Articles

Risk factors and vectors for SARS-CoV-2 household transmission: a prospective, longitudinal cohort study

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Summary

Background Despite circumstantial evidence for aerosol and fomite spread of SARS-CoV-2, empirical data linking either pathway with transmission are scarce. Here we aimed to assess whether the presence of SARS-CoV-2 on frequently-touched surfaces and residents' hands was a predictor of SARS-CoV-2 household transmission.

Methods In this longitudinal cohort study, during the pre-alpha (September to December, 2020) and alpha (B.1.1.7; December, 2020, to April, 2021) SARS-CoV-2 variant waves, we prospectively recruited contacts from households exposed to newly diagnosed COVID-19 primary cases, in London, UK. To maximally capture transmission events, contacts were recruited regardless of symptom status and serially tested for SARS-CoV-2 infection by RT-PCR on upper respiratory tract (URT) samples and, in a subcohort, by serial serology. Contacts' hands, primary cases' hands, and frequently-touched surface-samples from communal areas were tested for SARS-CoV-2 RNA. SARS-CoV-2 URT isolates from 25 primary case-contact pairs underwent whole-genome sequencing (WGS).

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Findings From Aug 1, 2020, until March 31, 2021, 620 contacts of PCR-confirmed SARS-CoV-2-infected primary cases were recruited. 414 household contacts (from 279 households) with available serial URT PCR results were analysed in the full household contacts' cohort, and of those, 134 contacts with available longitudinal serology data and not vaccinated pre-enrolment were analysed in the serology subcohort. Household infection rate was $28 \cdot 4\%$ (95% CI $20 \cdot 8 - 37 \cdot 5$) for pre-alpha-exposed contacts and $51 \cdot 8\%$ ($42 \cdot 5 - 61 \cdot 0$) for alpha-exposed contacts (p=0 \cdot 0047). Primary cases' URT RNA viral load did not correlate with transmission, but was associated with detection of SARS-CoV-2 RNA on their hands (p=0 \cdot 031). SARS-CoV-2 detected on primary cases' hands, in turn, predicted contacts' risk of infection (adjusted relative risk [aRR]= $1 \cdot 70$ [95% CI $1 \cdot 24 - 2 \cdot 31$]), as did SARS-CoV-2 RNA presence on household surfaces (aRR= $1 \cdot 66$ [$1 \cdot 09 - 2 \cdot 55$]) and contacts' hands (aRR= $2 \cdot 06$ [$1 \cdot 57 - 2 \cdot 69$]). In six contacts with an initial negative URT PCR result, hand-swab (n=3) and household surface-swab (n=3) PCR positivity preceded URT PCR positivity. WGS corroborated household transmission.

Interpretation Presence of SARS-CoV-2 RNA on primary cases' and contacts' hands and on frequently-touched household surfaces associates with transmission, identifying these as potential vectors for spread in households.

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Introduction

SARS-CoV-2 transmission mostly occurs in households,¹ yet our understanding of the routes of transmission remains an area of ongoing scientific debate. Despite public guidance on self-isolation, facemask wearing, handwashing, and surface and air decontamination, there is very little empirical evidence for the pathways through which the virus spreads. Modelling and epidemiological observations implicate viral spread through large respiratory droplets that settle on surfaces from where virus is transferred to contacts' mucosal membranes, known as fomite spread, as well as airborne spread through smaller, aerosolised droplets directly inhaled by contacts.²

Although previous studies have documented the presence of SARS-CoV-2 RNA on household surfaces,^{3,4} as well as in exhaled and ambient air from hospitalised patients with COVID-19,^{5,6} its detection on potential vectors has not hitherto been shown to correlate with transmission. This gap in knowledge hampers accurate modelling of disease transmission and development of evidence-based public health guidance,^{7,8} which remain important given the ability of SARS-CoV-2 to spread through highly vaccinated populations.^{9,10} As isolation restrictions for household contacts have now been lifted, understanding the risk factors and pathways of transmission within households has become increasingly important to prevent the spread of SARS-CoV-2.

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Research in context

Evidence before this study

Extensive literature on SARS-CoV-2 transmission has been published, with different types of studies (observational prospective cohort studies, retrospective cohort studies, modelling, case-control, etc) evaluating transmission in different scenarios and populations. Although most transmission globally occurs in households, the vectors and pathways of household SARS-CoV-2 transmission remain poorly understood and there is ongoing scientific debate as to the predominant modes of transmission. We searched for the terms "SARS-CoV-2" AND "household" AND "transmission" with "pathways" OR "surface" OR "hand" OR "fomite" OR "droplet" in PubMed for articles published from database inception up until Nov 2, 2022, in English. After omitting studies with different settings or study populations (hospitalised populations, health-care workers, schools, care homes, etc), we identified few studies assessing SARS-CoV-2 in household settings. Several systematic and nonsystematic reviews have been published on household infection rate, often denoted as secondary attack rate, and some studies have evaluated the contribution of demographic, clinical, and epidemiological factors to infection (sex, age, vaccinations status, co-infection, comorbidities, relation to the primary case, household characteristics, etc). At the time of writing, only six studies reported the presence of viral RNA on household surfaces, but none have correlated this presence to risk of infection among contacts. Additionally, we found no studies analysing the presence of viral RNA on primary cases' or contacts' hands, which might be important vectors of transmission.

