RESEARCH ARTICLE

Environmental screening for SARS-CoV-2 in long term care facilities: lessons from a pilot study [version 1; peer review: 1 not approved]

Rachel Kwiatkowska1-3, Nicola Yaxley4, Ginny Moore4, Allan Bennett4, Matthew Donati5,6, Nicola Love7, Roberto Vivancos8,9, Matthew Hickman2,3, Derren R Ready1,3,10

1Field Service, National Infection Service, Public Health England, Bristol, UK
2Population Health Sciences, University of Bristol, Bristol, UK
3NIHR Health Protection Research Unit in Behavioural Science and Evaluation, University of Bristol, Bristol, UK
5Severn Infection Sciences, North Bristol NHS Trust, Bristol, UK
6Public Health England South West Regional Laboratory, National Infection Service, Bristol, UK
7Field Service, National Infection Service, Public Health England, Newcastle, UK
8NIHR Health Protection Research Unit in Gastrointestinal Infections, University of Liverpool, Liverpool, UK
10Eastman Dental Institute, University College London, London, UK

Abstract

Background: The SARS-CoV-2 pandemic has highlighted the risk of infection transmission in long-term care facilities (LTCF) and the vulnerability of resident populations. It is essential to understand the environmental spread of the virus and risk of indirect transmission to inform Infection Prevention and Control (IPC) measures in these settings.

Methods: Upon notification of SARS-CoV-2 outbreaks, LTCF within a local authority in the South West of England were approached to take part in this pilot study. Investigators visited to swab common touchpoints and elevated ‘non-touch’ surfaces and samples were analysed for presence of SARS-CoV-2 genetic material (RNA). Data were collected regarding LTCF infrastructure, staff behaviours, clinical and epidemiological risk factors for infection (staff and residents), and IPC measures.

Criteria for success were: recruitment of three LTCF; detection of SARS-COV-2 RNA; variation in proportion of SARS-CoV-2 positive surfaces by sampling zone; potential to assess infection risk from SARS-CoV-2 positive surfaces.

Results: Three LTCFs were recruited, ranging in size and resident demographics. Outbreaks lasted 63, 50 and 30 days with resident attack rates of 53%, 40% and 8%, respectively. The proportion of
sample sites on which SARS-CoV-2 was detected was highest in rooms occupied by infected residents and varied elsewhere in the LTCF, with low levels in a facility implementing enhanced IPC measures. The heterogeneity of settings and difficulty obtaining data made it difficult to assess association between environmental contamination and infection. Elevated surfaces were more likely to test positive for SARS-CoV-2 RNA than common touch-points.

**Conclusions:** SARS-CoV-2 RNA can be detected in a variety of LTCF outbreak settings. We identified variation in environmental spread which could be associated with implementation of IPC measures, though we were unable to assess the impact on infection risk. Sampling elevated surfaces could add to ongoing public health surveillance for SARS-CoV-2 and other airborne pathogens in LTCF.

**Keywords**
infection control; infectious disease transmission; environmental exposure; fomites; disease outbreaks; long-term care; epidemiologic methods

---

**Corresponding author:** Rachel Kwiatkowska (rachel.kwiatkowska@bristol.ac.uk)

**Author roles:** **Kwiatkowska R:** Conceptualization, Data Curation, Formal Analysis, Investigation, Methodology, Writing – Original Draft Preparation, Writing – Review & Editing; **Yaxley N:** Formal Analysis, Methodology, Writing – Review & Editing; **Moore G:** Conceptualization, Formal Analysis, Investigation, Methodology, Resources, Supervision, Writing – Review & Editing; **Bennett A:** Conceptualization, Methodology, Supervision, Writing – Review & Editing; **Donati M:** Methodology, Resources, Writing – Review & Editing; **Vivancos R:** Methodology, Supervision, Writing – Review & Editing; **Hickman M:** Methodology, Supervision, Writing – Review & Editing; **Ready DR:** Conceptualization, Investigation, Methodology, Resources, Supervision, Writing – Original Draft Preparation, Writing – Review & Editing

**Competing interests:** No competing interests were disclosed.

**Grant information:** RK is funded through the Wellcome GW4 Clinical Academic Training programme [203918]

The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Copyright:** © 2021 Kwiatkowska R et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. The author(s) is/are employees of the US Government and therefore domestic copyright protection in USA does not apply to this work. The work may be protected under the copyright laws of other jurisdictions when used in those jurisdictions.

**How to cite this article:** Kwiatkowska R, Yaxley N, Moore G *et al.* Environmental screening for SARS-CoV-2 in long term care facilities: lessons from a pilot study [version 1; peer review: 1 not approved] Wellcome Open Research 2021, 6:235 https://doi.org/10.12688/wellcomeopenres.17047.1

**First published:** 15 Sep 2021, 6:235 https://doi.org/10.12688/wellcomeopenres.17047.1
Introduction

Long term care facilities (LTCF) are inadvertently ideal environments for the spread of pathogens. (Strausbaugh et al., 2003) Residents are often susceptible to infection or colonisation, and in frequent and close contact with staff who have links to the wider community. Outbreaks of infectious diseases are common in these settings (Inns et al., 2017; Inns et al., 2019) and the coronavirus 2019 (COVID-19) pandemic, caused by SARS-CoV-2 coronavirus, has highlighted the vulnerability of people in LTCF to infectious disease threats: there were an estimated 29,542 excess deaths among LTCF residents over the first 23 weeks of the epidemic in England. (Morciano et al., 2021)

If detected early enough transmission of pathogens within the LTCF can be curbed, (Inns et al., 2018) however SARS-CoV-2 infections are often asymptomatic or paucisymptomatic leading to large outbreaks. Regular testing of residents and staff helps identify cases early but is resource-intensive and unpleasant for frail individuals, so non-invasive surveillance strategies may be more sustainable in the long term. Swabbing touch-points and elevated surfaces (which airborne pathogens will settle on) could provide early warning of infection as well as providing insights into how the virus is transmitted, which can inform infection prevention and control (IPC) measures. According to a World Health Organization scientific brief on transmission of SARS-CoV-2 (2020), direct (droplet) transmission and indirect spread via fomites (contaminated surfaces) and long-distance aerosols are thought to occur; however there is no conclusive evidence for indirect transmission in LTCF. (Ben-Shmuel et al., 2020; Greenhalgh et al., 2021; Ong et al., 2020; Ye et al., 2020)

COVID-19: Detecting Indirect Transmission in Facilities for Enhanced Care sTudy (COVID-19: DISinFECT) aims to investigate the role of indirect transmission of SARS-CoV-2 in LTCF and evaluate the potential for environmental surveillance to inform IPC measures. We present findings from a pilot conducted between 14th January and 28th March 2021, during the second epidemic wave in South West England.

