Nonpharmacological interventions to reduce respiratory viral transmission: an evidence-based review with recommendations*

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Rhinology 59: 2, 114 - 132, 2021
https://doi.org/10.4193/Rhn20.563

*Received for publication: October 30, 2020
Accepted: December 15, 2020

Abstract

Background: Viral respiratory infections are a leading cause of worldwide mortality and exert the potential to cause global socioeconomic crises. However, inexpensive, efficacious, and rapidly deployable strategies to reduce viral transmission are increasingly important in the setting of an ongoing pandemic, though not entirely understood. This article provides a comprehensive review of commonly employed nonpharmacological interventions to interrupt viral spread and provides evidence-based recommendations for their use.

Methodology: A systematic review of three databases was performed. Studies with defined endpoints of subjects receiving one of five interventions (nasal washing, gargling, personal protective equipment (PPE), social distancing, and hand hygiene) were included. An evidence-based review of the highest level of evidence, with recommendations, was created in accordance with a previously described, rigorous, iterative process.

Results: Fifty-four primary studies were included. The most commonly studied intervention was hand hygiene, followed by PPE, gargling, saline nasal washing, and social distancing.

Conclusions: Mask use and hand hygiene are strong recommendations for prevention of viral transmission. Donning gloves, gowns, and eye protection are a recommendation in healthcare settings. Saline nasal washing and gargling are options in selected populations. Although an aggregate level of evidence is not provided, the authors recommend social distancing.

Key words: COVID-19, evidence-based medicine, viral infection, prophylaxis

Introduction

Although often self-limited, viral respiratory tract infections (VRTI) are associated with an enormous burden of disease. The total economic impact of non-influenza-related VRTIs approaches $40 billion in direct and indirect costs annually[1]. As highlighted by the current coronavirus disease of 2019 (COVID-19) pandemic, respiratory viruses are capable of causing vast morbidity and mortality in addition to social and economic crises. Such disease outbreaks over the past two decades have included Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV) in 2003, Middle Eastern Respiratory Syndrome Coronavirus (MERS-CoV) in 2012, and most recently, Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2).

Nonpharmacological interventions (NPI), which are often inexpensive and simple to implement for a wide-scale population, have enormous value in curtailing transmission in both healthcare and community settings. These methods can be instituted rapidly and demonstrate efficacy independent of the specific infectious pathogen, including novel viruses for which more targeted therapy may not be available[2]. Given the anatomic tropism to the sinonasal cavity and nasopharynx of these viruses, rhinologists are frequently positioned on the front-lines of viral outbreaks, and are potentially at increased risk of both infection and transmission based on their unique clinical exper-
NPIs to reduce respiratory viral transmission

Although the effectiveness of various NPIs to reduce viral transmission have been previously reported\(^2,3\), they have not to our knowledge been examined using an iterative evidence-based process.

The objective of this study is to thoroughly assess the current literature on the efficacy of five principal NPIs to reduce upper VRTI transmission using a structured and systematic review process, and provide evidence-based recommendations where possible: nasal washing, gargling, personal protective equipment (PPE), social distancing, and hand hygiene. Though recommendations are provided, this review is intended not to replace clinical judgment, but rather to inform the medical community, in particular rhinologists, about the evidence for behavioural strategies that may be employed to combat outbreaks caused by respiratory viruses and to provide foundation for future research.

**Materials and methods**

**Study design**

An evidence-based review with recommendations was prepared using an online iterative process, following the methodology described by Rudmik and Smith\(^4\). The Clinical Practice Guideline Manual\(^5\), Conference on Guideline Standardization (COGS)\(^6\), and the Appraisal of Guidelines and Research Evaluation (AGREE)\(^7\) instrument recommendations were followed to improve quality, transparency, and reporting of results in this review. This study was conducted according to Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines\(^8\).

**Literature search strategy**

To identify studies for inclusion, a research informationist (E.A.B.) developed detailed search strategies in the following three databases: PubMed (U.S. National Library of Medicine, National Institutes of Health), Scopus (Elsevier), and CINAHL (EBSCOhost). Databases were queried from date of inception through July 7, 2020, and English language filters were applied. The search strategies used a combination of subject headings (e.g., MeSH in PubMed) and keywords for all relevant concepts. The PubMed search strategy was modified for the other two databases, replacing MeSH terms with appropriate subject headings, when available, and maintaining similar keywords. The search strategies for each database are detailed in Appendix 1. In an effort to identify and provide the highest level of evidence, database searches were limited to systematic review and meta-analyses when available. For comprehensive review, and to identify additional articles, the reference lists of both systematic reviews and primary articles were hand-searched. References were exported into the Covidence review management software (Veritas Health Innovation Ltd, Melbourne, Australia) for study selection.

**Inclusion and exclusion criteria**

Studies investigating the effectiveness of the five aforementioned NPIs in preventing upper VRTI transmission were included. Data from primary studies was subsequently extracted and assessed for study eligibility. Original articles with clearly defined primary clinical endpoint(s) were included. Exclusion criteria included non-English language, non-human studies, duplicates, and case reports, case series, editorials, and practice guidelines. Studies that specified a clinical endpoint involving lower respiratory tract infections and those that solely examined the effect of multicomponent interventions, such as the concurrent use of face masks and hand hygiene, were excluded. Furthermore, studies that evaluated the role of these interventions in treating VRTIs and those in which data for outcomes of interest could not be extracted were not considered.

**Data extraction and collection**

After duplicates were removed, all abstracts and records were independently reviewed by two authors (E.Y. and C.S.). Following abstract review, ineligible articles were excluded while the remaining studies underwent full-text review. Any disagreements in inclusion were resolved by consensus, then outcomes data were independently extracted from each eligible individual study.

**Development of recommendations**

Summary tables were developed for the included articles. Aggregate grade of evidence (A to D), benefit-harm assessment, and value judgments were developed for each NPI studied. An
The results of the individual studies extracted from these reviews are summarized in Tables 3-10 and are discussed in the following sections. A summary of evidence and recommendations is provided in cases where sufficient data exists to support one. Regarding the quality of individual studies, 48 studies were level 2 while six studies were level 4.

Saline nasal washing

The literature search identified two case-control studies (level 4 evidence) evaluating the role of saline nasal washing in preventing SARS infection in healthcare workers (Table 3). Both studies retrospectively assessed the nasal washing habits of hospital personnel caring for SARS patients via questionnaires. Although Chen et al. (41) found that the frequency of nasal cavity washing was not significantly associated with reduced odds of infection, Liu et al. (42) reported that this strategy conferred a protective effect. The two studies are at high risk of bias. First, neither study tested a defined hypothesis; rather, both analysed a myriad of variables to identify associations in living healthcare workers. Second, there was no attempt at matching cases with controls, and whether interviewers were blinded to the subjects’ respective study groups was not addressed. A mortality rate of up to 20% of infected healthcare workers in the first weeks of the SARS epidemic (42) poses a potentially significant risk for selection bias. Finally, as with all retrospective surveys, recall bias is a concern. Consideration of these shortcomings was acknowledged during interpretation of the included data.

Summary: Saline nasal washing

1. Aggregate grade of evidence: C
2. Benefit: Reduced VRTI transmission
3. Harm: Physical discomfort, time-cost of performing washing
4. Cost: Low (cost of nasal saline bottle, and saline, in addition to short preparation and execution time)
5. Benefits-harm assessment: Preponderance of benefit over harm
6. Value judgments: With minimal potential harm and low cost, there may be moderate value to augment an innate

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Table 1. Quality rating according to Oxford centre for evidence-based medicine.

<table>
<thead>
<tr>
<th>Grade</th>
<th>Research quality</th>
<th>Preponderance of benefit over harm</th>
<th>Balance of benefit and harm</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Well-designed RCTs</td>
<td>Strong recommendation</td>
<td>Option</td>
</tr>
<tr>
<td>B</td>
<td>RCTs with minor limitations; overwhelming consistent evidence from observational studies</td>
<td>Strong recommendation/recommendation</td>
<td>Option</td>
</tr>
<tr>
<td>C</td>
<td>Observational studies (case control and cohort design)</td>
<td>Recommendation</td>
<td>Option</td>
</tr>
<tr>
<td>D</td>
<td>Expert opinion; case report; reasoning from first principles</td>
<td>Option</td>
<td>No recommendation</td>
</tr>
</tbody>
</table>

RCT: randomized controlled trials.
HCW: healthcare workers, LOE: level of evidence, OR: odds ratio, SARS: severe acute respiratory syndrome, VRTI: viral respiratory tract infection.

