherence: 29.4% Hard- ware provision: 87.8% or schools School- level adherence 61.4% Group compound cleaning 94.8%, toilet use: 75.5%, group toilet

(e.g. presence

with

ers: 90%

ed and

to local

months

of

checklist (Continued)			(1 bar of soap/ pupil)							tion of sup- plied	dex: stu- dent re- port of	cleaning: 68.3%, group	
			3 group hand- washing tables with soap and water							soap re- duced supply.	behav- iour- al out- comes index score (0 to 4)	hand- washing: 48.7%, indi- vidual hand- wash-	
			At least 1 drink- ing water filter per classroom								ŕ	ing with soap af- ter toi- let use: 23.9%.	
			Schedules of daily group hand-washing, compound and toilet cleaning										Further details (Chard 2018)
			Cost per school: USD 13,000 to 17,500										
Hartinger Inte- 2016 grat- ed en-	House- holds and	Re- duce infec-	Per household:	Community engage- ment with local and regional stakehold-	Health pro- moters	Face- to-face and to	House- holds in rur-	Stoves and sinks in-	Tai- lored to par-	Not de- scribed	Week- ly spot- check	SODIS use:	
viron- mental home- based	their house- hold- ers in-	tions and im- prove	"OPTIMA-im- proved stove": improved venti- lated solid-fuel	ers in design and development	hired local ele- men-	indi- vidual house- holds;	al com- muni- ties in Peru	stalled over ini- tial 3 months.	ticular house- hold facil-		observa- tions of house- hold hy-	60% ini- tially and 10% at end of	
inter- ven- tion pack- age	clud- ing chil- dren	child growth in house- holds	stove Kitchen sink	Provision of stoves, kitchen sinks, and plastic bottles for so- lar water treatment,	tary school teach- ers and imple-	mode of de- liv- ery of train-		Month- ly rein- force-	ities and envi- ron- ments		giene and en- viron- mental health	study Self-re-ported	
(IHIP)		in rur- al com-	with in-kitchen water connec-	and hygiene educa- tion	ment- ed and	ing as indi-		ment over 12	as need-		condi- tions	use by moth-	

moted

pro-

vid-

ual or

mu-

nities

tion providing

piped water

ipato-

ry ap-

proach

g		•
Training of mothers/caretakers in: - solar drinking-water disinfection (SODIS) ^[39] according to standard procedures - hand hygiene (washing own and children's hands with soap at critical times ^[40]) - advice to separate animals and their excreta from the kitchen environment	the interventions. 4 teams of field staff conducted spotcheck observations.	group not de- scribed
Project-initiated repairs		

SODIS,	beliefs	of SODIS	slight
child	and	bottles	decrease
and	cultur-	on the	at end
kitchen	al cus-	roof or	
hygiene	toms	kitchen)	
		using a checklist	Self-re- ported
Week-	Re-		stove
ly spot	pairs		use: 90%
checks	to	Monthly	daily
of com-	stoves	self-re-	
pliance	as	port by	
	need-	mothers	Sink use:
	ed and	of stove	66% dai-
Repairs	checked	and sink	ly
after 9	at 9 months	use	•
months	monus		
			35% of
			stoves
Environ-			needed
men-			minor
tal sam-			repairs,
ples test			, ,
middle			1%
and end			needed
of 12-			major re-
month			pairs.
surveil-			
lance.			

Bestfunctioning stoves achieved mean 45% and 27% reduction of $PM_{2.5}$ and CO, respectively, in



Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)

Com-

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10

checklist (Continued)

that addressed local beliefs and cultural views

Re-

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in their

house-

hold

mothers' personal exposure.

Vil-Huda Sanitation 2012 lages Hyand giene their house-Eduholds cation with a and Water child < Sup-5 years ply in old Bangladesh (SHE-WA-B)

training of comduce illness munity hygiene in chilpromoters and dren < promotion activities includ-5 years ing flip charts by improvand flash cards ing hywith messages giene alerting participants to pracpresence of tices, saniunobservable "germs" and tation practices to and water minimise germs supply and

Materials for

See Box 1 in paper for 11 key messages.[41]

Engaging local residents under guidance of local NGOs to develop community action plans addressing:

Latrine coverage and usage

Access to and use of arsenic-free water

Improved hygiene practices, especially hand-washing with soap

Recruitment and appointment of community hygiene promoters

Household visits. courtyard meetings, and social mobilisation activities (e.g. water, sanitation and hygiene fairs, village theatre, group discussions in tea stalls (the social meet-

Vil-18 Faceto-face months lages delivand overall ery to housegroups holds (vilin dis-Extricts lages pected of and Bangladesh house-hold vishouseholds) it and and incourtdividu-Comyard als munimeeting every 2 ty activities months held in villages. Handwashing op-Meetportunities: afings held in ter own or child's courtdefecayards of tion, groups prior to of preparhouseing and holds.

serving

food, pri-

or to eat-

HW: Structured Food-reobserlated: vation of hand-No sigwashnificant ing and differchild ence faeces from disposal basebehavline to 18 iour in months; households IG versus and spot CG checks of After type of anus housecleaning: hold wa-36% verter and sus 27% sanita-Defetion facation: cilities 30% ver-

sus 23%

No ac-

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR)
checklist (Continued)

led stu	dies, using the items	from the	Templat	e for Inte	rvention D	escriptio	n and Rej	olication (1	「IDieR)
	ing point for village men)) by community			House- hold	ing and feeding			10.3% to 6.8%.	
	promoters			visits	a child				
	Structured observa- tion in households								No significant improvement in access to improved latrines, solid waste disposal, drainage systems, and covered containers for water storage
nts: san alby, for na-	Collection and com- mercial cleaning of toys from nurseries: - linen and toys suit- able for washing ma-	Com- mer- cial clean- ing	Clean- ing com- panies col-	Day- care nurs- eries in Den-	2 to 3 months overall	Stag- gered clean- ing to ensure	None de- scribed.	None described.	None described.
Eco	chines were washed at 46 °C and subse- quently disinfected	com- pany: Berend-	lect- ed the toys	mark	Cleaning every 2	chil- dren had			

fection

ethoxylates) for

immersion or wiping

- toys not suitable for washing machines immersed in disinfectant or wiped with microfibre cloth

sen A/S, Søborg, Denmark

and Comlinen merand cial incleaned dusthem trial offsite, cleanthen ing farecility turned

them.

weeks toys to play with whilst others were being

cleaned

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued) treat-

		ment	
2 ac-	Low-	Pre-	A
tive in- terven-	in- come	vent or reduce	C
tions:	house-	trans-	_
	holds	mis-	S
	and	sion of	(: E
Δ	com-	•	n
	pounds	,	d
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'be-			С
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-		Water	
		Sani-	(
		tation	a
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		giene	t
CIOII			t
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	2	-	(
group			٧
		Hul-	t b
		land	h
		2013)	i
	tive interventions: A. Combined cholera vaccine and 'behaviour change communication' intervention B. Cholera vacc	tive in- terven- tions: house- holds and com- pounds Com- bined cholera vac- cine and 'be- hav- iour change com- muni- cation' inter- ven- tion B. Cholera vac- cine-alone	2 ac- tive in- terven- tions: A. Com- bined cholera vac- cine and iour change com- muni- cation' inter- ven- tion B. Cholera vac- cine and Com- muni- cation' inter- ven- tion B. Cholera vac- cine-alone group Com- MA. Com- pounds ratory illness based based on the var- ioural Behav- ioural Mod- el for Water Sani- tation and Hy- giene (IBM- WASH) theo- retical Cholera vac- cine-alone group bis 2013; Hul- land

	A. and B.	A. and B.
	Cholera vaccine ShanChol™ (Shantha Biotech-	Provision of cholera vaccine (2 doses at least 14 days apart)
	nics-Sanofi, India)	Provision of hand- washing hardware and behaviour change communica-
	A. Following hardware per compound:	tion activities
	a. Hand-wash- ing hardware:	Encouragement of hand-washing after defecation, after
	(i) Bucket with a tap (provided free of charge)	cleaning child's anus, and before preparing food
	(ii) Soapy water bottle (mixture of a commercially available sachet of powdered detergent	Encouragement to add chlorine to own water vessels
-	(~USD 0.03) with 1.5 L of water in a plastic	Benefits were again explained.
	hole punched in the cap) sup- plied by partic- ipating com- pounds	Follow-up visits by health promoters

Dushtha Shasthya Kendra (DSK), an NGO, deliv- ered the hard- ware and behav- iour- al in- terven- tion	Hand- wash- ing and water treat- ment hard- ware most- ly de- livered at the com- pound level in per- son.	House-holds and compounds (where several house-holds share a common water source, kitchen,	Behaviour change communication messages delivered first (within 3 months of cholera vaccination).
(through com- munity health pro- mot- ers).	Behaviour change communication mes-	and toi- lets) in Banglade	Point-of- use wa- ster hard- ware provid- ed 3 months later.
Separate data collec- tors ob- served soap avail- ability.	sages were deliv- ered both at com- pound and house- hold levels.		Fol- low-up health promot- er visits 3 times in 2 months after hard- ware instal-

lation,

e- ds re re- e- e- re- en,	Behaviour change communication messages delivered first (within 3 months of cholera vaccination). Point-ofuse waster hardware provided 3 months later.	Hard- ware-re- lated prob- lems (break- age/leak- age) were ad- dressed on health pro- mot- er fol- low-up visits.	None de- scribed.	Unan- nounced home visits by data col- lectors who ob- served presence of soap/ soapy water and wa- ter in most conve- nient place for hand- washing (either reserved in a con- tainer or avail- able at the tap)
	Follow-up health promoter visits 3 times in 2 months after hard-ware			Resid- ual chlo- rine was mea- sured in- dicating uptake of chlo- rine dis-

	Presence of soap / soapy water and wa- ter:
ò	A. Hand- washing group com- pounds: 45% (1729 / 3886);
	B. Vaccine-on- ly group com- pound: 22% (438 / 1965);
	C. Con- trol: 28% (556 / 1991)
	Residual chlorine

present

in stored

(160/3886)

drinking water of 4%

of house-

penser.



(iii) Bowl to col-
lect rinse water
after

washing hands (see photo in text or in Najnin 2017 doi.org/10.1093/ then 2 times/ month (over nearly 2 years). holds in the vac-cine-plus-behaviour-change compound and none in the other 2 compounds.

b. Water treatment hardware:

ije/dyx187)

Dispenser containing liquid sodium hypochlorite

See Figure 2 in Najnin 2017 for photos of both doi.org/10.1093/ ije/dyx187

and more details.

Participants own water vessels for water treatment

Print materials for behaviour change to compounds and households

Swarthou	ıt6 ac-	Resi-	lm-	Free technolo-	Provision and de-	Com-	Face to	8246	Installa-	Train-	None	Partici-	All in-
2020 (ad-	tive in-	dents	prove	gies as appro-	livery of supplies or	muni-	face in	house-	tion and	ing tai-	de-	pant re-	terven-
di-	terven-	of	envi-	priate to IG:	installations as de-	ty-based	groups	holds	supply	lored	scribed	ports	tions de-

Table 1.	Descripti	on of int	ervention	s in included stu	idies, using the items	from the	Templat	e for Inte	rvention D	escription and	Replication (1	ΓIDieR)
checklist	(Continued)											
tional	tions	house-	ron-		scribed in Materials	health	(e.g.	and	of ma-	for dif-	of visits	livered
sources:	Aronfolica-	holds	mental		column according to	pro-	house-	7960	terials	ferent	by pro-	within 3
2013, Ch	rister,	of vil-	condi-	W: water treat-	intervention type or	moters	holds	com-	before	inter-	moters	months
tensen	sanita-	lages	tions	ed with sodi-	combination	nom-	or	pounds	com-	ven-	in past	of enrol-
2015, De	nttion,	and for	to in-	um hypochlo-		inat-	com-	of rur-	muni-	tions	month	ment
2017, Nu	ll and	some	terrupt	rite (1.25% so-		ed by	pounds)	al vil-	ty meet-			
2018, Pic	k-hand-	inter-	trans-	lution / 2 mg/L)	D (their	or indi-	lages	ings			
ering	wash-	ven-	mis-	using chlorine	Provision of study	local	viduals	in Bun-	_	_		
2019)	ing	tions,	sion of	dispensers in-	materials to promot-	com-	(moth-	goma,		Trou-	Unan-	In-
	(WASH),	partic-	respi-	stalled at com-	ers	mu-	ers and	Kakameg	ga,	bleshoot-	nounced	creased
	and	ularly	ratory	munal water		nities	their	and Vi-		ing of	visits by	adher-
	nutri-	preg-	pathoger	_{IS} source collec-		and	chil-	higa	munity	solu-	staff to a	ence in-
	tion	nant	and	tion points or	Community meet-	trained	dren)	coun-	meeting	tions	random	dicators
	com-	women	im-	bottled chlo-	ings	in the	,	ties in	6 weeks	to bar-	sam-	of ≥ 30%
	ро-	(Ma-	prove	rine (1L for 333	83	rele-		west-	after en-	riers to	ple of	higher
	nents:	mas)	child	20-l jerry-cans		vant		ern	rolment	adher-	at least	in all IGs
		and	malnu-	worth) ^[45] pro-		inter-		Kenya		ence	20% of	relative
	A. Wa-	their	trition	vided to house-	Household and com-	ven-		,		by pro-	partic-	to the
	ter (W)	infants	there-	holds in com-	munity visits by pro-	tion			Month-	mot-	ipants	control
		and	by re-	pounds	moters who:	to be			ly visits	er and	in IGs at	in the
	B. San-	chil-	ducing			imple-			(45 to 60	partic-	2, 6, 10,	first year
	itation	dren	child-	Chlorine strips	- delivered interven-	ment-			min in	ipants	and 19	
	(S)	< 5	hood	to test chlorine	tion-specific behav-	ed			1 st year)	as	months	
	6	years;	respi-	levels	iour change mes-				by pro-	need-	after the	Adher-
	C.	Landowr	•		saging focusing on				moters	ed	interven-	ence was
	Hand-	ers of	illness		themes of nurture,				over 2		tions be-	compa-
	wash-	com-	and	S: installation	aspiration and self-	Field			years		gan to	rable be-
	ing (H)	munal	im-	of new or im-	efficacy, consider-	enu-			(2012 to	Nutri-	confirm	tween
	D.	water	prov-	provement of	ing convenience and	mera-			2014)	tion	delivery	the Indi-
	Com-	sources	ing	existing latrines	cultural norms to im-	tors as-			2014)	mes-	of mate-	vidual
	bined	and	child-	with plastic	prove adherence us-	sessed				saging	rials and	IGs com-
	(WSH)	com-	hood	slab latrines	ing scripts and visual	adher-				was	moni-	pared
	(1101.)	pound	mor-	with tight-fit-	aids;	ence in			Timing	tai-	tor avail-	with
	E. Nu-	heads	bidity	ting lids; plas-	- provided instruc-	com-			of visits	lored	ability	com-
	trition	for la-	based	tic potties and	tions on hardware	pounds			detailed	to be	of inter-	bined
	(N)	trine	on a	sani-scoops	use and consumable				in pro-	age-	vention	IGs.
		up-	litera-	sam-scoops					cedures	appro-	materi-	100.
	F.	grades	ture re-		supplies where ap- plicable	Study			provid-	priate	als and	
	Com-	and	view,		plicable	staff			ed at os-	priace	recom-	
	bined	con-	a theo-	H: 2 HW sta-	- advocated:	trained			f.io/7j9sk/		mended	W: 5
	(WSHN)	struc-	ry-based	tions (2-foot		pro-					behav-	chlo-
		tion	ap-	pedal-operat-	W: drinking water	mot-				Mate-	iours af-	rine dis-
		•	proach	ed jerry-cans	treatment with sodi-	ers,			147. 1 1	rials	ter the	pensers
			(health	that dispensed	um hypochlorite	provid-			W: 1 L	provid-	interven-	in-
			belief,	soapy and rinse		ed pe-			bottle	ed in	tions be-	stalled /
			•			ou po			of chlo-	both in		cluster

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

social cog- nitive theo-	water), 1 near food preparation, 1 near latrine.
ry and per- sua- sion theo- ry),[42], [43],[44]	Rinse water provided by households; bar soap for soapy water container
formative research and the WASH Benefits pilot RCT (Christensen 2015)	N: 2 x 10 g sachets / day / child of lipid-based nutrient supplementation (LNS) "Mwanzobora", (Nutriset, Malaunay, France) (118 kcal/day and 12 essential vitamins and 10 minerals)
	See Figure 2 of Christensen 2015 for photos of examples of some of the ma- terials
	Community meeting and household visit summa- ry sheets (in Kiswahili and

English) and

idies, using the items	from th
S: use of improved latrines for defecation and safe disposal of children's and animals' faeces and use of plastic potties by children < 3 years and sani-scoops for faeces removal H: HW with soap before food preparation and after defecating (including assisting child); helped participants identify compound members to refill taps and manage barriers to use such as running out of soap	riodic obser- vation and su- pervi- sion and month- ly phone calls
N: early initiation of breastfeeding, exclusive breastfeeding 0 to 6 months and continued till 24 months; at 6 months, introduction of appropriate and diverse complementary foods; feeding frequency and during illness; supply of LNS to children 6 to 24 months and instruction to mix it was foods twice/day	
Promoters used visual aids to promote messages:	

- cue cards provid-

ed to Mamas at ini-

rine / 6 months	Kiswahili and English	gan (Null 2018)	Year 1: 74%
H: bar soap provided every 3 months	Chlo- rine dis- pensers lo- cated	W: monthly tests of chlorine concen- tration in stored	Year 2: 37% house- holds were vis- ited by a pro-
N: LNS intro- duced at 6 months	based on list of sources partic-	water; negative results prompt- ed dis-	moter in previous month
of age of child	ipants report- ed (at base- line)	cussions to ad- dress chlorina- tion bar-	W: Year 1: 42%
Promot- er train- ing:	using for wa- ter col- lection	riers	Year 2: 21% had de-
6 days single IGs.	tection	S: partic- ipant re- port of	tectable total chlorine
7 days com-	Sani- scoops and	access to im- proved	CG: 3%
bined IGs.	potties were to be	latrine; field enumer-	S:
Refresher training at 6, 12 and 18 months after initial training	washed by care- givers with soap and wa- ter af-	ators ob- served if la- trine had plastic or ce- ment slab or venti-	Year 1 and 2: > 80% had latrine access CG: 20%
0	ter use	lation	

pipe;

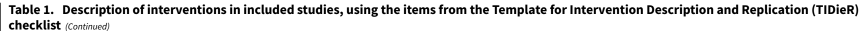
2018 for more details

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

list of materi- als provided	tial visits to hang on walls for reminders
as PDFs at os- f.io/7j9sk/	- picture sheets used by promoter to ex- plain key concepts or messages
Key messages and visual aids provided at os- f.io/7j9sk/	- calendars provided to households during first compound visit
Including ~6 primary key messages per intervention,	- stickers attached to LNS box
each with a se- ries of specif- ic topics, visu- al aids, and en-	Adherence checking unannounced visits
gagement activities (e.g. story- telling, mottos, etc.). Visual aids included:	Initial training on in- tervention-specific behaviour change messages and mate-
- cue card re- minders	rials
- picture sheets for use by pro- moters	Refresher training
- calendars for households with key mes- sages	Periodic observation and supportive su- pervision by study staff
- stickers for LNS box depict- ing appropri- ate feeding and	

storage

Supervi-	and	caregiv-	HW:
sion and obser- vation	tools kept out of	er re- port that child	Year 1: 77%
of pro- moter by study staff at 2, 4, 9, 14	reach of chil- dren (see the vi-	faeces safely disposed	Year 2: 21% had HW ma- terials
and 21 months and	sual aids provid-	H: field enumer- ator ob-	CG: 9%
month- ly phone calls	ed to partici- pants:	served if water	N:
calls	OS-	and soap available	Year 1: 95%
	f.io/9r4kg/ for	Ni rapart	Year 2: 115%
	potties and	N: report of LNS sachets	of ex- pected
	os- f.io/mz2c6/	con- sumed by child	sachets con-
	for sani-	in last week /	sumed
	scoops)	14	See Null



Promoter Training Materials for trainers and trainees for each intervention for initial training and for refresher training including detailed PDF training manuals available at osf.io/7j9sk/focusing on key hygiene messages, visitation scripts and visual aids and hardware for each intervention^[46]

Promoters' supplies:

Branded t-shirt, mobile phone, job aids and intervention materials, payment (\$US15/ month for first 6 months, then \$9/month thereafter), detailed plans for every visit (key messages, scripts for visual aids, instructions for activities)

Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist (Continued)