Methods

DISinFECT methods are detailed in the protocol which can be accessed online. (Kwiatkowska & Ready, 2021) LTCFs were eligible for inclusion if they provided residential care for older adults (>65 years), were within the boundaries of a selected local authority in the Public Health England (PHE) South West region, and experienced a COVID-19 outbreak, defined as two or more laboratory-confirmed cases among staff and/or residents within a 14-day period.

Recruitment

On notification of an outbreak, investigators contacted the LTCF manager with information about DISinFECT and offered environmental sampling as part of outbreak management. If managers expressed an interest they were asked to complete a consent form permitting the study team to conduct telephone interviews, collect information from care home records, sample the care home environment and approach residents and staff for involvement. Prior to the sampling visit, residents and staff were provided with written and pictorial leaflets describing the purpose of the investigations, sampling procedures and how their information would be processed. Each of the residents selected for sampling was consulted to make sure they understood this information and were happy to provide samples. Sampling was not carried out if the individual lacked capacity to complete a consent form. Staff were asked for consent to participate prior to accessing the electronic questionnaire.

Sampling

Settings varied in size and layout but sampling was done systematically, with a focus on common touch points (for example: door handles, light switches, television remote controls) and elevated surfaces onto which airborne virus might settle (for example: door sills, tops of wall-mounted cabinets).

Within each home, sampling sites were categorised into three ‘zones’: 1) rooms occupied by residents isolating with active SARS-CoV-2 infection, or equipment used by them, 2) areas/equipment used by both staff and residents such as lounges and dining areas, shared kitchen equipment, and 3) staff-only areas/equipment such as offices, recreation areas, and key cabinets (see Figure 1). Surfaces were sampled using wetted flocked swabs and sponges, and wetted swabs were also used to swab the fingertips of residents in isolation rooms. All samples were transported to a public health laboratory specialising in aerobiology, biocontainment and biosafety measures.

The full DISinFECT protocol includes sampling stool and saliva from resident cases and air and wastewater sampling from the facility but for logistical reasons, stool and saliva could not be taken during this pilot. Likewise, no air sampling was conducted, and wastewater was collected from only one facility; results will be reported separately.

Laboratory analysis

Each sample was analysed in duplicate using a Reverse Transcription Polymerase Chain Reaction (RT-PCR) assay targeting both the N gene and the ORF1ab gene of SARS-CoV-2 (Viasure, CerTest Biotec, Zaragoza). Results were reported in cycle threshold (Ct) numbers. A sample was classified as ‘positive’ for SARS-CoV-2 if amplification of one or both targets was detected in both duplicates; ‘suspect’ if detected in only one duplicate and ‘negative’ if no amplification was detected for either gene target (Ct cut-off was 39). RT-PCR was repeated (in duplicate) for ‘suspect’ samples and samples with an internal control Ct standard deviation of >0.5. If repeat RT-PCR detected amplification of a gene target in both duplicates then the ‘suspect’ sample was reclassified as ‘positive’; otherwise the sample remained ‘suspect’.

Where possible, diagnostic isolates were sent for whole genome sequencing (WGS) to identify SARS-CoV-2 variants and mutations of interest/concern.
Epidemiological data collection
In addition to sampling, the study team recorded details of LTCF layout, cleaning and IPC measures, and staff and resident behaviours. Clinical test results (nasopharyngeal swabs analysed with PCR) were obtained from the regional public health laboratory and LTCF managers provided additional information about clinical case notifications, resident risk factors for infection and IPC measures. Finally, staff members were sent an electronic questionnaire asking questions about exposures and risk factors for infection.

Criteria for success in the pilot are presented in Box 1.

Box 1. DISinFECT pilot criteria for success
- a) Recruitment of three LTCF and consent to participate from at least one resident per facility;
- b) Detection of SARS-CoV-2 RNA from surface sampling;
- c) Variation by sampling zone in the proportion of sample sites testing positive for SARS-CoV-2 RNA;
- d) Potential to assess transmission risk from environmental contamination, in the context of individual risk factors for infection.

Ethical considerations
These investigations were carried out as part of a public health response to the SARS-CoV-2 pandemic. Ethical approval was granted on 14th January 2021 by the Public Health England Research Ethics and Governance Group (PHE REGG: RD 415).

Results
Four LTCFs were approached on notification of an outbreak: one declined to participate on the grounds that they did not have capacity to consider the study information or arrange for residents to be consulted. Three LTCFs were sampled between 2nd February and 10th March 2021 and a total of 84 environmental swabs were taken (56 from common touch points, 28 from elevated sites). One home had two sampling visits, 14 days apart. For simplicity, we have labelled the homes A, B and C in order of sampling dates. Table 1 contains the full list of sampling sites.

Setting and population
LTCF sizes ranged in size: there were 40 beds in facility A, which was a self-contained unit within an 80-bedded LTCF, 16 beds in facility B and 13 beds in facility C. The number of occupants was 30 (A), 15 (B) and 12 (C) on the date of onset of the first case. Facility A was a short stay residential unit with clients aged between 55 and 98 years; facility B a residential home for older adults (65 and over) with and without dementia; facility C a residential home for adults with learning difficulties aged between 35 and 88 years. All residents were included in the epidemiological analysis, regardless of age. All residents slept in single occupancy rooms; residents in facilities A and C all had private bathrooms and 13/16 rooms in facility B were en-suite. Characteristics of the three facilities are summarised in Table 2.