Added value of this study

We prospectively enrolled symptomatic and asymptomatic household contacts recently exposed to their primary cases,

Risk factors, vectors, and pathways for transmission can only be rigorously assessed by maximally and prospectively capturing transmission to susceptible household contacts. However, large-scale data from national testing services rely on contacts requesting tests, usually because they are symptomatic.¹¹ Moreover, almost all contacts' infection rate estimates use upper respiratory tract (URT) PCR testing at a single timepoint to detect or exclude infection,¹² limiting the ability to detect new secondary infections.¹³

We therefore prospectively recruited susceptible household contacts recently exposed to PCR-positive COVID-19 primary cases. We serially sampled both symptomatic and asymptomatic contacts' URT for PCR testing and, when consented, blood for serial serology. We tested for the presence of SARS-CoV-2 RNA on primary cases' and contacts' hands and frequentlytouched surfaces. We aimed to assess whether the presence of SARS-CoV-2 on one or more of these candidate vectors would prospectively predict SARS-CoV-2 transmission. Methods

household spread.

Study design and participants

transmission in households.

Participants were enrolled in two longitudinal community-based observational studies in Greater London, UK: Integrated Network for Surveillance, Trials and Investigations into COVID-19 Transmission (INSTINCT), and Assessment of Transmission And Contagiousness of COVID-19 in Contacts (ATACCC). Ethics approval was obtained from the Health Research Authority (20/NW/0231).

and serially performed upper respiratory tract swabs RT-PCR

testing, and when consented, collected blood samples for serial serology. Additionally, we also took swabs from

frequently touched household surfaces, and primary cases'

SARS-CoV-2 RNA. Our prospective study design and dense

longitudinal testing of contacts enabled us to maximally

sequencing of primary and secondary cases' SARS-CoV-2

isolates confirmed household transmission. Presence of

SARS-CoV-2 RNA on primary cases' hands, household

surfaces, and contacts' hands strongly correlated with

contacts' risk of infection. To our knowledge, this study

SARS-CoV-2 on candidate vectors of transmission. This study

also identified primary cases' hands, household surfaces, and

contacts' hands as potential vectors of transmission. Aerosol

This study, to our knowledge, is the first to identify a correlation

of contacts. These longitudinal empirical data from a real-world

between SARS-CoV-2 RNA on candidate vectors and infection

community setting substantially advance our understanding

of SARS-CoV-2 household spread, which is the setting of most

transmission globally. Our findings support interventions such

distancing, reducing direct contact, and use of masks to reduce

as frequent handwashing, surface cleansing, physical

provides the first empirical data linking detection of

transmission was not investigated in this study and

therefore cannot be excluded as a potential route of

Implications of all the available evidence

and contacts' hands, and tested them for presence of

capture secondary COVID-19 cases among household

contacts. Epidemiological linkage and whole-genome

SARS-CoV-2 PCR-positive, symptomatic primary cases (defined as the first detected PCR-positive case in the household) and their contacts were identified through the national contact tracing system (National Health Service Test & Trace) and invited to participate by UK Health Security Agency (UKHSA) staff between Aug 1, 2020, and March 31, 2021. The study period included the pre-alpha (September to December 2020) and alpha (B.1.1.7; December, 2020, to April, 2021) waves of SARS-CoV-2, when vaccination coverage was low and social distancing policies were in place. Contacts were

deemed household contacts if resident at the same address as their primary case.

In the INSTINCT study, primary cases and their household contacts were enrolled within 5 days (median of 3 days [IQR 2-4]) of primary case symptom onset and visited by study staff on the day of enrolment (day 0), day 7, and day 28, and on day 14 if any contact had tested PCR positive. URT swabs were collected at all visits as shown in figure 1A. On study day 4, participants took an additional self-taken URT swab. In the ATACCC study, household and non-household contacts were recruited within 5 days (median of 3 days [IOR 2-4]) of their primary case symptom onset to perform self-taken daily URT swabs for 14 days and, if still PCR positive, up to day 28 after enrolment; self-taken swabs were delivered by courier to the UKHSA laboratory daily. Primary cases were not enrolled. Most ATACCC contacts were recruited remotely; and the remaining contacts who consented to household visits underwent additional sampling according to figure 1. In all INSTINCT and visited ATACCC households, surface swabs (sampled from the most frequently touched surfaces in communal areas as reported by the household residents) and hand swabs were taken by trained nurses at enrolment (day 0) and day 7. Serology samples were taken on day 0, day 7, day 14, and day 28. All participants provided sociodemographic, clinical, and epidemiological data via questionnaires on the day of enrolment.

Participants were free to leave the study prematurely if they wished or to omit some sociodemographic characteristic information. Among the visited participants, not all agreed to give all types of samples (serological, surface swabs, and hand swabs).

Two contacts cohorts were defined for analysis: the full household contacts' cohort, consisting of all household contacts with longitudinal (\geq 3) URT samples, and the serology subcohort, consisting of the contacts within the full cohort who also had longitudinal serology data (ie, serological data available at day 0 and at least one other timepoint), were known to be unvaccinated before enrolment, and were seronegative at enrolment. Households with co-primary cases (concurrent PCR positivity) were excluded.



ATACCC=Assessment of Transmission And Contagiousness of COVID-19 in Contacts. Ct=cycle threshold. INSTINCT=Integrated Network for Surveillance, Trials and Investigations into COVID-19 Transmission. URT=upper respiratory tract. WGS=whole-genome sequencing. *WGS was possible for 35 of the 152 PCR-positive household contacts, where URT swabs from both the contact and their respective epidemiologically-linked primary case were PCR positive, available, and had a viral load of more than 200 copies per mL. †URT swabs from 21 (36%) of 59 contacts in the serology subcohort deemed infected underwent WGS analysis alongside URT swabs from their primary case (appendix p 12). ‡Insufficient longitudinal serological data among contacts who were PCR positive (no serology after day 7).