Oral and	or nasal a	application	ns										
Alman- za-Reyes 2021	Mouth-wash and nose rinse with AR-GOVIT silver nanoparticles (Ag-NPs)	Health-care per-sonnel (doctors, nurses, administrative staff) of a metropolitan hospital caring for patients diagnosed with atypical pneumonia and/or COV-ID-19	Reduce morbidity in health-care professionals exposed to SARS-CoV-2 by inhibiting virus replication	Per participant: - 50 ml bottle of RGOVIT® AgNPs mouthwash and nasal rinse [Investigation and Production Center Vector-Vita Ltd., Novosibirsk, Russia] (metallic silver 0.06%, polyvinylpyrrolidone 0.63%, hydrolyzed collagen 0.31%, distilled water 99% wt.) - water - cotton swabs	Individuals provided with spray bottle containing AgNPs solution with 1 wt% concentration (0.6 mg/mL metallic silver) and instructed to do 1 of the following or a combination: a) mix 4 to 6 spray shots (~ 0.5 mL) with 20 mL of water and gargle solution for 15 to 30 seconds at least 3 times/day (gargle) or b) do not dilute with water and cover the oral cavity evenly with 1 to 2 direct spray shots (spray) c) apply the same solution to the inner part of the nasal alae and nasal passage with cotton swab twice a day (nasal rinse)	Re- searchers sup- plied mate- rials and in- struc- tions Partic- ipants self- ap- plied the mouth- wash and nasal rinse materi- als	Individually and face to face	Gener- al hos- pital in Ti- juana, Mexico	Over a 9 week period (April to June 2020) 4 to 6 spray shots of AgNP solution (0.5 mL) with 20 mL of water or 1 to 2 spray shots of solution without water for 15 to 30 seconds ≥ 3 times / day and 1 nasal lavage 2 times / day	Participants could choose application method	None de- scribed	Weekly self-re-port of number of: daily gargles; mouth-wash-es with spray; mouth-washes by gargle + spray; and nasal rinses	Mean applications/day: Gargle only: IG: 2 (n = 28 CG: 2.14 Spray only: IG: 2 (n = 34). Both gargle and spray: IG: 2 gargles, 4 sprays (n = 52) Nasal rinse: IG: 0.70 (n = 64) CG: 0.25
Gutiér- rez-Gar- cía 2022	Na- sopha- ryn- geal and	COV- ID-19 front- line med-	Re- duce risk of COV- ID-19	SES (pH 6.5 to 7.5; RE- DOX potential 750-950 mV;	Written instructions provided to follow a prophylactic rinse protocol with SES 3 times/day for 4	Not clearly spec- ified; lead-	Indi- vidual- ly and face to face	Mex- ican COV- ID-19 hospi-	4 nasal sprays (~ 0.4 mL) and 10 mL	None de- scribed	None de- scribed	None de- scribed	None de scribed

ers of

tal

mouth-

weeks with advice

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in



	- 4 plastic flasks of 240 mL oral SES (ESTERICIDE® Bucofaríngeo, COFEPRIS registration no. 1003C2013 SSA) with a graduated cap and - 4 plastic flasks of 30 mL nasal rinse (EsteriFlu®, COFEPRIS registration no. 308C2015 SSA), with a valve for spraying	vertical sprays in each nostril, inhaled deeply at the time of each spray b) oral cavity: mouthwash and gargle 10 mL for 60 seconds, then spit out In addition to standard COVID-19 safety protocols requiring wearing of adequate personal protection equipment at all times, [49] frequent handwashing [50] and disinfection of secondary uniform and footwear [51] and bath at end of working day	part- ment distrib- uted the study infor- mation and were the point of con- tact and moni- tored the proto- col so they may have distrib- uted inter- ven- tion		ber to No- vember 2020)		
			materi- als				



	(Continued) A. Vitamin D ₃ supplementation B. Gargling water		of URTI through in- creased vita- min D levels (asso- ciated with greater fre- quen- cy and sever- ity of URTI) and gar- gling (as pre- ven- tative mea- sure against URTI)	chased from Euro-Pharm International Canada Inc.) Weekly email reminder B. Gargling: 30 mL of tap water 2/day	B. Gargling: instructed to gargle twice daily for 30 seconds All participants received general lifestyle and health advice on sleep, nutrition, hand hygiene, and exercise.	ably the re- searchers includ- ing a study phar- macist	vidually, but, no further details. Method of lifestyle and health advice provision also not described.	ing (in resi- dences or off- cam- pus) in Cana- da	Vita- min D ₃ : weekly supple- menta- tion and email re- minder Gargling: 30 mL of wa- ter for 30 seconds twice daily				
Ide 2014	2 active interventions (no control): A. Green tea gargling B. Water gargling	High school stu- dents	Prevent influenza spread and infection in high school students who are at increased	A. Bottled green tea (500 mL) containing a catechin concentration of 37 ± 0.2 mg/dL, including approximately 18% (-)-epigallocatechin gallate (manufactured by the Kakegawa Tea Merchants Association).	A. Provision of green tea B. Advice to gargle with tap water and not to gargle green tea during study A. and B. Advice to gargle at least 3 times/day (after arriving at school, after lunch, and after school) Consumption of green tea and other	Materials supplied by researchers High schools' vice principals and head teachers as-	Green tea sup- plied individu- ally to stu- dents. Mode of gar- gling advice not de- scribed.	High schools in Japan	Gargling 3 times/ day for 90 days	None de- scribed.	None de- scribed.	Daily questionnaire included questions about daily adherence to gargling regimen. Adherence rate of	Gargling adher- ence rate: green tea group: 73.7%; water group: 67.2%

	from close inter- action through	measured by high-perfor- mance liquid	ed for either group.	with						at	
	inter- action	mance liquid		safety						or above	
	action		Safety monitoring	moni-						75%,	
	through	chromatogra-	carried out through-	toring.						and ab-	
			out the study (not	J						sence	
	gar-	the average	further described).							of green	
	gling	concentration								tea gar-	
	as a	in 10 bottles								gling	
	non-	from the same								when in	
	phar-	production								the	
	ma-	lot (September								water	
	ceuti-	2011) used for								gargling	
	cal in-	gargling in the								group.	
	terven-	study.									
	tion,	B. Tap water									
	specif-										
	ically										
	green										
	tea										
	con-										
	taining										
	highly bioac-										
	tive										
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	fluenza										
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	prop-										
	erties										
Sato- 2 ac-	Healthy Pre-	A. Water	Local administrators	Local	Not	18	60 days	If di-	3 par-	Comple-	9 part
mura tive in- 2005	adults vent URTIs	B. 15 to 30 times dilut-	instructed partici- pants to:	project admin-	spec- ified,	health-	overall	luted	tici-	tion of gargling	ipants did no

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Table 1. Description of interventions in included studies, using the items from the Template for Intervention Description and Replication (TIDieR) checklist

ist (Continued)										·	,
terven-	through	ed 7% povi-	- gargle dose of wa-	istra-	but	sites in	1. Water	done-io-	as-	diary:	com-
tions:	gar-	done-iodine (as	ter or povidone-io-	tors	likely	Japan	gargling:	dine	signed	frequen-	plete di-
	gling	indicated by	dine 3 times/day;	(18	to have	(4 in	20 mL	caused	to	cy of gar-	ary.
A. Wa-	water	manufacturer)	- maintain hand-	health-	been	north-	for 15 s	serious	povi-	gling	
ter gar-	alone,		washing routine;	care	face-	ern re-	at least	dis-	done-io-	and	Average
gling	which		- not change other	profes-	to-face	gion, 9	3 times/	com-	dine	hand-	frequen-
B. Povi-	may		hygiene habits;	sion-	and in-	in cen-	day	fort	gar-	washing	cy of gar- gling /
done-io-	wash		- not take any cold	als)	divid-	tral re-	2. Povi-	or was	gled	Weekly	person /
dine	out		remedies;	provid-	ually,	gion,	done-io-	not	with	monitor-	day:
gar-	pathogen from	15	 complete gargling diary. 	ed in- struc-	at least initially	5 in west-	dine gar- gling:	avail- able,	water instead	ing and encour-	auy.
gling	the		Weekly monitoring of	tions	for in-	ern re-	20 mL of	partic-	as the	agement	With wa-
0. 0	phar-		hygienic actions and	and	struc-	gion)	dilution	ipants	povi-	by local	ter:
	ynx		encouragement to	mon-	tions	gioni	3 times/	were	done-io-	adminis-	
	and		keep up assigned	itor-			day	al-	dine	trators	A: 3.6
	oral		intervention every	ing and			,	lowed	"did		B: 0.8
	cavity		week	en-				to gar-	not		2. 0.0
	through			cour-				gle	agree		Control:
	whirling			age-				with	with		0.9
	wa-			ment.				wa-	them".		14.51
	ter or							ter in-			With
	through							stead.			povi-
	chlo-										done-io- dine:
	rine, or										unie.
	povi-										A.: < 0.1
	done-io-										
	dine for its										B: 2.9
	for its per-										Control
	ceived										Control: 0.2
	viru-										0.2
	VII G										

ABH: alcohol-based rub

AGNPs: ARGOVIT silver nanoparticles ARI: acute respiratory infection

CDC: Centers for Disease Control and Prevention

cidal

properties

CG: control group

CHG: chlorhexidine gluconate CHW: community health worker

CO: carbon monoxide DCCs: daycare centres FM: face masks H: handwashing

HCP: healthcare personnel HCW: healthcare worker HH: hand hygiene

HSG: hand sanitiser group

HSW: hand-washing with soap and water

HW: hand-washing

HWWS: hand-washing with soap

IG: intervention group

IHIP: integrated environmental home-based intervention package

ILI: influenza-like illness IU: international units LFD: lateral flow device

LNS: lipid-based nutrient supplements

LTCFs: long-term care facilities

m: metre min: minute N: nutrition

NGOs: non-governmental organisations

NH: nursing home

NHS: National Health Service

no.: number

NPIs: non-pharmaceutical interventions

PCR: polymerase chain reaction

PM2.5: particulate matter of less than 2.5 microns

RAs: research assistants RIs: respiratory infections RTIs: respiratory tract infections

S: sanitation

SD: standard deviation SES: electrolysed water

SSTI: skin and soft-tissue infection SWG: soap-and-water group TCID: tissue-culture infectious dose URTI: upper respiratory tract infection

W: water

WHO: World Health Organization

WSH: combined water, sanitation and handwashing

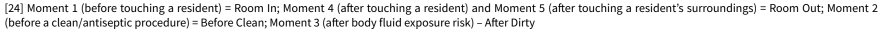
WSHN: combined water, sanitation, handwashing and nutrition

w/w: weight for weight

[1] Filtration efficiency testing was conducted using a Fluke 985 particle counter (volumetric sampling rate of 2.83 litres/ minute. The measurement was taken of particles 0.3-0.5 µm in diameter flowing through the material with a face velocity of 8.5 cm/s. Internal testing found that cloth masks with an external layer made of Pellon 931 polyester fusible

interface ironed onto interlocking knit with a middle layer of interlocking knit could achieve a 60% filtration efficiency. Upon discussions with the manufacturers, the researchers learned that those materials could not be procured. Using materials that were available, the highest filtration efficiency possible was 37%.

- [2] "the exterior and interiors were spunbond and the middle layer was meltblown"
- [3] 10 times with bar soap and water
- [4] Featured the Honorable Prime Minister of Bangladesh Sheikh Hasina, the head of the Imam Training Academy, and the national cricket star Shakib Al Hasan.
- [5] A grassroots organization with a network of volunteers across the country
- [6] "consistent with the WHO guideline that defines physical distancing as one meter of separation." www.who.int/westernpacific/emergencies/covid-19/information/physicaldistancing (accessed 13 June 2022).
- [7] Occupational Safety and Health Administration (OSHA). OSHA technical manual: section VIII: chapter 2: respiratory protection. US Department of Labor. www.osha.gov/dts/ osta/otm/otm viii/otm viii 2.html (accessed 21 April 2020).
- [8] Ministry of Health and Long-Term Care, Public Health Division, Provincial Infectious Diseases Advisory Committee. Preventing respiratory illnesses: protecting patient and staff: infection control and surveillance standards for febrile respiratory illness (FRI) in non-outbreak conditions in acute care hospitals [September 2005] http://www.health.gov.on.ca/ english/providers/program/infectious/diseases/best_prac/bp_fri_080406.pdf (accessed September 11 2009). [URL inactive]
- [9] Before eating, after sneezing, coughing, handling money, using restroom, returning to desk and interacting with others who may be sick
- [10] after coming into classroom, before and after lunch, after break, after physical education, when they went home and after coughing, sneezing or blowing their noses
- [11] after toileting and when visibly dirty plus a protocol for particular circumstances: after coming into the classroom; before and after lunch; after playing outside; when they went home; after coughing, sneezing, or blowing their noses; and after diapering
- [12] 1) when entering into the classroom; 2) after sneezing, coughing, or blowing their nose; 3) after using the toilet/washroom; 4) before eating any food; and 5) when leaving the school at the end of the day
- [13] what to do if hands were dirty, why students should wash their hands, benefits of washing hands and using hand sanitiser, procedure for washing hands using hand sanitiser, to cover mouth and nose with upper part of sleeve while coughing and/or sneezing
- [14] Boyce JM, Pittet D, Healthcare Infection Control Practices Advisory Committee, HICPAC/ SHEA/APIC/IDSA Hand Hygiene Task Force. Guideline for hand hygiene in healthcare settings. Recommendations of the Healthcare Infection Control Practices Advisory Committee and the HICPAC/SHEA/APIC/ IDSA Hand Hygiene Task Force. MMWR Recommendations and Reports 2002;51 (RR-16):1-45. www.cdc.gov/mmwr/preview/mmwrhtml/rr5116a1.htm (accessed 21 April 2020). International Bank for Reconstruction and Development/ World Bank, Bank-Netherlands Water Partnership, Water and Sanitation Program. Hand washing manual: a guide for developing a hygiene promotion program to increase handwashing with soap. http://go.worldbank.org/PJTS4A53C0 (Accessed 16 May 2007). [URL inactive] California State Department of Education. Techniques for Preventing the Spread of Infectious Diseases. Sacramento (CA): California State Department of Education, 1983. Geiger BF, Artz L, Petri CJ, Winnail SD, Mason JW. Fun with Handwashing Education. Birmingham (AL): University of Alabama, 2000. Roberts A, Pareja R, Shaw W, Boyd B, Booth E, Mata JI. A tool box for building health communication capacity. www.globalhealthcommunication.org/tools/29 (Accessed 10 October 2007), [URL inactive] Stark P. Handwashing Technique. Instructor's Packet, Learning Activity Package. Sacramento (CA): California State Department of Education, 1982.
- [15] DIN EN 1500: Chemische Desinfektionsmittel und Antiseptika, Hygienische Händedesinfektion, Prüfverfahren und Anforderungen (Phase 2/Stufe 2). Brüssel (Belgium): CEN, European Comittee for Standardization 1997;1-20.
- [16] DIN EN 12791: Chemische Desinfektionsmittel und Antiseptika, Chirugische Händedesinfektionsmittel Prüfverfahren und Anforderungen (Phase 2/Stufe 2). Brüssel (Belgium): CEN, European Comittee for Standardization 2005;1-31.
- [17] after defaecation, after cleaning an infant who had defaecated, before preparing food, before eating, and before feeding infants
- [18] non-governmental organisation that supports community-based health and development initiatives
- [19] "Healthy Hands" Rules (from Figure 3 in paper): Do use "special soap" when arrive to school, before lunch, after go to bathroom (only if soap and water not available), if rub nose or eyes or if fingers in mouth, if teacher asks. Do not: use "special soap" if hand dirt on them, put "special soap" on another student, play with 'special soap", put hands near eyes after using "special soap".
- [20] Calculated by subtracting each day's soap weight from the previous day's weight. Maximum number of grams of soap consumed for each compound was identified and the day on which the maximum soap consumption was recorded. A per capita estimate of daily soap consumption was calculated
- [21] National Health and Medical Research Council. Staying Healthy in Child Care. Canberra (Australia): Australian Government Publishing Service, 1994
- [22] upon arrival, before and after lunch, and prior to departure
- [23] World Health Organization. (2012). Hand hygiene in outpatient and home-based care and long-term care facilities: a guide to the application of the WHO multimodal hand hygiene improvement strategy and the "My Five Moments For Hand Hygiene" approach. World Health Organization. apps.who.int/iris/handle/10665/78060 (accessed 15 June 2022)



[25] Handsome: handhygiëne in verpleeghuizen.: Zorg voor beter; 2019 May 03. URL: www.zorgvoorbeter.nl/handsome (accessed 7 June 2022)

- [26] Veiligheid en Kwaliteit: Project Handen uit de Mouwen.: Stichting Samenwerkende Rijnmond Ziekenhuizen
- [27] Auditor training.: Hand Hygiene Australia URL: www.hha.org.au/audits/auditor-training (accessed 7 June 2022)
- [28] no long nails, acrylic nails, or polished nails and not wearing a ring, bracelet, wristwatch, brace, or long sleeves.
- [29] Persoonlijke hygiëne: Verpleeghuizen, woonzorgcentra, voorzieningen voor kleinschalig wonen voor ouderen.: Werkgroep Infectie Preventie; 2014. URL: tinyurl.com/wpfqr8p (accessed 7 June 2022)
- [30] knowledge and awareness of HH guidelines, perceived importance of performing HH, perceived behavioural control (i.e. perceived ease or difficulty of performing the behaviour), and habit
- [31] "According to the Dutch national guidelines, HH is mandatory for caregivers before touching/preparing food, before caregivers themselves ate or assisted children with eating, and before wound care; and after diapering, after toilet use/wiping buttocks, after caregivers themselves coughed/sneezed/wiped their own nose, after contact with body fluids (e.g. saliva, vomit, urine, blood, or mucus when wiping children's noses), after wound care, and after hands were visibly soiled." (p. 2495)
- [32] Having touched household items being used by the index patients and/or other symptomatic household contacts, and after coughing/sneezing, before meals, before preparing meals and when returning home
- [33] Which addresses "contextual, psychosocial, and technological factors at the societal, community, interpersonal, individual, and habitual levels". (Luby 2018)
- [34] Hussain F, Luby SP, Unicomb L, Leontsini E, Naushin T, Buckland AJ, et al. Assessment of the acceptability and feasibility of child potties for safe child feces disposal in rural Bangladesh. The American Journal of Tropical Medicine and Hygiene. 2017;97: 469-76.
- [35] Sultana R, Mondal UK, Rimi NA, Unicomb L, Winch PJ, Nahar N, et al. An improved tool for household faeces management in rural Bangladeshi communities. Tropical Medicine & International health 2013;18: 854-60.
- [36] Hulland KR. Leontsini E. Dreibelbis R. Unicomb L. Afroz A. Dutta NC. et al. Designing a handwashing station for infrastructure-restricted communities in Bangladesh using the integrated behavioural model for water, sanitation and hygiene interventions (IBM-WASH). BMC Public Health 2013; 13: 877.
- [37] Menon P, Nguyen PH, Saha KK, Khaled A, Sanghvi T, Baker J, et al. Combining intensive counseling by frontline workers with a nationwide mass media campaign has large differential impacts on complementary feeding practices but not on child growth; results of a cluster-randomized program evaluation in Bangladesh. The Journal of Nutrition 2016;146:2075-84.
- [38] comprised of: senior program manager-intervention delivery, senior program manager-operations, Sanitation Intervention Team leader, senior field research officer, training officer, field research officers, CHW supervisors and CHWs
- [39] SODIS: www.sodis.ch/index_EN.html
- [40] after defecation, after changing diapers, before food preparation and before eating
- [41] 1. Wash both hands with water and soap before eating / handling food 2. Wash both hands with water and soap/ash after defecation 3. Wash both hands with water and soap ash after cleaning baby's bottom 4. Use hygienic latrine by all family members including Children 5. Dispose of children's faeces into hygienic latrines 6. Clean and maintain latrine 7. Construct a new latrine if the existing one is full and fill the pit with soil/ash. 8. Safe collection and storage of drinking water 9. Draw drinking water from arsenic safe water point 10. Wash raw fruits and vegetables with safe water before eating and cover food properly 11. Manage menstruation period safely (p.605)
- [42] Rosenstock IM, Strecher VJ, Becker MH. Social learning theory and the Health Belief Model. Health Education Quarterly 1988;15:175–83.
- [43] Glanz K, Rimer BK, 2005. Theory at a Glance: A Guide for Health Promotion Practice. Washington, DC:US Department of Health and Human Services, Public Health Service, National Institutes of Health, National Cancer Institute.
- [44] Hovland CI, Janis IL, Kelley HH, 1953. Communication and Persuasion; Psychological Studies of Opinion Change. New Haven, CT: Yale University Press.
- [45] Based on family of five, consuming 2L of water per person per day, the bottle would last almost a year
- [46] W: key concepts for water treatment and contamination, procedures for refilling dispenser and distributing bottled chlorine, chlorine testing and reporting; H: HW with soap at critical times and creating supportive environment; S: contamination pathways; N: early initiation and exclusive breastfeeding, complementary and supplementary feeding, LNS procedures for collection from health facility and delivery tracking, teaching mamas how to feed Mwanzobora to the child, cooking demonstration, age-specific messaging about nutrition
- [47] Department of Health and Social Care. Lateral flow device performance data. July 7, 2021. www.gov.uk/government/publications/lateral-flow-device-performance-data (accessed 15 June 2022).
- [48] "applicable to schools as defined in national guidelines were, face to face contact (within 1 metre for any length of time) or skin to skin contact or someone the case coughed on; or within 1 metre for ≥1 minute; or within 1-2 metres for >15 minutes." P.2 of Supplementary appendix

[50] With liquid soap (2% chlorhexidine gluconate) and hand disinfection (0.05% chlorhexidine gluconate and 60-80% ethyl alcohol).

