A Hierarchical Framework for Assessing Transmission Causality of Respiratory Viruses

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Abstract
We propose a hierarchical framework based on our experience of systematically reviewing and synthesizing 378 primary studies for an evidence-based update of the modes of transmission for SARS-CoV-2. These studies revealed significant methodological shortcomings with a lack of standardization in the design, conduct, testing and reporting of SARS-CoV-2 transmission. While this situation is in part excusable at the outset of a pandemic, evidence rules of proof for assessing the transmission of this virus are needed for this and future pandemics of viral respiratory pathogens. We review the history of causality assessment related to microbial etiologies with a focus on respiratory viruses and suggest a hierarchy of evidence to integrate clinical, epidemiologic, molecular and laboratory perspectives on transmission. The hierarchy, if applied to future studies, should narrow the uncertainty over the twin concepts of causality and transmission of human respiratory viruses. We attempt to address the translational gap between the current research evidence and the assessment of causality in the transmission of respiratory viruses with a focus on SARS-CoV-2. Experimentation, consistency and independent replication of research alongside our proposed framework provide a chain of evidence that can reduce the uncertainty over the transmission of respiratory viruses and increase the level of confidence in specific modes of transmission and the measures that should be undertaken to prevent transmission.

Keywords: viral transmission; causation; evidence hierarchy; SARS-CoV-2; respiratory pathogens
Introduction

“our increasing proficiency in demonstrating viruses has produced a disconcerting but not entirely unwelcome paradox—the spectacle of new information leading to confusion.”

Robert J. Huebner. 1957

The COVID-19 pandemic has necessitated widespread global action, and research to inform evidence-based policy decisions has been urgently needed. Understanding the circumstances of SARS-CoV-2 transmission has been a public health priority. Yet, a consensus is lacking, partly due to the suboptimal quality of the available research to date.\textsuperscript{1-6} The lack of high-quality evidence has highlighted the pressing need for a contemporary conceptual framework to assess causality in the transmission of respiratory viruses in humans, especially SARS-CoV-2.

The background to this work was a series of systematic reviews funded by the WHO on modes of transmission of SARS-CoV-2 including airborne, fomite, orofecal and the association of close contact with transmission.\textsuperscript{7-10} The reviews revealed significant methodological shortcomings in the included studies with a lack of standard methods in the design, conduct, testing and reporting of SARS-CoV-2 transmission and a consistent lack of proof of replicability of the results.

Here we do not argue for or against any particular mode of transmission. Rather, we briefly review the history of causality assessment related to microbial etiologies with a focus on viral etiologies and suggest a hierarchy of evidence to integrate clinical, epidemiologic, molecular and laboratory perspectives on transmission. The hierarchy, if applied to future studies, should narrow the uncertainty over the twin concepts of causality and transmission of human respiratory viruses. These distinct concepts have become intertwined by the advent of genomics, as our historical and evidenciary narrative will show.

From bacteriology to virology

The first formulation of principles of causality between a microbe and disease is that by the German physician and microbiologist\textsuperscript{11} (Table 1). Although dated, the postulates are an essential first attempt at setting out attribution rules and are the definitive move away from miasma theory prevalent previously.\textsuperscript{12}

Table 1. \textit{Koch and Loeffler 1884 postulates in their original formulation:}

1. The microorganism must be found in abundance in all organisms suffering from the disease, but should not be found in healthy organisms.
2. The microorganism must be isolated from a diseased organism and grown in pure culture.
3. The cultured microorganism should cause disease when introduced into a healthy organism.
4. The microorganism must be reisolated from the inoculated, diseased experimental host and identified as being identical to the original specific causative agent.

The next significant contribution was by the American bacteriologist and virologist Thomas Rivers, who in 1936 analysed the implications on the postulates in the emergent fields of virology and immunology.\textsuperscript{13} He formulated two simple postulates: 1) a specific virus must be
found associated with a disease with a degree of regularity, and 2) the virus must be shown to occur in the unwell individual not as an incidental or accidental finding but as the cause of the disease under investigation. Rivers addressed the historical shortcomings of Koch’s work - and their subsequent distortions - by recognising the viral agent may not necessarily be present each time a case of the particular “malady” was observed. The originality of Rivers’ contribution lay in separating causality from association and recognising the possibility of contamination or co-infection (a point he makes in an elegant discussion of the aetiology of encephalitis lethargica or Von Economo’s disease, and the coincidental finding of herpes virus in some of the slides from those cases)\(^\text{14}\).