*Defined using criteria provided by the China Health Ministry: Travel to a SARS epidemic area in the 2 weeks before the onset of symptoms or close contact with a probable SARS patient, fever of ≥ 38°C, chest x-ray abnormalities, normal or decreased leukocyte count, and no response to treatment by antimicrobial drugs. *Defined by World Health Organization's criteria: Documented fever (> 38°C), presence of cough or breathing difficulty, AND a significant history of exposure to a SARS patient not more than 10 days prior to onset of symptoms OR a suspect case with radiographic evidence of infiltrates consistent with pneumonia or respiratory distress syndrome on chest x-ray.

### Table 3. Summary of nasal washing studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Study Design</th>
<th>LOE</th>
<th>Definition of VRTI</th>
<th>No. of Subjects</th>
<th>Study Group(s)</th>
<th>Study Protocol</th>
<th>Primary endpoint(s)</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chen et al.</td>
<td>2009</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Nasal/pharyngeal symptoms AND 2. Increased symptom severity by 2 grades3, AND 3. Symptom worsening of ≥1 increment for ≥3 days</td>
<td>748 HCWs</td>
<td>1. SARS IgG positive 2. SARS IgG negative 3. Control (maintain previous gargling habits)</td>
<td>Questionnaire: Nasopharyngeal rinse after attending to patients performed?</td>
<td>1. SARS infection rate</td>
<td>Odds of infection not reduced with nasal cavity washing (OR 3.21 [0.98-10.53]).</td>
</tr>
<tr>
<td>Liu et al.</td>
<td>2009</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Nasal/pharyngeal symptoms AND 2. Increased symptom severity by 2 grades3, AND 3. Symptom worsening of ≥1 increment for ≥3 days</td>
<td>477 HCWs</td>
<td>1. SARS IgG positive 2. SARS IgG negative 3. Control (maintain previous gargling habits)</td>
<td>Questionnaire: Nasopharyngeal rinse after attending to patients performed?</td>
<td>1. SARS infection rate</td>
<td>2.4x higher likelihood of infection without nose washing.</td>
</tr>
</tbody>
</table>

### Table 4. Summary of gargling studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Study Design</th>
<th>LOE</th>
<th>Definition of VRTI</th>
<th>No. of Subjects</th>
<th>Study Group(s)</th>
<th>Study Protocol</th>
<th>Primary endpoint(s)</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sato-mura et al.</td>
<td>2005</td>
<td>Non-blinded RCT</td>
<td>3</td>
<td>1. Nasal/pharyngeal symptoms AND 2. Increased symptom severity by 2 grades3, AND 3. Symptom worsening of ≥1 increment for ≥3 days</td>
<td>387 healthy volunteers</td>
<td>1. Tap water gargling4 2. PVP-I gargling5 3. Control (maintain previous gargling habits)</td>
<td>Daily frequency of gargling and URI complaints recorded.</td>
<td>1. URI incidence rate</td>
<td>Only tap water gargling associated with URI prevention (IRR 0.64 [0.41-0.99]).</td>
</tr>
<tr>
<td>Toyoi-zumi et al.</td>
<td>2013</td>
<td>RCT</td>
<td>2</td>
<td>1. Positive assay for influenza virus antigen OR 2. Fever (37.8°C) and 2. Cough, sore throat, headache, and myalgia</td>
<td>308 high school students</td>
<td>1. Green tea gargling6 2. Tap water gargling7 3. Control (maintain previous gargling habits)</td>
<td>Symptoms reported to physician or school nurse.</td>
<td>1. Incidence of influenza</td>
<td>No significant difference between green tea (7.1%) and water (7.9%) gargling.</td>
</tr>
<tr>
<td>Ide et al.</td>
<td>2014</td>
<td>Non-blinded RCT</td>
<td>3</td>
<td>1. Lab-confirmed influenza</td>
<td>757 high school students</td>
<td>1. Green tea gargling6 2. Tap water gargling7 3. Control (maintain previous gargling habits)</td>
<td>Questionnaire: Occurrence of influenza infection?</td>
<td>1. Incidence of lab-confirmed influenza</td>
<td>No significant difference between green tea (4.9%) and water (6.9%) groups.</td>
</tr>
<tr>
<td>Yamada et al.</td>
<td>2007</td>
<td>Double-blinded RCT</td>
<td>2</td>
<td>1. Positive rapid assay for influenza virus antigens. Assay performed if subject had ILI8 2. URI9</td>
<td>404 healthy volunteers</td>
<td>1. Tea catechin extract gargling6 2. Without tea catechin extract gargling7 3. Control (maintain previous gargling habits)</td>
<td>Presence/severity of cold-related symptoms recorded.</td>
<td>1. Incidence of URI</td>
<td>No significant difference between catechin (1%) and control (2%) groups.</td>
</tr>
</tbody>
</table>