Outbreak trajectory and control measures
SARS-CoV-2 attack rates among residents were highest in facility A: 16/30 (53%), of which 15/16 (94%) were symptomatic,
Table 1. Sampling sites and SARS-CoV-2 RNA positivity by facility.

<table>
<thead>
<tr>
<th>Facility A: sampling sites</th>
<th>*</th>
<th>PCR for SARS-CoV-2</th>
<th>Facility B: sampling sites</th>
<th>*</th>
<th>PCR for SARS-CoV-2</th>
<th>2nd visit PCR result</th>
<th>Facility C: sampling sites</th>
<th>*</th>
<th>PCR for SARS-CoV-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female staff changing room</td>
<td>E</td>
<td>Negative</td>
<td>Staff toilet</td>
<td>E</td>
<td>Negative</td>
<td></td>
<td>Staff office</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>1. Top of lockers (next to door)</td>
<td>T</td>
<td>Negative</td>
<td>1. Air vent/extractor</td>
<td>E</td>
<td>Negative</td>
<td></td>
<td>1. Top of key cupboard</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>2. Toilet door: inside handle and lock</td>
<td>T</td>
<td>Negative</td>
<td>2. Soap dispenser lever</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>2. Computer mouse</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>3. Air vent/extractor (toilet)</td>
<td>E</td>
<td>Negative</td>
<td>3. Door ledge (outer)</td>
<td>E</td>
<td>Positive (Ct N 36.47 ± 0.46)</td>
<td>Negative</td>
<td>3. Top of message board</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>4. Bench top</td>
<td>T</td>
<td>Negative</td>
<td>Manager’s office</td>
<td>E</td>
<td>Negative</td>
<td></td>
<td>4. Telephone receiver</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>5. Soap dispenser lever</td>
<td>T</td>
<td>Negative (Ct N 39.06)</td>
<td>4. Computer mouse</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>Shared toilet (staff/residents)</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>Dining room (staff)</td>
<td>T</td>
<td>Negative</td>
<td>Shared toilet (staff/residents)</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>7. Front door keypad (inside)</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>6. Top of TV</td>
<td>E</td>
<td>Negative</td>
<td>5. Air vent/extractor</td>
<td>E</td>
<td>Positive (Ct N 37.43 ± 0.01)</td>
<td>Negative</td>
<td>6. Soap dispenser lever</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>7. Top of trolley (for lunch trays)</td>
<td>E</td>
<td>Negative</td>
<td>6. Glove box (size small)</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>N/A Reception area</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sluice room</td>
<td>T</td>
<td>Negative</td>
<td>7. Toilet booster handles</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>8. Top of kitchen door frame (outside)</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>9. Code pad (door exterior)</td>
<td>T</td>
<td>Negative</td>
<td>8. Air vent/extractor</td>
<td>E</td>
<td>Positive (Ct N 36.22 ± 0.35)</td>
<td>Negative</td>
<td>Staff break area</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breakout room (staff)</td>
<td>T</td>
<td>Negative</td>
<td>9. Toilet booster underside</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>9. Air vent/extractor</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>10. Chair arms</td>
<td>T</td>
<td>Negative</td>
<td>Dining area (residents &amp; staff)</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>10. Fire exit sign</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>11. Air vent/extractor</td>
<td>E</td>
<td>Negative</td>
<td>10. Tablecloth</td>
<td>T</td>
<td>Suspect (Ct N 36.99)</td>
<td>Negative</td>
<td>11. Door handle (exit to reception)</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>Corridor</td>
<td>T</td>
<td>Negative</td>
<td>12. Armchair handles</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>Lounge/ shared kitchen area</td>
<td></td>
<td></td>
</tr>
<tr>
<td>13. UV cabinet handle</td>
<td>T</td>
<td>Negative</td>
<td>13. Top of picture frame</td>
<td>E</td>
<td>Suspect (Ct N 37.05)</td>
<td>Negative</td>
<td>13. Top of kitchen hatch</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>Photocopyer room</td>
<td>T</td>
<td>Negative</td>
<td>14. TV remote control</td>
<td>T</td>
<td>Positive (Ct N 37.54 ± 0.12)</td>
<td>Negative</td>
<td>14. Chair arms</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>Bedroom/ bathroom #1</td>
<td>T</td>
<td>Negative</td>
<td>15. Top of wardrobe</td>
<td>E</td>
<td>Positive (Ct N 33.80 ± 0.18)</td>
<td>Positive (Ct N 35.25 ± 0.35)</td>
<td>16. Microwave handle &amp; dial</td>
<td>T</td>
<td>Negative</td>
</tr>
<tr>
<td>15. Bedrails (unoccupied)</td>
<td>T</td>
<td>Suspect (Ct N 37.43)</td>
<td>16. Fingertips (L hand)</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>17. Air vent/extractor</td>
<td>E</td>
<td>Negative</td>
</tr>
<tr>
<td>16. Fingertips (R hand)</td>
<td>-</td>
<td>Negative</td>
<td>17. Fingertips (R hand)</td>
<td>T</td>
<td>Negative</td>
<td></td>
<td>Corridor</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Facility: A</td>
<td>Sampling sites</td>
<td>PCR for SARS-CoV-2</td>
<td>2nd visit PCR result</td>
<td>Designated common touch point/T or elevated site/E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>------------</td>
<td>----------------</td>
<td>---------------------</td>
<td>----------------------</td>
<td>--------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>17. Host handle</td>
<td>Neg/Pos/CI 37.75 ± 0.03</td>
<td>-</td>
<td>-</td>
<td>T</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>18. Glove box - size small</td>
<td>Neg/Pos/CI 37.44 ± 0.31</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>19. Lip of PPE cabinet</td>
<td>Neg/Pos/CI 37.72 ± 0.21</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20. Air vent/extractor (bedroom #2)</td>
<td>Neg/Pos/CI 37.40</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>21. Bed remote control (occupied)</td>
<td>Neg/Pos/CI 37.78 ± 0.21</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>22. Air vent/extractor (bedroom #2)</td>
<td>Neg/Pos/CI 37.77 ± 0.03</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>23. Wardrobe handle</td>
<td>Neg/Pos/CI 37.50</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>24. TV control (in use)</td>
<td>Neg/Pos/CI 37.66 ± 0.41</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>25. Table top</td>
<td>Neg/Pos/CI 37.78 ± 0.21</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>26. Bed remote control (bedroom #2)</td>
<td>Neg/Pos/CI 37.48</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>27. Fingers tips (right hand)</td>
<td>Neg/Pos/CI 37.67 ± 0.75</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>28. Top of light above bathroom sink</td>
<td>Neg/Pos/CI 37.40</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>29. Oxygen saturations probe (outside room)</td>
<td>Neg/Pos/CI 37.59 ± 0.03</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30. Wheelbarrow handles</td>
<td>Neg/Pos/CI 37.70 ± 0.30</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>31. Air vent/extractor (bathroom)</td>
<td>Neg/Pos/CI 37.75</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>32. Host rail (bedroom)</td>
<td>Neg/Pos/CI 37.50</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>33. Toilet handle</td>
<td>Neg/Pos/CI 37.46 ± 0.00</td>
<td>-</td>
<td>T</td>
<td>E</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**
- Designated common touch point/T or elevated site/E
- Samples are analysed in duplicate: a positive result means the target gene is detected in both replicates (reported as cycle threshold (Ct) number with 95% confidence intervals). If the target gene is identified in only one sample, it is reported as suspect. Ct values reported for N gene target: the higher the Ct value, the lower the concentration of viral RNA. Limit of detection set at Ct 39.
- NB not all sites were re-sampled on the second visit to facility B (labelled N/A), and some sample sites were combined (indicated by merged cells in the table).
Table 2. Characteristics of DISinFECT pilot LTCFs and resident populations.