As virology developed further, the issue of differentiating between causation and association led the American physician and virologist, Robert Huebner, in 1957, to issue a “Bill of Rights for Prevalent Viruses, a guarantee against the imputation of guilt by simple association”\(^\text{15}\). Huebner pointed out the paradox of the advancement of virological knowledge exemplified by noting, “our increasing proficiency in demonstrating viruses has produced a disconcerting but not entirely unwelcome paradox-the spectacle of new information leading to confusion”\(^\text{15}\).

Huebner asked the fundamental question of how can we tell what is causing what? He proposed eight points to assess viral causality (Table 2). The development of virological techniques is reflected in the detailed and multidisciplinary approach to the exclusion of a simple association. Each of his points per se is insufficient to prove or disprove anything. However, if taken in their entirety, they would narrow the chance of an association and strengthen the case for causality.

Table 2. Rober Huebner’s 1957 Bill of Rights for Prevalent Viruses\(^\text{15}\)

1) Isolation of a virus in culture;
2) Repeated recovery of the virus from human specimens;
3) Antibody response to the virus;
4) Characterization and comparison with known pathogenic viruses;
5) Constant association of the virus with specific illness;
6) Reproduction of clinical illness in volunteer challenge studies;
7) Epidemiologic studies (with controlled longitudinal studies offering the greatest value); and
8) Prevention of disease by vaccination.

In 1973, the Yale epidemiologist Alfred Evans reformulated Rivers’ postulates based on up to date immunology theory and the discovery of antigens and listed seven immunologic criteria, related to proof of causality which complemented Rivers' postulates\(^\text{16}\). Meanwhile, the statistician Austin Bradford Hill in 1965 had introduced what he called criteria for causality in another President’s address\(^\text{17}\).

Hill’s criteria
Hill, a British statistician by training - famous for his work on tobacco and lung cancer - proposed nine criteria to assess the nature of the association between two variables: Strength, Consistency, Specificity, Temporality, Biological gradient, Plausibility, Coherence, Experiment, and Analogy\(^\text{17}\). Hill stated there was no magic formula to assess causality and his criteria were a guide or “viewpoints” and did not provide “indisputable evidence for or against the cause-and-effect hypothesis.” A degree of subjectivity was necessary in interpreting the evidence and no viewpoint was a “sine qua non”.


Gwaltney and Hendley postulates
In 1978 Jack Gwaltney and J Owen Hendley, based at the University Of Virginia, proposed postulates based on their work with rhinovirus and other respiratory viruses to address respiratory virus causality (Table 3)\(^\text{18}\). The first four postulates can be considered a practical application of Hill’s criteria of strength and temporality, but the fifth is Hill’s experiment: “because of an observed association some preventive action is taken. Does it in fact prevent? Is the frequency of the associated events affected?”\(^\text{18}\)

<table>
<thead>
<tr>
<th>Postulate Number One</th>
<th>Microbial growth at the proposed anatomic site of origin.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Postulate Number Two</td>
<td>Microbes present in secretions or tissues shed from the site of origin.</td>
</tr>
<tr>
<td>Postulate Number Three</td>
<td>Microbes contaminate and survive in or on environmental substance or object.</td>
</tr>
<tr>
<td>Postulate Number Four</td>
<td>Contaminated substance or object reaches portal of entry of new host.</td>
</tr>
<tr>
<td>Postulate Number Five</td>
<td>Interruption of transmission by hypothesized route reduces incidence of natural infection</td>
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</tbody>
</table>

The postulates present the first convergence of causality and transmission as they represent a detailed checklist of the chain of transmission of respiratory viruses and require an experiment to test whether the hypothesised mode of transmission can be interrupted.