ILI: influenza-like illness, IRR: incidence rate ratio, LOE: level of evidence, PVP-I: povidone-iodine, RCT: randomized controlled trial, URI: upper respiratory infection, VRTI: viral respiratory tract infection. *Symptoms classified into four grades according to the Jackson method (none, mild, moderate, and severe). *Gargling regimen: 20 mL for 15 seconds 3x consecutively for at least 3x per day. *Gargled 3x daily for 90 days. *Temperature of ≥37.8°C and a recent or aggravated cough plus suggestive symptoms. *Presence of cold-related symptoms but not ILI. *Gargled for 15 seconds, 3 times consecutively, 3 times daily for 90 days.
Table 5. Summary of wearing mask studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Study Design</th>
<th>LOE</th>
<th>Definition of VRTI</th>
<th>No. of Subjects</th>
<th>Study Group(s)</th>
<th>Study Protocol</th>
<th>Primary endpoint(s)</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>MacIntyre et al. (58)</td>
<td>2009</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. ILI or 2. Symptom + lab confirmation of viral infection</td>
<td>290 adults</td>
<td>1. Surgical mask 2. P2 (N95) mask 3. No mask</td>
<td>Masks worn when in the same room as index child. Daily symptom assessment.</td>
<td>1. ILI HR 2. Lab-confirmed viral infection</td>
<td>75% less likely to develop ILI with mask use. Unable to determine difference between masks.</td>
</tr>
<tr>
<td>MacIntyre et al. (65)</td>
<td>2016</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. CRI 2. ILI 3. LabConfirmed viral respiratory infection</td>
<td>245 index cases in 597 households</td>
<td>1. Surgical mask (for index cases only) 2. No mask</td>
<td>Symptomatic individuals meeting CRI definition were tested for VRTI.</td>
<td>In contacts, RR of: 1. CRI 2. ILI 3. Lab-confirmed viral respiratory infection</td>
<td>No significant difference in RR of: CRI: 0.61 [0.18-2.13] ILI: 0.32 [0.03-3.13] Lab: 0.97 [0.06-15.54]</td>
</tr>
<tr>
<td>Cowling et al. (66)</td>
<td>2008</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. Positive viral culture or PCR for influenza 2. Clinical definitions</td>
<td>198 index subjects in 128 households</td>
<td>1. Surgical mask 2. Hand hygiene 3. Control (education)</td>
<td>Respiratory illness symptoms were documented.</td>
<td>1. SAR among household contacts</td>
<td>SAR for mask (7%), hand hygiene (6%), and control (6%) groups did not differ.</td>
</tr>
<tr>
<td>Aiello et al. (58)</td>
<td>2010</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. Cough and ≥1 constitutional symptom</td>
<td>1297 students in 7 resident halls</td>
<td>1. Face masks only 2. No intervention</td>
<td>1. ILI incidence</td>
<td>No significant incidence reduction during total study period (IRR 0.90 [0.77-1.05])</td>
<td></td>
</tr>
<tr>
<td>Aiello et al. (58)</td>
<td>2012</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. ILI 2. Lab-confirmed influenza A/B</td>
<td>1,178 young adults in 37 residence houses</td>
<td>1. Facemask only 2. Control</td>
<td>Symptoms surveyed. Throat swabs obtained for symptomatic patients.</td>
<td>1. ILI incidence 2. Lab-confirmed influenza</td>
<td>No significant incidence reduction as measured by ILI (IRR 1.10 [0.88-1.38]) and lab-confirmation (IRR 0.92 [0.59-1.42])</td>
</tr>
<tr>
<td>Barasheed et al. (58)</td>
<td>2014</td>
<td>Non-blinded cluster RCT</td>
<td>2</td>
<td>1. Lab-confirmed influenza A/B and other respiratory viruses</td>
<td>164 Haj pilgrims</td>
<td>1. Surgical masks 2. No masks</td>
<td>1. ILI symptoms 2. Virology lab results</td>
<td>Masks protective against ILI when compared to control (31% vs 53%, p= 0.04).</td>
<td></td>
</tr>
<tr>
<td>Suess et al. (58)</td>
<td>2012</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. Lab-confirmed influenza</td>
<td>218 contacts in 84 households</td>
<td>1. Surgical mask 2. No mask</td>
<td>1. Lab-confirmed influenza in household contact 2. ILI occurrence</td>
<td>Odds of infection did not differ with mask use, as measured by lab (OR 0.39 [0.13-1.19]) and ILI (OR 0.56 [0.18-1.68]).</td>
<td></td>
</tr>
<tr>
<td>Canini et al. (58)</td>
<td>2010</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. Index case: Positive rapid influenza A 2. Contact: ILI</td>
<td>306 contacts in 105 homes</td>
<td>1. Surgical mask (for index cases only) 2. Control</td>
<td>Daily symptom questionnaire. 1. ILI incidence</td>
<td>No significant difference between mask (16.2%) and control (15.8%) arms. Study prematurely terminated.</td>
<td></td>
</tr>
<tr>
<td>Jacobs et al. (58)</td>
<td>2009</td>
<td>RCT</td>
<td>2</td>
<td>1. Clinical symptoms</td>
<td>32 HCWs</td>
<td>1. Surgical masks 2. No surgical masks</td>
<td>Daily symptom diary. 1. URI incidence</td>
<td>No significant difference between mask (5.9%) and control (6.7%) arms.</td>
<td></td>
</tr>
<tr>
<td>Radonovich et al. (58)</td>
<td>2019</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. Lab-confirmed influenza</td>
<td>4,051 HCWs in 7 medical centres</td>
<td>1. N95 respirator 2. Surgical mask</td>
<td>Nasal and throat swabs collected.</td>
<td>1. Incidence of lab-confirmed influenza</td>
<td>No significant difference between N95 (8.2%) vs. surgical masks (7.2%).</td>
</tr>
<tr>
<td>Loeb et al. (58)</td>
<td>2009</td>
<td>RCT</td>
<td>2</td>
<td>1. Clinical symptoms</td>
<td>446 nurses</td>
<td>1. N95 respirator 2. Surgical mask</td>
<td>Nasal specimens collected for newly reported symptoms.</td>
<td>1. Incidence of lab-confirmed influenza</td>
<td>Use of surgical mask compared with N95 resulted in non-inferior rates (23.6% vs 22.9%) of infection.</td>
</tr>
</tbody>
</table>
A gargling diary daily, documenting the frequency of gargling and any URTI complaints. Incident rates were found to be lower in the water gargling cohort (0.17 episode/30 person-days) and in the PVP-I gargling cohort (0.24 episode/30 person-days) compared with control (0.26 episode/30 person-days). However, on multivariate analysis, gargling with tap water, but not with PVP-I, was found to significantly reduce VRTI incidence (36% decrease). Although participants were not blinded to the intervention, disease incidence was determined by one study physician who was blinded to the results of assignment. Notably, despite the potential irritant nature of PVP-I, only 2% of subjects complained of discomfort or difficulties in gargling and withdrew.

Three RCTs assessed the role of green tea gargling in reducing the incidence of infection during the influenza season. Each study utilized similar gargling regimens (three times daily for 90 days) and the incidence of laboratory-confirmed influenza as the primary endpoint. None of these studies found benefit in

**Table 4**

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
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<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Macintyre et al.</td>
<td>2011</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. CRI or ILI 1. Lab-confirmed viral respiratory infections</td>
<td>1,441 HCWs in 15 hospitals</td>
<td>1. Medical mask 2. Fit-tested N95 3. Non-fit-tested N95 4. No mask</td>
<td>Daily contact to identify incident cases of respiratory infection.</td>
<td>1. CRI and ILI incidence 2. Lab-confirmed respiratory virus or influenza incidence</td>
<td>1. CRI (3.9% vs 6.7%), ILI (0.3% vs 0.6%), lab-confirmed virus (1.4% vs 2.6%) and influenza (0.3% vs 1%) rates lower for N95 compared to medical masks. 2. Non-fit-tested N95 more protective (3.3% vs 6.7%) than medical masks against CRI only.</td>
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<td>61</td>
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</tr>
<tr>
<td>Macintyre et al.</td>
<td>2013</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. CRI or ILI 1. Lab-confirmed viral respiratory infections</td>
<td>1,669 HCWs in 19 hospitals</td>
<td>1. Medical mask 2. N95 3. Targeted use of N95 during high-risk procedures</td>
<td>Daily assessment for respiratory infections. Swabbed if symptomatic.</td>
<td>1. CRI or ILI incidence 2. Lab-confirmed VRTI incidence</td>
<td>1. CRI incidence significantly lower with N95 (7.2%) vs. medical mask (17.1%). 2. Rate of lab-confirmed VRTI did not differ between medical mask (33%), N95 (22%), and targeted N95 (33%) arms.</td>
</tr>
<tr>
<td>61</td>
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</tr>
<tr>
<td>Macintyre et al.</td>
<td>2015</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. CRI or ILI 1. Lab-confirmed viral respiratory infections</td>
<td>1,607 HCWs working in high-risk wards across 14 hospitals</td>
<td>1. Medical mask 2. Cloth mask</td>
<td>Daily assessment for respiratory infections. Swabbed if symptomatic.</td>
<td>1. CRI or ILI incidence 2. Lab-confirmed VRTI incidence</td>
<td>1. ILI rate higher (RR 13 [1.69-100.07]) with cloth vs medical mask. 2. Lab-confirmed VRTI rate higher (RR 1.72 [1.01-2.94]) with cloth vs medical mask.</td>
</tr>
<tr>
<td>62</td>
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</table>


1. Temperature ≥ 37.8°C, history of fever or feeling feverish, ≥2 symptoms (arthralgia, sore throat, cough, sneezing, runny nose, nasal congestion, headache).
2. Influenza A and B, RSV, PIV Types 1-3, enteroviruses, rhinoviruses, adenoviruses, coronavirus 229E and OC43, and hMPV.
3. Two or more respiratory or one respiratory symptom and a systemic symptom.
4. Fever ≥38°C plus one respiratory symptom (i.e. cough, runny nose, sore throat, etc.).
5. Detection of adenovirus, human metapneumovirus, coronavirus 229E/NL63/OC43/HKU1, parainfluenza viruses 1, 2, and 3, influenza viruses A and B, respiratory syncytial virus A and B, rhinovirus A/B.
6. Presence of cough and ≥1 of fever/feverishness, chills, or body aches.
7. Subjective (or proven) fever plus one respiratory symptom (e.g. dry or productive cough, runny nose, sore throat, shortness of breath).
8. Temperature >37.8°C or at least two of the following: sore throat, cough, runny nose, or fatigue.

immune mechanism in healthcare workers and patients who tolerate saline irrigations or are at especially high risk.

**Recommendation level**: Option

**Intervention**: Prophylactic nasal washing with saline solution. Optimal frequency, technique, and solution undetermined.