<table>
<thead>
<tr>
<th>ID</th>
<th>CQC* rating</th>
<th>No. residents/ no. beds (% occupancy*)</th>
<th>No. floors</th>
<th>Private/ shared bathroom</th>
<th>Care provision</th>
<th>Agency staff</th>
<th>Dependent/ independent</th>
<th>Age range (years)</th>
<th>Walk with purpose†</th>
<th>Prevalence comorbidities†</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>O</td>
<td>30/40 (75.0%)</td>
<td>1</td>
<td>Private</td>
<td>Residential: short stay</td>
<td>Yes - block</td>
<td>Mixed</td>
<td>55–98</td>
<td>No</td>
<td>23/30 (76.7%)</td>
</tr>
<tr>
<td>B</td>
<td>G</td>
<td>15/16 (93.8%)</td>
<td>2</td>
<td>Mixed</td>
<td>Residential: older adults (+/- dementia)</td>
<td>Yes</td>
<td>Mixed</td>
<td>71–97</td>
<td>Yes</td>
<td>13/15 (86.7%)</td>
</tr>
<tr>
<td>C</td>
<td>G</td>
<td>12/13 (92.3%)</td>
<td>2</td>
<td>Private</td>
<td>Residential: adults with Learning Difficulties</td>
<td>No</td>
<td>Mixed</td>
<td>35–88</td>
<td>Yes</td>
<td>data unavailable</td>
</tr>
</tbody>
</table>

*registered occupancy at time of sampling visit; †CQC = Care Quality Commission, O = Outstanding, G = Good; ††one or more resident unable to adhere to self-isolation within private room; †††presence of one or more chronic condition (overweight/obese; chronic respiratory disease; chronic heart disease; dementia; diabetes; hypertension; immunocompromised/ cancer) among registered occupants over the course of the outbreak; Dependent/ independent relates to resident mobility.

followed by facility B: 6/15 (40%, of which 3/6 (50%) symptomatic). Only one resident tested positive in LTCF C (8%); this individual was asymptomatic and had received the first vaccine dose four weeks beforehand. They also had a history of laboratory-confirmed COVID-19 a year previously. A repeat sample taken 10 days after the most recent diagnosis was PCR negative for SARS-CoV-2, nonetheless the individual remained in isolation for 14 days as a precaution. Non-agency staff attack rates were also highest in facility A: 16/60 (27%) followed by facility B: 4/22 (18%) and C: 1/35 (3%). Numbers of agency staff were not available. Duration of outbreak (calculated from the date of first illness onset to 28 days after onset of the final case) was 63, 50 and 30 days for facilities A, B and C respectively. Facilities A and B had residents admitted to hospital (n=5 symptomatic cases and n=3 of which one was symptomatic, respectively). Sadly there were COVID-19-related deaths among residents (facility A: n=2, both receiving end-of-life care, one hospitalised; facility B: n=1, hospitalised). Figure 2 illustrates outbreak trajectories in facilities A and B.

All homes adopted enhanced cleaning protocols in response to the COVID-19 epidemic, with increased frequency and a focus on common touch points. In addition, facility A provided fresh uniforms for staff at the beginning of each shift (laundered on site) and had installed a UV cabinet for treating phones and keys prior to handover. All LTCF managers stated that personal protective equipment (PPE) was available to staff in line with national guidance.

In facilities B and C, 100% of residents had received the first dose of a COVID-19 vaccine more than two weeks prior to outbreak onset. In facility A, 83% of residents had received the first dose of vaccine but just four days before onset of the outbreak. In facility A, 80% of non-agency staff were reported to have received a vaccine, though these figures related to the wider facility and exact vaccination dates were not provided. Facility B reported 55% and facility C 96% of non-agency staff vaccinated with at least one dose more than two weeks prior to the outbreak.

Observations

Facility A was a modern building with spacious, uncluttered rooms of a uniform layout. Signage was in place to remind staff to clean surfaces and socially distance, and the sampling team observed good adherence to PPE donning and doffing protocols. All residents were isolated in their rooms, and staff wore ‘scrubs’.

Facility B was an older building, once a large house. Residents’ rooms were small and somewhat cluttered with several sampling sites visibly soiled. Several residents were observed using the dining area and lounge (unmasked); staff wore their own clothing.