[51] With 80% ethyl alcohol



Table 2. Results from trials of hand hygiene compared to control

Study	Comparison (see Table 1 for details of interventions)	Reported outcomes	Results
Alzaher 2018	Hand-washing workshop and posters versus usual practice	% absence days due to URI	0.39% and 0.72% in intervention group schools; 0.86% and 1.39% in control schools
cluster-RCT			
Saudi Arabia			
Arbogast 2016	Hand sanitiser + wipes + hand foam versus none Both groups received education + signage about hand-washing	Health insurance claims for preventable illnesses per employee	 0.30 claims in intervention; 0.37 in control (27% relative reduction; P = 0.03) 1.45 in intervention; 1.53 in control (5.0% relative reduction in intervention; P = 0.30)
cluster-RCT			
USA		2. Absences per employee	
Ashraf 2020	6 intervention arms: water quality, sanitation, hand washing, combined WSH, nutrition, nutrition + WSH	7-day prevalence of acute respiratory illness (ARI).	Hand washing reduced ARI cases by 32% (RR 0.68, 95% CI 0.52 to 0.88)
cluster-RCT			
Bangladesh			
Azor-Martinez 2016	Hand-washing with soap and- water plus hand sanitiser versus usual hand-washing practices	% absence days due to URI	1.15% in intervention; 1.68% in control. Significantly lower in intervention (P < 0.001)
RCT			
Spain			
Azor-Martinez 2018	Education and hand hygiene with soap and water versus hand hy- giene with sanitiser versus usual hand-washing procedures	URI incidence rate ratio (primary) Percentage difference in absenteeism days	1. HH soap versus control 0.94 (95% CI 0.82 to 1.08); HH sanitiser versus control 0.77 (95% CI 0.68 to 0.88); HH soap versus HH sanitiser 1.21 (95% CI 1.06 to 1.39)
cluster-RCT			
Spain			
			2. HH soap 3.9% versus control 4.2% (P < 0.001); HH sanitiser 3.25% versus control 4.2% (P = 0.026); HH soap 3.9% versus HH sanitiser 3.25% (P < 0.001)
Biswas 2019	Hand sanitiser and respiratory hygiene education and cough/ sneeze hygiene versus no inter- vention	I. ILI incidence rate (at least 1 episode) Laboratory-confirmed influenza	1. 22 per 1000 student-weeks in interven- tion; 27 per 1000 student-weeks in con- trol, not statistically significantly different
cluster-RCT			
Bangladesh			2. 3 per 1000 student-weeks in intervention; 6 per 1000 student-weeks in control, P = 0.01
Correa 2012	Alcohol-based hand sanitiser in addition to hand-washing versus usual hand-washing practice	ARIs in 3rd trimester of follow-up	Hazard ratio for intervention to control 0.69 (95% CI 0.57 to 0.83)
cluster-RCT			
Colombia			
Cowling 2008	Hand hygiene (36 households) versus face mask (mask) versus education (control)	Secondary attack rate for:	1. HH 0.06; mask 0.07; control 0.06
cluster-RCT		1. laboratory-confirmed in-	2. HH 0.18; mask 0.18; control 0.18
Hong Kong		fluenza;	3. HH 0.11; mask 0.10; control 0.11
riong nong		2. ILI definition 1;	



Table 2.	Results from trials of hand hygiene compared to control (Cont	tinued)

Cowling 2009	Hand hygiene (HH) versus face	Secondary attack rate for:	1. HH 5; HH + mask 7; control 10
cluster-RCT	mask + hand hygiene (HH + mask) versus education (control)	1. laboratory-confirmed in-	2. HH 16; HH + mask 21; control 19
Hong Kong		fluenza;	3. HH 4; HH + mask 7; control 5
		2. ILI definition 1;	
		3. ILI definition 2.	
DiVita 2011 (conference abstract)	Hand-washing stations with soap and motivation vs none	1. SAR for laboratory-confirmed influenza	1. SAR higher in intervention group (11.0% versus 7.5%)
RCT		2. SAR for ILI	2. SAR higher in intervention group
Bangladesh			(14.2% versus 11.9%)
Feldman 2016	Hand disinfection + soap and wa-	1. Number of respiratory in-	1. 11 in each group
cluster-RCT	ter installed versus none	fections	2. 112 in intervention; 104 in control
Israel		2. Number of off-duty days	
Gwaltney 1980	Virucidal hand wash versus	1. Number with illness after	1. 0 of 8 in intervention; 7 of 7 in control
RCT	placebo immediate exposure		2. 1 of 10 in intervention; 6 of 10 in contro
USA		2. Number with illness after 2-hour delay in exposure	,
Hubner 2010	Hand disinfection provided versus none	Odds ratios (95% CI) (inter-	1. 1.02 (0.20 to 5.23)
RCT		vention:control) 1. Influenza 2. Common cold 3. Sinusitis	2. 0.35 (0.17 to 0.71)
Germany			3. 1.87 (0.52 to 6.74)
			4. 0.62 (0.31 to 1.25)
			5. 0.38 (0.14 to 0.99)
		4. Sore throat	6. 0.45 (0.22 to 0.91)
		5. Fever	
		6. Cough	
Ladegaard 1999	Hand hygiene and education ver-	Sick days during the "effect	22 days/child in the intervention group
RCT	sus none	period"	versus 36 days/child in the control group
Denmark			
Larson 2010	Education versus education with	Incidence rate ratios	1. HS 29; HS + masks 39; control 35
cluster-RCT	alcohol-based hand sanitiser versus education with hand sanitiser	(episodes per 1000 per- son-weeks) for:	2. HS 1.9; HS + masks 1.6; control 2.3
USA	and face masks	1. URI; 2. ILI;	3. HS 0.6; HS + masks 0.5; control 2.3
		3. influenza.	4. HS 0.14; HS + masks 0.12; control 0.14
		Secondary attack rates for: 4. URI/ILI/influenza; 5. ILI/influenza.	5. HS 0.02; HS + masks 0.02; control 0.02



Little 2015	Bespoke automated web-based hand hygiene motivational inter-	Number of participants with 1 or more episodes of URI	Risk ratio for intervention to control 0.86 (95% CI 0.83 to 0.89; P < 0.001)	
RCT	vention with tailored feedback	1 of more episodes of oki	(95% C1 0.65 to 0.69, F < 0.001)	
England	versus none			
Luby 2005	Antibacterial soap and education	1. Cough or difficulty	All outcomes significantly lower than cor	
RCT	about hand-washing versus plain soap and education versus none	breathing in children < 15 yrs (episodes/100 per- son-weeks)	trol	
Pakistan			1. 4.21 in antibacterial soap group; 4.16 in plain soap group; 8.50 in control group	
		 Congestion or coryza in children < 15 yrs (episodes/100 per- son-weeks) 	2. 7.32 in antibacterial soap group; 6.87 in plain soap group; 14.78 in control group	
		3. Pneumonia in children < 5 yrs (episodes/100 per- son-weeks)	3. 2.42 in antibacterial soap group; 2.20 in plain soap group; 4.40 in control group	
Millar 2016 cluster-RCT	Standard educational promotion of hand-washing versus en-	Incidence rates of ARI over 20 months	37.7 enhanced + body wash; 29.3 enhanced; 35.3 standard; RR for enhanced	
USA	hanced promotion versus promo- tion plus a once-weekly applica- tion of chlorhexidine-based body wash	20 months	body wash to standard 1.07 (95% CI 1.0 to 1.11); RR for enhanced to enhanced body wash 0.78 (95% CI 0.75 to 0.81)	
Morton 2004	Alcohol gel plus education versus	Absence due to infectious	Results not stated numerically	
cluster-RCT	regular hand-washing	illness		
cross-over study				
USA				
Nicholson 2014	Combination hand-washing pro-	Target children:	1. 16 in intervention; 19 in control	
cluster-RCT	motion with provision of free soap versus none	1. Episodes of ARI (per 100 person-weeks)	2. 1.2 in intervention; 1.7 in control	
India		2. School absence episodes (per 100 person-days)	3. 10 in intervention; 11 in control	
		Families: 3. Episodes of ARI		
Priest 2014	Hand hygiene education and hand sanitiser versus education	1. % absence days due to respiratory illness	1. 0.84% in intervention group; 0.80% in control (P = 0.44)	
cluster-RCT	alone	2. % absence days due to	2. 1.21% in intervention group; 1.16% in	
New Zealand		any illness	control (P = 0.35)	
Ram 2015	Education to promote intensive	1. Secondary attack ratio for	1. 1.24 (95% CI 0.93 to 1.65)	
RCT	hand-washing in households plus soap provision versus none	intervention to control for ILI	2. 2.40 (95% CI 0.68 to 8.47)	
Bangladesh		2. Laboratory-confirmed in- fluenza		
Roberts 2000 cluster-RCT	Hand-washing programme with training for staff and children versus none	Incidence rate ratio for ARI	IRR 0.92 for intervention to control (95% CI 0.86 to 0.99)	



Table 2. Results from trials of hand hygiene compared to control (Continued) Australia

Sandora 2008	Hand sanitiser and education	Incidence rates for	0.43 in intervention; 0.42 in control	
uster-RCT versus none		ARI (episodes per per- son-month)	, , , , , , , , , , , , , , , , , , ,	
USA		Son moneny		
Savolainen-Kopra 2012	Hand hygiene with soap and water (IR1 group) versus with alco	1. Number of respiratory infection episodes/week	1. 0.076 in IR1; 0.085 in IR2; 0.080 in control, NS	
cluster-RCT Finland	versus control (none); interven- tion groups also received educa-		2.0.097in IR1;0.107in IR2;0.104in control,NS	
Timanu	tion	3. Number of reported sick leave episodes/week	3. 0.042 in IR1; 0.035 in IR2; 0.035 in control. Significantly higher in IR1 compared with control	
Simmerman 2011	Hand-washing (HW) versus hand- washing plus paper surgical face	Odds ratios for secondary attack rates for influenza	OR for HW: control 1.20 (95% CI 0.76 to 1.88)	
cluster-RCT Thailand	masks (HW + FM) versus control (none)		OR for HW + masks: control 1.16 (95% CI 0.74 to 1.82)	
			OR for HW + masks: HW 0.72 (95% CI 0.21 to 2.48)	
Stebbins 2011	Training in hand and respiratory	Incidence rate ratios for in-	1. IRR 0.81 (95% CI 0.54 to 1.23)	
cluster-RCT	(cough) hygiene + hand sanitiser versus none	tervention to control for: 1. laboratory-confirmed influenza (RT-PCR); 2. influenza-A; 3. absence.	2. IRR 0.48 (95% CI 0.26 to 0.87)	
USA			3. IRR 0.74 (95% CI 0.56 to 0.97)	
Swarthout 2020	There were 6 intervention	orinated drinking dren 0.90 to 1 mproved sanitation	No evidence of an effect: RR 0.97, 95% CI	
cluster-RCT	groups: chlorinated drinking water (W), improved sanitation		0.90 to 1.04.	
Kenya	(S), handwashing with soap (H), combined WSH, improved nutrition (N) through counselling lipid based nutrient supplementation (LNS) combined WSHN There were 2 control groups passive control (no promotional visits), a double-sized active control (monthly visits to measure midupper arm circumference)			
Talaat 2011	Mandatory hand-washing intervention + education versus none	Number of absence days due to ILI	1. 917 in intervention; 1671 in control (P < 0.001)	
cluster-RCT	vention - education versus notice	2. Number of absence days	2. 13,247 in intervention; 19,094 in control	
Egypt		2. Number of absence days	(P < 0.001)	
Teesing 2021	Hand hygiene enhancement ac-	Incidence of gastroenteritis,	Hand hygiene reduced risk of ILI (RR 0.51,	
cluster-RCT	tivities versus no activities.	influenza-like illness (ILI), assumed pneumonia, uri-	95% CI 0.31 to 0.83)	
Netherlands		nary tract infections (UTIs), and infections caused MRSA in residents		



Temime 2018	Hand hygiene with alcohol-based	Incidence rate of ARI clus-	2 ARI clusters in intervention; 1 in contro	
cluster-RCT	hand rub, promotion, staff educa- tion, and local work groups ver-	ters (5 or more people in same nursing home)		
France	sus none			
Turner 2012	Antiviral hand treatment versus no treatment	Number of rhinovirus in- fections	1. 49 in intervention; 49 in control, NS	
RCT	no deadnent		2. 56 in intervention; 72 in control, NS	
USA		2. Common cold infections	3. 26 in intervention; 24 in control, NS	
		3. Rhinovirus-associated ill- nesses		
White 2001	Hand rub with benzalkonium chloride (hand sanitiser) versus	ARI symptoms	30% to 38% decrease of illness and absenteeism (RR for illness absence inci-	
DB-RCT	placebo	Laboratory: testing of viru- cidal and bactericidal activi-	dence 0.69; RR for absence duration 0.	
USA		ty of the product		
Yeung 2011	Alcohol-based hand gel + mate-	Difference between pre-	0.63/1000 reduction in intervention	
cluster-RCT	rials + education versus control (basic life support workshop)	in pneumonia infections	group; 0.16/1000 increase in control	
Hong Kong		recorded in residents		
Zomer 2015 cluster-RCT	4 components:	Incidence rate ratio for in-	IRR 1.07 (95% CI 0.97 to 1.19)	
	 Hand hygiene products, pa- per towel dispensers, soap, al- 	tervention to control for common cold	8.2 episodes per child-year in interven-	
Netherlands	cohol-based hand sanitiser, and hand cream provided for 6 months		tion; 7.4 episodes per child-year in control	
	2. Training and booklet			
	3. 2 team training sessions aimed at hand hygiene improvement			
	Posters and stickers for care- givers and children as reminders.			
	Combination versus usual practice			

ARI: acute respiratory infection

CI: confidence interval

cluster-RCT: cluster-randomised controlled trial DB-RCT: double-blind randomised controlled trial

HH: hand hygiene
HS: hand sanitiser
HW: hand-washing
ILI: influenza-like illness
IRR: incidence rate ratio
NS: non-significant
OR: odds ratio

RCT: randomised controlled trial

RR: risk ratio

RT-PCR: reverse-transcriptase polymerase chain reaction

SAR: secondary attack rate



URI: upper respiratory infection yrs: years

Table 3. Results from trials of hand hygiene + medical/surgical masks compared to control

Study	Comparison (see Table 1 for details of interventions)	Reported outcomes	Results	
Aelami 2015 (conference abstract)	Hand hygiene education + al- cohol-based hand rub + soap +	Proportion with ILI (de- fined as presence of ≥ 2 of	52% in intervention; 55.3% in control (P < 0.001)	
RCT	surgical masks vs none	the following during their stay: fever, cough, and sore		
Saudi Arabia		throat)		
Aiello 2010	Face mask use (FM) vs face	1. ILI	Significant reduction in ILI cases in both in-	
cluster-RCT	masks + hand hygiene (FM + HH) vs control	2. Laboratory-confirmed influenza A or B	tervention groups compared with control over weeks 3 to 6	
USA	Note that this study is not included in meta-analysis as each treatment group included only 1 cluster.		No significant differences between FM and FM + HH	
Aiello 2012	Face mask use (FM) vs face	1. Clinical ILI	1. Non-significant reductions in FM group	
cluster-RCT	masks + hand hygiene (FM + HH) vs control	2. Laboratory-confirmed influenza A or B	compared with control over all weeks. Sig nificant reduction in FM + HH group com-	
USA			pared with control in weeks 3 to 6	
			2. Non-significant reductions in both intervention groups compared with control	
Cowling 2009	Hand hygiene (HH) vs hand hy-	Secondary attack ratio for:	1. HH 5; HH + mask 7; control 10	
cluster-RCT	giene plus face masks (HH + mask) vs control	1. laboratory-confirmed in- fluenza;	2. HH 16; HH + mask 21; control 19 3. HH 4; HH + mask 7; control 5	
Hong Kong		2. ILI definition 1;3. ILI definition 2.		
Larson 2010	Education (control) vs educa-	Incidence rate ratios	1. HS 29; HS + mask 39; control 35	
cluster-RCT	tion with alcohol-based hand sanitiser (HS) vs education +	(episodes per 1000 per- son-weeks) for:	2. HS 1.9; HS + mask 1.6; control 2.3 3. HS 0.6; HS + mask 0.5; control 2.3	
USA	HS + face masks (HS + mask)	1. URI; 2. ILI; 3. influenza.	4. HS 0.14; HS + mask 0.12; control 0.14 5. HS 0.02; HS + mask 0.02; control 0.02	
		Secondary attack rates for: 4. URI/ILI/influenza; 5. ILI/influenza.		
Simmerman 2011	Control vs hand-washing (HW)	Odds ratio for secondary at-	OR for HW: control 1.20 (95% CI 0.76 to 1.88)	
cluster-RCT	vs hand-washing + paper sur- gical face masks (HW + mask)	tack rates for influenza	OR for HW + mask: control 1.16 (95% CI 0.74 to 1.82)	
Thailand			OR for HW + mask: HW 0.72 (95% CI 0.21 to 2.48)	
Suess 2012	Face mask + hand hygiene	Secondary attack rates in	1. Mask 9; mask + HH 15; control 23	
cluster-RCT	(mask + HH) vs face masks on- ly (mask) vs none (control)	household contacts: 1. Laboratory-confirmed in-	2. Mask 9; mask + HH 9; control 17	
Germany		fluenza 2. ILI		



CI: confidence interval

cluster-RCT: cluster-randomised controlled trial

FM: face mask HH: hand hygiene HS: hand sanitiser HW: hand-washing ILI: influenza-like illness

OR: odds ratio

RCT: randomised controlled trial URI: upper respiratory infection

vs: versus

Table 4. Results from trials of soap + water compared to hand sanitisers

Study	Comparison (see Table 1 for details of interventions)	Reported out- comes	Results
Azor-Martinez 2018 cluster-RCT Spain	Education and hand hygiene with soap and water (HH soap) vs hand hygiene with sanitiser (HH sanitiser) vs usual hand-washing procedures	URI incidence rate ratio (primary) Percentage dif- ference in absenteeism days	1: HH soap vs control 0.94 (95% CI 0.82 to 1.08); HH sanitiser vs control 0.77 (95% CI 0.68 to 0.88); HH soap vs HH sanitiser 1.21 (95% CI 1.06 to 1.39) 2: HH soap 3.9% vs control 4.2% (P < 0.001); HH sanitiser 3.25% vs control 4.2% (P = 0.026); HH soap 3.9% vs HH sanitiser 3.25% (P < 0.001)
Pandejpong 2012 cluster-RCT Thailand	Alcohol hand gel applied every 60 minutes vs every 120 minutes vs once before lunch (3 groups).	Absent days due to confirmed ILI/ present days	0.017 in every hour group; 0.025 in every 2 hours group; 0.026 in before lunch group. Statistically significant difference between every hour group and before lunch group, and between every hour and every 2 hours groups
Savolainen-Kopra 2012 cluster-RCT Finland	Hand hygiene with soap and water (IR1 group) vs with alco- hol-based hand rub (IR2 group) vs control (none); intervention groups also received education	1. Number of respiratory infection episodes/week 2. Number of reported infection episodes/week 3. Number of reported sick leave episodes/week	1. 0.076 in IR1; 0.085 in IR2; 0.080 in control, NS 2: 0.097 in IR1; 0.107 in IR2; 0.104 in control, NS 3: 0.042 in IR1; 0.035 in IR2; 0.035 in control. Significantly higher in IR1 compared with control
Turner 2004a and- Turner 2004b RCT Canada	Study 1. Ethanol vs salicylic acid 3.5% vs salicylic acid 1% and pyroglutamic acid 3.5% Study 2. Skin cleanser wipe vs ethanol (control)	% of volunteers infected with rhi-novirus	7% in each intervention group; 32% in control (study 1) 22% in intervention, 30% in control (study 2)

CI: confidence interval

cluster-RCT: cluster-randomised controlled trial

HH: hand hygiene ILI: influenza-like illness NS: non-significant

RCT: randomised controlled trial URI: upper respiratory infection

vs: versus



Table 5. Results from trials of surface/object disinfection (with or without hand hygiene	e) compared to control
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Study	Comparison (see Table 1 for details of interventions)	Reported outcomes	Results	
Ban 2015	Hand hygiene products, surface	1. Respiratory illness	1. OR 0.47 for intervention to control (95%	
cluster-RCT	cleaning and disinfection provided to families and kindergartens vs none	Cough and expectoration	CI 0.38 to 0.59) 2. OR 0.56 (95% CI 0.48 to 0.65)	
China				
Carabin 1999	One-off hygiene education and disin-	Difference in inci-	0.28 episodes per 100 child-days lower in	
cluster-RCT	fection of toys with bleach vs none	dence rate for URTI (cluster-level result)	intervention group (95% CI 1.65 lower to 1.08 higher); URTI incidence rate IRR 0.80	
Canada			(95% CI 0.68 to 0.93)	
Ibfelt 2015	Disinfectant washing of linen and	Presence of respirato-	Statistically significant reduction in inter-	
cluster-RCT	toys by commercial company every 2 weeks vs usual care	ry viruses on surfaces	vention group in adenovirus, rhinovirus, RSV, metapneumovirus, but not other	
Denmark			viruses including coronavirus	
Kotch 1994	Training in hand-washing and dia-	Respiratory illness in-	1. 14.78 episodes per child-year in inter-	
RCT	pering and disinfection of surfaces vs none	cidence rate in: 1. children < 24	vention; 15.66 in control	
USA		months;	2. 12.87 in intervention; 11.77 in control	
		2. children >= 24 months.		
McConeghy 2017	Staff education, cleaning products,	Infection rates	Upper respiratory infections not reliably	
RCT	and audit of compliance and feedback vs none		recorded or reported.	
USA				
Sandora 2008	Hand sanitiser and disinfection of	Absence due to respi-	Rate ratio 1.10 for intervention to control	
cluster-RCT	classroom surfaces vs materials about good nutrition (control)	ratory illness (multi- variable analysis)	(95% CI 0.97 to 1.24)	
USA				

CI: confidence interval

cluster-RCT: cluster-randomised controlled trial

IRR: incident rate ratio

OR: odds ratio

RCT: randomised controlled trial RSV: respiratory syncytial virus URTI: upper respiratory tract infection

vs: versus

Table 6. Results from trials of complex interventions compared to control

Study	Comparison (see Table 1 for details of interventions)	Reported out- comes	Results
Complex hygien	e and sanitation interventions compar	ed to control	
Chard 2019	Complex sanitation intervention	Pupil-reported	NS difference between groups. 29% of interven-
cluster-RCT	and education vs none	symptoms of res-	tion group; 32% control group; adjusted risk ratio 1.08 (95% CI 0.95 to 1.23)



Table 6. R	esults from trials of	f complex interventions compared	d to control (Continued)
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Laos		piratory infection over 1 week	
Hartinger 2016	Cooking and sanitation provision and education vs none	Number of ARI	NS difference between groups. Risk ratio for intervention to control 0.95 (95% CI 0.82 to 1.10)
cluster-RCT	and education vs none	episodes per child- year	vention to control 0.95 (95% Cl 0.82 to 1.10)
Peru			
Huda 2012	Sanitation provision and educa-	Respiratory illness	12.6% in intervention group; 13.0% in control
cluster-RCT	tion vs none er-RCT		group. Not adjusted for multiple outcome measurements. No CIs reported.
Bangladesh			
Najnin 2019	Sanitation and behaviour change	Respiratory illness	2.8% in intervention group; 2.9% in control group
cluster-RCT	intervention (plus cholera vac- cine) vs none	in past 2 days	
Bangladesh			

ARI: acute respiratory infection

CI: confidence interval

cluster-RCT: cluster-randomised controlled trial

NS: non-significant

RCT: randomised controlled trial

vs: versus

Table 7. Results from trials of virucidal tissues compared to control

Study	Comparison	Reported outcomes	Results	
Virucidal tissues cor	Virucidal tissues compared with placebo or no tissues			
Farr 1988a and Farr 1988b	Trial 1. Virucidal nasal tissues vs placebo vs none	Respiratory illnesses per person over 24 weeks Trial 1	Trial 1: 3.4 in tissues group; 3.9 in placebo group; 3.6 in no-tissues	
cluster-RCT	Trial 2. Virucidal nasal tis-	Trial 2	group Trial 2: 3.4 in tissues group; 3.6 in	
USA Trial 1 and Trial 2	sues vs placebo		placebo group NS	
Longini 1988	Virucidal nasal tissues vs	Secondary attack rate of viral infec-	10.0 in intervention; 14.3 in placebo;	
DB-PC RCT	placebo	tions (number of infections in house- hold members of index case)	NS	
USA				

cluster-RCT: cluster-randomised controlled trial DB-PC: double-blind, placebo-controlled

NS: non-significant

RCT: randomised controlled trial

vs: versus

Table 8. Summary of main results of the review for the primary outcomes

Interventions	RCT/cluster-RCT (N = 78)
Medical/surgical masks	Masks (medical/surgical) compared to no masks



Table 8. Summary of main results of the review for the primary outcomes (Continued)

9 trials in the community showed no effect on ILI (RR 0.95, 0.84 to 1.09) (Abaluck 2022; Aiello 2010; Alfelali 2020; Barasheed 2014; Canini 2010; Cowling 2008;; MacIntyre 2009;; MacIntyre 2016; Suess 2012); and 6 trials in the community showed no effect on laboratory-confirmed influenza 95% CI RR 1.01 (0.72 to 1.42) (Aiello 2012; Alfelali 2020; Bundgaard 2021; Cowling 2008; MacIntyre 2009; Suess 2012). Two trials in health care workers where the control group wore masks if they were required provided inconclusive results with very wide confidence intervals (Jacobs 2009; MacIntyre 2015).