**Gargling**

Four unique RCTs evaluating the role of gargling in preventing upper VRTIs were identified (Table 4). Only one was a double-blinded trial, in part due to the difficulty of masking or mimicking the tastes of povidone iodine (PVP-I) and green tea. Satomura et al. conducted a trial in which 387 participants, followed for 60 days, were randomly assigned to three treatment arms: tap water gargling, PVP-I gargling, and usual care (control). Participants in the two intervention arms were instructed to gargle three times per day. All subjects were asked to maintain a gargling diary daily, documenting the frequency of gargling and any URTI complaints. Incident rates were found to be lower in the water gargling cohort (0.17 episode/30 person-days) and in the PVP-I gargling cohort (0.24 episode/30 person-days) compared with control (0.26 episode/30 person-days). However, on multivariate analysis, gargling with tap water, but not with PVP-I, was found to significantly reduce VRTI incidence (36% decrease). Although participants were not blinded to the intervention, disease incidence was determined by one study physician who was blinded to the results of assignment. Notably, despite the potential irritant nature of PVP-I, only 2% of subjects complained of discomfort or difficulties in gargling and withdrew. Three RCTs assessed the role of green tea gargling in reducing the incidence of infection during the influenza season. Each study utilized similar gargling regimens (three times daily for 90 days) and the incidence of laboratory-confirmed influenza as the primary endpoint. None of these studies found benefit in
gargling with green tea. Each study, however, had noteworthy limitations. Yamada et al. recruited healthy adults inoculated with the influenza vaccine prior to study participation, which may lower the incidence of influenza below what is required to obtain statistical power. Toyozumi et al. and Ide et al. suffered from low adherence rates (<75%) among high school students. When both studies performed a per protocol set analysis, in which non-adherent participants were excluded, the reduction in influenza infection was greater, suggesting that a significant difference in outcomes between the green tea and water gargling groups may be detected in future investigations with greater compliance. In addition, as all four trials were conducted with healthy subjects in Japan, additional studies including different populations with varying sociodemographic characteristics may improve the generalizability of the results.

Summary: Gargling

1. **Aggregate grade of evidence:** C
2. **Benefit:** Reduced VRTI transmission
3. **Harm:** Physical discomfort and potentially taste associated with gargling
4. **Cost:** Low
5. **Benefits-harm assessment:** Preponderance of benefit over harm
6. **Value judgments:** None
7. **Recommendation level:** Option
8. **Intervention:** Gargling. Population, optimal frequency, technique, and solution undetermined

### Masks

The efficacy of surgical masks, N95 respirators, and cloth masks in preventing upper VRTI transmission were reviewed (Table 5). The literature search identified 14 unique RCTs for evaluation. All 14 RCTs (level 2 evidence) assessed the use of masks, including surgical masks, N95 respirators, and cloth masks, as physical barriers. These studies were conducted either in community or healthcare settings.

#### Surgical masks

The 8 studies that evaluated the use of surgical masks in the community setting failed to clearly demonstrate any benefit. Several studies were underpowered to detect any statistical difference in primary endpoints between the study groups and...
Table 7. Summary of wearing gowns studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Study Design</th>
<th>LOE</th>
<th>Definition of VRTI</th>
<th>No. of Subjects</th>
<th>Study Group(s)</th>
<th>Study Protocol</th>
<th>Primary endpoint(s)</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chen et al.</td>
<td>2004</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Probable SARS cases(^a), 2. SARS IgG positivity</td>
<td>748 HCWs</td>
<td>1. SARS IgG positive 2. SARS IgG negative</td>
<td>Questionnaire: Quantify gowns worn while caring for SARS patients.</td>
<td>1. SARS infection rate</td>
<td>Wearing double gowns not significantly more protective than wearing a single one.</td>
</tr>
<tr>
<td>Nishiura et al.</td>
<td>2005</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Lab-confirmed SARS infection</td>
<td>115 HCWs</td>
<td>1. SARS cases 2. SARS exposure</td>
<td>Survey: Gowns worn when in contact with SARS patients?</td>
<td>1. SARS infection rate</td>
<td>Use of gowns lowered rate by 80%.</td>
</tr>
<tr>
<td>Seto et al.</td>
<td>2003</td>
<td>Case-control study</td>
<td>4</td>
<td>1. SARS infection(^b), confirmed by positive serology results</td>
<td>254 HCWs</td>
<td>1. SARS cases 2. SARS exposure</td>
<td>Questionnaire: Were gowns used during patient care?</td>
<td>1. SARS infection rate</td>
<td>Gown use significantly differed between infected (0%) and non-infected (34%) staff.</td>
</tr>
<tr>
<td>Teleman et al.</td>
<td>2004</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Probable SARS cases(^c), confirmed by positive serology results</td>
<td>86 HCWs</td>
<td>1. HCWs with probable SARS 2. HCWs with SARS exposure</td>
<td>Questionnaire: Gown compliance during patient contact?</td>
<td>1. SARS infection rate</td>
<td>Gowns not significantly protective (OR 0.5 [0.4-6.9]).</td>
</tr>
<tr>
<td>Yin et al.</td>
<td>2004</td>
<td>Case-control study</td>
<td>4</td>
<td>Limited information obtained from partial translation</td>
<td>257 HCWs</td>
<td>1. HCWs with SARS 2. HCWs without SARS</td>
<td>Questionnaire: Gown wearing?</td>
<td>1. SARS infection rate</td>
<td>Wearing a gown provided significant protection against SARS infection.</td>
</tr>
</tbody>
</table>

HCW: healthcare workers, LOE: level of evidence, OR: odds ratio, SARS: severe acute respiratory syndrome, VRTI: viral respiratory tract infection. \(^a\)Defined using criteria provided by the China Health Ministry. \(^b\)Defined as fever of ≥38°C, radiological infiltrates compatible with pneumonia, and two of: chills, new cough, malaise, and signs of consolidation. \(^c\)Defined by WHO's criteria.

thus failed to report conclusive findings on their efficacy in preventing viral transmission in community settings. Of note, the trial conducted by Canini et al.\(^{50}\) was severely underpowered due to premature termination of the study. MacIntyre et al.\(^{57}\) cited poor compliance (<50%) as one of the main limitations of the study, concluding that without the threat of an epidemic or pandemic to encourage greater adherence, household use of face masks would be ineffective for controlling seasonal respiratory disease.

In contrast to earlier studies that sought to ascertain the impact of mask use in preventing secondary viral transmission, Aiello et al. designed two studies\(^{52,53}\) to examine their effectiveness when implemented prior to the onset of influenza-like illness symptoms. In both trials, students living in residence houses were assigned to one of three arms: face mask and hand hygiene, face mask only, or control. Using the rate of VRTI as the primary endpoint, Aiello et al. reached the same conclusion in both studies: neither intervention was associated with a significant reduction cumulatively during the 6-week study period. Despite these findings, the significance of these two trials lies in their study design, which more accurately represents guidelines that recommend the use of NPIs before susceptible individuals become infected and an outbreak ensues.

In healthcare settings, evidence on the efficacy of surgical masks in preventing upper VRTI transmission was similarly inconclusive. In two studies\(^{61,63}\), although the incidence of VRTI was higher among healthcare workers in the no-mask group, the difference was not significant. However, both studies reported limitations, including lack of randomization of the control arm\(^{61}\) and a small sample size \((n = 32)\).\(^{63}\)

**N95 Respirators**

Evidence supporting the use of N95 respirators for preventing viral transmission was stronger in healthcare than in community settings. Among the studies conducted in the community, only one\(^{37}\) assessed its effectiveness relative to surgical masks and no masks. Although the secondary attack rate in exposed adults did not significantly differ among the three study arms, the trial suffered from low compliance as previously mentioned. In addition, a small sample size precluded any conclusive comparison of the relative efficacy of N95 respirators and surgical masks. Four RCTs assessed the efficacy of N95 respirators when utilized by healthcare workers. MacIntyre et al. demonstrated in two separate trials\(^ {60,61}\) that N95 respirators were superior to medical masks in preventing clinical respiratory illness despite the greater discomfort and lower adherence associated with respirator use. Moreover, to observe this benefit, healthcare workers had to wear the respirator consistently during the entirety of their shifts rather than intermittently. In contrast, two studies\(^ {64,65}\) reported no significant difference in VRTI incidence between
the two interventions. The study conducted by Loeb et al.\textsuperscript{44} had several limitations, including small sample size (n = 446), lack of control arm, and the use of serology to confirm influenza infection. Despite recruiting more patients (n = 4,051) and utilizing both clinical symptoms and laboratory confirmation to reach the diagnosis, Radonovich et al.\textsuperscript{65} similarly found no difference in primary endpoints.

**Cloth masks**

The efficacy of cloth masks was evaluated in one trial\textsuperscript{62}, which found that the rate of VRTI was significantly higher among healthcare workers wearing cloth masks compared with those using medical masks, likely because the penetration of particles through the cloth masks was found to be very high (97%).