Contacts within the full household contacts' cohort were defined as being PCR positive (positive URT swab within the first 14 days after enrolment) or PCR negative if all URT swabs (\geq three) were negative throughout the first 14 days after enrolment. Hand-swab and surfaceswab PCR positivity was identically assigned (ie, 5 RNA copies per mL). Contacts within the serology subcohort were classified as infected if they were PCR positive or seroconverted within the first 14 days after enrolment, or both, and uninfected if they remained both PCR negative and seronegative. Further details on laboratory results definitions are available in the appendix (pp 2–4).

See Online for appendix

Study covariates considered were sex assigned at birth, age (continuous and grouped in categories), ethnicity (grouped as White and non-White), BMI (grouped as <18.5, 18.5 to <25.0, 25.0 to <30.0, and ≥ 30.0 kg/m²), presence of one or more comorbidities, and COVID-19 vaccination pre-enrolment. COVID-19 vaccination preenrolment was defined as receiving one or more doses before enrolment. Household contacts were defined as contacts who were co-resident with the primary case. Relationship status to the primary case was assessed with predetermined categories: partner, parent to child aged 16 years or younger, child aged 16 years or younger, or sibling in same bedroom; parent to child older than 16 years, child older than 16 years, or sibling without shared bedroom; and housemate or residential employee. We used the 2019 Index of Multiple Deprivation (IMD), the UK Government's official measure of relative deprivation at a small area level (average population of 1500) across England,¹⁴ to assess household deprivation according to postcodes. Participants gave written informed consent.

Procedures

URT swabs were analysed by RT-PCR at Imperial College London, London, UK (INSTINCT)15 and UKHSA Colindale, London, UK (ATACCC) with comparable sensitivity and specificity (appendix pp 2-3). Serology was performed using a hybrid double antigen binding assay (DABA), as previously described (appendix pp 3-4).16 A DABA level of less than 1 indicated seronegativity and a DABA level of 1 or more indicated seropositivity. A DABA level of less than 1 at day 0 that increased by 3 in the binding ratio in the first 28 days after participant enrolment (if PCR positive) or in the first 14 days after enrolment (if PCR negative) indicated seroconversion. Variant status was assigned through S-gene target failure status (ATACCC; appendix p 5) and whole-genome sequencing (WGS; INSTINCT; appendix pp 5-6). Libraries for WGS were generated with the EasySeq RT-PCR SARS-CoV-2 (novel coronavirus) WGS kit v1 or v2 (Nimagen, Nijmegen, Netherlands) and an iSeq 100 device was used for sequencing (Illumina, San Diego, CA, USA). Hand swabs and surface swabs were tested for RNA presence at the Department of Infectious Disease, Imperial College London, London, UK (appendix p 4).³ PCR-positive hand and surface swabs with a cycle threshold (Ct) of less than 33 were subjected to viral culture in a mammalian cell line-based in vitro assay (appendix pp 4–5). Swabs were maintained at 4°C for 36–72 h until culture was performed, reflecting the time for transportation from residents' homes and laboratory storage while RT-PCR was performed. Further assay details are found in the appendix (pp 2–6).

Statistical analysis

The primary outcome was infection among contacts, defined as contacts becoming PCR positive in the full household contact cohort, or infected in the serology subcohort, as defined earlier. All analyses were performed with the data available for each covariate used. To identify confounding risk factors for contacts' infection, univariable associations between PCR positive or infected versus PCR negative or uninfected cases according to different covariates were evaluated using a modified χ^2 test accounting for contacts' household clustering.¹⁷

Household infection rate was calculated as the proportion of contacts who became PCR positive or infected among all those tested.

To estimate the contribution of viral RNA presence on candidate vectors to contacts' risk of infection, adjusted relative risks (aRR) from multivariable, modified Poisson regression analyses were estimated, with an implemented generalised estimating equation model to account for household clustering and obtain robust standard errors.¹⁸ Binary indicators for infection or PCR positive versus PCR negative candidate vectors were used. The model was adjusted for significant demographic covariates and covariates associated with household characteristics identified with univariable analyses.

Analyses were performed on R version 4.0.5 (appendix p 8).

Role of the funding source

The funder of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report.

Results

620 contacts of PCR-confirmed SARS-CoV-2-infected primary cases were recruited in London from Aug 1, 2020, to March, 31, 2021. After excluding non-household contacts (n=138), contacts without URT samples for three or more timepoints (n=53), and contacts with a PCR-positive swab at a single time-point with high Ct value between consecutive PCR-negative swabs (n=15), 414 household contacts (from 279 households) with available serial URT PCR results were included in our analyses (full household contacts' cohort; figure 1; appendix p 9). 134 household contacts within the full cohort also had longitudinal serology data and were included in the serology subcohort.

No significant differences were observed in sex, age, ethnicity, or BMI between the 152 PCR-positive and 262 PCR-negative contacts from the full household contacts' cohort (table 1); although men had an increased risk of infection in the serology subcohort (p=0.047). In this predominantly vaccine-naive cohort, vaccinated contacts comprised a significantly higher proportion of PCR-negative cases (p=0.015) and participants with comorbidities were less likely to be PCR positive than those without (p=0.017; table 1).

Household-visited participants and remotely recruited participants had similar characteristics (appendix pp 13–15). The characteristics of primary cases who infected one or more of their recruited contacts were also similar to participants who did not infect their household contacts, except for age, with older primary cases resulting in more secondary infection events (appendix pp 16–17).