The Gwaltney-Hendley postulates, however, omit Hill’s criteria of “specificity” and “consistency.” Specificity is a particular problem concerning viral respiratory infections. Since the clinical presentation cannot distinguish those viral infections that present with similar clinical features, laboratory identification of the responsible agent becomes necessary. Evans pointed this out in 1991 in his “Five Realities of Acute Respiratory Disease” (Table 4)\(^\text{16}\). As any practising physician knows, it is impossible to distinguish with absolute confidence an acute respiratory infection caused by rhinovirus from one caused by adenovirus, or indeed to be able to rule out any co-infection with two or more viruses, based on clinical presentation alone. Proof requires microbiological identification and evidence for the chain of transmission, which is now possible with the advent of genomics.\(^\text{19}\)

Table 4. Evans’ 1991 list of “Five Realities of Acute Respiratory Disease”\(^\text{16}\)

1) the same syndrome could be produced by several agents;
2) The same virus could produce several clinical syndromes;
3) the cause of the syndrome varied by geographic area, age, and other factors;
4) the causes of only about half of the common acute respiratory and intestinal syndromes and of about one-quarter of acute viral infections of the central nervous system have been identified; and
5) diagnosis of the etiological agent could rarely be made on clinical grounds alone and required laboratory methods such as isolation of the virus and/or demonstration of an antibody response.
The genomics era
As knowledge of genetics grew, Fredricks and Relman in 1996 proposed reformulation of the original Koch’s postulates incorporating the state of knowledge at the time (Table 5). This list contains many previously raised points, and crucially, includes the 7th postulate: the issue of consistency or reproducibility of results. Further reformulations were proposed by Lipkin in 2013 in the Levels of Certainty in pathogen discovery and Byrd and Segre in 2016 to advocate genetic sequencing of all members of a “microbial community” for classification and definitive identification.

Table 5. Fredricks & Relman’s 1996 proposed reformulation of the original Koch’s postulates

1) A nucleic acid sequence belonging to a putative pathogen should be present in most cases of an infectious disease. Microbial nucleic acids should be found preferentially in those organs or gross anatomic sites known to be diseased (i.e., with anatomic, histologic, chemical, or clinical evidence of pathology) and not in those organs that lack pathology.
2) Fewer, or no, copy numbers of pathogen-associated nucleic acid sequences should occur in hosts or tissues without disease.
3) With resolution of disease (for example, with clinically effective treatment), the copy number of pathogen-associated nucleic acid sequences should decrease or become undetectable. With clinical relapse, the opposite should occur.
4) When sequence detection predates disease, or sequence copy number correlates with severity of disease or pathology, the sequence-disease association is more likely to be a causal relationship.
5) The nature of the microorganism inferred from the available sequence should be consistent with the known biological characteristics of that group of organisms. When phenotypes (e.g., pathology, microbial morphology, and clinical features) are predicted by sequence-based phylogenetic relationships, the meaningfulness of the sequence is enhanced.
6) Tissue-sequence correlates should be sought at the cellular level: efforts should be made to demonstrate specific in situ hybridization of microbial sequence to areas of tissue pathology and to visible microorganisms or to areas where microorganisms are presumed to be located.
7) These sequence-based forms of evidence for microbial causation should be reproducible.

A Proposed Framework for assessing transmission causality of respiratory viruses

Certainty (or diminution of uncertainty) is essential for all Gwaltney’s postulates, especially on the crucial issues of correct identification of those who have an active infection and are contagious, i.e. capable of infecting contacts. This simple concept has been lost in a plethora of information overload of hurried low-quality studies in the current pandemic (in line with Huebner’s prescient observations). Clinical details of the exposed patient and a chain of transmission with laboratory confirmation of active infection in the exposed susceptible person are necessary to reduce the uncertainty over transmission.

Across 378 studies, we identified laboratory confirmation of active infection as a critical result for interpreting the circumstances and finality of transmission. However, our reviews show very heterogeneous methods of confirmation with low quality of evidence: from binary PCR to serial viral culture done with little context and even less rigour, reported in the studies to assess the transmission. Few studies reported testing within a clinical context. For example, time from onset of symptoms and their severity are intimately tied to the likelihood of contagiousness. Another review of studies of viral cultures for SARS-CoV-2 infectious
potential assessment found that only 11/29 studies reported date of symptom onset in relation to the date of specimen collection.