**Summary: Masks**

**For Community Settings:**

1. **Aggregate grade of evidence:** B
2. **Benefit:** Reduced VRTI transmission
3. **Harm:** Discomfort while wearing, more protective masks may lead to increased user discomfort.
4. **Cost:** Variable; low cost of materials in setting of normal demand, but may increase substantially in setting of reduced supply. Fit testing for respirators.
5. **Benefits-harm assessment:** Preponderance of benefit over harm
6. **Value judgments:** High value to prevent community spread in setting of high risk of exposure, with partially diminished value in lower risk, community settings
7. **Recommendation level:** Recommendation
8. **Intervention:** Donning masks. Optimal mask type undetermined due to varying efficacy.

For Healthcare Settings:

1. **Aggregate grade of evidence:** A
2. **Benefit:** Reduced VRTI transmission
3. **Harm:** Discomfort while wearing; lower compliance of respirators as compared to other masks
4. **Cost:** Variable; low cost of materials in setting of normal demand, but may increase substantially in setting of reduced supply
5. **Benefits-harm assessment:** Preponderance of benefit over harm
6. **Value judgments:** Potential exceedingly high value in high risk, healthcare settings
7. **Recommendation level:** Strong recommendation
8. **Intervention:** Donning masks. Optimal mask type undetermined. Respirators are warranted for high risk scenarios, though continuous use is associated with discomfort and risk of lower compliance. Cloth masks are not recommended in the clinical setting.

**Gloves, Gowns, and Eye Protection**

Because no RCTs assessing the use of gloves, gowns, and eye protection in preventing upper VRTI transmission were identified, available evidence derived from case-control studies was evaluated (Tables 6-8). All six studies recruited healthcare workers who cared for SARS patients during the outbreak in 2003, retrospectively assessing their compliance with PPE usage via questionnaires. The incidence of SARS infection among participants was used as the measured outcome. Of the six case-control\textsuperscript{41-46} studies on glove-wearing, half of them reported no significant benefit in donning gloves to reduce viral transmission. Five case-control studies\textsuperscript{41,43-46} on gown-wearing simi-

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### Table 8. Summary of wearing eye protection (mask/goggles) studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>LOE</th>
<th>Definition of VRTI</th>
<th>No. of Subjects</th>
<th>Study Group(s)</th>
<th>Study Protocol</th>
<th>Primary endpoint(s)</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chen et al.</td>
<td>2009</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Probable SARS cases\textsuperscript{a} 2. SARS IgG positivity</td>
<td>748 HCWs</td>
<td>1. SARS IgG positive 2. SARS IgG negative</td>
<td>Questionnaire: Quantify frequency of wearing goggles while caring for SARS patients.</td>
<td>1. SARS infection rate</td>
<td>Frequency of wearing goggles did not significantly reduce infection rate.</td>
</tr>
<tr>
<td>Liu et al.</td>
<td>2009</td>
<td>Case-control study</td>
<td>4</td>
<td>1. Probable SARS cases\textsuperscript{b} 2. SARS IgG positivity</td>
<td>477 HCWs</td>
<td>1. SARS IgG positive 2. SARS IgG negative</td>
<td>Questionnaire: Did participants wear goggles during patient care?</td>
<td>1. SARS infection rate</td>
<td>Wearing goggles significantly reduced rate (7.7% vs. 13.3%).</td>
</tr>
<tr>
<td>Yin et al.</td>
<td>2004</td>
<td>Case-control study</td>
<td>4</td>
<td>[Limited information obtained from partial translation]</td>
<td>257 HCWs</td>
<td>1. HCWs with SARS 2. HCWs without SARS</td>
<td>Questionnaire: Goggle wearing?</td>
<td>1. SARS infection rate</td>
<td>Wearing goggles reduced odds of infection (OR 0.20 [0.10-0.41]).</td>
</tr>
</tbody>
</table>

HCW: healthcare workers, LOE: level of evidence, OR: odds ratio, SARS: severe acute respiratory syndrome, VRTI: viral respiratory tract infection. \textsuperscript{a}Defined using criteria provided by the China Health Ministry. \textsuperscript{b}Defined by WHO's criteria.
larly provided conflicting evidence, with three demonstrating benefit. Finally, of three case-control studies\(^ {51, 64, 65}\) on utilizing eye protection, two determined that the use of eye masks or goggles conferred a protective effect against viral infection for healthcare workers. As previously mentioned, nuanced and cautious interpretation of the relatively lower quality evidence derived from case-control studies was undertaken.

**Summary: Gloves, gowns, and eye protection**

1. **Aggregate grade of evidence:** C
2. **Benefit:** Reduced VRTI transmission
3. **Harm:** Discomfort, minimal
4. **Cost:** Low, cost of materials, which may be fluid during supply shortages of large viral outbreaks
5. **Benefits-harm assessment:** Preponderance of benefit over harm, as there is (limited) evidence to support benefit, but definitely zero evidence to support harm
6. **Value judgments:** With limited evidence to support benefit of use, and no evidence of potential harm, this intervention has potential for high value in the analysed healthcare setting
7. **Recommendation level:** Recommendation in healthcare setting
8. **Intervention:** Donning PPE for healthcare workers. Differences in efficacy of gloves, gowns, and eye protection could not be determined.

**Social distancing**

One quasi-cluster RCT (level 2 evidence) was reviewed (Table 9). Miyaki et al.\(^ {51}\) evaluated the effectiveness of social distancing in workplaces, utilizing the incidence of influenza A H1N1 as the primary endpoint. Company employees in the intervention arm were asked to remain home if a co-habitation family member developed a VRTI while those in the control group reported to work as usual if the same situation occurred. Following the 233-day study period, the authors demonstrated that implementation of a stay-at-home policy significantly reduced the overall risk of influenza infection in the workplace by 20%. Full compliance was achieved among those following social distancing protocol, likely because employees continued to receive full pay while at home and were explained about its public health benefit. Therefore, the authors regarded promoting a public health mindset and offering financial support as important measures to maintain high adherence to social distancing policies.

An aggregate level of evidence is not provided for this level 2 study, however, its findings are notable, including the potential for reduced VRTI transmission. Potential harm and cost include reduced productivity, especially if workers lose compensation or cannot work from home. Despite potential financial and psychosocial risks when implemented over long periods, there appears to be high value of this intervention, especially when employed in the acute infectious period. Based on the available evidence, the authors recommend social distancing protocols with measures to mitigate productivity and compensation loss (e.g. ability to work remotely).

**Hand hygiene**

The literature search identified 29 unique RCTs (level 2 evidence) for review (Table 10). Heterogeneity of settings, hand hygiene interventions, and primary endpoints existed across studies. Nine studies\(^ {66-74}\) were conducted in elementary schools, of which six\(^ {66-71}\) reported a significant reduction in VRTI-associated absenteeism in the intervention arm compared with control.