Medical/surgical masks versus other (non-N95) masks: 1 trial showed more ILI with cloth mask (RR 13.25, 1.74 to 100.97) (MacIntyre 2015); 1 trial showed no effect of catechin-treated masks on influenza (adjusted OR 2.35, 0.40 to 13.72) (Ide 2016).

N95 respirator

N95 respirators compared to medical/surgical masks

3 trials showed no difference for clinical respiratory illness (RR 0.70, 0.45 to 1.10) (MacIntyre 2011; MacIntyre 2013; Radonovich 2019);

4 trials showed no difference for ILI (95% CI RR 0.81, 0.62 to 1.05) (Loeb 2009; MacIntyre 2009; MacIntyre 2011; Radonovich 2019); and 4 trials showed no difference for laboratory-confirmed influenza (95% CI RR 1.06, 0.81 to 1.38) (Loeb 2009; MacIntyre 2009; MacIntyre 2011; Radonovich 2019).

4 trials conducted in HCWs: 3 trials showed no difference for clinical respiratory illness (RR 0.70, 0.45 to 1.10) (MacIntyre 2011; MacIntyre 2013; Radonovich 2019); 3 trials showed no difference for ILI (RR 0.64, 0.32 to 1.31) (Loeb 2009; MacIntyre 2011; Radonovich 2019); and 3 trials showed no difference for laboratory-confirmed ILI (RR 1.02, 0.73 to 1.43) (Loeb 2009; MacIntyre 2011; Radonovich 2019).

Hand hygiene

Hand hygiene compared to control

19 trials found an effect on combined outcome (ARI or ILI or influenza) (RR 0.89, 0.83 to 0.94) (Ashraf 2020; Azor-Martinez 2018; Biswas 2019; Correa 2012; Cowling 2008; Cowling 2009; Hubner 2010; Larson 2010; Little 2015; Millar 2016; Nicholson 2014; Ram 2015; Roberts 2000; Sandora 2005; Simmerman 2011; Stebbins 2011; Swarthout 2020; Teesing 2021; Zomer 2015); 9 trials showed an effect on ARI (RR 0.86, 0.81 to 0.90) (Ashraf 2020; Azor-Martinez 2018; Correa 2012; Larson 2010; Little 2015; Millar 2016; Nicholson 2014; Sandora 2005; Swarthout 2020); 11 trials showed no effect on ILI (RR 0.94, 0.81 to 1.09) (Biswas 2019; Cowling 2008; Cowling 2009; Hubner 2010; Larson 2010; Little 2015; Ram 2015; Roberts 2000; Simmerman 2011; Teesing 2021; Zomer 2015); and 8 trials no effect on laboratory-confirmed influenza (RR 0.91, 95% CI 0.63 to 1.30) (Biswas 2019; Cowling 2008; Cowling 2009; Hubner 2010; Larson 2010; Ram 2015; Simmerman 2011; Stebbins 2011).

Hand hygiene + medical/surgical masks

Hand hygiene + medical/surgical masks compared to control

7 trials showed no effect on ILI (95% CI RR 0.97, 0.80 to 1.19) (Aelami 2015; Aiello 2010; Aiello 2012; Cowling 2009; Larson 2010; Simmerman 2011; Suess 2012); and 4 trials showed no effect on laboratory-confirmed influenza (RR 0.97, 0.69 to 1.36) (Cowling 2009; Larson 2010; Simmerman 2011; Suess 2012).

Hand hygiene + medical/surgical masks compared to hand hygiene

3 trials showed no effect on ILI (RR 1.03, 0.69 to 1.53) or laboratory-confirmed influenza (RR 0.99, 0.69 to 1.44) (Cowling 2009; Larson 2010; Simmerman 2011).

Soap + water compared to sanitiser, and comparisons of different types of sanitiser

Soap + water compared to sanitiser, and comparisons of different types of sanitiser

1 trial hand sanitiser was more effective than soap and water (Azor-Martinez 2018); 1 trial there was no difference (Savolainen-Kopra 2012).

2 trials in children antiseptic was more effective (Morton 2004; White 2001); 1 trial in children antiseptic = soap (Luby 2005).

1 trial hand sanitisers were better than placebo, but no difference between sanitisers (Turner 2004a); 1 trial no difference between different wipes (Turner 2004b).



Surface/object disinfection (with or without hand hygiene) compared to control	Surface/object disinfection compared to control 2 trials were effective on ARI (Ban 2015; Carabin 1999); 1 trial was effective for viruses detected on surfaces (Ibfelt 2015); 2 trials showed no difference in ARIs (Kotch 1994; McConeghy 2017).
Disinfection of living quarters	-
Complex interventions	Complex interventions compared to control
	4 trials in low-income countries found no effect on respiratory viral illness (Chard 2019; Hartinger 2016; Huda 2012; Najnin 2019).
Physical interventions (masks, gloves, gowns combined)	-
Gloves	-
Gowns	-
Physical distancing	Physical distancing compared to self-isolation
	1 trial reported 1 positive SARS-CoV-2 case in the fitness centre access arm versus 0 in the no access arm (risk difference 0.05%, 95% CI – 0.05 to 0.16%) (Helsingen 2021)
Quarantine in the community	Quarantine compared to control
	1 trial effective for influenza (Cox hazard ratio 0.799, 95% CI 0.66 to 0.97) (Miyaki 2011).
	Daily contact testing compared to self-isolation
	1 trial showed non-inferiority of daily contact testing of school-based contacts compared to self-isolation for SARS-CoV-2 (RR 0.96, 95% CI 0.75 to 1.22) (Young 2021)
Eye protection	Glasses compared to no glasses 1 pragmatic RCT conducted in Norway wearing any type of eyeglasses when close to other people outside their home (on public transport, in shopping malls etc.), over a 14-day period. Positive COVID-19 tests based on self-reporting were 9.6% and 11.5% (RR 0.83, 95% CI 0.69 to 1.00) (Fretheim 2022a).
Gargling	Gargling compared to control 1 trial gargling with tap water was effective, povidone-iodine was not effective (Satomura 2005); 1 trial gargling with green tea was not more effective than tap water (Ide 2014); 1 trial gargling with water was not effective (Goodall 2014); pooling of 2 trials showed no effect of gargling (RR 0.91, 95% CI 0.63 to 1.31) (Goodall 2014; Satomura 2005).
	Mouth/nose rinse compared to control
	2 trials found a large protective effect on SARS-CoV-2 (RR 0.07, 0.01 to 0.23) (Almanza-Reyes 2021; Gutiérrez-García 2022).
Virucidal tissues	Virucidal tissues compared to control
	1 trial had a small effect (Farr 1988a) ("The study authors conclude that virucidal tissues have only a small impact upon the overall rate of natural acute respiratory illnesses"); 2 trials showed a non significant difference (Farr 1988b; Longini 1988).
Nose wash	-



HCW: healthcare worker ILI: influenza-like illness

OR: odds ratio

RCT: randomised controlled trial

RR: risk ratio

Table 9. Trial authors' outcome definitions

Study Outcome definitions		
Masks (n = 16)		
Abaluck 2022	COVID-19 symptoms as per the WHO case definition of probable COVID-19 given epidemiologi-	
cluster-RCT	cal risk factors: (i) fever and cough; (ii) 3 or more of the following symptoms (fever, cough, general weakness and/or fatigue, headache, myalgia, sore throat, coryza, dyspnoea, anorexia, nausea,	
Bangladesh	and/or vomiting, diarrhoea, and altered mental status); or (iii) loss of taste or smell. The owner of the household's primary phone completed surveys by phone or in-person at weeks 5 and 9 after the start of the intervention. They were asked to report symptoms experienced by any household member consistent with the WHO. COVID-19 case definition.	
	Laboratory: seropositivity was defined by having detectable IgG antibodies in blood samples against SARS-CoV-2, using the SCoV-2 Detect™ IgG ELISA kit (InBios, Seattle, Washington). This assay detects IgG antibodies against the spike protein subunit (S1) of SARS-CoV-2.	
	Safety: harms were not directly assessed in this study, but it is stated no adverse events were reported.	
Alfelali 2020	Laboratory: swabs were placed it into UTM™ (COPAN) viral transport media. Swabs labelled with	
cluster-RCT	the participant's unique barcode number were stored in an icebox at –20°C before being re-stored by day's end in a –80°C freezer at the laboratory of the Hajj Research Center at Umm Al-Qura Uni-	
Haj in Makkah, Saudi Arabia	versity, Makkah. After Hajj, these swabs were shipped in refrigerated or cold containers to the Centre for Infectious Disease and Microbiology Laboratory Services, Westmead Hospital, NSW, Australia. There, nucleic acid was extracted with the Qiagen bioROBOT EZ instrument (Qiagen, Valencia, CA), and amplification was performed using the Roche LC 480 (Roche Diagnostics GmbH, Mannheim, Germany) instrument. Respiratory viruses were detected using a real-time, multiplex reverse transcription polymerase chain reaction assay targeting human coronaviruses (OC43, 229E and NL63), influenza A and B viruses, respiratory syncytial virus (RSV), parainfluenza viruses 1 to 3, human metapneumovirus, rhinovirus, enterovirus and adenovirus. Middle East respiratory syndrome coronavirus (MERS-CoV) assay targeting the upstream region of the E gene (upE) was also performed.	
	Safety: harms of using face masks were difficulty in breathing (26.2%); discomfort (22%); and a small minority (3%) reported feeling hot, sweating, a bad smell or blurred vision with eyeglasses.	
Bundgaard 2021	Laboratory: viral RNA was extracted from swab samples in DNA/RNA Shield (Zymo Research) using Quick-RNA Microprep Kit (Zymo Research) with the below modifications. 200 µl samples were	
RCT	incubated for 1 min with proteinase K (Qiagen) in a final concentration of 0.2 $\mu g/\mu l$ prior to treat-	
Denmark	ment with lysis buffer (Quick-RNA Microprep Kit). Only a single washing step using 400 µl RNA Wash Buffer (Quick-RNA Microprep Kit) was performed before elution in 15µl RNase free water.	
	Participants tested for SARS-CoV-2 IgM and IgG antibodies in whole blood using a point-of-care test (Lateral Flow test [Zhuhai Livzon Diagnostics]) according to the manufacturer's recommendations. After puncturing a fingertip with a lancet, they withdrew blood into a capillary tube and placed 1 drop of blood followed by 2 drops of saline in the test chamber in each of the 2 test plates (IgM and IgG).	
	Safety: harms were not mentioned as an outcome in the methods, but psychological adverse effects were mentioned, and 14% reported adverse reactions from other people regarding wearing a face mask.	



Cowling 2008

cluster-RCT

Hong Kong

Laboratory:

QuickVue Influenza A+B rapid test

Viral culture on MDCK (Madin-Darby canine kidney cells)

Samples were harvested using NTS, but the text refers to a second procedure from June 2007 onwards with testing for influenza viruses on index participants with a negative QuickVue result but a fever \geq 38 °C who were also randomised and further followed up. Data on clinical signs and symptoms were collected for all participants, and an additional NTS was collected for later confirmation of influenza infection by viral culture. It is noteworthy that dropout was higher in households of index participants who had a negative result on the rapid influenza test (25/44, 57%) compared to those who had a positive result (45/154, 29%).

Effectiveness: secondary attack ratios (SAR): SAR is the proportion of household contacts of an index case who subsequently were ill with influenza (symptomatic contact individuals with at least 1 NTS positive for influenza by viral culture or PCR)

3 clinical definitions were used for secondary analysis:

- 1. fever ≥ 38 °C or at least 2 of the following symptoms: headache, coryza, sore throat, muscle aches and pains;
- 2. at least 2 of the following S/S: fever ≥ 37.8 °C, cough, headache, sore throat and muscle aches and pains; and
- 3. fever of ≥ 37.8 °C plus cough or sore throat.

Safety: harms were not mentioned as an outcome in the methods, but it was reported in the results that there were no adverse events.

Jacobs 2009

RCT Japan Laboratory-confirmation not reported.

Effectiveness: URTI is defined on the basis of a symptom score with a score > 14 being a URTI according to Jackson's 1958 criteria ("Jackson score"). These are not explained in text, although the symptoms are listed in Table 3 (any, sore throat, runny nose, stuffy nose, sneeze, cough, headache, earache, feel bad) together with their mean and scores (SD) by intervention arm.

Safety: the text does not mention or report harms. These appear to be indistinguishable from URTI symptoms (e.g. headache, which is reported as of significantly longer duration in the intervention arm). Compliance is self-reported as high (84.3% of participants).

Loeb 2009

cluster-RCT HCW Canada Clinical respiratory illness, influenza-like illness, and laboratory-confirmed respiratory virus infection.

- 1. Clinical respiratory illness, defined as 2 or more respiratory symptoms or 1 respiratory symptom and a systemic symptom.
- 2. Influenza-like illness, defined as fever ≥ 38 °C plus 1 respiratory symptom.
- 3. Laboratory-confirmed viral respiratory infection. Laboratory confirmation was by nucleic acid detection using multiplex RT-PCR for 17 respiratory viruses.

Safety: harms were not mentioned as an outcome in the methods, but it is stated in the results that no adverse events were reported by participants.

MacIntyre 2009 cluster-RCT Australia

Eligibility criteria were stipulated as follows:

- 1. the household contained > 2 adults > 16 years of age and 1 child 0 to 15 years of age;
- 2. the index child had fever (temperature > 37.8 °C) and either a cough or sore throat;
- 3. the child was the first and only person to become ill in the family in the previous 2 weeks;
- 4. adult caregivers consented to participate in the study; and
- 5. the index child was not admitted to the hospital.

Definitions used for outcomes:



- 1. ILI defined by the presence of fever (temperature > 37.8 °C), feeling feverish or a history of fever, > 2 symptoms (sore throat, cough, sneezing, runny nose, nasal congestion, headache), or 1 of the symptoms listed plus laboratory confirmation of respiratory viral infection.
- 2. Laboratory confirmation: multiplex RT-PCR tests to detect influenza A and B and RSV, PIV types 1 to 3, picornaviruses (enteroviruses or rhinoviruses), adenoviruses, coronaviruses 229E and OC43, and hMPV plus > 1 symptom

Effectiveness: presence of ILI or a laboratory diagnosis of respiratory virus infection within 1 week of enrolment.

Safety: harms not mentioned as an outcome in the methods, but it is reported in the results that more than 50% of participants reported concerns with mask wearing, mainly that wearing a face mask was uncomfortable, but there were no significant differences between the P2 (N95) and surgical mask groups. Other concerns were that the child did not want the parent wearing a mask.

Aiello 2010

cluster-RCT

USA

Laboratory details are described in appendix.

Effectiveness: ILI, defined as cough and at least 1 constitutional symptom (fever/feverishness, chills, headache, myalgia). ILI cases were given contact nurses phone numbers to record the illness and paid USD 25 to provide a throat swab. 368 participants had ILI, 94 of which had a throat swab analysed by PCR. 10 of these were positive for influenza (7 for A and 3 for B), respectively by arm 2, 5 and 3 using PCR, 7 using cell culture.

Safety: no outcomes on harms planned or reported.

Canini 2010

cluster-RCT USA The primary endpoint was the proportion of household contacts who developed an ILI during the 7 days following inclusion. Exploratory cluster-level efficacy outcome, the proportion of households with 1 or more secondary illness in household contacts.

A temperature over 37.8 °C with cough or sore throat was used as primary clinical case definition.

The authors also used a more sensitive case definition based on a temperature over 37.8 °C or at least 2 of the following: sore throat, cough, runny nose, or fatigue.

Safety: adverse reactions due to mask wearing were reported, with 38 (75%) participants in the intervention arm experiencing discomfort with mask use due to warmth (45%), respiratory difficulties (33%), and humidity (33%). Children wearing children face masks reported feeling pain more frequently than other participants wearing adult face masks (P = 0.036).

Aiello 2012

cluster-RCT in halls of residence in the USA

Clinically verified ILI - case definition (presence of cough and at least 1 or more of fever/feverishness, chills, or body aches)

Laboratory-confirmed influenza A or B. Throat swab specimens were tested for influenza A or B using real-time PCR.

Safety: no outcomes on harms planned or reported.

Barasheed 2014

cluster-RCT Saudi Arabia Laboratory: 2 nasal swabs from all ILI cases and contacts. 1 for influenza POCT using the QuickVue Influenza (A+B) assay (Quidel Corporation, San Diego, USA) and 1 for later NAT for influenza and other respiratory viruses. However, there was a problem with getting POCT on time during Hajj.

Effectiveness: to assess the effectiveness of face masks in the prevention of transmission of ILI. ILI was defined as subjective (or proven) fever plus 1 respiratory symptom (e.g. dry or productive cough, runny nose, sore throat, shortness of breath).

Safety: no outcomes on harms planned or reported.

MacIntyre 2011

cluster-RCT China Clinical respiratory illness

Influenza-like illness

Laboratory-confirmed viral respiratory infection



Laboratory-confirmed influenza A or B

- Clinical respiratory illness, defined as 2 or more respiratory or 1 respiratory symptom and a systemic symptom.
- 2. Influenza-like illness, defined as fever ≥ 38 °C plus 1 respiratory symptom (i.e. cough, runny nose, etc.)
- 3. Laboratory-confirmed viral respiratory infection (detection of adenoviruses, human metapneumovirus, coronavirus 229E/NL63, parainfluenza viruses 1, 2, and 3, influenza viruses A and B, respiratory syncytial virus A and B, rhinovirus A/B and coronavirus OC43/HKU1 by multiplex PCR).
- 4. Laboratory-confirmed influenza A or B.
- 5. Adherence with mask/respirator use.

Safety: adherence and adverse effects of mask wearing were collected at exit interviews 4 weeks' post study. Significantly higher adverse events with N95 respirator compared to medical mask for discomfort, headache, difficulty breathing, nose pressure, trouble communicating, not wearing, and unspecified "other" side effects. Over 50% of those wearing N95 respirators reported adverse events. Of those wearing medical masks versus N95 respirators, 85.5% (420/491) versus 47.4% (447/943) reported no adverse events (P < 0.001), respectively.

MacIntyre 2013 cluster-RCT China

Laboratory:

- 1. Laboratory-confirmed viral respiratory infection in symptomatic participants, defined as detection of adenoviruses; human metapneumovirus; coronaviruses 229E/NL63 and OC43/HKU1; parainfluenza viruses 1, 2, and 3; influenza viruses A and B; respiratory syncytial viruses A and B; or rhinoviruses A/B by NAT using a commercial multiplex PCR (Seegen, Inc., Seoul, Korea).
- 2. Laboratory-confirmed influenza A or B in symptomatic participants.
- 3. Laboratory-confirmed bacterial colonisation in symptomatic participants, defined as detection of *Streptococcus pneumoniae*, *Legionella*, *Bordetella pertussis*, *Chlamydia*, *Mycoplasma pneumoniae*, *or Haemophilus influenzae* type B by multiplex PCR (Seegen, Inc.).

Effectiveness: clinical respiratory illness defined as 2 or more respiratory symptoms or 1 respiratory symptom and a systemic symptom. ILI defined as fever (38 °C) plus 1 respiratory symptom.

Safety: adverse effects measured using a semi-structured questionnaire. Investigators stated that there was higher reported adverse effects and discomfort of N95 respirators compared with the other 2 arms. In terms of comfort, 52% (297 of 571) of the medical mask arm reported no problems, compared with 62% (317 of 512) of the targeted arm and 38% (217 of 574) of the N95 arm (P < 0.001).

MacIntyre 2015

cluster-RCT Vietnam

Clinical respiratory illness, influenza-like illness, and laboratory-confirmed respiratory virus infection

- 1. Clinical respiratory illness, defined as 2 or more respiratory symptoms or 1 respiratory symptom and a systemic symptom.
- 2. Influenza-like illness, defined as fever ≥ 38 °C plus 1 respiratory symptom.
- 3. Laboratory-confirmed viral respiratory infection. Laboratory confirmation was by nucleic acid detection using multiplex RT-PCR for 17 respiratory viruses.

Safety: adverse events associated with face mask use were reported in 40.4% (227/562) of HCWs in the medical/surgical mask arm and 42.6% (242/568) in the cloth mask arm (P = 0.45). The most frequently reported adverse events were: general discomfort (35.1%; 397/1130) and breathing problems (18.3%; 207/1130). The rate of ILI was higher in the cloth mask arm compared to medical/surgical masks (RR 13.25, 95% CI 1.74 to 100.97).

MacIntyre 2016 cluster-RCT China

Clinical respiratory illness, influenza-like illness, and laboratory-confirmed viral respiratory infection.

1. Clinical respiratory illness, defined as 2 or more respiratory symptoms (cough, nasal congestion, runny nose, sore throat, or sneezes) or 1 respiratory symptom and a systemic symptom (chill, lethargy, loss of appetite, abdominal pain, muscle or joint aches).