To efficiently use all possible information and reduce uncertainty over the interpretation of the mode of transmission, we built a four-level hierarchical structure, similar to the levels of evidence used in evidence-based medicine, and carefully building on the work of the many scientists mentioned above who have contributed so much to causality and transmission.

**Level 1**: Binary PCR or nucleic acid antigen result in the absence of clinical data.
Lack of agent specificity means there are often no distinctive signs or symptoms readily attributable to a specific respiratory virus: reporting only PCR or other nucleic acid amplification tests as binary positive/negative does not identify current active infection. At the lowest level of confirmation, residual non-infectious RNA or DNA of the responsible viruses can be harboured and shed for weeks to months.

**Level 2**: Quantitative PCR result
Adding further information, such as log concentrations of virus or PCR cycle threshold (Ct) values, reduces some of the uncertainty for identifying those with active infection but is not definitive. Current evidence shows that the lower the Ct representing a higher concentration of virus particles, the more likely a "positive" has infectious potential. Contagiousness is also highest during the first few days from the onset of symptoms for many respiratory viruses, including SARS-CoV-2. In addition, immune-suppressing treatments or conditions may prolong infectiousness time. Underpinning why we need clinical anamnestic data to interpret the results of PCR.

**Level 3**: Cytopathic Effects (CPE)
More certainty is provided by cell culture and observation of the effects of inoculation of specimen washouts on the cells. Observed structural changes in host cells caused by viral invasion leading to visible cell lysis and other cytopathic phenomena are further indicative of viral replication. Clinically the patient may be still symptomatic, but the viral concentration may be too low to infect, and the CPE may be classed as weak. Therefore, coinfection or contamination cannot be fully ruled out. For example, Lednicky et al. reported general virus-induced CPE within two days post-inoculation of specimens from a COVID-19 patient. However, PCR tests for SARS-CoV-2 vRNA from the cell culture were negative; three other respiratory viruses were identified: Influenza A H1N1, H3N2, and human coronavirus OC43. The exclusion of contaminants or co-infections in the plaques and serial culturing of similar specimens with the same results across different laboratories is required to reduce uncertainty further.

**Level 4**: Serial Viral Culture and genome sequencing
Immunohistochemistry directed at specific viruses and genome sequencing excludes co-infection and indicates a higher probability of the correct identification of the agent and viral similarity between donor and recipient, which are crucial for determining proof of transmission. Here again, accurate contact history and the clinical picture will provide chronological evidence of the likely chain of transmission. Patients within 5-7 days of symptom onset, severe symptoms or pre-existing pathologies fall in this category. Sequencing does not require the presence of a viable virus, so only its coupling with repeated serial cultures will diminish the uncertainty around both transmission and
causation. The presence of replicating virus, the absence of co-infection, and the similarity of the viral lineage of the presumed source gives rise to the strongest evidence of transmission and causation.

To prove transmission from one person to another requires two viral sequences to share an otherwise statistically improbable number of identical mutations. Genome sequences are required to exclude the possibility that the second person was infected by another circulating virus from elsewhere in the environment. The quantity of genome needing sequencing depends upon the spontaneous mutation rate (the rate of drift of the sequences). Ideally, these would be “silent” mutations (random sequence noise) unconnected to phenotypes subject to selection (for example, by the administration of antivirals).

To document how the second person got infected requires evidence they were exposed in their environment (the route) and that the source was sufficiently contaminated (low Ct) with infectious material (cultivatable virus) to transmit an infection to another human. Availability of all such evidence provides a high standard of proof of transmission in the age of genomics.

Our hierarchical Framework for Assessing Transmission Causality of Respiratory Viruses is synthesized in Figure 1.