To elucidate the reported effect of hand hygiene on reducing viral transmission, the interventions studied were stratified into three broad categories to allow for comparison: those that mainly promoted hand sanitizer use, those that emphasized soap use, and those that provided education only. Four trials studied the effect of hand sanitizer (alcohol-based or alcohol-free) use alone but because this intervention was offered at different times throughout the school day, a comparison of outcomes was challenging. Of the three studies that tested the effect of an alcohol-based handrub alone, two\(^ {67, 70}\) reported a significant re-
Table 10. Summary of hand hygiene studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Design</th>
<th>LOE</th>
<th>Definition of VRTI</th>
<th>No. of Subjects</th>
<th>Study Group(s)</th>
<th>Study Protocol</th>
<th>Primary endpoint(s)</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cowling et al.</td>
<td>2009</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>1. Lab evidence of influenza OR 2. &gt;2 of the following: temperature &gt;37.8°C, cough, headache, sore throat, myalgia OR 3. Temperature &gt;37.8°C plus cough or sore throat</td>
<td>1,201 (407 index patients, 794 household contacts)</td>
<td>1. Hand hygiene + education 2. Lifestyle education</td>
<td>All household contacts kept daily symptom diaries. Nasal and throat swabs collected at home visits.</td>
<td>1. SAR among household contacts</td>
<td>No significant difference in SAR between intervention (14%) and control (24%).</td>
</tr>
<tr>
<td>Simmerman et al.</td>
<td>2011</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>1. ILI: Lab-confirmed influenza 2. URI: Not specified</td>
<td>885 subjects</td>
<td>1. Handwashing 2. Education</td>
<td>Respiratory swabs and serum collected from all household members.</td>
<td>1. SAR among household contacts</td>
<td>SAR not reduced by promotion of handwashing (23%) vs. control (19%).</td>
</tr>
<tr>
<td>Larson et al.</td>
<td>2010</td>
<td>Cluster block RCT</td>
<td>2</td>
<td>1. ILI: Temperature of &gt;37.8°C and cough and/or sore throat 2. URI: Not specified</td>
<td>2788 participants in 617 households</td>
<td>1. Alcohol-based HS + education 2. Education</td>
<td>Presence or absence of symptoms reported for every household member.</td>
<td>1. Incidence and secondary transmission of VRTI</td>
<td>No reduction in incidence (29% vs. 35%) or SAR (14.4% vs. 13.7%) with HS.</td>
</tr>
<tr>
<td>Priest et al.</td>
<td>2014</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>Runny, stuffy, or blocked nose, noisy breathing, cough, fever, sore throat or sneezing*</td>
<td>2443 students</td>
<td>1. Alcohol-based HS + education 2. Hand hygiene education</td>
<td>Absence information of children collected.</td>
<td>1. Absence episodes per 100 child-days.</td>
<td>Absence episodes not significantly different with HS (1.21) vs. control (1.16).</td>
</tr>
<tr>
<td>Sandora et al.</td>
<td>2008</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>Runny, stuffy, or blocked nose, cough, fever or chills, sore throat, or sneezing.</td>
<td>285 students</td>
<td>1. Alcohol-based HS 2. Usual practices</td>
<td>Reason for absence recorded on a standardized form.</td>
<td>1. Rate of school absenteeism</td>
<td>No significant impact on rate ratio (1.07 [0.92-1.24]) with HS.</td>
</tr>
<tr>
<td>Stebbins et al.</td>
<td>2011</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>ILI: fever &gt;38°C with sore throat or cough</td>
<td>3360 students</td>
<td>1. HS + education 2. Standard practice</td>
<td>Student absenteeism recorded.</td>
<td>1. Absence episodes</td>
<td>No significant effect of intervention (IRR 0.81 [0.54-1.23]).</td>
</tr>
<tr>
<td>Bowen et al.</td>
<td>2007</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>Conjunctivitis, otalgia, rhinorrhea, sore throat, or cough.</td>
<td>3962 students in 87 groups</td>
<td>1. Soap + education 2. Education 3. Standard practice</td>
<td>Symptoms or signs of illness identified by teachers.</td>
<td>1. ARI-related absenteeism 2. ARI illness rate</td>
<td>Significant reductions in absenteeism (1.2 vs. 2.6 d) and rate (1.2 vs. 2 episodes) with soap use + education only.</td>
</tr>
<tr>
<td>Patel et al.</td>
<td>2012</td>
<td>Cluster RCT 2</td>
<td>2</td>
<td>ARI: reported fever and cough or difficulty breathing</td>
<td>43 schools and 643 households</td>
<td>1. Handwashing + education 2. Standard practice</td>
<td>ARI assessed for and recorded.</td>
<td>1. ARI illness rate</td>
<td>Significant rate reduction with handwashing vs. control (2% vs. 3%).</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Study Design</td>
<td>LOE</td>
<td>Definition of VRTI</td>
<td>No. of Subjects</td>
<td>Study Group(s)</td>
<td>Study Protocol</td>
<td>Primary endpoint(s)</td>
<td>Conclusion</td>
</tr>
<tr>
<td>---------------</td>
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<td>---------------------------------------------------------------------------</td>
</tr>
<tr>
<td>White et al.</td>
<td>2001</td>
<td>Double-blinded, placebo-controlled cluster RCT</td>
<td>2</td>
<td>Symptoms of cough, sneezing, sinus trouble, bronchitis, fever, pink-eye, headache, mononucleosis</td>
<td>769 students</td>
<td>1. Alcohol-free HS</td>
<td>Daily attendance and nature of student's absence recorded.</td>
<td>1. Illness absence incidence</td>
<td>HS significantly decreased likelihood of absence by 31%.</td>
</tr>
<tr>
<td>Talaat et al.</td>
<td>2011</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. ILI: fever &gt;38°C + cough or sore throat 2. Lab-confirmed influenza</td>
<td>44,451 students</td>
<td>1. Soap + education 2. Standard practice</td>
<td>Absences due to illness recorded.</td>
<td>1. Absences due to illness</td>
<td>Absenteeism significantly reduced by 40-50%.</td>
</tr>
<tr>
<td>Pandejpong et al.</td>
<td>2012</td>
<td>Cluster RCT</td>
<td>2</td>
<td>Runny/stuffy nose, cough, fever, chills, sore throat, headache, hand-foot-mouth ulcers</td>
<td>1,437 students</td>
<td>1. HS every 60 min. 2. HS every 120 min. 3. HS before lunch</td>
<td>Absences caused by ILI recorded.</td>
<td>1. Change in absenteeism rate</td>
<td>Rate significantly lowered with HS every 60 min. (1.7%), but not 120 min. (2.5%), vs. control (2.6%)</td>
</tr>
<tr>
<td>Kotch et al.</td>
<td>1994</td>
<td>Cluster RCT</td>
<td>2</td>
<td>Symptoms of coughing, runny nose, wheezing, sore throat, or earache</td>
<td>389 children in 24 childcare centres</td>
<td>1. Handwashing training 2. No training</td>
<td>Respiratory symptoms elicited.</td>
<td>1. Incidence of respiratory episodes</td>
<td>No reduction in incidence (RR 0.94 [-2.43-0.66]).</td>
</tr>
<tr>
<td>Sandora et al.</td>
<td>2005</td>
<td>Cluster RCT</td>
<td>2</td>
<td>2 of the following: runny nose, stuffy/ blocked nose, noisy breathing, cough, fever/chills, sore throat, sneezing</td>
<td>292 families with children enrolled in childcare centres</td>
<td>1. Alcohol-based HS + education 2. Education only</td>
<td>Self-reported symptoms collected.</td>
<td>1. Weekly prevalence of RTI symptoms</td>
<td>No risk reduction with soap (RR 0.8, [0.66-1.03]) or alcohol rub (RR 0.96, [0.76-1.20]) vs. control.</td>
</tr>
<tr>
<td>Hovi et al.</td>
<td>2017</td>
<td>Cluster RCT</td>
<td>2</td>
<td>&quot;Symptoms typical of acute RTI&quot;</td>
<td>683 office employees</td>
<td>1. Plain soap 2. Antibacterial soap</td>
<td>Self-reported symptoms collected.</td>
<td>1. Weekly prevalence of RTI symptoms</td>
<td>No risk reduction with soap (RR 0.8, [0.66-1.03]) or alcohol rub (RR 0.96, [0.76-1.20]) vs. control.</td>
</tr>
<tr>
<td>Luby et al.</td>
<td>2005</td>
<td>Cluster RCT</td>
<td>2</td>
<td>Symptoms of cough, difficulty breathing, congestion, or coryza</td>
<td>4,691 participants and 36 households</td>
<td>1. Plain soap 2. Antibacterial soap 3. Standard practice</td>
<td>Symptoms in all households recorded.</td>
<td>1. Incidence of ARI in children</td>
<td>1. 50% reduction with both soaps. 2. Incidence between 2 soaps did not significantly differ.</td>
</tr>
<tr>
<td>Larson et al.</td>
<td>2004</td>
<td>Double-blind RCT</td>
<td>2</td>
<td>Symptoms of fever, sore throat, cough, rhinorrhea, conjunctivitis</td>
<td>1,178 subjects in 238 households</td>
<td>1. Antibacterial products 2. Non-antibacterial products</td>
<td>Symptoms in individual household members assessed.</td>
<td>1. Risk for symptoms of ARI</td>
<td>Risk not significantly different (RR 0.96 [0.82-1.12]).</td>
</tr>
<tr>
<td>Little et al.</td>
<td>2015</td>
<td>RCT</td>
<td>2</td>
<td>1. RTIc 2. ILId</td>
<td>20,066 subjects</td>
<td>1. Education 2. No intervention</td>
<td>Nature and frequency of symptoms documented.</td>
<td>1. RTI/ILI incidence</td>
<td>Decreased risk of RTI (RR 0.86 [0.83-0.89]) or ILI (RR 0.8 [0.72-0.92]) vs. control.</td>
</tr>
<tr>
<td>Carabin et al.</td>
<td>1999</td>
<td>Cluster RCT</td>
<td>2</td>
<td>1. Nasal discharge + fever, sneezing, cough, sore throat, ear pain, malaise, or irritability</td>
<td>1,729 children in 47 daycare centres</td>
<td>1. Education 2. No education</td>
<td>Absences and daily occurrence of colds recorded.</td>
<td>1. Incidence of URI</td>
<td>Intervention reduced incidence of URI (IRR 0.80 [0.68, 0.93]).</td>
</tr>
<tr>
<td>Study</td>
<td>Year</td>
<td>Study Design</td>
<td>LOE</td>
<td>Definition of VRTI Cluster</td>
<td>No. of Subjects</td>
<td>Study Group(s)</td>
<td>Study Protocol</td>
<td>Primary endpoint(s)</td>
<td>Conclusion</td>
</tr>
<tr>
<td>-------------------</td>
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<tr>
<td>Ban et al.</td>
<td>2015</td>
<td>Cluster RCT</td>
<td>2</td>
<td>ARI'</td>
<td>466 children</td>
<td>1. Soap, alcohol-based HS 2. Standard practice</td>
<td>Daily data on illness symptoms collected.</td>
<td>1. ARI illness reduced with intervention (OR 0.47 [0.38-0.59]).</td>
<td></td>
</tr>
<tr>
<td>Correa et al.</td>
<td>2012</td>
<td>Cluster RCT</td>
<td>2</td>
<td>ARI'</td>
<td>1,727 students in 42 childcare centres</td>
<td>1. Alcohol-based HS 2. Standard practice</td>
<td>ARI symptoms recorded.</td>
<td>1. ARI illness Significant risk reduction with intervention (HR 0.69 [0.57-0.83]).</td>
<td></td>
</tr>
<tr>
<td>Hubner et al.</td>
<td>2010</td>
<td>RCT</td>
<td>2</td>
<td>Common cold, sore throat, fever, cough, sinusitis, influenza</td>
<td>129 employees</td>
<td>1. Alcohol-based HS 2. Standard practice</td>
<td>Respiratory symptoms and days of work missed recorded.</td>
<td>1. Odds of common cold 2. Days of work missed</td>
<td>HS reduced odds (OR 0.35 [0.17-0.71]) but not days absent (OR 0.50 [0.22-1.17]).</td>
</tr>
<tr>
<td>Ram et al.</td>
<td>2015</td>
<td>RCT</td>
<td>2</td>
<td>ILLP</td>
<td>3,421 participants within 377 households</td>
<td>1. Handwashing 2. Standard practice</td>
<td>Daily surveillance and collection of oropharyngeal specimens from contacts.</td>
<td>1. SAR among household contacts No difference in SARs for ILL (ratio 1.24 [0.92-1.65]) or influenza (2.4 [0.68-8.47]).</td>
<td></td>
</tr>
<tr>
<td>Nicholson et al.</td>
<td>2014</td>
<td>Cluster RCT</td>
<td>2</td>
<td>ARI'</td>
<td>11,725 participants within 70 neighbourhoods</td>
<td>1. Education + plain bar soap 2. Standard practice</td>
<td>Illnesses and school absences recorded.</td>
<td>1. ARI incidence rate 2. School absenteeism Increased risk of illness (by 15%) and school absence (by 31%) without soap use.</td>
<td></td>
</tr>
<tr>
<td>Slayton et al.</td>
<td>2016</td>
<td>Cluster RCT</td>
<td>2</td>
<td>ARI (no definition provided)</td>
<td>369 participants within 33 geographical areas</td>
<td>1. Education + antimicrobial hand towels 2. Education</td>
<td>Information on illness in the past 48 hours collected.</td>
<td>1. ARI illness rate per 100 person-visits No significant difference between intervention (1.38) vs. control (1.48).</td>
<td></td>
</tr>
<tr>
<td>Shafi-que et al.</td>
<td>2016</td>
<td>Cluster RCT</td>
<td>2</td>
<td>URI'</td>
<td>227 participants among 48 clusters</td>
<td>1. Benzalkonium chloride-based HS 2. Education</td>
<td>URI symptoms recorded daily.</td>
<td>1. Incidence of URI No significant difference between HS (30.2%) and control (34.4%).</td>
<td></td>
</tr>
<tr>
<td>Morton et al.</td>
<td>2004</td>
<td>Cluster randomized study</td>
<td>2</td>
<td>Symptoms of URI, such as nasal congestion, cough, or sore throat, with or without fever</td>
<td>253 children</td>
<td>1. Alcohol gel + handwashing 2. Handwashing</td>
<td>Crossover study, with 1-week washout period. Symptoms recorded.</td>
<td>1. School absenteeism for respiratory illness Alcohol gel reduced odds of absenteeism by 43%.</td>
<td></td>
</tr>
</tbody>
</table>