- 2. Influenza-like illness, defined as fever ≥ 38 °C plus 1 respiratory symptom.
- 3. Laboratory-confirmed viral respiratory infection, defined as detection of adenoviruses, human metapneumovirus, coronaviruses 229E/NL63 and OC43/HKU1, parainfluenza viruses 1, 2, and 3, influenza viruses A and B, respiratory syncytial virus A and B, or rhinovirus A/B by NAT using a commercial multiplex PCR.

Safety: no outcomes on harms planned or reported.

Radonovich 2019

cluster-RCT USA Laboratory. Primary outcome: incidence of laboratory-confirmed influenza, defined as:

- 1. detection of influenza A or B virus by RT-PCR in an upper respiratory specimen collected within 7 days of symptom onset;
- 2. detection of influenza from a randomly obtained swab from an asymptomatic participant; and
- 3. influenza seroconversion (symptomatic or asymptomatic), defined as at least a 4-fold rise in haemagglutination inhibition antibody titres to influenza A or B virus between pre-season and postseason serological samples deemed not attributable to vaccination.

Effectiveness. Secondary outcomes: incidence of 4 measures of viral respiratory illness or infection as follows:

- 1. acute respiratory illness with or without laboratory confirmation;
- 2. laboratory-detected respiratory infection, defined as detection of a respiratory pathogen by PCR or serological evidence of infection with a respiratory pathogen during the study surveillance period(s), which was added to the protocol prior to data analysis; and
- 3. laboratory-confirmed respiratory illness, identified as previously described (defined as self-reported acute respiratory illness plus the presence of at least PCR-confirmed viral pathogen in a specimen collected from the upper respiratory tract within 7 days of the reported symptoms and/ or at least a 4-fold rise from pre-intervention to postintervention serum antibody titres to influenza A or B virus).

Influenza-like illness, defined as temperature of at least 100 °F (37.8 °C) plus cough and/or a sore throat, with or without laboratory confirmation.

Safety: 19 participants reported skin irritation or worsening acne during years 3 and 4 at 1 site in the N95 respirator group.

Hand and hygiene (n = 35)			
Alzaher 2018	Episode of URI was defined as having 2 of the following symptoms for a day or 1 of the symptoms		
cluster-RCT	for 2 or more consecutive days: 1) a runny nose, 2) a stuffy or blocked nose or noisy breathing, 3) sneezing, 4) a cough, 5) a sore throat, and 6) feeling hot, having a fever or a chill.		
Saudi Arabia			
Arbogast 2016	ICD-9 used: 46611: acute bronchiolitis due to respiratory syncytial virus, 46619: acute bronchioli-		
cluster-RCT	tis due to other infectious organisms, 4800: pneumonia due to adenovirus, 4809: viral pneumonia, unspecified, 4870: influenza with pneumonia, 07999: unspecified viral infection, 4658: acute upper		
USA	respiratory infections of other multiple sites, 4659: acute upper respiratory infections of unspecified site, 4871: influenza with other respiratory manifestations.		
Ashraf 2020	Main outcome: 7-day prevalence of acute respiratory infection (ARI), defined as caregiver-reported		
cluster-RCT	symptoms of persistent cough or panting, wheezing, or difficulty breathing (1 or 2) in the 7 days before the interview.		
Bangladesh			
Azor-Martinez 2016	Upper respiratory illness was defined as 2 of the following symptoms during 1 day, or 1 of the		
RCT	symptoms for 2 consecutive days: (1) runny nose; (2) stuffy or blocked nose or noisy breathing; (3) cough; (4) feeling hot or feverish or having chills; (5) sore throat; or (6) sneezing.		
Spain			



Azor-Martinez 2018	Respiratory illness (RI) was defined as the presence of 2 of the following symptoms during 1 day or		
RCT	the presence of 1 of the symptoms for 2 consecutive days: (1) runny nose, (2) stuffy or blocked nose or noisy breathing, (3) cough, (4) feeling hot or feverish or having chills, (5) sore throat, or (6) sneezing. ICD-10 and ICD-9 diagnosis codes used: nonspecific upper respiratory tract infection (465.9), otitis media (382.9), pharyngotonsillitis (463), lower respiratory tract infections (485 and 486), acute bronchitis (490), and bronchiolitis (466.19). Study authors combined the bronchopneumonia code (485) and pneumonia code (486) under the label "lower respiratory tract infections." If > 1 antibiotic was prescribed during an episode, they used the first prescription for analysis. The final diagnosis was done by the medical researchers on the basis of the symptoms described above and a review of the medical history of children with RIs.		
Spain			
Biswas 2019	Influenza-like illness: an ILI episode was defined as measured fever > 38 °C or subjective fever and		
cluster-RCT	cough.		
Bangladesh	Laboratory-confirmed influenza		
Ü	Nasal swabs for real-time RT-PCR.		
Correa 2012	Acute respiratory infection was defined as 2 or more of the following symptoms for at least 24		
cluster-RCT	hours, lasting at least 2 days: runny, stuffy, or blocked nose or noisy breathing; cough; fever, hot sensation, or chills; and/or sore throat. Ear pain alone was considered ARI alternately.		
Colombia			
Cowling 2009	Laboratory-confirmed of influenza virus infection by RT-PCR for influenza A and B virus.		
cluster-RCT	Clinical influenza-like illness: used 2 clinical definitions of influenza based on self-reported data		
Hong Kong	from the symptom diaries as secondary analyses. The first definition of clinical influenza was at least 2 of the following signs and symptoms: temperature 37.8 °C or greater, cough, headache, sor throat, and myalgia; the second definition was temperature 37.8 °C or greater plus cough or sore throat.		
DiVita 2011 (conference abstract)	Influenza-like illness was defined as fever in children < 5 years old and fever with cough or sore throat in individuals > 5 years old.		
RCT			
Bangladesh			
Feldman 2016	Infectious diseases grouped into diarrhoeal, respiratory, and skin infection. Based on ICD-9, but		
cluster-RCT	no supplementary material was accessible for further definition (Supplementary Material C lists al ICD-9 diagnoses tallied in this "outcome").		
Israel			
Gwaltney 1980 RCT	Viral cultures and serology if rhinovirus in laboratory-inoculation		
USA			
Hubner 2010	Assessing illness rates due to common cold and diarrhoea. Collecting data on illness symptoms		
RCT	(common cold, sinusitis, sore throat, fever, cough, bronchitis, pneumonia, influenza, diarrhoea) and associated absenteeism at the end of every month.		
Germany	Definitions of symptoms were given to the participants as part of the individual information at the beginning of the study. Whilst most symptoms are quite self-explanatory, "influenza" and "pneumonia" are specific diagnoses that were confirmed by professional diagnosis only. Similarly, (self-) diagnosis of "fever" required objective measurement with a thermometer.		
Ladegaard 1999	Laboratory: serological evidence		
Ladegaard 1999			



п	r	т
ĸ	ι.	

Denmark

Effectiveness: influenza-like illness (described as fever, history of fever or feeling feverish in the past week, myalgia, arthralgia, sore throat, cough, sneezing, runny nose, nasal congestion, headache). However, a positive laboratory finding for influenza converts the ILI definition into one of influenza.

Larson 2010

cluster-RCT

USA

Study goals: rates of symptoms and secondary transmission of URIs, incidence of virologically confirmed influenza, knowledge of prevention and treatment strategies for influenza and URIs, and rates of influenza vaccination.

- 1. Laboratory-confirmed influenza: nasal swabs to test for influenza types A and B as well as other common respiratory viruses by rapid culture (R-Mix, Diagnostic Hybrids, Inc., Athens, OH, USA). PCR and subtyping of the samples was done during the second half of the second year of the study.
- 2. Influenza-like illness: CDC definition of ILI from the Sentinel Physicians' Network was used to determine when masks should be worn: "temperature of ≥37.8°C and cough and/or sore throat in the absence of a known cause other than influenza".
- 3. Episodes of URI = upper respiratory infection: not clear, no explicitly stated definition, reported that the most commonly reported URI symptoms are cough or rhinorrhoea.

Little 2015

RCT

England

Respiratory tract infections defined as 2 symptoms of an RTI for at least 1 day or 1 symptom for 2 consecutive days. For reported ILI, study authors did not use WHO or CDC definitions because these definitions require measured temperature, and thus were not appropriate (participants were not included after a clinical examination), and they did not use the European Centre for Disease Prevention and Control definition (1 systemic and 1 respiratory symptom) because, according to the international influenza collaboration, this definition does not necessarily differentiate ILI from a common cold. Influenzanet suggests making high temperature a separate element. Their pragmatic definition of ILI therefore required a high temperature (feeling very hot or very cold; or measured temperature > 37.5 °C), a respiratory symptom (sore throat, cough, or runny nose), and a systemic symptom (headache, severe fatigue, severe muscle aches, or severe malaise).

Luby 2005

RCT

Defined pneumonia in children according to the WHO clinical case definition: cough or difficulty breathing with a raised respiratory rate (> 60 per minute in individuals younger than 60 days old, > 50 per minute for those aged 60 to 364 days, and > 40 per minute for those aged 1 to 5 years)

Pakistan

Millar 2016 cluster-RCT

USA

Medically attended, outpatient cases of acute respiratory infection in the study population. The case definition was any occurrence of the following International Classification of Disease, 9 Revision, Clinical Modification (ICD-9) symptom or disease-specific codes: 460 to 466, 480 to 488, and specifically 465.9, 482.9, 486, and 487.1.

Acute respiratory infections (460 to 466)

460 Acute nasopharyngitis (common cold)

461 Acute sinusitis

462 Acute pharyngitis

463 Acute tonsillitis

464 Acute laryngitis and tracheitis

465 Acute upper respiratory infections of multiple or unspecified sites

466 Acute bronchitis and bronchiolitis

Pneumonia and influenza (480 to 488)

480 Viral pneumonia

481 Pneumococcal pneumonia (Streptococcus pneumoniae pneumonia)

482 Other bacterial pneumonia



Table 9. Trial authors' outco	ome definitions (Continued) 483 Pneumonia due to other specified organism
	484 Pneumonia in infectious diseases classified elsewhere
	485 Bronchopneumonia, organism unspecified
	486 Pneumonia, organism unspecified
	487 Influenza
	488 Influenza due to identified avian influenza virus
	465.9 Acute upper respiratory infections of unspecified site
	482.9 Bacterial pneumonia NOS
	487.1 Diagnosis of influenza with other respiratory manifestations
Morton 2004	Respiratory illnesses defined by symptoms of upper respiratory infections such as nasal conges-
cluster-RCT	tion, cough, or sore throat, in any combination, with or without fever
Cross-over study	
USA	
Nicholson 2014	Acute respiratory infections
cluster-RCT	Operational definitions for all the illnesses were taken from Black's Medical Dictionary. ARIs de-
India	fined as "Pneumonia, cough, fever, chest pain and shortness of breath, cold, inflammation of any or all of the airways, that is, nose, sinuses, throat, larynx, trachea and bronchi".
Pandejpong 2012	Influenza-like illness defined if 2 or more symptoms of stuffy nose, cough, fever or chills, sore
cluster-RCT	throat, headache, diarrhoea, presence of hand, foot, or mouth ulcers.
Thailand	
Priest 2014	Respiratory illness was defined as an episode of illness that included at least 2 of the following
cluster-RCT	caregiver-reported symptoms for 1 day, or 1 of these symptoms for 2 days (but not fever alone): runny nose, stuffy or blocked nose or noisy breathing, cough, fever, sore throat, or sneezing.
New Zealand	
Ram 2015	Influenza-like illness
RCT	Age-specific definitions of ILI. For individuals ≥ 5 years old, ILI was defined as history of fever with
Bangladesh	cough or sore throat. For children < 5 years old, ILI was defined as fever; study authors used this relatively liberal case definition in order to include influenza cases with atypical presentations in children.
	Laboratory-confirmed influenza infection
	Oropharyngeal swabs from index case patients for laboratory testing for influenza. All swabs were tested by PCR for influenza A and B, with further subtyping of influenza A isolates.
Roberts 2000	The symptoms of acute upper respiratory illness elicited from parents were: a runny nose, a
cluster-RCT	blocked nose, and cough. Study authors used a definition of colds based on a community intervention trial of virucidal impregnated tissues.
Australia	A cold was defined as either 2 symptoms for 1 day or 1 of the respiratory symptoms for at least 2 consecutive days, but not including 2 consecutive days of cough alone. Study authors defined a



Table 9. Trial authors' ou	new episode of a cold as the occurrence of respiratory symptoms after a period of 3 symptom-free days.		
Sandora 2005 cluster-RCT	The overall rates of secondary respiratory and GI illness.		
USA	Respiratory illness was defined as 2 of the following symptoms for 1 day or 1 of the symptoms for 2 consecutive days: (1) runny nose; (2) stuffy or blocked nose or noisy breathing; (3) cough; (4) fever, feels hot, or has chills; (5) sore throat; and (6) sneezing. An illness was considered new or separate when a period of at least 2 symptom-free days had elapsed since the previous illness. An illness was defined as a secondary case when it began 2 to 7 days after the onset of the same illness type (respiratory or GI) in another household member.		
Savolainen-Kopra 2012	Nasal and pharyngeal stick samples from participants with respiratory symptoms		
cluster-RCT			
Finland			
Simmerman 2011	Influenza-like illness defined by WHO as fever plus cough or sore throat, based on self-reported		
cluster-RCT	symptoms.		
Thailand	Laboratory-confirmed secondary influenza virus infections amongst household members described as the secondary attack rate. The secondary influenza virus infection was defined as a positive rRT-PCR result on days 3 or 7 or a four-fold rise in influenza HI antibody titres with the virus type and subtype matching the index case.		
Stebbins 2011 cluster-RCT	The primary outcome was an absence episode associated with an influenza-like illness that was subsequently laboratory-confirmed as influenza A or B. The following CDC definition for ILI was		
USA	used: fever ≥ 38 °C with sore throat or cough.		
Swarthout 2020	The primary outcome in this study is ARI symptoms - defined as having caregiver-reported cough or		
cluster-RCT	difficulty breathing, including panting or wheezing, within 7 days before the interview - in children younger than 3 years. Prespecified secondary outcomes in this study include difficulty		
Kenya	breathing, including panting or wheezing, in the past 7 days (a more specific indicator of respiratory infection than a cough alone); ARI symptoms presenting with fever in the past 7 days (a potentially more severe infection); and enumerator-observed runny nose (an objective outcome).		
Talaat 2011	Nasal swab for QuickVue test for influenza A and B viruses.		
cluster-RCT	Influenza-like illness (defined as fever > 38 °C and either cough or sore throat).		
Egypt			
Teesing 2021	Incidence of gastroenteritis, ILI, assumed pneumonia, UTIs using the McGeer criteria, and infec-		
cluster-RCT	tions caused by MRSA.		
The Netherlands			
Temime 2018	ARIs were defined as the combination of at least 1 respiratory symptom and 1 symptom of systemic		
cluster-RCT	infection.		
France			
Turner 2004b	Virologic assays		
RCT			
Canada			



Tahle 9	Trial author	s' outcome definitions	(Continued)
Iable 3.	iiialauliivi	3 Outcome deminitions	(Continuea)

Turner 2012	Laboratory-confirmed rhinovirus infection by PCR assay.		
RCT	Common cold illness was defined as the presence of any of the symptoms of nasal obstruction,		
USA	rhinorrhoea, sore throat, or cough on at least 3 consecutive days. Illnesses separated by at least 3 symptom-free days were considered as separate illnesses.		
Yeung 2011	Pneumonia		
cluster-RCT			
Hong Kong			
Zomer 2015 cluster-RCT	Incidence of gastrointestinal and respiratory infections in children monitored by parents. The common cold was defined as a blocked or runny nose with at least 1 of the following symptoms: coughing speezing fever sore throat or earache		
Netherlands	ing, sneezing, fever, sore throat, or earache.		
Hand hygiene and masks (n =	6)		
Aelami 2015 (conference abstract)	Influenza-like illness was defined as the presence of at least 2 of the following during their stay: fever, cough, and sore throat.		
RCT	Safety: no outcomes on harms planned or reported.		
Saudi Arabia			
Aiello 2010	Influenza-like illness case definition (presence of cough and at least 1 constitutional symptom		
cluster-RCT	(fever/feverishness, chills, or body aches).		
USA	Safety: no outcomes on harms planned or reported.		
Cowling 2009	2 clinical definitions of influenza. First definition was at least 2 of the following signs and symp-		
cluster-RCT	toms: temperature 37.8 °C or greater, cough, headache, sore throat, and myalgia. The second was temperature 37.8 °C or greater plus cough or sore throat.		
Hong Kong	Safety: no outcomes on harms planned or reported.		
Larson 2010	Study goals: rates of symptoms and secondary transmission of URIs, incidence of virologically-con-		
cluster-RCT	firmed influenza, knowledge of prevention and treatment strategies for influenza and URIs, and rates of influenza vaccination.		
USA	 Laboratory-confirmed influenza: nasal swabs to test for influenza types A and B as well as other common respiratory viruses by rapid culture (R-Mix, Diagnostic Hybrids, Inc., Athens, OH, USA). PCR and subtyping of the samples was done during the second half of the second year of the study. 		
	2. Influenza-like illness: CDC definition of ILI from the Sentinel Physicians' Network was used to determine when masks should be worn: "temperature of ≥37.8°C and cough and/or sore throat in the absence of a known cause other than influenza".		
	3. Episodes of URI = upper respiratory infection: not clear, no explicitly stated definition, reported that the most commonly reported URI symptoms are cough or rhinorrhoea.		
	Safety: no outcomes on harms planned or reported.		
Simmerman 2011	Laboratory-confirmed secondary influenza virus infections amongst household members de-		
cluster-RCT	scribed as the secondary attack rate. The secondary influenza virus infection was defined as a positive rRT-PCR result on days 3 or 7 or a four-fold rise in influenza HI antibody titres with the virus		
Thailand	type and subtype matching the index case.		
	Influenza-like illness defined by WHO as fever plus cough or sore throat, based on self-reported symptoms.		



Table 9.	Trial authors'	outcome de	finitions (Continued)
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Safety: no outcomes on harms planned or reported.

Suess 2012 Quantitative RT-PCR for samples of nasal wash.

cluster-RCT Influenza virus infection as a laboratory-confirmed influenza infection in a household member who developed fever (> 38.0 °C), cough, or sore throat during the observation period. Also secondary outcome measure of the occurrence of ILI as defined by WHO as fever plus cough or sore throat.

Safety: the study reported that the majority of participants (107/172, 62%) did not report any problems with mask wearing. This proportion was significantly higher in the group of adults (71/100, 71%) compared to the group of children (36/72, 50%) (P = 0.005). The main problem stated by participants (adults and children) was "heat/humidity" (18/34, 53% of children; 10/29, 35% of adults) (P = 0.1), followed by "pain" and "shortness of breath" when wearing a face mask.

Surface/object disinfection (with or without hand hygiene)(n = 8)

Ban 2015	Acute respiratory illness classified as the appearance of 2 or more of the following symptoms: fever, cough and expectoration, runny nose and nasal congestion.	
cluster-RCT		
China		
Carabin 1999	The presence of nasal discharge (runny nose) accompanied by 1 or several of the following symp-	
cluster-RCT	toms: fever, sneezing, cough, sore throat, ear pain, malaise, irritability. A URTI was defined as a cold for 2 consecutive days.	
Canada		
Chard 2019	Pupils were considered to have symptoms of respiratory infection if they reported cough, runny	
cluster-RCT	nose, stuffy nose, or sore throat.	
Laos		
Ibfelt 2015	Laboratory confirmation of 16 respiratory viruses: influenza A; influenza B; coronavirus NL63229E,	
cluster-RCT	OC43 and HKU1; parainfluenza virus 1, 2, 3, and 4; rhinovirus; RSV A/B; adenovirus; enterovirus; parechovirus; and bocavirus using quantitative PCR	
Denmark		
Kotch 1994	Respiratory symptoms include coughing, runny nose, wheezing or rattling in the chest, sore throat,	
RCT	or earache.	
USA		
McConeghy 2017	Classified infections as lower respiratory tract infections (i.e. pneumonia, bronchitis, or chronic	
RCT	structive pulmonary disease exacerbation) or other.	
USA		
Sandora 2008	RI was defined as an acute illness that included > 1 of the following symptoms: runny nose, stuffy or	
cluster-RCT	blocked nose, cough, fever or chills, sore throat, or sneezing.	
USA		
White 2001	RI was defined as: cough, sneezing, sinus trouble, bronchitis, fever alone, pink-eye, headache,	
DB-RCT	mononucleosis, and acute exacerbation of asthma.	
USA		



Other (miscellaneous) interventions (n = 5)

other (miscetaineous) meer ventions (ii = 5)			
Fretheim 2022a	Respiratory infection was defined as having 1 respiratory symptom (stuffed or runny nose, sore		
pragmatic RCT Norway	throat, cough, sneezing, heavy breathing) and fever, or 1 respiratory symptom and at least 2 more symptoms (body ache, muscular pain, fatigue, reduced appetite, stomach pain, headache, loss of smell.		
Hartinger 2016	ARI was defined as a child presenting cough or difficulty breathing, or both. ALRI was defined as a		
cluster-RCT	child presenting cough or difficulty breathing, with a raised respiratory rate > 50 per minute in children aged 6 to 11 months and > 40 per minute in children aged > 12 months on 2 consecutive mea-		
Peru	surements. An episode was defined as beginning on the first day of cough or difficulty breathing and ending with the last day of the same combination, followed by at least 7 days without those symptoms.		
Huda 2012	Study authors classified acute respiratory illness as having cough and fever or difficulty breathing		
cluster-RCT	and fever within 48 h prior to interview.		
Bangladesh			
Najnin 2019	Classified participants as having respiratory illness if they reported having fever plus either cou		
cluster-RCT	or nasal congestion or fever plus breathing difficult.		
Bangladesh			
Satomura 2005	Upper respiratory tract infection defined as all of the following conditions:		
RCT	1. both nasal and pharyngeal symptoms;		
Japan	2. severity of at least 1 symptom increased by 2 grades or more; and		
oupun	worsening of a symptom of 1 increment or more for > 3 days.		
	Because of the difference in the mode of transmission, study authors excluded influenza-like diseases featured by moderate or severe fever; anti-influenza vaccination in the preseason and arthralgia, and treated them separately. The incidence was determined by 1 study physician who was blinded to group assignment.		
Virucidal tissues (n = 2)			
Farr 1988a	RI defined as: occurrence of at least 2 respiratory symptoms on the same day or the occurrence of a		
cluster-RCT	single respiratory symptom on 2 consecutive days (except for sneezing). The respiratory symptoms were as follows: sneezing, nasal congestion, nasal discharge, sore throat, scratchy throat, hoarse-		
USA trial 1 and trial 2	ness, coughing, malaise, headache, feverishness, chilliness and myalgia.		
Longini 1988	Respiratory illness defined as 1 or more of the following symptoms occurring during the course of acute episode: coryza, sore throat or hoarseness, earache, cough, pain on respiration, wheezy breathing or phlegm from the chest.		
DB-PC RCT			
USA			