Facility C was a relatively modern building; rooms were small but uncluttered with fewer soft furnishings than LTCFs
A and B. Two residents were observed walking with purpose (unmasked), accompanied by carers; staff wore their own clothing.

Proportion of sites testing positive and distribution of SARS-CoV-2 RNA
Facility B had the highest proportion of sampling sites testing positive/ suspect for SARS-CoV-2 RNA (PCR positive on one or both duplicates): 17/28 (61%), followed by LTCF A: 6/27 (22%). In LTCF C, all environmental swabs were negative for SARS-CoV-2 except for one suspect positive from an air extractor in the index case’s bathroom: positivity 1/31 (3%).

A repeat visit to facility B two weeks after the initial sampling visit yielded a much lower proportion of SARS-CoV-2 positive/ suspect sampling sites (4/19; 21%).

Concentrations of SARS-CoV-2 RNA, as inferred from Ct values, were very low for all positive and suspect samples. The lowest Ct value found was 31.8, and only four samples had a Ct value of below 35. Amplification was below the limit of detection in two samples (see Table 1).

SARS-CoV-2 positive/ suspect surfaces were most common in zone 1 (rooms occupied by residents with active SARS-CoV-2 infection, and equipment used by them), as illustrated in Figure 3.

Proportion of SARS-CoV-2 positive sites in proximity to a COVID-19 case
Within zone 1, there was significant variation in the proportion of sample sites testing positive/ suspect for SARS-CoV-2 RNA. For example, in facility A two residents’ rooms were sampled: in the first room, 1/5 (20%) of sample sites was ‘suspect positive’ for SARS-CoV-2 RNA and in the second room 6/8 (75%) of sample sites tested positive. Both rooms were similar in size and layout, and subject to the same cleaning protocols. The first room was occupied by an individual who was bed/chair bound, and who had tested positive for the virus 11 days previously and had fever and a slight cough. The occupant of the second room spent much of their time confined to bed, though was mobile with a wheelchair. This individual tested positive for SARS-CoV-2 infection seven days prior to the visit and had a cough. Figure 4 shows the environmental sample site positivity in relation to the time from onset of illness for the room occupant.

Proportion of common touch points testing positive for SARS-CoV-2 vs elevated surfaces
Overall, the proportion of common touch points testing positive/ suspect for SARS-CoV-2 RNA (15/56; 27%) was slightly lower than the proportion of sites testing positive/ suspect (10/28; 36%), though this was not consistent between LTCFs (see Table 4). Where the proportion of positive common touch points was highest (facility B), a large proportion of elevated sites were also tested positive/ suspect for SARS-CoV-2 RNA.

Fingertip samples
None of the swabs taken from fingertips of residents with confirmed COVID-19 were PCR positive for SARS-CoV-2 RNA.

WGS of clinical isolates
A small number of outbreak isolates were sequenced by the COVID-19 Genomics UK Consortium (COG-UK) (facility A: n=3; facility B: n=3). All were identified as B.1.1.7 (VOC-202012/01), with no E484K substitution detected.

Staff survey results
Response rates to the electronic staff survey were poor: 11/60 (18%) from facility A, 6/22 (27%) from facility B and 0/35
Table 3. Outbreak metrics and control measures for DISinFECT pilot LTCFs.

<table>
<thead>
<tr>
<th>ID</th>
<th>Outbreak metrics</th>
<th>Infection Prevention &amp; Control</th>
<th>COVID-19 vaccine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Attack rate residents</td>
<td>Attack rate staff</td>
</tr>
<tr>
<td>A</td>
<td></td>
<td>16/30 (53%)</td>
<td>16/60 (27%)</td>
</tr>
<tr>
<td>B</td>
<td></td>
<td>6/15 (40%)</td>
<td>4/22 (18%)</td>
</tr>
<tr>
<td>C</td>
<td></td>
<td>1/12 (8%)</td>
<td>1/35 (3%)</td>
</tr>
</tbody>
</table>

*Approximate staff attack rates, based on numbers of non-agency staff. *Period between onset/ swab dates of index case, and 28 days after illness onset for the final case; With the exception of end of life visits. *as per PHE guidelines: daily Lateral Flow Tests (LFT) for at least 5 days followed by twice weekly LFT and once weekly PCR testing. *Personal Protective Equipment; *Receipt of 1 or more dose >=14 days prior to onset in the index case; *Staff vaccination figures provided for whole facility, not just the short stay unit. No dates reported so figures may not accurately represent vaccine-induced immunity.
(0%) from facility C, meaning there was insufficient data to assess clinical and epidemiological risk factors for infection. None of the respondents raised concerns about access to PPE as recommended in national guidelines for working in care homes.

**Discussion**

The greatest proportions of SARS-CoV-2 positive/suspect sample sites were found in the immediate vicinity of laboratory-confirmed COVID-19 cases, which is consistent with findings from other studies and indicates that environmental swabbing...
can detect the presence of an infected individual. (Onakpoya et al., 2021) The proportion of SARS-CoV-2 positive/suspect sites varied considerably within one LTCF, even between rooms with similar layouts and cleaning regimes occupied by individuals with comparable symptom profiles. The exact timing of cleaning was not captured and may have affected these results, though variation was also observed in elevated sites which were unlikely to have been cleaned as part of daily routine. Equally, the sensitivity of surface swabbing may have differed between rooms though the same sampler swabbed both. Environmental sampling around COVID-19 patients evacuated from the Diamond Princess cruise ship demonstrated a similar lack of correlation between clinical signs of illness and levels of environmental contamination. (Santarpia et al., 2020) It is likely that our findings illustrate the myriad of environmental, clinical, behavioural and pathogen factors affecting dispersion of the virus, (Moore et al., 2021) which must be controlled for in any analysis of infection risk by indirect transmission.

Survey response rates were poor, which is likely to reflect the pressures staff were under at the time. Consequently, there was insufficient epidemiological information to assess whether transmission occurred via fomites or long-distance aerosol during these outbreaks. Concentrations of SARS-CoV-2 RNA were very low for all positive and suspect samples, therefore it is highly unlikely that viable virus was present. (Huang et al., 2020; Transmission of Covid-19 in the Wider Environment Group (TWEG) reporting to UK Scientific Advisory Group on Emergencies (SAGE) 2020) This is in keeping with results from environmental surveillance in Canadian care homes. (Nelson et al., 2021)

Table 4. Proportion of common touch point and elevated sampling sites testing positive for SARS-CoV-2 RNA, by LTCF.