ARI: Acute respiratory illness, CI: confidence interval, d: day, HR: hazard ratio, HS: hand sanitizer, ILL: influenza-like illness, IRR: incidence rate ratio, OR: odds ratio, RCT: randomized controlled trial, RTI: respiratory tract infection, SAR: secondary attack rate/ratio, URI: upper respiratory infection, VRTI: viral respiratory tract infection. ^<2 years of age: Fever >38°C and ≥1 of the following: nasal discharge/congestion, cough, conjunctivitis, respiratory distress (tachypnea, retractions), sore throat, and new seizure; >2 years of age: fever >38°C and cough or sore throat in the absence of another explanation. ^At least two caregiver-reported symptoms for 1 day, or one of the following symptoms for 2 days (but not fever alone). ^Two of the following symptoms for 1 day or 1 of the following symptoms for 2 consecutive days, not including 2 consecutive days of cough alone, sneezing alone, or fever alone: runny nose, stuffy/blacked nose or noisy breathing, cough, feeling hot/feverish or having chills, sore throat, or sneezing. ^High temperature (>37.5°C), a respiratory symptom, and a systemic symptom. ^1) sudden appearance of symptoms; (2) at least 1 of the following 4: fever/feeling feverish, body aches, headache, muscle aches, and (3) at least 1 of the following 3: cough, sore throat, difficulty breathing; and (4) absence of other suspected diagnosis. ^Symptoms of fever (>37.3°C), cough and expectoration, rhinorrhoea and nasal congestion. ^≥2 of the following symptoms for ≥24 hours, lasting ≥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. ^<5 years old: fever; ≥5 years old: fever with cough or sore throat. Pneumonia, cough, fever, chest pain and shortness of breath, cold, inflammation of any or all of the airways. ^Reported symptoms of stuffy or runny nose.
duction in VRTI-associated absenteeism, as high as 34.6%, while the other did not (79). Three other trials (74-76) conducted in the community setting concurred that the effect of alcohol-based hand sanitizer use (combined with education) was weak. One study (77) that examined the use of an alcohol-free hand sanitizer alone found a significant reduction in VRTI-associated absenteeism (31.7%) in the intervention groups.

Two studies evaluating the effect of handwashing with soap compared with usual handwashing practices (control) provided conflicting findings. Hovi et al. (78) found that while the use of soap and water significantly lowered the incidence of acute gastrointestinal infections (AGI) among office employees, that of respiratory tract infections was not, potentially due to different transmission routes. In contrast, Talaat et al. (79) found significant reductions in both outcomes associated with AGI (33%) and VRTI (40%) in the intervention group. Of note, two studies (78,79) that compared the effect of antibacterial soap versus non-antibacterial soap on respiratory illness rates in community settings both found no significant difference.

Of the two studies that examined education alone, both observed positive results, reporting either a significant decrease in risk (80) or incidence (81) of reported VRTI among those in the intervention arms. When all three hand hygiene interventions (i.e. hand sanitizer use, soap use, and education) were employed in the experimental group, two studies found a significant reduction in school absenteeism due to respiratory illness relative to the control group, which followed usual handwashing procedures (82,83).

In addition to schools, implementation of hand hygiene interventions was studied in other settings, including childcare centres, workplaces, and households. In one cluster RCT, Roberts et al. (84) demonstrated high-quality evidence that the practice of hand hygiene in childcare centres significantly reduced the incidence of colds in children under 24 months of age and those who adhered best to the intervention. Two other trials provided evidence supporting its role in reducing VRTI incidence (85,86). In a small individually randomized trial carried out in the workplace, Hubner et al. (87) reported that employees who used an alcohol-based hand gel had a significantly lower risk of a cold compared with control. In contrast, evidence of a protective effect when implemented in households is lacking. One study (88) found no benefit of hand hygiene in reducing VRTI rates compared with education alone. To ascertain the effect of introducing hand hygiene interventions to prevent secondary household influenza transmission from an index case, three studies (89-91) were executed. All found that the secondary attack rate did not significantly differ across the intervention arms. This observation may be explained by the relatively short study period in all three trials (≥21 days), as participants require time to learn and adopt new hygiene behaviours before viral transmission can be interrupted. Furthermore, Sandora et al. (76) reported that even when the intervention was implemented prior to any index case, no demonstrable benefit in preventing secondary household transmission of respiratory tract illnesses was seen.

