Reducing risks from coronavirus transmission in the home—the role of viral load

Paul Little and colleagues call for better promotion of simple measures that can help reduce the spread and severity of infection among those living with people who have covid-19

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Most people with covid-19 are cared for at home, increasing the likely exposure of household members. Although the evidence is limited, high infection rates among health workers have been attributed to more frequent contact with infected patients, and higher viral load—the size of the infecting dose of virus. This has led to demands for better personal protection equipment (PPE). Less attention, however, has been given to family members and others caring for people with covid-19 in the community. Providing them with the same level of PPE as in hospitals is not practicable, but promotion of simple evidence based interventions may lower the risk of infection transmission and help reduce morbidity and demand on hospitals.

Transmission in home and community

The long incubation and high presymptomatic infectivity of covid-19 makes transmission between family members a particular risk. Modelling of viral shedding in 94 patients with covid-19 and 77 transmission pairs suggests that the highest viral load is at or just before symptom onset, with 44% of transmission occurring before symptoms.1

Public health advice recommends isolation of symptomatic household members, but this can be difficult, particularly in small flats with shared facilities. Motivation to overcome these difficulties may not be high enough if members of the public are sceptical about reducing transmission in the home and unaware that the illness of other family members may be more severe if they do not reduce their level of exposure.

The medical community is commendably reluctant to make recommendations in the absence of evidence. An expert team that reviewed the evidence for viral load concluded that until the evidence is more conclusive: “As our grandparents used to say, when you do not know what is going on, do nothing.”2 However, given that measures to reduce the viral load from exposure to symptomatic household members have little risk of harm, the precautionary principle suggests that we should be promoting them. It is difficult to get good dose-response data, but it seems prudent not to treat absence of direct evidence in the pandemic as evidence of absence, particularly given evidence from other viral infections from both animal and human models, and trial evidence for reducing the transmission among families in non-pandemic years. Greater awareness of the potential risk from viral load may help motivate family members to pay sufficient attention to protecting themselves despite their concern for sick family members.

Evidence for viral load effect

It is intuitive that viral load should influence the incidence and severity of disease. The main problem is that measuring the viable infecting dose of the virus in people is extremely challenging: contemporary measures of viral density, viability, and viral contamination are all hard to obtain. Establishing the relation between infecting dose and the likelihood of developing disease is therefore difficult.

The challenge of establishing the infecting dose is complicated by environmental contamination. Experience with Middle East respiratory syndrome (MERS), caused by another coronavirus (MERS-CoV), suggests that environmental contamination with SARS-CoV-2 is likely to be high,3 and this is supported by recent case reports of extensive environmental contamination from patients with covid-19.4 Indirect evidence, including from
animal models and epidemiological studies also provides support.

### Animal models

Although the infecting dose from a combination of droplets and environmental contamination cannot be easily measured, high quality experiments under controlled conditions in animal models can provide indirect evidence. We are not aware of infecting dose experiments with animal models of covid-19, but animal models of other viral infections show that variation in the infecting dose determines how many animals get infected and how severe the illness is.

A model of African swine flu virus shows a clear dose-response relation between the infecting dose and disease in the animal. Likewise, a strong dose-response effect is found in the animal model for haemopoietic necrosis virus. A dose-response has also been shown in mouse models with several strains of SARS-CoV-1, which is closely related to the virus causing SARS-CoV-2. The infectivity varies between different strains of the virus, which modifies the shape of the dose-response curve, but nevertheless consistent dose-response relations are observed with the severity of the infection.

### Defective viral genomes

One of the key factors in determining how severe an infection becomes is also the extent to which defective viral genomes are produced. These effectively reduce the infecting doses during the early part of an infection by competing with non-defective genomes. The greater the abundance of viruses with defective genomes within an infecting inoculum, the better the clinical outcome: genomic analysis of viruses isolated from previously healthy people requiring admission to the intensive care unit with influenza A infection, those not requiring intensive care, and those who died (who also had underlying medical conditions) showed that defective genomes were associated with fewer severe or fatal outcomes.

### Epidemiology of serious viral infections

The evidence from other similar serious viral infections also suggests the infecting viral load may be important. A retrospective study of survivors of Ebola from the Kerry Town treatment centre in Sierra Leone, investigated disease in more than 933 family members (those who had died, those surviving, and those not infected). The severity of infecting dose was graded according to the history of exposure. Although there was no clear relation with mortality, perhaps because of the mediating effect of treatment at the treatment centre, the study found a very strong linear relation with the likelihood of infection developing, ranging from 80% likelihood of getting the disease with the highest infecting dose (direct contact with body of someone who had died) and 10% with the lowest dose (no contact).