The hierarchy allows an understanding of transmission evidence across our reviews and its uncertainties. Across our (four) reviews, the results interpreted by using our proposed framework show a lack of higher-level evidence. Overall, we found that none of the 378 studies reached the level of serial viral culture; of the 276 studies which used PCR as a diagnostic tool, 79 reported a cycle threshold (Table 6).
Table 6. Virological and genomic evidence reported in 378 studies included in 4 systematic reviews of transmission causality of SARS-CoV-2. Key: CT=Cycle threshold; CPE=cytopathic effect

<table>
<thead>
<tr>
<th>Review</th>
<th>Transmission causality</th>
<th>Primary studies</th>
<th>Report PCR result (% of studies)</th>
<th>Report Cycle threshold (% of studies)</th>
<th>CT&lt;25 (% of studies)</th>
<th>Attempted Viral Culture (% of studies)</th>
<th>CPE (% of studies)</th>
<th>Genome Sequencing (% of studies)</th>
<th>Serial viral culture positive (% of studies)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Airborne Transmission in COVID-19: a systematic review</td>
<td>Airborne</td>
<td>67</td>
<td>53 (79.1%)</td>
<td>12 (17.9%)</td>
<td>2 (3.0%)</td>
<td>10 (14.9%)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>SARS-CoV-2 and the Role of Fomite Transmission: Systematic Review</td>
<td>Fomites</td>
<td>63</td>
<td>51 (81.0%)</td>
<td>13 (20.6%)</td>
<td>3 (4.8%)</td>
<td>11 (17.5%)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>SARS-CoV-2 and the Role of Orofecoal Transmission: Systematic Review</td>
<td>Orofecoal</td>
<td>77</td>
<td>46 (59.7%)</td>
<td>22 (28.6%)</td>
<td>7 (9.1%)</td>
<td>6 (7.8%)</td>
<td>1*</td>
<td>1</td>
<td>0**</td>
</tr>
<tr>
<td>SARS-CoV-2 and the Role of Close Contact in Transmission: A Systematic Review</td>
<td>Close contact</td>
<td>171</td>
<td>126 (73.7%)</td>
<td>32 (18.7%)</td>
<td>5 (2.9%)</td>
<td>3 (1.8%)</td>
<td>2</td>
<td>10</td>
<td>2</td>
</tr>
<tr>
<td>% of primary studies</td>
<td>378</td>
<td>276 (73.0%)</td>
<td>79 (20.9%)</td>
<td>17 (4.5%)</td>
<td>30 (7.9%)</td>
<td>3 (0.3%)</td>
<td>11</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
- *CPE does not show plaques and is not immunostained,
- **Isolated the virus from the stools of one severe hospitalised COVID-19 case. Evidence of infectious material unclear.

One study observed presumed virus-induced CPE.
Forward
Assigning causation relies on a consistent body of evidence which excludes other biologically plausible alternative explanations for the events or makes them unlikely. Experimentation and consistency alongside our proposed framework provide a chain of evidence that can reduce the uncertainty over the transmission of respiratory viruses and increase the level of confidence in specific modes of transmission. The addition of a sixth postulate to Gwaltney’s work requires the validation of the transmission chain by independent replication of results. However, experimentation in the case of SARS-CoV-2 is unlikely to be ethical unless an intervention to interrupt the chain of transmission is tested and challenge studies are ethically difficult to justify.

Evidence that follows Bradford Hill’s principles for assessing causation: Strength, Temporality and Experimentation (Gwaltney’s fifth postulate) along with independent replication and the hierarchy of transmission outcome events provides a coherent framework for assessing causation for the transmission of respiratory viruses. Continued uncertainty and lack of understanding of the major mode(s) of transmission and their respective frequency in any given setting with respect to SARS-CoV-2, underscores the critical importance of applying a framework for assessing causality in transmission of respiratory viruses. Given the current global COVID-19 pandemic and that a year since its historic arrival as a “modern day plague” has gone by, we still do not have the fundamental questions definitively answered regarding the mode(s) and frequency of transmission, nor the minimal infective dose or the serological correlates of infection. It is imperative to provide high quality evidence so that optimal intervention measures may be applied to achieve prevention of lives lost and protection against global economic and social demise.

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Authors’ contributions
All authors contributed in equal part to the conceptualisation and development of the content. TJ and CH wrote the first draft. All authors contributed to the subsequent drafts and approved the final version.