Lastly, studies conducted in urban and rural domestic settings produced vastly different outcomes. Two trials (79,92) performed in urban settlements provided high-quality evidence of reduced VRTIs in children after hand hygiene education and soap were introduced. Both studies promoted hand hygiene practices through regular group training sessions and household visits over a 10- to 12-month period. In contrast, two studies (93,94) conducted in rural settings observed no such reduction with intervention.

Summary: Hand hygiene

1. Aggregate grade of evidence: A
2. Benefit: Decreased VRTI
3. Harm: Minimal: inconvenience, time, dry skin on hands
4. Cost: Low (costs of soap and hand sanitizers)
5. Benefits-harm assessment: Preponderance of benefit over harm
6. Value judgments: Low cost intervention with significant improvements in outcome of interest, gives high value for intervention
7. Recommendation level: Strong recommendation
8. Intervention: Employ hand hygiene strategies in healthcare and community settings.

Table 11. Summary of evidence for interventions to reduce VRTI transmission.

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Aggregate grade of evidence</th>
<th>Balance of benefit to harm</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal washing</td>
<td>C</td>
<td>Preponderance of benefit to harm</td>
<td>Option</td>
</tr>
<tr>
<td>Gargling</td>
<td>C</td>
<td>Preponderance of benefit to harm</td>
<td>Option</td>
</tr>
<tr>
<td>Masks (community settings)</td>
<td>B</td>
<td>Preponderance of benefit to harm</td>
<td>Recommendation</td>
</tr>
<tr>
<td>Masks (healthcare settings)</td>
<td>A</td>
<td>Preponderance of benefit to harm</td>
<td>Strong recommendation</td>
</tr>
<tr>
<td>Other personal protective equipment (gloves, gowns, eye protection)</td>
<td>C</td>
<td>Preponderance of benefit to harm</td>
<td>Recommendation in healthcare settings</td>
</tr>
<tr>
<td>Social distancing</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Hand hygiene</td>
<td>A</td>
<td>Preponderance of benefit to harm</td>
<td>Strong recommendation</td>
</tr>
</tbody>
</table>
Discussion
This review provides evidence-based recommendations of five broad nonpharmacological interventions (Table 11) aimed to reduce upper VRTI transmission amid an ongoing respiratory virus pandemic, with an impending cold and flu season in the Northern Hemisphere. With potential spread of disease through both air and physical contact, the nonpharmacological options assessed herein may be particularly meaningful in situations where efficacious antivirals and/or vaccines are in short supply or non-existent.

Through the iterative process, mask use and hand hygiene have emerged as the most efficacious strategies to impede viral transmission. Their adoption in healthcare and community settings is supported by a high level of evidence, a favourable safety profile, and a preponderance of benefit over harm. However, as described in multiple included studies, their effectiveness is contingent upon user compliance. This review was unable to determine the optimal mask type for use in either setting, which represents a study limitation. Nevertheless, based upon the available evidence, cloth masks should not be used in healthcare settings when surgical masks or N95 respirators are available. In addition, although the evidence supporting the use of gloves, gowns, and eye protection is weak, there is no data to suggest significant harm, beyond added cost and potential scarcity. Due to their potential benefit in high-risk situations, donning gloves, gowns, and eye protection are recommended in healthcare settings.

Saline nasal washing and gargling are two low-cost interventions with minimal potential harm. Although the available evidence suggests potential benefit for both, the limited available data suggest that these behavioural strategies are an option. Consideration of the potential risks of nasal washing and gargling, such as Eustachian tube dysfunction symptoms and throat irritation, should be considered. Individuals who are more susceptible to VRTIs and can tolerate nasal saline irrigation and/or gargling may derive greater benefit from these interventions. Notably, in response to the COVID-19 pandemic, novel therapeutic and preventative strategies to combat SARS-CoV-2 are currently under investigation. Further research is required to ascertain the optimal dosing for prophylaxis and the associated risks of these interventions.

Although a summary of findings for social distancing measures is not possible due to limited data from one study, some points are worth noting, particularly in the COVID-19 era. During the early stages of an outbreak or pandemic, the goal of social distancing is to reduce and delay the peak attack rate, thereby allowing time to accumulate resources, distribute antivirals, and administer vaccines if available. In situations where antivirals and/or vaccines do not exist for a novel virus, such as the current predicament with SARS-CoV-2, these measures, which can be readily available and activated, may represent the first line of defence and play a more critical role in reducing the spread of the infectious virus.

This review is not intended to replace clinical judgment, but rather to aid clinicians, in particular rhinologists, in comprehending the available evidence and develop an evidence-based approach to identify efficacious, rapidly deployed NPIs to reduce viral transmission. Although the scope of this review is broad, we acknowledge that by limiting the inclusion criteria to studies with the highest level of evidence, emerging and anecdotal interventions with potentially meaningful clinical impact may have inadvertently been excluded. Furthermore, we chose to only examine studies that specified a clinical endpoint involving upper respiratory tract infections. Therefore, caution must be exercised when considering these recommendations in the context of preventing lower airway infections.

Conclusion
The five NPIs examined represent a small fraction of the armamentarium currently under investigation. The current pandemic has catalysed research efforts to reexamine existing methods and propose novel strategies to reduce viral exposure, such as modified protective equipment or topical nasal and oral applications of PVP-I. As the pandemic continues to unfold, these NPIs may serve as the only viable strategies to contain the outbreak until an efficacious vaccine is developed.

Acknowledgement
N/A

Authorship contribution
DAG, RJS, NRR: Concept and design; EY, JF, CS, EAB: Acquisition, analysis, or interpretation of data; EY, JF: Drafting of Manuscript; DAG, RJS, SAN, NRR: Critical revision of manuscript; DAG, RJS, SAN, NRR: Supervision.

Conflict of interest
RJS is a consultant for Stryker, Optinose, GSK, a medical director for Healthy Humming and has received grant support from Stryker, Optinose and Healthy Humming. NRR has received grant funding from Optinose US, Inc.

Financial disclosure
No funding has been received to support the development of this article.
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NPIs to reduce respiratory viral transmission


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APPENDIX

Appendix 1. Detailed search strategy.

PubMed (U.S. National Library of Medicine, National Institutes of Health) search strategy:


Scopus (Elsevier) search strategy:


CINAHL (EBSCOhost) search strategy:

(MH "Respiratory Tract Infections” OR respiratory OR URI OR VURI OR MH "Common Cold” OR MH "Influenza+” OR MH "Laryngitis+” OR MH "Group” OR "Laryngotracheitis” OR MH "Pharyngitis” OR nasopharyngitis OR MH "Rhinitis+" OR MH "Severe Acute Respiratory Syndrome” OR SARS OR MH "Sinusitis+” OR supraglottitis OR MH “Epiglottitis” OR tracheitis OR MH “Middle East Respiratory Syndrome Coronavirus”) OR (MH “Middle East Respiratory Syndrome” OR MERS OR MH "Coronavirus+” OR MH "Coronavirus Infections+” OR COVID-19 OR MH "Otitis Media+” OR MH "Mastoiditis” OR petrositis) AND (MH "Handwashing+” OR gargle) OR MH "Nasal Lavage” OR MH "Mouthwashes+” OR "social distancing” OR "physical distancing” OR MH "Social Isolations+” OR MH "Quarantine” OR "personal protective” OR MH "Protective Clothing+” OR MH "Masks” OR facemasks OR N95 OR MH "Respiratory Protective Devices” OR "respirator protection” OR MH "Gloves” AND (prevent* OR MH "Respiratory Tract Infections+/PC” OR MW "PC" OR risk OR transmission)) • Number of records identified: 51

Limiters: Publication Types: Meta-Analysis, Meta Synthesis, Systematic Review • Expanders: Apply equivalent subjects • Search modes: Boolean/Phrase • Note: The search strategy below has to be pasted one line at a time (one concept on each line). If the entire search strategy is pasted onto the top row of the search box, the results will not be accurate.