<table>
<thead>
<tr>
<th>LTCF</th>
<th>No. (proportion) common touch sites positive for SARS-CoV-2</th>
<th>No. (proportion) elevated sites positive for SARS-CoV-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>5/17 (29%)</td>
<td>2/10 (20%)</td>
</tr>
<tr>
<td>B</td>
<td>10/20 (50%)</td>
<td>6/7 (86%)</td>
</tr>
<tr>
<td>C</td>
<td>0/19 (0%)</td>
<td>1/10 (10%)</td>
</tr>
<tr>
<td>Total</td>
<td>15/56 (27%)</td>
<td>10/28 (36%)</td>
</tr>
</tbody>
</table>

The fact that fingertip swabs were all PCR negative for SARS-CoV-2, despite observing infected residents touch various items which returned a positive result, was surprising: people typically touch their nose, eyes and mouth more than 20 times per hour and an experimental study suggests that the virus can persist on skin for at least 8 hours at body temperature. (Harbourt et al., 2020; Kwok et al., 2015) Sampling may have occurred too late to detect viral shedding, since participants were between seven and 14 days of diagnosis, and the concentration of virus on fomites was low which might reduce the likelihood of transfer to fingers. It is also possible that residents applied hand sanitiser unobserved, or that the sampler did not apply sufficient pressure or friction to pick up viral RNA. (Mbithi et al., 1992)

Notably, facility A experienced the highest attack rates despite implementing more comprehensive IPC and cleaning measures compared to facilities B and C. Residents in facility A were more susceptible to infection than those in facilities B and C, having only received the first dose of vaccine shortly before the outbreak onset. (Shrotri et al., 2021; Tenforde et al., 2021) Facility A also had a higher rate of admissions from the local hospitals and these factors, as well as the relatively large size of the facility, may have increased the probability of multiple introductions of the virus to the premises. (Burton et al., 2020; Shallcross et al., 2021) Unfortunately staff survey response rates were too low to enable a comparative analysis of other infection risk factors within and outside the facility.

High attack rates may also have reflected community case rates, which peaked at the end of December and remained high through January (see Figure 2 for pilot LTCF outbreak trajectories). This surge in case rates was fuelled by emergence of the more transmissible B.1.1.7 Alpha variant, which quickly entered English LTCFs. (Krutikov et al., 2021) At this time a substitution at the E484K location of the receptor binding domain also emerged, raising concerns that the virus might evade the host immune response. (Wise, 2021) Six clinical samples were sequenced and were all of the B.1.1.7 Alpha variant with no E484K substitution, however we cannot exclude the possibility that multiple strains of the virus were contributing to these outbreaks.

Results from surface swabbing provided some reassurance that facility A (with staff cohorting and enhanced IPC measures) was effectively containing the environmental spread of the virus, in contrast to LTCF B (without cohorting) in which viral RNA was widely distributed. Repeat sampling 14 days after the initial visit to facility B yielded a much lower proportion of SARS-CoV-2 positive/suspect sites. Since the first visit corresponded with the end of the final case’s infectious period and no further cases of COVID-19 were identified it is reasonable to assume that nobody in the facility was actively shedding virus at the second sampling visit. Our observations could reflect the effectiveness of cleaning protocols introduced between sampling visits or of swabbing at the first round of sampling, or degradation of viral RNA over a 14 day period. (Onakpoya et al., 2021; van Doremalen et al., 2020) Facility C was the only one to isolate its resident case on a separate floor/wing which may have helped reduce egress to other areas within the home. However, this individual’s history of vaccination and prior infection, and a negative repeat PCR test suggest that the diagnosis was a false positive and they were not shedding SARS-CoV-2 at the time of sampling.

Elevated sampling sites, being cleaned less regularly, may be a more pragmatic means of SARS-CoV-2 detection than common touch points: of the four sites that remained positive/suspect
for SARS-CoV-2 on a repeat visit to facility B, three were elevated. Reactive sampling, as applied in this pilot, will not distinguish between historic and current viral shedding but there is evidence that levels of surface contamination with SARS-CoV-2 RNA mirror contemporaneous levels of airborne SARS-CoV-2 RNA. (Cherrie et al., 2021; Dumont-Leblond et al., 2021) Air vents may be useful sentinel sampling points since three of four air vents in facility B tested positive for SARS-CoV-2 RNA, and in facility C the air vent was the only sampling site that tested suspect positive. Similar observations are reported from sampling ventilation grates in the Diamond Princess COVID-19 quarantine rooms, and respiratory viruses have been isolated from air filters in aeroplanes and large public buildings. (Goyal et al., 2011; Korves et al., 2011; Santarpia et al., 2020) It is interesting that none of the air vents sampled from facility A tested positive for SARS-CoV-2 RNA, including one in a room that was otherwise quite heavily contaminated. This could have been an artefact of different sampling techniques, or reflect the design of the air vents, which were circular with a single ring opening rather than a slatted grate, though the vent that tested suspect positive in facility C was of the same circular design. Facility A may also have been better ventilated than the other facilities, however this seems unlikely given positive results in other elevated sites in the building.

The uncluttered environment in facility A, in reducing build-up of dust, may also have helped limit environmental spread of SARS-CoV-2. Evidence that respiratory droplets containing SARS-CoV-2 are adsorbed to dust and particulate matter, creating ‘aerosolised fomites’, is emerging (Andree, 2020; Conticini et al., 2020; Qu et al., 2020; Renninger et al., 2021; Setti et al., 2020; Travaglio et al., 2021) and in healthcare settings we have observed that elevated surfaces accumulate greater quantities of SARS-CoV-2 RNA in dusty environments such as changing rooms, bathrooms, and cluttered spaces (unpublished data). This merits further investigation.