The overall household infection rate was 36.7% (95% CI 32.2-41.5) in the full cohort (table 2). 31 of the 152 PCR-positive contacts were defined contacts with incident infection as they became PCR positive after enrolment (as opposed to contacts with prevalent infection, who were PCR positive at enrolment), hence the household infection rate at the day of enrolment was lower than the overall HIR (29.2%, 25.0-33.8; data not shown). The number of days from primary case symptom onset to recruitment did not differ between contacts with prevalent and incident infection (median 3.00 days [IQR 2.00-4.00] vs 3.00 days [3.00-4.00]); Mann-Whitney U test p=0.98; data not shown). In the serial serology subcohort, after removal of 17 previously

	Full household	contacts' coho	rt (n=414)		Serial serology	/ subcohort (n=1	34)	
	Total (n=414)	PCR-positive (n=152)	PCR-negative (n=262)	χ² test p value*	Total (n=134)	Infected with SARS-CoV-2† (n=59)	Not infected with SARS-CoV-2 (n=75)	χ² test p value*
Sex assigned at birth				0.77				0.047
Male	200 (48%)	75 (49%)	125 (48%)		70 (52%)	37 (63%)	33 (44%)	
Female	214 (52%)	77 (51%)	137 (52%)		64 (48%)	22 (37%)	42 (56%)	
Median age, years (IQR)	34 (25–50)	37 (25-49)	33 (24–51)	0.90	36 (24–51)	36 (25-47)	36 (24–52)	0.54
Age group				0.36				0.36
≤16 years	45 (11%)	19 (12%)	26 (10%)		12 (9%)	5 (8%)	7 (9%)	
17–29 years	124 (30%)	39 (26%)	85 (32%)		43 (32%)	19 (32%)	24 (32%)	
30–49 years	141 (34%)	60 (39%)	81 (31%)		43 (32%)	23 (39%)	20 (27%)	
50–69 years	100 (24%)	33 (22%)	67 (26%)		35 (26%)	11 (19%)	24 (32%)	
≥70 years	4 (1%)	1 (1%)	3 (1%)		1(1%)	1(2%)	0 (0%)	
Ethnicity				0.53				0.17
White	325 (78%)	122 (80%)	203 (77%)		112 (84%)	51 (86%)	61 (81%)	
Non-White	73 (18%)	24 (16%)	49 (19%)		16 (12%)	4 (7%)	12 (16%)	
Data missing	16 (4%)	6 (4%)	10 (4%)		6 (4%)	4 (7%)	2 (3%)	
BMI				0.12				0.28
<18.5 kg/m²	8 (2%)	2 (1%)	6 (2%)		1(1%)	0	1(1%)	
18·5 to <25·0 kg/m²	163 (39%)	47 (31%)	116 (44%)		48 (36%)	16 (27%)	32 (43%)	
25·0 to <30·0 kg/m ²	111 (27%)	45 (30%)	66 (25%)		41 (31%)	21 (36%)	20 (27%)	
≥30·0 kg/m²	59 (14%)	26 (17%)	33 (13%)		13 (10%)	4 (7%)	9 (12%)	
Data missing	73 (18%)	32 (21%)	41 (16%)		31 (23%)	18 (30%)	13 (17%)	
Presence of comorbidities‡				0.017				0.0038
Yes	105 (25%)	28 (18%)	77 (29%)		22 (16%)	3 (5%)	19 (25%)	
No	298 (72%)	121 (80%)	177 (68%)		107 (80%)	54 (92%)	53 (71%)	
Data missing	11 (3%)	3 (2%)	8 (3%)		5 (4%)	2 (3%)	3 (4%)	
COVID-19 vaccination pre-enrolment§				0.015				NA
Yes	25 (6%)	3 (2%)	22 (8%)		0	0	0	
No	382 (92%)	144 (95%)	238 (91%)		134 (100%)	59 (100%)	75 (100%)	
Data missing	7 (2%)	5 (3%)	2 (1%)					

Data are n (%) unless otherwise indicated. NA=not applicable. *p values are given for each covariate; χ^2 test accounting for household clustering was used for all categories for which data are available, except for median age, for which the Mann-Whitney test was used. †Contacts were considered infected if PCR-positive or, in the serology subcohort, if PCR-positive or seroconverted. ‡Presence of comorbidities was considered if the participant had one or more conditions (appendix p 7). \$Vaccination pre-enrolment was defined as having received one or more doses before day 0.

Table 1: Baseline demographic and clinical characteristics of household contacts exposed to PCR-confirmed SARS-CoV-2 primary cases, in the full household contacts' cohort and in the serial serology subcohort.

	Full household contac	ts' cohort (n=414)		Serial serology sub-col	hort (n=134)	
	Number of PCR- positive/total number	Household infection rate (95% CI)	χ² test p value*	Number infected/total number	Household infection rate (95% CI)	χ² test p value*
Overall	152/414	36.7% (32.2-41.5)		59/134	44.0% (35.9–52.5)	
Contacts' exposure to the primary case						
Relationship status to the primary case			<0.0001			0.0016
Partner, parent to child aged ≤16 years, child aged ≤16 years, or sibling in same bedroom	85/174	48.8% (41.5–56.2)		36/56	64·3% (51·2–75·5)	
Parent to child aged >16 years, child aged >16 years, or sibling without shared bedroom	39/126	30.9% (23.5–39.5)		11/44	25.0% (14.6–39.4)	
Housemate or residential employee	12/71	16.9% (9.9–27.2)		6/21	28.6% (13.8–50.0)	
Data missing	16/43			6/13		
Primary case SARS-CoV-2 variant			0.0047			0.0016
Alpha (B.1.1.7)	56/108	51.8% (42.5-61.0)		25/34	73.5% (56.9–85.4)	
Pre-alpha	31/109	28.4% (20.8-37.5)		11/38	28.9% (17.0-44.8)	
Data missing	65/197			23/62		
Primary case maximum-measured URT RNA viral load, viral copy number per mL			0.83			0.64
5 to ≤4500	36/82	43.9% (33.7-54.7)		24/47	51.1% (37.2-64.7)	
4500 to ≤12 000 000	35/76	46.0% (35.3-57.2)		25/44	56.8% (42.2-70.3)	
Data missing	81/256			10/43		
Contacts' household exposure characteristics						
Number of people resident in the household, including the primary case			0.22			0.44
2 people	25/66	37.9% (27.1–49.9)		12/21	57.1% (36.5-75.5)	
3-4 people	90/224	40.2% (34.0-46.7)		29/71	40.8% (30.2-52.5)	
5 or more people	26/96	27.1% (19.2–36.7)		16/35	45.7% (30.5–61.8)	
Data missing	11/28			2/7		
Ratio of the number of people resident in the household per number of bedrooms			0.065			0.23
≤1 resident per bedroom	70/217	32.2% (26.4–38.7)		27/69	39.1% (28.5–50.9)	
>1 resident per bedroom	70/161	43.5% (36.1–51.2)		30/58	51.7% (39.2–64.1)	
Data missing	12/36			2/7		
By household decile of Index of Multiple Deprivation			0.68			0.51
More deprived deciles, 1 to 5	71/200	35.5% (29.2-42.3)		29/71	40.8% (30.2-52.5)	
Least deprived deciles, 6 to 10	81/214	37.8% (31.6-44.5)		30/63	47.6% (35.8-59.7)	