ALRI: acute lower respiratory infection ARIs: acute respiratory infections

CDC: Centers for Disease Control and Prevention

CI: confidence interval

cluster-RCT: cluster-randomised controlled trial

CRI: clinical respiratory illness

DB-PC: double-blind, placebo-controlled

DB-RCT: double-blind randomised controlled trial

DNA: deoxyribonucleic acid



ELISA: enzyme-linked immunosorbent assay

GI: gastrointestinal

h: hours

HCW: healthcare workers HI: haemagglutinin

hMPV: human metapneumo virus

ICD-9: International Classification of Disease, 9th Revision, Clinical Modification ICD-10: International Classification of Disease, 10th Revision, Clinical Modification

IgG: immunoglobulin G IgM: immunoglobulin M ILI: influenza-like illness

min: minutes

MRSA: methicillin-resistant Staphylococcus aureus

NAT: nucleic acid testing NOS: not otherwise specified NTS: nasal and throat swab PCR: polymerase chain reaction

PIV: parainfluenza virus POCT: point-of-care testing RCT: randomised controlled trial

RI: respiratory infection RNA: ribonucleic acid

RR: risk ratio

rRT-PCR: real-time reverse transcriptase polymerase chain reaction

RSV: respiratory syncytial virus RTI: respiratory tract infection

RT-PCR: reverse transcriptase polymerase chain reaction

SAR: secondary attack ratios
SD: standard deviation
S/S: signs and symptoms
URI: upper respiratory infection
URTI: upper respiratory tract infection

UTI: urinary tract infection WHO: World Health Organization

APPENDICES

Appendix 1. Cochrane Central Register of Controlled Trials (CENTRAL) search string

([mh "Influenza, Human"] OR [mh "Influenzavirus A"] OR [mh "Influenzavirus B"] OR [mh "Influenzavirus C"] OR Influenza:ti,ab OR [mh "Respiratory Tract Diseases"] OR Influenzas:ti,ab OR "Influenza-like":ti,ab OR ILI:ti,ab OR Flu:ti,ab OR Flu:ti,ab OR [mh ^"Common Cold"] OR "common cold":ti,ab OR colds:ti,ab OR coryza:ti,ab OR [mh coronavirus] OR [mh "sars virus"] OR coronavirus:ti,ab OR Coronavirus:ti,ab OR [mh "coronavirus infections"] OR [mh "severe acute respiratory syndrome"] OR "severe acute respiratory syndrome":ti,ab OR "severe acute respiratory syndromes":ti,ab OR [mh "respiratory syncytial viruses"] OR [mh "respiratory syncytial virus, human"] OR [mh "Respiratory Syncytial Virus Infections"] OR "respiratory syncytial virus":ti,ab OR "respiratory syncytial viruses":ti,ab OR roughing OR Sneezing)) OR ((respiratory:ti,ab AND Tract) AND (infection:ti,ab OR Infections:ti,ab OR illness:ti,ab)))

([mh "Hand Hygiene"] OR handwashing:ti,ab OR "hand-washing":ti,ab OR ((Hand:ti,ab OR Alcohol:ti,ab) AND (wash:ti,ab OR Washing:ti,ab OR Cleansing:ti,ab OR Rinses:ti,ab OR hygiene:ti,ab OR rub:ti,ab OR Rubbing:ti,ab OR sanitizer:ti,ab OR sanitizer:ti,ab OR cleanser:ti,ab OR disinfected:ti,ab OR Disinfectant:ti,ab OR Disinfect:ti,ab OR antiseptic:ti,ab OR virucid:ti,ab) OR [mh "gloves, protective"] OR Glove:ti,ab OR Gloves:ti,ab OR [mh Masks] OR [mh "respiratory protective devices"] OR facemask:ti,ab OR Facemasks:ti,ab OR mask:ti,ab OR Masks:ti,ab OR respirator:ti,ab OR respirators:ti,ab OR [mh ^"Protective Clothing"] OR [mh "Protective Devices"] OR "patient isolation":ti,ab OR ((school:ti,ab OR Schools:ti,ab) AND (Closure:ti,ab OR Closures:ti,ab OR Closed:ti,ab)) OR [mh Quarantine] OR quarantine:ti,ab OR "Hygiene intervention":ti,ab OR [mh Mouthwashes] OR gargling:ti,ab OR "nasal tissues":ti,ab OR [mh "Eye Protective Devices"] OR Glasses:ti,ab OR Goggle:ti,ab OR "Eye protection":ti,ab OR "Face shield:ti,ab OR Faceshields:ti,ab OR Visors:ti,ab)

AND



([mh "Communicable Disease Control"] OR [mh "Disease Outbreaks"] OR [mh "Disease Transmission, Infectious"] OR [mh "Infection Control"] OR "Communicable Disease Control":ti,ab OR "Secondary transmission":ti,ab OR ((Reduced:ti,ab OR Reduce:ti,ab OR Reduce:ti,ab OR Reduce:ti,ab OR Cocurrence:ti,ab OR Transmission:ti,ab OR Secondary:ti,ab))

Appendix 2. PubMed search string

("Influenza, Human" [Mesh] OR "Influenzavirus A" [Mesh] OR "Influenzavirus B" [Mesh] OR "Influenzavirus C" [Mesh] OR Influenza [tiab] OR "Respiratory Tract Diseases" [Mesh] OR "Bacterial Infections/transmission" [Mesh] OR Influenzas [tiab] OR "Influenza-like" [tiab] OR ILI [tiab] OR Flu [tiab] OR Flu [tiab] OR "Common Cold" [Mesh: No Exp] OR "common cold" [tiab] OR colds [tiab] OR coryza [tiab] OR coronavirus [Mesh] OR "savere acute respiratory syndrome" [Mesh] OR "severe acute respiratory syndrome" [Mesh] OR "severe acute respiratory syndrome" [tiab] OR "respiratory syndromes" [tiab] OR "respiratory syncytial virus [tiab] OR "Respiratory [tiab] OR [ti

AND

("Hand Hygiene"[Mesh] OR handwashing[tiab] OR hand-washing[tiab] OR ((Hand[tiab] OR Alcohol[tiab]) AND (wash[tiab] OR Washing[tiab] OR Cleansing[tiab] OR Rinses[tiab] OR hygiene[tiab] OR rub[tiab] OR Rubbing[tiab] OR sanitizer[tiab] OR sanitizer[tiab] OR cleanser[tiab] OR disinfected[tiab] OR Disinfectant[tiab] OR Disinfect[tiab] OR antiseptic[tiab] OR virucid[tiab])) OR "gloves, protective"[Mesh] OR Glove[tiab] OR Gloves[tiab] OR Masks[Mesh] OR "respiratory protective devices"[Mesh] OR facemask[tiab] OR Facemasks[tiab] OR mask[tiab] OR masks[tiab] OR masks[tiab] OR "Protective Clothing"[Mesh:NoExp] OR "Protective Devices"[Mesh] OR "patient isolation"[tiab] OR ((school[tiab] OR Schools[tiab]) AND (Closure[tiab] OR Closures[tiab] OR Closures[tiab] OR "Mouthwashes"[Mesh] OR gargling[tiab] OR "nasal tissues"[tiab] OR "Eye Protective Devices"[Mesh] OR Glasses[tiab] OR Goggles[tiab] OR Goggles[tiab] OR Faceshield[tiab] OR "Face shields"[tiab] OR Visors[tiab])

("Communicable Disease Control"[Mesh] OR "Disease Outbreaks"[Mesh] OR "Disease Transmission, Infectious"[Mesh] OR "Infection Control"[Mesh] OR Transmission[sh] OR "Prevention and control"[sh] OR "Communicable Disease Control"[tiab] OR "Secondary transmission"[tiab] OR ((Reduced[tiab] OR Reduce[tiab] OR Reduction[tiab] OR Reducing[tiab] OR Lower[tiab]) AND (Incidence[tiab] OR Occurrence[tiab] OR Transmission[tiab] OR Secondary[tiab])))

ΔΝΝ

(Randomized controlled trial[pt] OR controlled clinical trial[pt] OR randomized[tiab] OR randomised[tiab] OR placebo[tiab] OR "drug therapy"[sh] OR randomly[tiab] OR trial[tiab] OR groups[tiab])

NOT

(Animals[Mesh] not (Animals[Mesh] and Humans[Mesh]))

NOT

("Case Reports"[pt] OR Editorial[pt] OR Letter[pt] OR Meta-Analysis[pt] OR "Observational Study"[pt] OR "Systematic Review"[pt] OR "Case Report"[ti] OR "Case series"[ti] OR Meta-Analysis[ti] OR "Meta Analysis"[ti] OR "Systematic Review"[ti])

Appendix 3. Embase (Elsevier) search string

('influenza'/exp OR Influenza:ti,ab OR 'Respiratory Tract Disease'/exp OR Influenzas:ti,ab OR Influenza-like:ti,ab OR ILI:ti,ab OR Flus:ti,ab OR 'Common Cold'/de OR "common cold":ti,ab OR colds:ti,ab OR coryza:ti,ab OR 'coronavirus'/exp OR 'SARS coronavirus'/exp OR coronavirus:ti,ab OR Coronaviruses:ti,ab OR 'coronavirus infection'/exp OR 'severe acute respiratory syndrome'/exp OR "severe acute respiratory syndrome":ti,ab OR "severe acute respiratory syndromes":ti,ab OR 'Pneumovirus'/exp OR 'Human respiratory syncytial virus'/exp OR "respiratory syncytial viruses":ti,ab OR rsv:ti,ab OR parainfluenza:ti,ab OR "Respiratory illness":ti,ab OR ((Transmission) AND (Coughing OR Sneezing)) OR ((respiratory:ti,ab AND Tract) AND (infection:ti,ab OR Infections:ti,ab OR illness:ti,ab)))

AND

('hand washing'/exp OR handwashing:ti,ab OR hand-washing:ti,ab OR ((Hand:ti,ab OR Alcohol:ti,ab) AND (wash:ti,ab OR Washing:ti,ab OR Cleansing:ti,ab OR Rinses:ti,ab OR hygiene:ti,ab OR rub:ti,ab OR Rubbing:ti,ab OR sanitizer:ti,ab OR sanitizer:ti,ab OR cleanser:ti,ab OR disinfected:ti,ab OR Disinfect:ti,ab OR antiseptic:ti,ab OR virucid:ti,ab)) OR 'protective glove'/exp OR Glove:ti,ab OR Gloves:ti,ab OR 'mask'/exp OR 'gas mask'/exp OR facemask:ti,ab OR Facemasks:ti,ab OR mask:ti,ab OR Masks:ti,ab OR respirator:ti,ab OR respirators:ti,ab OR 'protective clothing'/de OR 'protective equipment'/exp OR "patient isolation":ti,ab OR ((school:ti,ab OR Schools:ti,ab) AND (Closure:ti,ab OR Closure:ti,ab OR Closed:ti,ab)) OR 'Quarantine'/exp OR quarantine:ti,ab OR "Hygiene intervention":ti,ab OR 'mouthwash'/exp OR gargling:ti,ab OR "nasal tissues":ti,ab OR 'eye protective device'/exp OR Glasses:ti,ab OR Goggle:ti,ab OR "Eye protection":ti,ab OR Faceshield:ti,ab OR Faceshields:ti,ab OR Goggles:ti,ab OR "Face shield":ti,ab OR "Face shields":ti,ab OR Visors:ti,ab)

('Communicable Disease Control'/exp OR 'epidemic'/exp OR 'disease transmission'/exp OR 'Infection Control'/exp OR "Communicable Disease Control":ti,ab OR "Secondary transmission":ti,ab OR ((Reduced:ti,ab OR Reduce:ti,ab OR Reduction:ti,ab OR Reducing:ti,ab OR Lower:ti,ab) AND (Incidence:ti,ab OR Occurrence:ti,ab OR Transmission:ti,ab OR Secondary:ti,ab)))

AND



(random* OR factorial OR crossover OR placebo OR blind OR blinded OR assign OR assigned OR allocate OR allocated OR 'crossover procedure'/exp OR 'double-blind procedure'/exp OR 'randomized controlled trial'/exp OR 'single-blind procedure'/exp NOT ('animal'/exp NOT ('animal'/exp AND 'human'/exp)))

Appendix 4. CINAHL (EBSCO) search string

((MH "Influenza, Human+") OR (MH "Orthomyxoviridae+") OR TI Influenza OR AB Influenza OR (MH "Respiratory Tract Diseases+") OR TI Influenzas OR AB III OR AB II

((MH "Handwashing+") OR TI handwashing OR AB handwashing OR TI hand-washing OR AB hand-washing OR ((TI Hand OR AB Hand OR TI Alcohol OR AB Alcohol) AND (TI wash OR AB wash OR TI Washing OR AB Washing OR TI Cleansing OR AB Cleansing OR TI Rinses OR AB Rinses OR TI hygiene OR AB hygiene OR TI rub OR AB rub OR TI Rubbing OR AB Rubbing OR TI sanitizer OR AB sanitiser OR TI sanitizer OR AB sanitiser OR TI cleanser OR AB cleanser OR TI disinfected OR AB disinfected OR TI Disinfectant OR AB Disinfectant OR TI Disinfect OR AB Disinfect OR TI antiseptic OR AB antiseptic OR TI virucid OR AB virucid)) OR (MH "gloves+") OR TI Glove OR AB Glove OR Gloves OR (MH "Masks+") OR (MH "respiratory protective devices+") OR TI facemask OR AB facemask OR TI Facemasks OR AB Facemasks OR TI mask OR AB mask OR TI Masks OR AB Masks OR TI respirator OR AB respirators OR (MH "Protective Clothing") OR (MH "Protective Devices+") OR TI "patient isolation" OR AB "patient isolation" OR ((TI school OR AB school OR TI Schools OR AB Schools) AND (TI Closure OR AB Closure OR TI Closures OR AB Closures OR TI Closed OR AB Closed)) OR (MH "Quarantine+") OR TI quarantine OR AB quarantine OR TI "Hygiene intervention" OR AB "Hygiene intervention" OR (MH "Mouthwashes+") OR TI gargling OR AB gargling OR TI "nasal tissues" OR AB "nasal tissues" OR (MH "Eye Protective Devices+") OR TI Glasses OR AB Glasses OR TI Goggle OR AB Goggle OR TI "Eye protection" OR AB "Face shield" OR AB "Face shield" OR AB "Face shields" OR AB "Face shields" OR AB Visors)

AND

((MH "Infection Control+") OR (MH "Disease Outbreaks+") OR (MH "Infection Control+") OR TI "Communicable Disease Control" OR AB "Communicable Disease Control" OR TI "Secondary transmission" OR AB "Secondary transmission" OR ((TI Reduced OR AB Reduced OR TI Reduce OR AB Reduce OR TI Reducion OR AB Reduction OR TI Reducing OR AB Reducing OR TI Lower OR AB Lower) AND (TI Incidence OR AB Incidence OR TI Occurrence OR AB Occurrence OR TI Transmission OR AB Transmission OR TI Secondary OR AB Secondary)))

AND

((MH "Clinical Trials+") OR (MH "Quantitative Studies") OR TI placebo* OR AB placebo* OR (MH "Placebos") OR (MH "Random Assignment") OR TI random* OR AB random* OR TI ((singl* or doubl* or tripl* or trebl*) W1 (blind* or mask*)) OR AB ((singl* or doubl* or tripl* or trebl*) W1 (blind* or mask*)) OR TI clinic* trial* OR AB clinic* trial* OR PT clinical trial)

Appendix 5. Previous search strategies (pre-2010)

Details of the 2010 update and the search strategy used in the original review and the 2009 search strategy updates for MEDLINE, CENTRAL, EMBASE and CINAHL

In the 2010 update we searched, as we have done previously, the Cochrane Central Register of Controlled Trials (CENTRAL) 2010, Issue 3, which includes the Acute Respiratory Infections Group's Specialised Register, MEDLINE (April 2009 to October week 2, 2010), EMBASE (April 2009 to October 2010) and CINAHL (January 2009 to October 2010). Details of previous searches are in Appendix 1. In addition, to include more of the literature of low-income countries in this update, we ran searches in LILACS (2008 to October 2010), Indian MEDLARS (2008 to October 2010) and IMSEAR (2008 to October 2010).

We used the following search strategy (updated to include new and emerging respiratory viruses) to search MEDLINE and CENTRAL. We combined the MEDLINE search strategy with the Cochrane Highly Sensitive Search Strategy for identifying randomised trials in MEDLINE: sensitivity- and precision-maximising version (2008 revision) (Ovid format) (Lefebvre 2011). We also included an additional search strategy based on the work of Fraser, Murray and Burr (Fraser 2006) to identify observational studies.

1 Influenza, Human/

2 exp Influenzavirus A/

3 exp Influenzavirus B/

4 Influenzavirus C/

5 (influenza* or flu).tw.

6 Common Cold/

7 common cold*.tw.



- 8 Rhinovirus/
- 9 rhinovir*.tw.
- 10 adenoviridae/ or mastadenovirus/ or adenoviruses, human/
- 11 adenoviridae infections/ or adenovirus infections, human/
- 12 adenovir*.tw.
- 13 coronavirus/ or coronavirus 229e, human/ or coronavirus oc43, human/ or infectious bronchitis virus/ or sars virus/
- 14 coronavir*.tw.
- 15 coronavirus infections/ or severe acute respiratory syndrome/
- 16 (severe acute respiratory syndrome* or sars).tw.
- 17 respiratory syncytial viruses/ or respiratory syncytial virus, human/
- 18 Respiratory Syncytial Virus Infections/
- 19 (respiratory syncytial virus* or rsv).tw.
- 20 Pneumovirus Infections/
- 21 parainfluenza virus 1, human/ or parainfluenza virus 3, human/
- 22 parainfluenza virus 2, human/ or parainfluenza virus 4, human/
- 23 (parainfluenza* or para-influenza* or para influenza).tw.
- 24 enterovirus a, human/ or exp enterovirus b, human/ or enterovirus c, human/ or enterovirus d, human/
- 25 Enterovirus Infections/
- 26 enterovir*.tw.
- 27 Human bocavirus/
- 28 bocavirus*.tw.
- 29 Metapneumovirus/
- 30 metapneumovir*.tw.
- 31 Parvovirus B19, Human/
- 32 parvoviridae infections/ or erythema infectiosum/
- 33 parvovirus*.tw.
- 34 Parechovirus/
- 35 parechovirus*.tw.
- 36 acute respiratory tract infection*.tw.
- 37 acute respiratory infection*.tw.
- 38 or/1-37
- 39 Handwashing/
- 40 (handwashing or hand washing or hand-washing).tw.
- 41 hand hygiene.tw.
- 42 (sanitizer* or sanitiser*).tw.
- 43 (cleanser* or disinfectant*).tw.
- 44 gloves, protective/ or gloves, surgical/
- 45 glov*.tw.
- 46 masks/ or respiratory protective devices/
- 47 (mask or masks or respirator or respirators).tw.
- 48 Protective Clothing/
- 49 Protective Devices/
- 50 Patient Isolators/
- 51 Patient Isolation/
- 52 patient isolat*.tw.
- 53 (barrier* or curtain* or partition*).tw.
- 54 negative pressure room*.tw.
- 55 ((reverse barrier or reverse-barrier) adj3 (nurs* or unit or isolation)).tw.
- 56 Cross Infection/pc [Prevention & Control]
- 57 (cross infection* adj2 prevent*).tw.
- 58 Communicable Disease Control/
- 59 Infection Control/
- 60 (school* adj3 (clos* or dismissal*)).tw.
- 61 temporary closur*.tw.
- 62 mass gathering*.tw.
- 63 (public adj2 (gathering* or event*)).tw.
- 64 (bans or banning or banned or ban).tw.
- 65 (outbreak adj3 control*).tw.
- 66 distancing*.tw.
- 67 Quarantine/
- 68 quarantine*.tw.
- 69 (protective adj2 (cloth* or garment* or device* or equipment)).tw.