Limitations
This pilot has several limitations, not least the small sample size, lack of control sites and heterogeneity of LTCFs. Our interpretation of results is speculative and intended to generate hypotheses rather than answer questions.

Between-site variation in layout and infrastructure means that sampling frames cannot be entirely standardised, and there may be a tendency to oversample areas that are visibly soiled. Sampling technique may also vary between samplers. Results from surface swabs represent a snapshot in time and cover a fraction of the LTCF environment so we may not have accurately captured overall levels of environmental contamination. Among other things these may have been influenced by trends in community prevalence of COVID-19 (affecting risk of importation), expansion of new variants, and the effects of vaccination rollout (affecting viral shedding and transmissibility).

We were unable to confirm whether the diagnostic test for the single resident case in facility C was a true positive, therefore the SARS-CoV-2 RNA detected in this individual’s rooms may have been residual from previous occupants or their carers. This bias also applies to the other facilities to some extent, since all were likely to have been exposed to the virus (whether or not it manifested clinically) at some point prior to the outbreak.

Sequencing data were only available for minority of outbreak samples therefore we were unable to assess whether new or multiple strains were responsible for the outbreaks in question.

Conclusions and recommendations
We have successfully recruited three pilot LTCF to the Disinfect study and observed SARS-CoV-2 RNA on a high proportion of surfaces around individuals with a laboratory-confirmed infection, though this varied considerably within and between settings.

This pilot demonstrates that surface swabbing can provide reassurance that IPC measures such as self-isolation, staff cohorting and enhanced cleaning collectively reduce egress of the virus from quarantine rooms to the wider care home environment. The heterogeneity of settings and situations means that we cannot assess impact of environmental contamination or individual IPC measures on transmission risk, however. An analysis of national LTCF-level data would be beneficial to assess which, if any, IPC measures influence attack rates during outbreaks of SARS-CoV-2.

Results also highlight that LTCF staff can be overburdened with information requests during outbreaks. There is a need for efficient and parsimonious data collection tools, using routine sources of intelligence wherever possible, to gather sufficient epidemiological information for the interpretation of environmental surveillance data.

Finally, sampling frameworks focussing on elevated surfaces/those which accumulate particulate matter may be less susceptible to the effects of cleaning regimes and thus a useful tool for detecting outbreaks and evaluating IPC measures.

In summary: this pilot demonstrates the potential utility of surface swabbing in LTCF to assess environmental spread and effectiveness of IPC measures, and monitor for outbreaks of infectious disease. At present, the potential to assess SARS-CoV-2 infection risk via indirect routes is limited by the heterogeneity of LTCFs and their populations, and challenges around data collection.

Data availability
Underlying data
To preserve anonymity of LTCF residents and staff, the study data are stored on a secure drive hosted by Public Health...
England (PHE) Field Services South West. Access to personal identifiable data is restricted to personnel responsible for outbreak investigations. Non-identifiable data may be made available to others upon formal request (please contact the corresponding author for an information request form) and subject to approval from the PHE Office for Data Release.

Extended data

This project contains the following files:
- DISeNFECT_Protocol_OSF.pdf. The protocol for this research study.
- DISeNFECT_staffsurvey.pdf. The electronic survey distributed to staff to collect information on epidemiological and clinical risk factors for SARS-CoV-2 infection.
- DISeNFECT_Tools_line list_v6.xlsx. The data collection template for LTCF residents: clinical and epidemiological risk factors for SARS-CoV-2 infection.
- DISeNFECT_Tools_setting_log_v2.xlsx. The data collection template for details of LTCF layout and staffing arrangements.

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

Acknowledgements
The authors would like to express thanks to the staff and residents of care homes who supported these investigations during a very challenging time. Also to acknowledge the hard work of regional Health Protection Teams and local authorities throughout the SARS-CoV-2 epidemic, along with laboratory staff without whom this work would not have been possible.

References


Onakpoya IJ, Heneghan CJ, Spencer EA, et al.: SARS-CoV-2 and the Role of
Open Peer Review

Current Peer Review Status: X

Version 1

Reviewer Report 15 October 2021

https://doi.org/10.21956/wellcomeopenres.18828.r45883

© 2021 Julian T. This is an open access peer review report distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Timothy R. Julian
Eawag, Swiss Federal Institute of Aquatic Science and Technology, Dübendorf, Switzerland

Overview:

The study describes epidemiology of SARS-CoV-2 outbreaks in three long term care facilities (LTCF) during SARS-CoV-2 outbreaks in the South West of England from January to March 2021, with a focus on the results of surface sampling. The topic is importantly and timely, highlighting the potential use of environmental sampling to both inform disease prevalence (surveillance) and to identify risks of transmission (fomite-mediated transmission).

The study identifies extensive contamination of SARS-CoV-2 RNA in one of the three sites, and almost no detectable contamination at one of the sites. Further, the study concluded that elevated surfaces – or surfaces with sufficient height they would not be expected to be cleaned regularly – were more likely to have detectable RNA than other sites. The study is well-written, describing a case study of contamination in LTCFs which are an important site of concern for transmission of SARS-CoV-2. Further, the study is embedded within a larger, more extensive study which may provide important insights on transmission routes of SARS-CoV-2 in these settings.

The manuscript should minimally be updated to include more details on methodology of surface sampling, as the text description is limited and hinders interpretation of the results. Additionally, laboratory controls (field blank, extraction blank, no template controls, positive controls, inhibition testing, recovery efficiency experiments) are not described, making it unclear the extent to which results may be due to sample contamination (or cross-contamination) as well as efficacy of the protocol. Given this is environmental sampling typically characterized by low concentrations and low positivity rates (as reflected in the results), adequate controls are particularly important.

Additional improvements (or discussion at least) of a lack of a standard curve is warranted, given that it hinders experimental interpretation, particularly discussions of concentrations of SARS-CoV-2 on surfaces and the associated risks of contacts. Synthetic SARS-CoV-2 RNA is commercially available, and would provide a sufficient standard for quantification of samples.
An additional update is that authors should consider modifying the manuscript discussion and conclusions to more closely align with the results. A specific example is that the study investigated three LTCFs and attempts to draw conclusions on results as a function of cleaning regime: such inference would require a larger sample size with multiple facilities with variable cleaning and quarantining regimes to rule out location-specific effects not associated with cleaning.