Household infection rate is given as the proportion of PCR-positive contacts in the full household contact cohort (n=414), and as the proportion of infected (PCR-positive or seroconverted) contacts in the serial serology subcohort (n=134). Primary case maximum-measured URT RNA viral load categories were divided to give similar number of contacts. URT=upper respiratory tract. *p values for each covariate were calculated with the χ^2 test accounting for household clustering, applying for all categories where data was available.

Table 2: SARS-CoV-2 household infection rate in the full household contacts' cohort and in the serial serology subcohort, stratified by type of exposure to the primary case and household characteristics

infected contacts (baseline seropositive) and reclassifying four PCR-negative contacts as incident infections (seroconverters between day 0 and day 14), the overall household infection rate was $44 \cdot 0\%$ ($35 \cdot 9 - 52 \cdot 5$; table 2). Contacts' risk of infection increased with a closer relationship to the primary case, with the household infection rate ranging from $16 \cdot 9\%$ (95% CI $9 \cdot 9 - 27 \cdot 2$) for non-related housemates to $48 \cdot 8\%$ ($41 \cdot 5 - 56 \cdot 2$) for primary-case partners or bedroom-sharing relatives ($p < 0 \cdot 0001$; table 2). The total number of residents per household was not associated with increased risk of infection; however, the point estimate proportion of infected contacts was higher in households with more than one resident per bedroom than in those with one resident per bedroom, although this difference was not significant (p=0.065; table 2). Household deprivation index (IMD deciles) did not significantly differ between the 152 PCR-positive and 262 PCR-negative contacts. Household infection rate was higher in contacts exposed to alpha (B.1.1.7) SARS-CoV-2 versus pre-alpha SARS-CoV-2 variants (table 2; 51.8% [42.5-61.0] *vs* 28.4% [20.8-37.5] in the full household contacts' cohort; p=0.0047 and 73.5% [56.9-85.4] *vs* 28.9% [17.0-44.8] in the serology subcohort; p=0.0016).

All PCR-positive surface swabs and primary cases' hand swabs, and most of the PCR-positive hand swabs from the contacts (16 [72%] of 22), were positive on the day of enrolment. This RNA detection on surface and

hand swabs was simultaneous in time with the first detection of SARS-CoV-2 in prevalent PCR-positive contacts' respective URT swabs, at day 0. Of the 31 PCR-positive contacts with incident infection (ie, with initially PCR-negative URT swabs), 14 had household surface samples collected, of which three were positiveall were positive at day 0, before the respective contacts' URT swabs becoming PCR positive. 19 of 31 contacts with incident infection had hand swabs collected, of which seven were positive. In three contacts, hand swabs were already PCR positive before their respective URT swabs becoming positive. In the remaining four contacts, hand swabs were positive at the second time-point (day 7); hence, it is not possible to temporally resolve whether these contacts' hands became PCR positive before or after their URT swabs became PCR positive.

Although the maximum URT RNA viral load measured in primary cases did not correlate with the household infection rate (p=0.83; table 2; appendix p 10), primary cases with SARS-CoV-2 RNA detected on hand swabs (21 [24%] of 88 primary cases with hand swabs collected) had significantly higher URT RNA viral load than did those whose hand swabs were SARS-CoV-2 PCR-negative (p=0.031; appendix p 11). Presence of SARS-CoV-2 RNA on primary cases' hands was associated with SARS-CoV-2 infection in their contacts (aRR 1.70, 95% CI 1.24-2.31; table 3). Presence of SARS-CoV-2 RNA on primary cases' hands was also associated with viral RNA detection on their corresponding contacts' hands (3.11, 1.21-8.05; appendix pp 18-19), and presence of SARS-CoV-2 on contacts' hands was associated with their infection status—ie, having a PCR-positive UTR sample (2.06, 1.57-2.69; table 3). Among the three contacts with incident infection (with a PCR-negative initial URT sample) and positive hand swabs at day 0 before URT PCR positivity, SARS-CoV-2 RNA was first detected on their hands 1, 2, and 3 days before their first PCR-positive URT sample. One of these hand swabs was also positive on viral culture (attempted on ten hand swabs) and yielded sufficient RNA for WGS, which revealed indistinguishable sequence from that from the PCRpositive URT swab collected 1 day later. A primary case sample was unavailable for comparison.

33 (10%) of 344 surface samples from 103 households were PCR positive on day 0 (appendix p 20). SARS-CoV-2 presence on one or more household surfaces was associated with increased risk of infection among contacts (aRR 1.66, 95% CI 1.09–2.55; table 3). Contacts were also substantially more likely to have detectable SARS-CoV-2 RNA on their hands if household surfaces were contaminated (3.77, 1.27–11.14; appendix pp 18–19). However, presence of viral RNA on surfaces did not correlate with its detection on primary cases' hands (1.30, 0.53–3.15; data not shown). No household surface swabs were positive for viral culture.