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Conflict of interest statements
TJ was in receipt of a Cochrane Methods Innovations Fund grant to develop guidance on the use of regulatory data in Cochrane reviews (2015-018). In 2014–2016, he was a member of three advisory boards for Boehringer Ingelheim. TJ was a member of an independent data monitoring committee for a Sanofi Pasteur clinical trial on an influenza vaccine. TJ is occasionally interviewed by market research companies about phase I or II pharmaceutical products for which he receives fees (current). TJ was a member of three advisory boards for Boehringer Ingelheim (2014-16). TJ was a member of an independent data monitoring committee for a Sanofi Pasteur clinical trial on an influenza vaccine (2015-2017). TJ is a relator in a False Claims Act lawsuit on behalf of the United States that involves
sales of Tamiflu for pandemic stockpiling. If resolved in the United States favor, he would be entitled to a percentage of the recovery. TJ is coholder of a Laura and John Arnold Foundation grant for development of a RIAT support centre (2017-2020) and Jean Monnet Network Grant, 2017-2020 for The Jean Monnet Health Law and Policy Network. TJ is an unpaid collaborator to the project Beyond Transparency in Pharmaceutical Research and Regulation led by Dalhousie University and funded by the Canadian Institutes of Health Research (2018-2022). TJ consulted for Illumina LLC on next generation gene sequencing (2019-2020). TJ was the consultant scientific coordinator for the HTA Medical Technology programme of the Agenzia per i Servizi Sanitari Nazionali (AGENAS) of the Italian MoH (2007-2019). TJ is Director Medical Affairs for BC Solutions, a market access company for medical devices in Europe. TJ is funded by NIHR UK and the World Health Organization (WHO) to update Cochrane review A122, Physical Interventions to interrupt the spread of respiratory viruses. TJ is funded by Oxford University to carry out a living review on the transmission epidemiology of COVID-19. Since 2020, TJ receives fees for articles published by The Spectator and other media outlets. TJ is part of a review group carrying out Living rapid literature review on the modes of transmission of SARS-CoV-2 (WHO Registration 2020/1077093-0). He is a member of the WHO COVID-19 Infection Prevention and Control Research Working Group.

CJH holds grant funding from the NIHR, the NIHR School of Primary Care Research, the NIHR BRC Oxford and the World Health Organization for a series of Living rapid review on the modes of transmission of SARs-CoV-2 reference WHO registration No2020/1077093. He has received financial remuneration from an asbestos case and given legal advice on mesh and hormone pregnancy tests cases. He has received expenses and fees for his media work including occasional payments from BBC Radio 4 Inside Health and The Spectator. He receives expenses for teaching EBM and is also paid for his GP work in NHS out of hours (contract Oxford Health NHS Foundation Trust). He has also received income from the publication of a series of toolkit books and for appraising treatment recommendations in non-NHS settings. He is Director of CEBM and is an NIHR Senior Investigator.

DE holds awards from the Canadian Institutes for Health Research and the Natural Sciences and Engineering Research Council of Canada to support poxvirus research as well as research relating to Covid-19 vaccine development and aerosol transmission. Funding from Alberta Health Services, Alberta Innovates, and the World Health Organization supports research concerning the mechanism of Covid-19 transmission and inactivation technologies. He also holds research contracts and consults with Tonix Pharmaceuticals in support of the development of poxvirus-based Covid-19 vaccines.

JMC holds grants from the Canadian Institutes for Health Research on acute and primary care preparedness for COVID-19 in Alberta, Canada 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 also received support from the Centers for Disease Control and Prevention (CDC) to attend an Infection Control Think Tank Meeting. He is a member of the WHO Infection Prevention and Control Research and Development Expert Group for COVID-19 and the WHO Health Emergencies Programme (WHE) Ad-hoc COVID-19 IPC Guidance Development Group, both of which provide multidisciplinary advice to the WHO and for which no funding is received.

IJO, EAS, JB, and AP have no interests to disclose.

**Ethics committee approval.**

No approval was necessary

**Data Availability**

All data included in the review are provided in the tables and text.
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