**Detailed Comments:**

**Abstract:**
- “potential to assess infection risk from SARS-CoV-2 positive surfaces” - should be connected to methods for this approach, unclear how this would be done.
- Define “elevated” here and in Methods.
- “could be associated with implementation” – this is speculation, and was not specifically tested, so suggest authors remove this from the abstract.

**Introduction:**
- “touch-points and elevated surfaces” – how did authors conclude they should test elevated surfaces? Is prior literature available here that could be referenced?
- Author discussion of COVID-19: DISinFECT suggests a project larger than the study results presented here, but it is unclear. Author clarification on the project DISinFECT and its relationships to this study would be helpful and interesting.

**Methods:**
- “complete a consent form” – suggest “written consent form”.
- Substantially more information is recommended on collecting samples. Detection/Non-detection of viral RNA on surfaces is influenced by the sensitivity of the methods, which includes sample collection, RNA extraction, and RNA detection. Authors should consider reporting material used (not only swabs, but also eluent volume and material, volume of material subsequently processed, RNA extraction methods, including volumes eluted (if applicable), and full details of the QPCR methodology. Inhibition may also influence sample detection, and some indication of qPCR inhibition testing should be included. Indeed, less frequently contacted surfaces (like elevated surfaces) may have more dust and thus may be more inhibited?
- Bustin et al. 2009 MIQE Guidelines\(^1\) provide a motivation and framework for reporting on data relying on qPCR. Additionally, authors should report all controls used, including positive controls (source and concentration), field blanks, extraction blanks, and no template controls. Authors should also conduct, or report on previously studies that conducted, SARS-CoV-2 or surrogate virus RNA from surfaces to provide an estimate of recovery efficiency. Environmental sampling relying on detection of low concentrations should demonstrate sufficient use of controls to rule out contamination and cross contamination of samples, and recovery efficiency to provide insights into relative magnitude of surface contamination.
- How was a Ct value of 39 chosen as a cut-off? How does this value compare to no template controls? See MIQE guidelines\(^1\) for guidance on Limits of Detection. Authors could consider
motivating why no standard curve was included to estimate concentrations.

- On mention of WGS, it was unclear in the methods that this meant clinical isolates and not environmental samples. Suggest authors clarify. Also, authors should specify directly, for example through referencing, what protocols were followed, what primer sets were used, for sequencing.

- Figure 1 provides minimal novel insights into the study, authors could consider replacing with more descriptive figure covering all specific locations tested, with further information on their classification (elevated or not).

### Results
- Term “home” is used and could be replaced with LCTF.

- Table 1 – in one negative sample, a threshold Ct value is reported. Similar, positive Ct is reported for all samples. Consider reporting Ct for all samples as a separate column.

- Table 2 – a helpful table for comparing locations, is it possible to add the percent of surfaces that were positive at each site?

- Figure 2- why is C not included in this trajectory?

- “Concentrations of SARS-CoV-2 RNA” – in the absence of standard curves and recovery efficiency experiments, there is an obscure relationship between Ct values and true (unobserved) concentrations on surfaces. Suggest authors avoid discussing or inferring concentrations and instead focus on detection/non-detection.

- “lower than the proportion of sites” – should this say elevated sites?

- Fingertip samples – repeat how many samples were tested here.

- Figure 3 – suggest updating legend descriptions from Zone 1-3 to more detailed descriptions provided in legend to make it easier to read.

- Figure 4 – unclear why there are two y-axes here since there are only four data points. Suggest authors place days since illness as a number above the bar charts. An alternative, if the point is to visualize the relationship (or lack of) between days since illness and positivity rate, is to plot a scatter plot with days since illness on x-axis and positivity rate on y-axis and label each point with A1, A2, B, or C with text labels.

### Discussion
- “with findings from other studies” – authors should reference these studies in this sentence, only one study is currently referenced.

- “can detect the presence of an infected individual” – although conceptually the authors demonstrate detection in rooms with infected patients, the study does not evaluate the specificity or sensitivity of the approach: how often in an infected patients room or other building area is SARS-CoV-2 detected in the room? How often is SARS-CoV-2 detected in uninfected patients rooms or other building areas? What are the false negative/false positive rates? These values are necessary to determine the feasibility of this approach in
environmental surveillance.

○ “Consequently, there was insufficient epidemiological... “ – I am a bit skeptical that survey results could provide insight into the role of fomites vs. long range aerosol vs. close contact in disease transmission. Can the authors further describe how this could be done?

○ “The fact that fingertip swabs...”, could also be very low methodological recovery. This should be tested or at least discussed in the context of other work on recovery efficiency. An alternative is spatial heterogeneity (in addition to temporal heterogeneity discussed by authors).

○ “may have been residual” – or analytical false negatives.

○ “This pilot demonstrates...” - the efficacy of surface swabbing to determine efficacy of IPC measures should rely on substantially more than three LCTFs, given that the variation in site-specific contamination may be caused by factors other than IPC. The sample size is too small to clearly demonstrate this. Suggest authors rephrase this point to highlight that future studies could use these methodology to explore impacts of IPC on surface contamination.

Comments on specific elements of the review:

Technically Sound / Methods and Analysis:
○ As discussed, more data on methods and associated results of controls would be beneficial.

Statistical Analysis:
○ The analysis presented here is primarily qualitative given the pilot nature, small sample size, and heterogeneity in results.

Data Availability:
○ The authors present much of their data in the manuscript, and offer the rest (non-identifiable) on contact. Although this is generally sound, suggest authors make non-identifiable data freely available, in particular a modified version of Table 1 that includes all Ct values (not the mean, SD).

References

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly

If applicable, is the statistical analysis and its interpretation appropriate?
Not applicable

Are all the source data underlying the results available to ensure full reproducibility?
No

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Environmental Microbiology, Environmental Virology, Exposure Assessment, Risk Assessment

I confirm that I have read this submission and believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.