Each of the vectors of transmission identified in the full household contacts' cohort (primary cases' hands,

	Full household	contacts' cohor	t (n=414)			Serial serology	subcohort (n=134)			
	Household cont	tacts			Multivariable analysis	Household con	tacts			Multivariable analysis
	Total (n=414)	PCR positive (n=152)	PCR negative (n=262)	χ² test p value*	Adjusted relative risk (95% CI)	Total (n=134)	Infected with SARS-CoV-2† (n=59)	Not infected with SARS-CoV-2 (n=75)	χ² test p value*	Adjusted relative risk (95% CI)
rry cases' hand-swab ·CoV-2 PCR status	:	:	:	0.075	:	:	:	:	0.0037	:
itive	33 (8%)	20 (13%)	13 (5%)	:	1.70 (1.24-2.31)	19 (14%)	17 (29%)	2 (3%)	:	1.67 (1.23–2.28)
jative	123 (30%)	49 (32%)	74 (28%)	:	1 (ref)	65 (49%)	31 (52%)	34 (45%)	:	1 (ref)
a missing	258 (62%)	83 (55%)	175 (67%)	:	:	50 (37%)	11 (19%)	39 (52%)	:	:
tcts' hand-swab SARS-CoV-2 tatus	:	:	:	<0.0001	:	:	÷	÷	<0.0001‡	:
itive	22 (5%)	20 (13%)	2 (1%)	:	2.06 (1.57-2.69)	16 (12%)	16 (27%)	0 (%0) 0	:	1.51 (1.07-2.14)
jative	189 (46%)	59 (39%)	130 (50%)	:	1 (ref)	103 (77%)	40 (68%)	63 (84%)	:	1 (ref)
a missing	203 (49%)	73 (48%)	130 (50%)	:	:	15 (11%)	3 (5%)	12 (16%)	:	:
onmental surface-swabs .CoV-2 PCR status	:	:	:	0.044	:	:	÷	÷	0.0069	:
itive	23 (5%)	13 (9%)	10 (4%)	:	1.66 (1.09–2.55)	15 (11%)	12 (20%)	3 (4%)	:	1.78 (1.21–2.62)
jative	160 (39%)	50 (33%)	110 (42%)	:	1 (ref)	94 (70%)	34 (58%)	60 (80%)	:	1 (ref)
a missing	231 (56%)	89 (59%)	142 (54%)	:	:	25 (19%)	13 (22%)	12 (16%)	:	:
e n (%) unless otherwise indicat d at birth, comorbidity status, v ting for household clustering, a	ed. Relative risk est accination pre-enr pplying for all cate <u>c</u>	timates from the r olment status, reli gories where data	nultivariable analys. ationship status to t are available. †Cont	es were calculat. the primary case acts in the seria	ed with a modified Poisson re , and ratio number of resider serology subcohort were de	egression using ro ats in the househc emed infected if F	bust standard errors, acco old to number of bedroom CR-positive or if they sero	unting for household clu s. *p values for each cove converted. ‡Fisher's exac	stering, and adj ariate were calcu ct test was used i	usting for contact's sex lated with the χ^2 test to calculate p values.
:: Detection of SARS-CoV-2 (*** subcohort	on environmenta	al surfaces, and c	on the hands of th	ie contact and	primary case, by PCR-sta	tus in the full h	ousehold contacts' coh	ort, and by infection s	tatus in the se	rology analysis

Po: Ne Dai Conta Pos Neg Dat Envir household surfaces, and contacts' hands) was corroborated in the serology subcohort, with similar correlations with transmission (table 3; appendix pp 18–19).

Of the 262 contacts whose URT samples remained PCR negative, 20 were linked to one or more PCR-positive swabs from household surfaces or residents' hands. Of the 152 contacts with PCR-positive URT samples, 42 were linked to one or more positive swabs from household surfaces or residents' hands (table 3, figure 2). We found no significant differences in the quantified RNA viral load in samples from any of the candidate vectors between the contacts with PCR-positive and PCR-negative URT samples (Figure 2).

18 of the 20 contacts with PCR-negative URT swabs that were linked to a PCR-positive household surface or residents' hand sample had serological data available, of whom 12 (67%) were seropositive at baseline (half of whom were recently vaccinated, including the two contacts whose URT samples remained PCR negative despite having PCR-positive hand swabs). By contrast, of the 37 (88%) of 42 contacts with PCR-positive URT samples who had serological results available, only five (13%) were seropositive at baseline (figure 2), significantly fewer than the contacts with PCR-negative URT samples linked to PCR-positive household surface or residents' hand samples (χ^2 test p=0.0002).

Sensitivity analysis assessed the effect of incomplete recruitment of contacts within households on household infection rate (appendix p 9). 200 contacts were from 115 fully recruited households (ie, households where all



Figure 2: Measured SARS-CoV-2 RNA viral load in PCR-positive environmental samples, stratified by contacts' URT PCR status and baseline serological status

Only contacts linked to one or more PCR-positive environmental samples are depicted. (A) Primary cases' handswab RNA viral load (median 82-09 copies per mL, IQR 46-09–1082-29). (B) Household surface-swab RNA viral load (226-58, 101-38–602-51). (C) Contacts' hand-swab RNA viral load (117-66, 21-75–244-39). The measured environmental viral loads do not differ by contacts' URT PCR status, as shown by the p values in each panel (Mann-Whitney U test). All positive primary case hand swabs and surface swabs were collected on day 0, whereas contacts' hand swabs were positive on day 0 (n=16) and day 7 (n=6). Environmental samples with more than 5 RNA copies per mL were considered PCR positive. URT=upper respiratory tract.