- 70 ((protective or preventive) adj2 (procedure* or behaviour* or behavior*)).tw.
- 71 personal protect*.tw.
- 72 (isolation room* or isolation strateg*).tw.
- 73 (distance adj2 patient*).tw.
- 74 ((spatial or patient) adj separation).tw.
- 75 cohorting.tw.
- 76 or/39-75
- 77 38 and 76
- 78 (animals not (animals and humans)).sh.
- 79 77 not 78

Ovid MEDLINE

- 1 Influenza, Human/
- 2 exp Influenzavirus A/
- 3 exp Influenzavirus B/
- 4 Influenzavirus C/
- 5 (influenza* or flu).tw.
- 6 Common Cold/
- 7 common cold*.tw.
- 8 Rhinovirus/
- 9 rhinovir*.tw.
- 10 adenoviridae/ or mastadenovirus/ or adenoviruses, human/
- 11 adenoviridae infections/ or adenovirus infections, human/
- 12 adenovir*.tw.
- 13 coronavirus/ or coronavirus 229e, human/ or coronavirus oc43, human/ or infectious bronchitis virus/ or sars virus/
- 14 coronavir*.tw.
- 15 coronavirus infections/ or severe acute respiratory syndrome/
- 16 (severe acute respiratory syndrome* or sars).tw.
- 17 respiratory syncytial viruses/ or respiratory syncytial virus, human/
- 18 Respiratory Syncytial Virus Infections/
- 19 (respiratory syncytial virus* or rsv).tw.
- 20 Pneumovirus Infections/
- 21 parainfluenza virus 1, human/ or parainfluenza virus 3, human/
- 22 parainfluenza virus 2, human/ or parainfluenza virus 4, human/
- 23 (parainfluenza* or para-influenza* or para influenza).tw.
- 24 enterovirus a, human/ or exp enterovirus b, human/ or enterovirus c, human/ or enterovirus d, human/
- 25 Enterovirus Infections/
- 26 enterovir*.tw.
- 27 Human bocavirus/
- 28 bocavirus*.tw.
- 29 Metapneumovirus/
- 30 metapneumovir*.tw.
- 31 Parvovirus B19, Human/
- 32 parvoviridae infections/ or erythema infectiosum/
- 33 parvovirus*.tw.
- 34 Parechovirus/
- 35 parechovirus*.tw.
- 36 acute respiratory tract infection*.tw.
- 37 acute respiratory infection*.tw.
- 38 or/1-37
- 39 Handwashing/
- 40 (handwashing or hand washing or hand-washing).tw.
- 41 hand hygiene.tw.
- 42 (sanitizer* or sanitiser*).tw.
- 43 (cleanser* or disinfectant*).tw.
- 44 gloves, protective/ or gloves, surgical/
- 45 glov*.tw.
- 46 masks/ or respiratory protective devices/
- 47 (mask or masks or respirator or respirators).tw.
- 48 Protective Clothing/
- 49 Protective Devices/



- 50 Patient Isolators/
- 51 Patient Isolation/
- 52 patient isolat*.tw.
- 53 (barrier* or curtain* or partition*).tw.
- 54 negative pressure room*.tw.
- 55 ((reverse barrier or reverse-barrier) adj3 (nurs* or unit or isolation)).tw.
- 56 Cross Infection/pc [Prevention & Control]
- 57 (cross infection* adj2 prevent*).tw.
- 58 Communicable Disease Control/
- 59 Infection Control/
- 60 (school* adj3 (clos* or dismissal*)).tw.
- 61 temporary closur*.tw.
- 62 mass gathering*.tw.
- 63 (public adj2 (gathering* or event*)).tw.
- 64 (bans or banning or banned or ban).tw.
- 65 (outbreak adj3 control*).tw.
- 66 distancing*.tw.
- 67 Quarantine/
- 68 quarantine*.tw.
- 69 (protective adj2 (cloth* or garment* or device* or equipment)).tw.
- 70 ((protective or preventive) adj2 (procedure* or behaviour* or behavior*)).tw.
- 71 personal protect*.tw.
- 72 (isolation room* or isolation strateg*).tw.
- 73 (distance adj2 patient*).tw.
- 74 ((spatial or patient) adj separation).tw.
- 75 cohorting.tw.
- 76 or/39-75
- 77 38 and 76
- 78 (animals not (animals and humans)).sh.
- 79 77 not 78

Embase.com search strategy, October 2010

The search strategy was broadened in 2010 to be more inclusive of new and emerging viruses.

- #3 #1 AND #25899
- #2 766172
- #2.8 #2.3 NOT #2.7766172
- #2.7 #2.4 NOT #2.6
- #2.6 #2.4 AND #2.5
- #2.5 'human'/de AND [embase]/lim
- #2.4 'animal'/de OR 'nonhuman'/de OR 'animal experiment'/de AND [embase]/lim
- #2.3 #2.1 OR #2.2
- #2.2 random*:ab,ti OR placebo*:ab,ti OR crossover*:ab,ti OR 'cross over':ab,ti OR allocat*:ab,ti OR trial:ti OR (doubl* NEXT/1 blind*):ab,ti AND [embase]/lim
- #2.1 'randomized controlled trial'/exp OR 'single blind procedure'/exp OR 'double blind procedure'/exp OR 'crossover procedure'/exp AND [embase]/lim
- #1 74545
- #1.65 #1.28 AND #1.6474545
- $\pm 1.64 \pm 1.29$ OR ± 1.30 OR ± 1.31 OR ± 1.32 OR ± 1.33 OR ± 1.34 OR ± 1.35 OR
- #1.36 OR #1.37 OR #1.38 OR #1.39 OR #1.40 OR #1.41 OR #1.42 OR #1.43
- OR #1.44 OR #1.45 OR #1.46 OR #1.47 OR #1.48 OR #1.49 OR #1.50 OR
- #1.51 OR #1.52 OR #1.53 OR #1.54 OR #1.55 OR #1.56 OR #1.57 OR #1.58
- OR #1.59 OR #1.60 OR #1.61 OR #1.62 OR #1.63
- #1.63 cohorting:ab,ti OR 'cohort isolation':ab,ti AND [embase]/lim
- #1.62 ((spatial OR patient*) NEAR/2 separation):ab,ti AND [embase]/lim
- #1.61 (distance NEAR/2 patient*):ab,ti AND [embase]/lim
- #1.60 (isolation NEXT/1 (room* OR strateg*)):ab,ti AND [embase]/lim
- #1.59 'personal protection':ab,ti AND [embase]/lim
- $\verb|#1.58| ((protective OR preventive) NEAR/2 (procedure* OR behaviour* OR behavior*)): ab, ti AND [embase]/limeline (protective OR preventive) (procedure* OR behaviour* OR behavior*)): ab, ti AND [embase]/limeline (procedure* OR behavior*)): ab, ti AND [embase]/limeline (proce$
- #1.57 (protective NEAR/2 (cloth* OR garment* OR device* OR equipment)):ab,ti AND [embase]/lim
- #1.56 quarantin*:ab,ti AND [embase]/lim



- #1.55 distancing:ab,ti AND [embase]/lim
- #1.54 ((outbreak* OR transmission OR infection*) NEAR/2 control):ab,ti AND [embase]/lim
- #1.53 bans:ab,ti OR banning:ab,ti OR banned:ab,ti OR ban:ab,ti AND [embase]/lim
- #1.52 (public NEAR/2 (gathering* OR event*)):ab,ti AND [embase]/lim
- #1.51 'mass gathering':ab,ti OR 'mass gatherings':ab,ti AND [embase]/lim
- #1.50 (temporar* NEAR/2 closur*):ab,ti AND [embase]/lim
- #1.49 (school* NEAR/3 (clos* OR dismissal*)):ab,ti AND [embase]/lim
- #1.48 'infection control'/de AND [embase]/lim
- #1.47 'epidemic'/dm_pc AND [embase]/lim
- #1.46 (('cross infection' OR 'cross infections') NEAR/2 prevent*):ab,ti AND [embase]/lim
- #1.45 'cross infection'/dm_pc AND [embase]/lim
- #1.44 (('reverse barrier' OR 'reverse-barrier') NEAR/3 (nurs* OR unit OR isolat*)):ab,ti AND [embase]/lim
- #1.43 'negative pressure room':ab,ti OR 'negative pressure rooms':ab,ti AND [embase]/lim
- #1.42 barrier*:ab,ti OR curtain*:ab,ti OR partition*:ab,ti AND [embase]/lim
- #1.41 (patient* NEAR/2 isolat*):ab,ti AND [embase]/lim
- #1.40 'patient isolator'/de AND [embase]/lim
- #1.39 'protective equipment'/de AND [embase]/lim
- #1.38 'protective clothing'/de AND [embase]/lim
- #1.37 facemask*:ab,ti OR mask:ab,ti OR masks:ab,ti OR goggles:ab,ti
- OR respirator*:ab,ti OR respirators:ab,ti AND [embase]/lim
- #1.36 'face mask'/exp OR 'mask'/de OR 'surgical mask'/de AND [embase]/lim
- #1.35 glov*:ab,ti AND [embase]/lim
- #1.34 'surgical glove'/de AND [embase]/lim
- #1.33 cleanser*:ab,ti OR disinfect*:ab,ti OR antiseptic*:ab,ti OR virucid*:ab,ti AND [embase]/lim
- #1.32 sanitizer*:ab,ti OR sanitiser*:ab,ti AND [embase]/lim
- #1.31 (alcohol NEAR/2 rub*):ab,ti AND [embase]/lim
- #1.30 handwash*:ab,ti OR (hand* NEAR/2 (wash* OR cleans* OR hygiene)):ab,ti AND [embase]/lim
- #1.29 'hand washing'/de AND [embase]/lim
- #1.28 #1.1 OR #1.2 OR #1.3 OR #1.4 OR #1.5 OR #1.6 OR #1.7 OR #1.8 OR #1.9 OR #1.10 OR #1.11 OR #1.12 OR #1.13 OR #1.14 OR #1.15 OR
- #1.16 OR #1.17 OR #1.18 OR #1.19 OR #1.20 OR #1.21 OR #1.22 OR #1.23
- OR #1.24 OR #1.25 OR #1.26 OR #1.27
- #1.27 (respiratory NEAR/2 (infect* OR illness* OR virus* OR pathogen* OR acute)):ab,ti AND [embase]/lim
- #1.26 parechovirus*:ab,ti AND [embase]/lim
- #1.25 'parechovirus'/de AND [embase]/lim
- #1.24 parvovirus*:ab,ti AND [embase]/lim
- ${\tt \#1.23~'parvovirus~infection'/de~OR~'erythema~infectiosum'/exp~AND~[embase]/lim}$
- #1.22 'parvovirus'/de OR 'human parvovirus b19'/de AND [embase]/lim
- #1.21 'human metapneumovirus'/de OR 'human metapneumovirus infection'/de AND [embase]/lim
- #1.20 'bocavirus'/de OR 'bocavirus infection'/de AND [embase]/lim
- #1.19 enterovir*:ab,ti AND [embase]/lim
- #1.18 'enterovirus infection'/de OR 'coxsackie virus infection'/de OR 'echovirus infection'/de AND [embase]/lim
- #1.17 'enterovirus'/de OR 'coxsackie virus'/exp OR 'echo virus'/de AND [embase]/lim
- #1.16 parainfluenza:ab,ti OR 'para influenza':ab,ti OR 'para-influenza':ab,ti AND [embase]/lim
- #1.15 'parainfluenza virus'/exp AND [embase]/lim
- #1.14 'pneumovirus infection'/de AND [embase]/lim
- #1.13 'respiratory syncytial virus':ab,ti OR 'respiratory syncytial viruses':ab,ti OR rsv:ab,ti AND [embase]/lim
- #1.12 'respiratory syncytial pneumovirus'/de OR 'respiratory syncytial virus infection'/exp AND [embase]/lim
- #1.11 coronavir*:ab,ti OR sars:ab,ti OR 'severe acute respiratory syndrome':ab,ti AND [embase]/lim
- #1.10 'coronavirus infection'/de OR 'severe acute respiratory syndrome'/de AND [embase]/lim
- #1.9 'coronavirus'/de OR 'human coronavirus nl63'/de OR 'sars coronavirus'/de OR 'transmissible gastroenteritis virus'/de
- #1.8 adenovir*:ab,ti AND [embase]/lim
- #1.7 'adenovirus infection'/de OR 'human adenovirus infection'/de OR 'human adenovirus'/exp AND [embase]/lim
- #1.6 rhinovir*:ab,ti AND [embase]/lim
- #1.5 'rhinovirus infection'/de OR 'human rhinovirus'/de AND [embase]/lim
- #1.4 'common cold':ab,ti OR 'common colds':ab,ti OR coryza:ab,ti OR colds:ab,ti AND [embase]/lim
- #1.3 'common cold'/de OR 'common cold symptom'/de AND [embase]/lim
- #1.2 influenza*:ab,ti OR flu:ab,ti AND [embase]/lim
- #1.1 'influenza'/exp AND [embase]/lim

CINAHL (EBSCO) search strategy, October 2010

The search strategy was broadened in 2010 to be more inclusive of new and emerging viruses.



S54 S32 and S53

S53 S44 or S52

S52 S45 or S46 or S47 or S48 or S49 or S50 or S51

S51 TI observational stud* or AB observational stud*

S50 TI cohort stud* or AB cohort stud*

S49 (MH "Cross Sectional Studies")

S48 (MH "Nonconcurrent Prospective Studies")

S47 (MH "Correlational Studies")

S46 (MH "Case Control Studies+")

S45 (MH "Prospective Studies")

S44 S33 or S34 or S35 or S36 or S37 or S38 or S39 or S40 or S41 or S42 or S43

S43 TI allocat* N1 random* or AB allocat* N1 random*

S42 (MH "Quantitative Studies")

S41 TI placebo* or AB placebo*

S40 (MH "Placebos")

S39 TI random* allocation* or AB random* allocation*

S38 (MH "Random Assignment")

S37 TI (randomised control* trial* or randomized control* trial*) or AB (randomised control* trial* or randomized control* trial)

S36 TI ((singl* W1 blind*) or (singl* W1 mask*) or (doubl* W1 blind*) or (doubl* W1 mask*) or (trebl* W1 blind*) or (trebl* W1 mask*) or (tripl* W1 blind*) or (tripl* W1 mask*) or (doubl* W1 blind*) or (trebl* W1 mask*) or (trebl* W1 mask*) or (trebl* W1 blind*) or (tripl* W1 blind*) or (tripl* W1 blind*) or (tripl* W1 mask*))

S35 TI clinic* W1 trial* or AB clinic* W1 trial*

S34 PT clinical trial

S33 (MH "Clinical Trials+")

S32 S15 and S31

S31 S16 or S17 or S18 or S19 or S20 or S21 or S22 or S23 or S24 or S25 or S26 or S27 or S28 or S29 or S30

S30 TI (bans or banning or banned or ban or "outbreak control" or "outbreak controls" or distancing* or quarantine* or "protective clothing" or "protective garment" or "protective garments" or "protective gown" or "protective gowns" or "protective device" or "protective devices" or "protective equipment" or "protective behaviour" or "protective behavior" or "protective behaviors" or "protective behaviors" or "protective procedure" or "protective procedures" or "preventive behaviours" or "preventive behaviors" or "preventive behaviors" or "preventive behaviors" or "preventive procedures" or "patient distance" or "patient distance" or "patient distance" or "patient distance" or "patient distances" or "patient separation" or "spatial separation") or AB (handwashing or "hand washing" or hand-washing or "hand hygiene" or sanitizer or sanitizer or cleanser* or disinfectant* or glov* or mask or masks or respirator or respirators or "patient isolation" or "patient isolators" or barrier* or curtain* or partition* or "negative pressure rooms" or "reverse barrier nursing" or "reverse barrier unit" or "reverse barrier isolation" or "cross infection" or "infection control" or "disease control" or "school closure" or "school closures" or "school dismissals" or "temporary closures" or "temporary closures" or "mass gathering" or "mass gatherings" or "public gatherings" or "public events")

S29 TI (handwashing or "hand washing" or hand-washing or "hand hygiene" or sanitizer or sanitiser or cleanser* or disinfectant* or glov* or mask or masks or respirator or respirators or "patient isolation" or "patient isolators" or barrier* or curtain* or partition* or "negative pressure rooms" or "reverse barrier nursing" or "reverse barrier unit" or "reverse barrier isolation" or "cross infection" or "infection control" or "disease control" or "school closure" or "school closures" or "school dismissal" or "school dismissals" or "temporary closures" or "mass gathering" or "mass gatherings" or "public gathering" or "public gathering" or "public events") or AB (handwashing or "hand washing" or hand-washing or "hand hygiene" or sanitizer or sanitizer or cleanser* or disinfectant* or glov* or mask or masks or respirator or respirators or "patient isolation" or "patient isolators" or barrier* or curtain* or partition* or "negative pressure room" or "negative pressure rooms" or "reverse barrier nursing" or "reverse barrier unit" or "reverse barrier isolation" or "cross infection" or "infection control" or "disease control" or "school closure" or "school closures" or "school dismissals" or "temporary closure" or "temporary closures" or "mass gatherings" or "public gatherings" or "public events")

S28 (MH "Sterilization and Disinfection")

S27 (MH "Quarantine")

S26 (MH "Area Restriction (Iowa NIC)") OR (MH "Infection Protection (IowaNIC)")

S25 (MH "Infection Control")

S24 (MH "Cross Infection/PC")

S23 (MH "Isolation, Reverse")

S22 (MH "Patient Isolation") S21 (MH "Protective Devices")

S20 (MH "Protective Clothing")

S19 (MH "Respiratory Protective Devices")

S18 (MH "Masks")

S17 (MH "Gloves")

S16 (MH "Handwashing+")



S15 S1 or S2 or S3 or S4 or S5 or S6 or S7 or S8 or S9 or S10 or S11 or S12 or S13 or S14

S14 TI ("acute respiratory tract infection" or "acute respiratory tract infections" or "acute respiratory infections") or AB (influenza* or flu or "common cold" or "common colds" or rhinovir* or adenovir* or coronavir* or sars or "severe acute respiratory syndrome" or "respiratory syncytial virus" or "respiratory syncytial viruses" or rsv or pneumovir* or parainfluenza* or "para influenza" or para-influenza or enterovir* or bocavir* or metapneumovir* or parvovir* or parechovir*)

S13 TI (influenza* or flu or "common cold" or "common colds" or rhinovir* or adenovir* or coronavir* or sars or "severe acute respiratory syndrome" or "respiratory syncytial virus" or "respiratory syncytial viruses" or rsv or pneumovir* or parainfluenza* or "para influenza" or para-influenza or enterovir* or bocavir* or metapneumovir* or parvovir* or parechovir*) or AB (influenza* or flu or "common cold" or "common colds" or rhinovir* or adenovir* or coronavir* or sars or "severe acute respiratory syndrome" or "respiratory syncytial viruses" or rsv or pneumovir* or parainfluenza* or "para influenza" or para-influenza or enterovir* or bocavir* or metapneumovir* or parvovir* or parechovir*)

S12 (MH "Respiratory Tract Infections+")

S11 (MH "Parvovirus Infections+")

S10 (MH "Enterovirus Infections+")

S9 (MH "Enteroviruses+")

S8 (MH "Respiratory Syncytial Virus Infections")

S7 (MH "Respiratory Syncytial Viruses")

S6 (MH "SARS Virus")

S5 (MH "Severe Acute Respiratory Syndrome")

S4 (MH "Coronavirus Infections+")

S3 (MH "Coronavirus+") OR (MH "Coronavirus Infections")

S2 (MH "Common Cold")

S1 (MH "Influenza+") OR (MH "Influenza A H5N1") OR (MH "Influenza A

LILACS (Latin America and Caribbean) search strategy

(mh:"Influenza, Human" OR "Gripe Humana" OR "Influenza Humana" OR influenza* OR flu OR grippe OR gripe OR mh:"Influenzavirus A" OR mh:b04.820.545.405* OR mh:b04.909.777.545.405* OR mh:"Influenzavirus B" OR mh:b04.820.545.407* OR mh:b04.909.777.545.407* OR "influenzavirus B" OR mh: "Influenzavirus C" OR "Influenzavirus C" OR mh: "Common Cold" OR "common cold" OR "common cold" OR "Resfriado Común" OR "Resfriado Comum" OR coryza OR "Coriza Aguda") AND (mh:handwashing OR "Lavado de Manos" OR "Lavagem de Mãos" OR "Desinfección de Manos" OR "Desinfecção de Mãos" OR "Higienização de Mãos Pré-Cirúrgica" OR handwash* OR "hand washing" OR "hand hygiene" OR "hand cleaning" OR "hand cleanse" OR "hand cleansing" OR higiene OR sanitizer* OR sanitiser* OR cleanser* OR disinfect* OR esteriliza* OR desinfectar* OR virucid* OR antiseptic* OR mh:"Gloves, Protective" OR "protective glove" OR "protective gloves" OR "Guantes Protectores" OR "Luvas Protetoras" OR mh:e07.700.600.400* OR mh:j01.637.215.600.400* OR mh:j01.637.708.600.400* OR glov* OR guantes OR luvas OR mh:masks OR mask* OR máscaras OR mascarillas OR facemask* OR goggles OR respirator* OR mh: "Respiratory Protective Devices" OR "Dispositivos de Protección Respiratoria" OR "Dispositivos de Proteção Respiratória" OR mh: "Protective Clothing" OR "Ropa de Protección" OR "Roupa de Proteção" OR mh:e07.700.600* OR mh:j01.637.215.600* OR mh:j01.637.708.600* OR mh:"Protective Devices" OR "Equipos de Seguridad" OR "Equipamentos de Proteção" OR mh:e07.700* OR mh:j01.637.708* OR mh:vs2.006.001.001* OR mh:vs4.002.001.001.007.002.002* OR mh:"Patient Isolation" OR "patient isolation" OR "Aislamiento de Pacientes" OR "Isolamento de Pacientes" OR mh: "Patient Isolators" OR "patient isolators" OR "Aisladores de Pacientes" OR "Isoladores de Pacientes" OR barrier* OR curtain* OR partition* OR barrera OR barreira OR cortina OR tabique OR mh:"Cross Infection" OR "cross infection" OR "Infección Hospitalaria" OR "Infecção Hospitalar" OR "Infecciones en Hospitales" OR "Infecciones Nosocomiales" OR "Infecções Nosocomiais" OR mh: "Infection Control" OR mh:n06.850.780.200.450* OR "Control de Infecciones" OR "Controle de Infecções" OR mh: "Communicable Disease Control" OR "Control de Enfermedades Transmisibles" OR "Controle de Doenças Transmissíveis" OR mh:n06.850.780.200* OR mh:sp8.946.819.811* OR mh:"Disease Outbreaks/prevention & control" OR mh:quarantine OR cuarentena OR quarentena OR "personal protection" OR "isolation room" OR "sala de aislamiento" OR "quarto de isolamento" OR "patient distance" OR "distancia del paciente" OR "spatial separation" OR cohort* OR ban OR bans OR banning OR banned OR prohibici* OR proibi* OR "outbreak control" OR distanc* OR "school closure" OR "school closures" OR "temporary closure" OR "temporary closures" OR "cierre de la escuela" OR "fechamento da escola" OR "public gathering" OR "public gatherings" OR "reunion publica" OR "reverse barrier nursing" OR "reverse barrier unit" OR "reverse barrier isolation" OR "negative pressure room" OR "negative pressure rooms" OR "patient separation") AND db: ("LILACS") AND type_of_study:("clinical_trials" OR "cohort" OR "case_control")

Indian MEDLARS search strategy

(influenza\$ or flu or common cold\$ or rhinovir\$ or coronavir\$ or adenovir\$ or severe acute respiratory syndrome\$ or sars or respiratory syncytial virus\$ or rsv or parainfluenza\$ or enterovir\$ or metapneumovir\$ or parvovir\$ or bocavir\$ or parechovir\$) and (handwashing or hand washing or mask\$ or glov\$ or protect\$ or isolat\$ or barrier\$ or curtain\$ or partition\$ or cross infection\$ or infection control\$ or disease control\$ or school\$ or quarantine\$ or ban\$ or cohort\$ or distanc\$ or spatial separation\$)

IMSEAR (Index Medicus for the South East Asia Region) search strategy

(influenza or flu or common cold or rhinovirus or coronavirus or adenovirus or severe acute respiratory syndrome or sars or respiratory syncytial virus or rsv or parainfluenza or enterovirus or bocavirus or metapneumovirus or parvovirus or parechovirus) and (handwashing or hand washing or hand hygiene or sanitizer or sanitizer or cleanser or disinfectant or gloves or masks or mask or protective clothing or protective devices or patient isolation or barrier or curtain or partition or cross infection or disease control or infection control or school or schools or bans or banning or banned or ban or distancing or quarantine or isolation or spatial separation or cohorting or cohort isolation)



In the first publication of this review we searched the Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library 2006, issue 4); MEDLINE (1966 to November 2006); OLDMEDLINE (1950 to 1965); EMBASE (1990 to November 2006) and CINAHL (1982 to November 2006). The MEDLINE search terms were modified for OLDMEDLINE, EMBASE and CINAHL.