During the 2003 SARS outbreak older age, comorbidities (adjusted hazard ratio (HR) 3.36, 95% confidence interval 1.44 to 7.82), and higher initial viral levels in nasopharyngeal specimens (adjusted HR 1.21 per log increase in number of RNA copies/mL, 95% CI 1.06 to 1.39) were associated with worse survival. Viral level 10 days after the onset of symptoms was associated with a series of poor clinical markers (oxygen desaturation, mechanical ventilation) and death. Recent data from covid-19 have shown that those with severe infections had viral levels 60 times higher at presentation than those with mild disease. Although the levels of virus once the disease has started will be in part a function of the immune response of the patient, the size of the initial viral load is likely to be a contributing factor, allowing immune defences to be more easily over-run.

The difference in case fatality rates in the three waves of the Spanish flu pandemic of 1918-19 can be explained by the number of simultaneous contacts a susceptible person had with infected people (the more contact the higher the infectious doses). However, in a detailed study modelling influenza virus transmission within households in 2008-12, infectivity was proportional to viral load but viral load alone provided a poor fit to the models. Clearly we need to better understand the relation between infecting dose and other prognostic factors in modifying the immune response and clinical outcome (age, comorbidity, etc).

### Pragmatic evidence that could help carers

We are aware of only one behavioural intervention that is proved to reduce virus transmission within households and is suitable for rapid dissemination in a pandemic. Two of us (PL and LY) were involved in a randomised trial of Germ Defence, a website that provides advice on infection control measures and helps users think about when and how to carry out key infection control behaviours such as handwashing and cleaning, avoiding sharing rooms and surfaces, managing incoming deliveries, and ventilating rooms. This could supplement public health advice on infection control in the home since it uses behaviour change techniques to help people implement this advice (box 1).

Germ Defence was trialled in 20 066 people during the H1N1 pandemic and subsequent seasonal flu years and reduced the number of respiratory infections (mean number of infections 0.84 y 1.09 in the control group, hazard ratio 0.75, 95% confidence intervals 0.72 to 0.79). Infection transmission among family members was lower in the intervention group (hazard ratio 0.79, 95% confidence interval 0.74 to 0.83), and there was a modest reduction in severity of infections (median number of days of moderately bad illness 3.9 (median 2 days) in intervention versus 4.5 (3 days) in the control group). Reductions were also observed in gastrointestinal infections, GP consultations, and antibiotic prescriptions.

The team has been funded by UK Research and Innovation to adapt this intervention for covid-19 and disseminate it nationally and internationally. It has already been translated into over 20 languages for this purpose (see www.germdefence.org/). Germ Defence may help limit transmission of covid-19 as well as the other viruses that are still causing the majority of respiratory illnesses in the current pandemic, even in secondary care settings. Other viruses may also be important given recent evidence that coinfection with other viruses occurs in more than 20% of cases.
Conclusion

Care is needed when extrapolating evidence from other disease, but viral load is likely to be important for covid-19. The precautionary principle suggests that people caring for household members who are unwell should be encouraged to take measures to reduce infecting viral load in order to reduce the incidence and severity of infection. Promoting infection control measures in the community is a priority for the UK government and will continue to be so as “stay at home” policies are lifted. Dissemination of evidence based behavioural interventions may help increase adoption of public health advice and reduce viral load.

Key messages

Government policy is aimed at reducing transmission of covid-19 between family units, but less attention has been given to transmission between family members.

Evidence from controlled experiments in animal models, viral genome studies, and other epidemics suggests the infecting viral load may be important.

A web based intervention has been shown to reduce incidence, transmission, and severity of seasonal flu.

Use of such behavioural interventions could support public health advice to improve infection control in families.

Contributors and sources: PL is a clinical GP academic with expertise in infections in primary care and the community. LY is a behavioural scientist with expertise in digital support for illness management. RCR is an infectious disease physician. TC has a split role as head of behavioural insights and evaluation lead at Public Health England and expert analyst in the Systems Unit at the Cabinet Office. Since surviving a stroke, CR has undertaken public involvement in a wide range of health research. JB is a public adviser to the covid-19 Germ Defence project, a member of the Healthcare Infection Society and led the public involvement in the evaluation of the implementation of the UK’s antimicrobial resistance strategy. RA is coeditor of the NIHR Health Protection Research Unit in Behavioural Science and Evaluation of Interventions and a visiting professor of practice in the RA is codirector of the NIHR Health Protection Research Unit in Behavioural trial of Germ Defence funded by the Medical Research Council. All other authors are part of the team that is expanding and disseminating Germ Defence for covid-19, which LY is leading.

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