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residents were recruited), of which 72 were PCR positive and 128 PCR negative, giving a household infection rate of $36 \cdot 0\%$ (95% CI 29.7–42.9), very similar to that for the full household contacts' cohort ($36 \cdot 7\%$; table 2).

URT samples were collected from 72 of 152 PCR-positive household contacts alongside their epidemiologically linked primary case. Of these, 35 contacts and their 25 respective primary cases had adequate RNA viral load in their URT samples (ie, >200 genome copies per mL of virus transport media) to undergo WGS (figure 3; appendix p 12). Phylogenetic analysis indicated that all isolates clustered within their respective households (figure 3). In 20 of the 25 households, isolates from primary cases and contacts isolates were indistinguishable on WGS (figure 3). In three households, isolates from contacts had one or two additional non-synonymous mutations compared with isolates from the primary cases, and in two households, the isolates from the primary cases isolate had one non-synonymous mutation not detected in their contacts. These data indicate that the primary case-contact pairs were most likely infected with the same SARS-CoV-2 strain,19 strongly supporting household transmission as the source of infection in all infected contacts sequenced.

Discussion

Rigorous assessment of risk factors and vectors of transmission necessitates maximising the capture of transmission events prospectively in epidemiologically well defined households. Our study design maximally captured transmission events by enrolling asymptomatic and symptomatic, recently exposed household contacts of newly diagnosed primary cases with COVID-19 and proactively detecting infection by repeated URT PCR testing. In a subcohort, longitudinal serology was conducted to detect contacts whose URT samples were PCR negative and seroconverted and exclude baseline seropositive, previously infected contacts. We also simultaneously tested for the presence of SARS-CoV-2 RNA on frequently touched household surfaces and residents' hands. Although other studies, with smaller sample sizes, have reported the presence of SARS-CoV-2 RNA on surfaces in households or other settings,^{3,4,20} this study is, to our knowledge, the first to identify correlations between SARS-CoV-2 RNA on candidate vectors with contacts' infection risk. These longitudinal empirical data from a real-world community setting with rigorously defined infection outcomes substantially advance our understanding of SARS-CoV-2 household spread, the setting of most transmission globally.

We report a household infection rate of 51.8% in the full household contacts' cohort and 73.5% in the serology subcohort for contacts exposed to alpha variant-infected primary cases, which is one of the highest overall household infection rates for SARS-CoV-2 reported in predominately unvaccinated and previously unexposed contacts.¹ This finding probably reflects our rigorous,



Figure 3: Genomic evidence of SARS-CoV-2 transmission among households

Genomic analysis of 25 households (comprising 35 contacts and 25 of their respective epidemiologically linked primary cases) of non-synonymous mutations within the SARS-CoV-2 genome, to assess whether the same or different strains are present within the household. Genomic analyses were performed where upper respiratory tract swabs with viral loads of more than 200 copies per mL were available for both primary cases and contact(s).

intensive sampling approach, which captured a higher proportion of transmission to susceptible contacts than previous studies.^{21–23}

Unlike some previous studies,^{22,24} but consistent with others,²⁵ primary case SARS-CoV-2 RNA viral load in URT samples did not correlate with contacts' risk of infection in our cohort. Interestingly, however, primary cases' RNA viral load in URT samples did predict their risk of having a SARS-CoV-2 PCR-positive hand swab, which in turn was associated with transmission to their respective contacts. Presence of viral RNA on frequently touched household surfaces strongly associated with contacts' risk of having a PCR-positive hand swab and predicted contacts' risk of infection. Moreover, presence of SARS-CoV-2 RNA on contacts' hands correlated with presence of viral RNA on their respective primary cases' hands and with contacts' own risk of SARS-CoV-2 infection.

Interestingly, in six contacts with incident infection, three had SARS-CoV-2 PCR-positive hand swabs, and another three had PCR-positive surfaces in their households, before their URT swabs became PCR positive. Because the collection of URT swabs involved sampling both the nasal turbinates and pharynx, it is unlikely that SARS-CoV-2 infection was missed on earlier samples due to anatomical compartmentalisation.²⁶ SARS-CoV-2 was cultured from one of these contacts' hand swabs, showing the presence of live, infectious virus on the hands of this individual. WGS confirmed that the viral isolates from the contact's hand and URT were indistinguishable. Conversely, in non-incident contacts who already had PCR-positive URT samples at enrolment, the corresponding PCR-positive hand and household surface swabs were also consistently positive at enrolment.

Among all contacts with one or more positive candidate vector samples, no differences were observed in the RNA viral load quantified on these vectors between contacts with PCR-positive and PCR-negative URT samples, suggesting that the source of the SARS-CoV-2 RNA on these vectors was unlikely to be retrograde contamination from the contacts' URT. Notably, contacts with PCRnegative URT samples and with viral RNA detected on their hands, their primary cases' hands, or household surfaces, predominantly had pre-existing immunity, as evidenced by being seropositive at enrolment. By contrast, very few contacts with PCR-positive URT samples linked to positive hand swabs or surface swabs in their households were seropositive at enrolment. This finding suggests that the presence of SARS-CoV-2 RNA on candidate vectors linked to uninfected contacts with PCR-negative URT samples might not have resulted in infection because of their pre-existing immunity.

The potential modes of SARS-CoV-2 transmission are the subject of ongoing scientific controversy.^{2,27} To our knowledge, our data are the first empirical evidence to correlate presence of SARS-CoV-2 RNA on candidate vectors with contacts' risk of infection. However, our findings do not prove causality. SARS-CoV-2 spread is believed to be predominantly through airborne transmission,28-30 which must be the sole route in several settings, including those involving long-range transmission.² The transmission we observed could therefore have been exclusively airborne and the presence of SARS-CoV-2 RNA on residents' hands and household surfaces might reflect bystander contamination by large respiratory droplets that might be generated alongside infectious aerosols that mediate transmission. We did not systematically sample air in the households so we cannot exclude this possibility.