In this 2009 update we searched the Cochrane Central Register of Controlled Trials (CENTRAL) (The Cochrane Library 2009, issue 2); Ovid MEDLINE (2006 to May Week 1 2009); OLDMEDLINE (1950 to 1965); Ovid EMBASE (2006 to Week 18, 2009) and Ovid CINAHL (2006 to May Week 1 2009).

Ovid MEDLINE

- 1 exp Influenza/
- 2 influenza.tw.
- 3 flu.tw.
- 4 exp Common Cold/
- 5 common cold.tw.
- 6 exp Rhinovirus/
- 7 rhinovirus*.tw.
- 8 exp Adenoviridae/
- 9 adenovirus*.tw.
- 10 exp Coronavirus/
- 11 exp Coronavirus Infections/
- 12 coronavirus*.tw.
- 13 exp Respiratory Syncytial Viruses/
- 14 exp Respiratory Syncytial Virus Infections/
- 15 respiratory syncytial virus*.tw.
- 16 respiratory syncythial virus.tw.
- 17 exp Parainfluenza Virus 1, Human/
- 18 exp Parainfluenza Virus 2, Human/
- 19 exp Parainfluenza Virus 3, Human/
- 20 exp Parainfluenza Virus 4, Human/
- 21 (parainfluenza or para-influenza or para influenza).tw.
- 22 exp Severe Acute Respiratory Syndrome/
- 23 (severe acute respiratory syndrome or SARS).tw.
- 24 acute respiratory infection*.tw.
- 25 acute respiratory tract infection*.tw.
- 26 or/1-25 (59810)
- 27 exp Hand Washing/
- 28 (handwashing or hand washing or hand-washing).tw.
- 29 hand hygiene.tw.
- 30 (sanitizer* or sanitiser*).tw.
- 31 (cleanser* or disinfectant*).tw.
- 32 exp Gloves, Protective/
- 33 exp Gloves, Surgical/
- 34 glov*.tw.
- 35 exp Masks/
- 36 mask*1.tw.
- 37 exp Patient Isolators/
- 38 exp Patient Isolation/
- 39 patient isolat*.tw.
- 40 (barrier* or curtain* or partition*).tw.
- 41 negative pressure room*.tw.
- 42 reverse barrier nursing.tw.
- 43 Cross Infection/pc [Prevention]
- 44 school closure*.tw.
- 45 (clos* adj3 school*).tw.
- 46 mass gathering*.tw.
- 47 public gathering*.tw.
- 48 (ban or bans or banned or banning).tw.
- 49 (outbreak* adj3 control*).tw.
- 50 distancing.tw.
- 51 exp Quarantine/
- 52 quarantine*.tw.
- 53 or/27-49



54 26 and 53

55 (animals not (humans and animals)).sh.

56 54 not 55

CENTRAL search strategy

#1 MeSH descriptor Influenza, Human explode all trees

#2 influenza:ti,ab,kw

#3 flu:ti,ab,kw

#4 MeSH descriptor Common Cold explode all trees

#5 "common cold":ti,ab,kw

#6 MeSH descriptor Rhinovirus explode all trees

#7 rhinovirus*:ti,ab,kw

#8 MeSH descriptor Adenoviridae explode all trees

#9 adenovirus*:ti,ab,kw

#10 MeSH descriptor Coronavirus explode all trees

#11 MeSH descriptor Coronavirus Infections explode all trees

#12 coronavirus*:ti,ab,kw

#13 MeSH descriptor Respiratory Syncytial Viruses explode all trees

#14 MeSH descriptor Respiratory Syncytial Virus Infections explode all trees

#15 respiratory syncytial virus*:ti,ab,kw

#16 respiratory syncythial virus*:ti,ab,kw

#17 MeSH descriptor Parainfluenza Virus 1, Human explode all trees

#18 MeSH descriptor Parainfluenza Virus 2, Human explode all trees

#19 MeSH descriptor Parainfluenza Virus 3, Human explode all trees

#20 MeSH descriptor Parainfluenza Virus 4, Human explode all trees

#21 (parainfluenza or para-influenza or para influenza):ti,ab,kw

#22 MeSH descriptor Severe Acute Respiratory Syndrome explode all trees

#23 (severe acute respiratory syndrome or SARS):ti,ab,kw

#24 acute respiratory infection*:ti,ab,kw

#25 acute respiratory tract infection*:ti,ab,kw

#26 (#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21 OR #22 OR #23 OR #24 OR #25)

#27 MeSH descriptor Handwashing explode all trees

#28 (handwashing or hand washing or hand-washing):ti,ab,kw

#29 hand hygiene:ti,ab,kw

#30 (sanitizer* or sanitiser*):ti,ab,kw

#31 (cleanser* or disinfectant*):ti,ab,kw

#32 MeSH descriptor Gloves, Protective explode all trees

#33 MeSH descriptor Gloves, Surgical explode all trees

#34 glov*:ti,ab,kw

#35 MeSH descriptor Masks explode all trees

#36 mask*:ti,ab,kw

#37 MeSH descriptor Patient Isolators explode all trees

#38 MeSH descriptor Patient Isolation explode all trees

#39 (barrier* or curtain* or partition*):ti,ab,kw

#40 negative NEXT pressure NEXT room*:ti,ab,kw

#41 "reverse barrier nursing":ti,ab,kw

#42 MeSH descriptor Cross Infection explode all trees with qualifier: PC

#43 school NEXT closure*:ti,ab,kw

#44 (clos* NEAR/3 school*):ti,ab,kw

#45 mass NEXT gathering*:ti,ab,kw

#46 public NEXT gathering*:ti,ab,kw

#47 ("ban" or "bans" or banned or banning):ti,ab,kw

#48 (outbreak* NEAR/3 control*):ti,ab,kw

#49 distancing:ti,ab,kw

#50 MeSH descriptor Quarantine explode all trees

#51 quarantine*:ti,ab,kw

#52 (#27 OR #28 OR #29 OR #30 OR #31 OR #32 OR #33 OR #34 OR #35 OR #36 OR #37 OR #38 OR #39 OR #40 OR #41 OR #42 OR #43 OR #44 OR #45 OR #46 OR #47 OR #48 OR #49 OR #50 OR #51)

#53 (#26 AND #52)

Ovid Embase search strategy



- 1 exp Influenza/
- 2 influenza.tw.
- 3 flu.tw.
- 4 exp Common Cold/
- 5 common cold.tw.
- 6 exp Human Rhinovirus/
- 7 rhinovirus*.tw.
- 8 exp Adenovirus/
- 9 adenovirus*.tw.
- 10 exp Coronavirus/
- 11 coronavirus*.tw.
- 12 exp Respiratory Syncytial Pneumovirus/
- 13 respiratory syncytial virus*.tw.
- 14 respiratory syncythial virus.tw.
- 15 (parainfluenza or para-influenza or para influenza).tw.
- 16 exp Severe Acute Respiratory Syndrome/
- 17 (severe acute respiratory syndrome or SARS).tw.
- 18 acute respiratory infection*.tw.
- 19 acute respiratory tract infection*.tw.
- 20 or/1-19
- 21 exp Hand Washing/
- 22 (handwashing or hand washing or hand-washing).tw.
- 23 hand hygiene.tw.
- 24 (sanitizer\$ or sanitiser\$).tw.
- 25 (cleanser\$ or disinfectant\$).tw.
- 26 exp Glove/
- 27 exp Surgical Glove/
- 28 glov*.tw.
- 29 exp Mask/
- 30 mask*1.tw.
- 31 patient isolat*.tw.
- 32 (barrier* or curtain* or partition*).tw.
- 33 negative pressure room*.tw.
- 34 reverse barrier nursing.tw.
- 35 Cross Infection/pc [Prevention]
- 36 school closure*.tw.
- 37 (clos* adj3 school*).tw.
- 38 mass gathering*.tw.
- 39 public gathering*.tw. (5)
- 40 (ban or bans or banned or banning).tw.
- 41 (outbreak* adj3 control*).tw.
- 42 distancing.tw.
- 43 quarantine*.tw.
- 44 or/21-43
- 45 20 and 44

EBSCO CINAHL search strategy

- S26 S10 and S24
- S25 S10 and S24
- S24 S11 or S12 or S13 or S14 or S15 or S16 or S17 or S18 or S19 or S20 or S21 or S22 or 23 or S24
- S23 TI outbreak* N3 control* or AB outbreak* N3 control*
- S22 TI (school closure* or mass gathering* or public gathering* or ban or bans or banned or banning or distancing or quarantine*) or AB (school closure* or mass gathering* or public gathering* or ban or bans or banned or banning or distancing or quarantine*)
- S21 TI (patient isolat* or barrier* or curtain* or partition* or negative pressure room* or reverse barrier nursing) or AB (patient isolat* or barrier* or curtain* or partition* or negative pressure room* or reverse barrier nursing)
- S20 TI (glov* or mask*) or AB (glov* or mask*)
- S19 TI (handwashing or hand washing or hand-washing or hand hygiene) or AB (handwashing or hand washing or hand-washing or hand hygiene)
- S18 (MH "Quarantine")
- S17 (MM "Cross Infection")
- S16 (MH "Isolation, Reverse")
- S15 (MH "Patient Isolation+")



S14 (MH "Respiratory Protective Devices")

S13 (MH "Masks")

S12 (MH "Gloves")

S11 (MH "Handwashing+")

S10 S1 or S2 or S3 or S4 or S5 or S6 or S7 or S8 or S9

S9 TI (influenza or flu or rhinovirus* or adenovirus* or coronavirus* or respiratory syncytial virus* or respiratory syncythial virus* or para-influenza or para-influenza or severe acute respiratory syndrome or SARS or respiratory viral infection* or viral respiratory infection*) or AB (influenza or flu or rhinovirus* or adenovirus* or coronavirus* or respiratory syncytial virus* or para-influenza or para-influenza or para-influenza or severe acute respiratory syndrome or SARS or respiratory viral infection* or viral respiratory

infection*)TI (influenza or flu or rhinovirus* or adenovirus* or coronavirus* or respiratory syncytial virus* or respiratory syncythial virus* or para-influenza or para-influenza or severe acute respiratory (syndrome or SARS or respiratory viral infection* or viral respiratory infection*) or AB (influenza or flu or rhinovirus* or adenovirus* or coronavirus* or respiratory syncytial virus* or respiratory syncytial virus* or para-influenza or para-influenza or severe acute respiratory syndrome or SARS or respiratory viral infection* or viral

respiratory infection*)

S8 (MH "SARS Virus")

S7 (MH "Severe Acute Respiratory Syndrome")

S6 (MH "Respiratory Syncytial Virus Infections")

S5 (MH "Respiratory Syncytial Viruses")

S4 (MH "Coronavirus+")

S3 (MH "Coronavirus Infections+")

S2 (MH "Common Cold")

S1 (MH "Influenza+")

WHAT'S NEW

Date	Event	Description
4 April 2023	Amended	John Conly's declaration of interest statement has been clarified in response to a feedback comment.

HISTORY

Protocol first published: Issue 4, 2006 Review first published: Issue 4, 2007

Date	Event	Description
27 January 2023	New citation required but conclusions have not changed	Our conclusions remain unchanged.
27 January 2023	New search has been performed	Searches updated. We included 11 new trials (Abaluck 2022; Alfelali 2020; Almanza-Reyes 2021; Ashraf 2020; Bundgaard 2021; Fretheim 2022a; Gutiérrez-García 2022; Helsingen 2021; Swarthout 2020; Teesing 2021; Young 2021), and excluded 20 new trials (Ahmadian 2022; Chen 2022; Costa 2021; Cyril Vitug 2021; Dalakoti 2022; Egger 2022; Ferrer 2021; Gharebaghi 2020; Giuliano 2021; Karakaya 2021; Kawyannejad 2020; Lim 2022; Malaczek 2022; Meister 2022; Montero-Vilchez 2022; Munoz-Basagoiti 2022; Sanchez Barrueco 2022; Seneviratne 2021; Sevinc Gul 2022). We identified two new ongoing trials (Brass 2021; NCT04471766), and five trials awaiting classification (Contreras 2022; Croke 2022; Delaguerre 2022; Loeb 2022; Varela 2022).



Date	Event	Description
1 April 2020	New search has been performed	Searches updated. In this 2020 update we only searched for RCTs and cluster-RCTs. We included 44 new trials (Aelami 2015; Aiello 2012; Alzaher 2018; Arbogast 2016; Azor-Martinez 2016; Azor-Martinez 2018; Ban 2015; Barasheed 2014; Biswas 2019; Canini 2010; Chard 2019; Correa 2012; DiVita 2011; Feldman 2016; Goodall 2014; Hartinger 2016; Hubner 2010; Huda 2012; Ibfelt 2015; Ide 2014; Ide 2016; Little 2015; MacIntyre 2011; MacIntyre 2013; MacIntyre 2015; MacIntyre 2016; McConeghy 2017; Millar 2016; Miyaki 2011; Najnin 2019; Nicholson 2014; Pandejpong 2012; Priest 2014; Radonovich 2019; Ram 2015; Savolainen-Kopra 2012; Simmerman 2011; Stebbins 2011; Suess 2012; Talaat 2011; Temime 2018; Turner 2012; Yeung 2011; Zomer 2015).
		We excluded 12 new trials (Azor-Martinez 2014; Bowen 2007; Chami 2012; Denbak 2018; Lennell 2008; Nandrup-Bus 2009; Patel 2012; Rosen 2006; Slayton 2016; Stedman-Smith 2015; Uhari 1999; Vessey 2007).
		We identified 5 new ongoing trials (NCT03454009; NCT04267952; NCT04296643; NCT04337541; Wang 2015) one of which – NCT04337541 - published as this review was going to press.
		We focused on RCTs and cluster-RCTs only and removed observational studies from this update.
1 April 2020	New citation required and conclusions have changed	There is now sufficient randomised controlled trial (RCT) evidence to show that hand hygiene is likely to provide a modest-benefit. Uncertainty remains for the other interventions. Further RCT evidence is needed.
22 October 2010	New citation required but conclusions have not changed	We updated the review again at the behest of the World Health Organization (WHO). External sources of support amended. External support from the WHO. The WHO interim guidelines document on 'Infection Prevention and Control of Epidemic and Pandemic Prone Acute Respiratory Diseases in Health Care' was published in 2007 to provide infection control guidance to help prevent the transmission of acute respiratory diseases in health care. The update of these guidelines will be evidence-based, and an update of this review was requested to assist in informing the evidence base for the revision of the WHO guidelines. Dr John Conly, Dr Mark Jones, and Sarah Thorning joined the review team.
22 October 2010	New search has been performed	Searches conducted. We included 7 new trials: 4 randomised controlled trials and 3 non-randomised comparative studies. We excluded 36 new trials.
7 May 2009	New search has been performed	For the 2009 update, we included 3 cluster-randomised controlled trials, Cowling 2009; MacIntyre 2009; Sandora 2008, and 1 individual randomised controlled trial (Satomura 2005, with its linked publication Kitamura 2007). We also included 1 retrospective cohort study (Foo 2006), 1 case-control study (Yu 2007), and 2 prospective cohort studies (Wang 2007; Broderick 2008).
		The content and conclusions of the 2007 review changed little, but the additional 8 studies add more information and certainty. Our meta-analysis remains unchanged as there were no new studies for pooling.



Date	Event	Description
30 April 2009	New citation required but conclusions have not changed	New author joined the review team.
8 July 2008	Amended	Converted to new review format.
20 August 2007	Amended	Review first published Issue 4, 2007.

CONTRIBUTIONS OF AUTHORS

For this 2022 update:

Co-ordinated the update: LD

Updated Background section: LD, MJ, LA

Updated searches: JC

Excluded irrelevant citations and disputed resolutions for trial registry searches: GB, LA

Screened titles and abstracts: EB, GB, LA, TJ

Selected studies: PG, GB, JMC

Extracted study data: MJ, TH, GB, JMC, EF, TJ Adjudicated data extraction: PG, JMC Assessed of risk of bias: MJ, GB, EF

Analysed data: MJ

Contributed to writing the update: PG, MJ, LD, TH, GB, JMC, JC, EF, MVD, LA, TJ

Approved final draft: EB, LD, PG, MJ, TH, GB, JMC, JC, EF, MVD, LA, TJ

DECLARATIONS OF INTEREST

LAA: has declared that they have no conflict of interest.

GAB: reports working at King Saud University, Medical City, Riyadh, Saudi Arabia as clinical faculty in the College of Pharmacy, collaborating with pharmacy services to provide clinical pharmacy services in primary care clinics (non-paid).

EMB: has declared that they have no conflict of interest.

JC: is an Information Specialist at the Cochrane Acute Respiratory Infections Group but was not involved in the editorial process for this review.

JMC: has held or holds peer reviewed grants from the Canadian Institutes for Health Research (CIHR) on acute and primary care preparedness for COVID-19 in Alberta, Canada and has received components of funding from a CIHR funded study via McMaster University for a randomised trial of medical masks versus N95 respirators for preventing COVID-19 amongst healthcare workers. He has also been engaged in WHO funded studies using integrated human factors and ethnography approaches to identify and scale innovative IPC guidance implementation supports in primary care with a focus on low-resource settings and using drone aerial systems to deliver medical supplies and PPE to remote First Nations communities during the COVID-19 pandemic and was the primary local Investigator for a Staphylococcus aureus vaccine study funded by Pfizer for which all funding was provided only to the University of Calgary. He has received travel support from the Centers for Disease Control and Prevention (CDC) to attend an Infection Control Think Tank Meeting and from bioMerieux Canada to speak at a symposium on antimicrobial resistance co-hosted by the University of Toronto and bioMerieux Canada. He also reports being a member and Chair of the WHO Infection Prevention and Control Research and Development Expert Group for COVID-19 and reports being a member of the WHO Health Emergencies Programme (WHE) Ad-hoc COVID-19 IPC Guidance Development Group, both of which provide multidisciplinary advice to the WHO, for which no funding is received and from which no funding recommendations are made for any WHO contracts or grants. He reports declaring an opinion on topics in this review in Clinical Microbiology and Infection and Antimicrobial Resistance and Infection Control; reports being engaged as a co-author on a randomised trial of medical masks versus N95 respirators for preventing COVID-19 amongst healthcare workers published in the Annals of Internal Medicine in 2022 and mentioned in this current Cochrane Review, but no extraction or risk of bias assessment or data pooling or other assessment was undertaken by him nor will it be in any future updates. He has also been a member of the W21C since 2004 and served as its Medical Director from 2012-2022. W21C is a notfor-profit healthcare systems research and innovation initiative based in the University of Calgary's O'Brien Institute for Public Health and the Calgary Zone of Alberta Health Services. He reports working as an Infectious Diseases Consultant at Alberta Health Services, Calgary, Canada.

LD: is a Managing Editor at the Cochrane Acute Respiratory Infections Group but was not involved in the editorial process for this review.

EF: has declared that they have no conflict of interest.

PG: reports a grant from the National Health and Medical Research Council, Australia.

TH: is a member of the Cochrane Stroke Group Editorial Board but was not involved in the editorial process for this review.



TJ: reports declaring an opinion on the topic of the review in articles for popular media. TJ is an Editor at the Cochrane Acute Respiratory Infections Group but was not involved in the editorial process for this review. See full statement here: https://restoringtrials.org/competing-interests-tom-jefferson/

MAJ: reports a grant from the National Institute for Health Research, UK. MAJ is Co-ordinating Editor at the Cochrane Acute Respiratory Infections Group but was not involved in the editorial process for this review.

MLvD: reports being a primary care panel member for the National COVID-19 Clinical Evidence Taskforce, Australia. MLvD is Deputy Coordinating Editor at the Cochrane Acute Respiratory Infections Group but was not involved in the editorial process for this review.

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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We changed the title of the review in 2010 (see Published notes below).

For the 2020 update, we added one additional outcome: adverse events related to the intervention, and we split the outcomes into primary and secondary outcomes. We also focused only on randomised controlled trials (RCTs) and cluster-RCTs and removed observational studies.

NOTES

In Issue 1, 2010, the title of the review was changed from 'Interventions for the interruption or reduction of the spread of respiratory viruses' to 'Physical interventions to interrupt or reduce the spread of respiratory viruses'.

The original review was subsequently published as Jefferson T, Foxlee R, Del Mar C, Dooley L, Ferroni E, Hewak B, Prabhala A, Nair S, Rivetti A. Physical interventions to interrupt or reduce the spread of respiratory viruses: systematic review. BMJ 2008;336:77-80 and Jefferson T, Del Mar C, Dooley L, Ferroni E, Al-Ansary LA, Bawazeer GA, van Driel ML, Foxlee R, Rivetti A. Physical interventions to interrupt or reduce the spread of respiratory viruses: systematic review. BMJ 2009;339:b3675. DOI: 10.1136/bmj.b3675.

INDEX TERMS

Medical Subject Headings (MeSH)

*Communicable Disease Control [methods]; COVID-19 [epidemiology] [prevention & control]; Global Health [statistics & numerical data]; Influenza A Virus, H1N1 Subtype; Influenza, Human [epidemiology] [prevention & control]; Randomized Controlled Trials as Topic; *Respiratory Tract Infections [epidemiology] [prevention & control]; SARS-CoV-2



MeSH check words

Aged; Child, Preschool; Humans