To demonstrate causality of either fomite or airborne transmission, a highly controlled artificial experiment would be required, in which SARS-CoV-2-naive, healthy, sentinel participants would be confined in an environment previously contaminated by an infectiouscase. Experimental interventions testing the effects of complete air exchange or decontamination versus surface decontamination before the participants' confinement would then be correlated with their subsequent infection status. Although such an experimental design might confirm the route of transmission, it would not reflect the real-world setting of community transmission with its natural spectrum of environmental conditions and host factors.

Only one of our candidate vector samples, a contact's hand swab, was positive for viral culture. The very high

proportion of SARS-CoV-2 RNA-positive but viral culture-negative swabs probably reflects viral degradation during the 36–72 h interval to transport the swabs from participants' homes and laboratory storage, while RT-PCR was performed, before they were subjected to in vitro cell culture. This explanation is supported by the recent observation that similar swabs processed in the same laboratory with the same cell culture assay by the same scientist, but within 12 h, yielded a much larger proportion of culture-positive samples.³¹

Our study has several limitations. We cannot exclude that in a proportion of primary case-contact pairs where the prevalent infected contact might have been asymptomatic or pre-symptomatic, the contact might have been the initial source of transmission, or that the contact might have been infected from outside of the household. These scenarios are unlikely for most of the households, because our assignment of casecontact pairs is underpinned by epidemiological linkage between each primary case and their respective contact(s), and further corroborated by WGS of SARS-CoV-2 isolates from primary cases and their contacts in all 25 households where this was possible. Similarly, although some of the contacts' infections could possibly have been tertiary rather than secondary to the primary case, we consider this unlikely because contacts were enrolled on average 3 days after symptom onset of the primary case. The probability of infection arising from outside the household was low because London was in a state of lockdown during our study, resulting in very limited social or occupational contact outside of households. A corollary of this point, however, is that our household infection rate estimate might not be generalisable to households in settings where societal non-pharmaceutical interventions are not in force with correspondingly less contact time between cohabitants.

We did not systematically collect behavioural information on hygiene practices and physical distancing, as preventative measures typically took place after primary case symptom onset, when exposure had already occurred, limiting the effect of such practices on reducing contacts' risk of infection. Additionally, the collection of such behavioural data would have been biased by the participants' conditioning of the underlying circumstances (eg, lockdown and participation in an observational study) and would not represent their usual practices.

The households that consented to participate might not be representative of the general population because non-White ethnicities and older age groups were underrepresented. In some households, only a proportion of cohabitants were recruited; however, infection rate in households where all contacts were recruited approximated the infection rate in the full household contact cohort. Although the study could be biased by the difference between household-visited contacts and remotely recruited contacts, these groups were similar. Our study began when the pre-alpha lineages were dominant in London and continued until the alpha variant became dominant. Our results cannot therefore be directly extrapolated to the more recent and more infectious variants. Interestingly, some recent evidence suggests that the omicron variant BA.1 is significantly more stable than ancestral SARS-CoV-2 on different surfaces, potentially increasing its propensity for transmission by the fomite route,^{27,32} whereas other evidence suggests increased likelihood for aerosol transmission relative to earlier variants.^{31,34}

Our study provides the first empirical evidence to correlate presence of SARS-CoV-2 on candidate vectors with risk of infection in household contacts. These findings from a real-world community setting substantially advance our understanding of SARS-CoV-2 household transmission, the setting of most transmission globally. Our results also have practical implications and support interventions such as frequent handwashing, surface cleansing, physical distancing, reducing direct contact, and use of masks to curb transmission in households. As governments worldwide develop policies to manage successive new COVID-19 variants that transmit widely through vaccinated populations,10 alternative interventions to impede transmission are urgently needed. The straightforward public health interventions and messaging underpinned by our evidence provide a timely, pragmatic component of the future toolkit for living safely with COVID-19.

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Contributors

RK, RV, JF, TDP, SHak, and AL designed the study and data collection methodology. RV, JD, JLB, JF, RK, SHak, EC, JSN, MRT-W, ND, AKoy, JC, VQ, SHam, EMcD, CL, KT, JS, SB, SE, LW, SN, MD, CT, GM, HC, AKon, RT, WSB, GPT, and MZ did administrative and technical study logistics. ND, AKov, CT, ME, GM, AVK, HC, MGW, KIM, and ILB acquired data. LM, CR, PW, AB, SD, PSF, JP, GPT, and MZ analysed the RT-PCR data. EP, IJDW, FM, MOM, and RT analysed the serology data. JZ and WSB analysed the environmental (surface swabs and hand swabs) data. CA, MAC, LM, GPT, and PSF analysed the sequencing data and S-gene target failure data. ND, NMF, and AL conceived the work and design of analyses. ND and AKoy analysed the data. ND, AKoy, TDP, SHak, AS, and AL interpreted the data. ND, AKoy, SHak, and AL drafted the manuscript. SHak, JF, TDP, HH, AS, EC, and AL reviewed the manuscript. AL is the principal investigator for INSTINCT. AL and JD are the principal investigators for ATACCC. ND and AKoy directly accessed and verified the underlying data of the study. All authors contributed important intellectual content during manuscript drafting or revision and accept accountability for the overall work by ensuring that questions pertaining to the accuracy or integrity of any portion of the work are appropriately investigated and resolved. All authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Declaration of interests

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Data sharing

An anonymised, deidentified version of the dataset can be made available upon request to the corresponding author to allow all results to be reproduced.

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