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A review on measurements of SARS-CoV-2 genetic material in air in outdoor and indoor environments: Implication for airborne transmission

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HIGHLIGHTS

- 73 papers furnished 77 datasets of detection/quantification of SARS-CoV-2 in air.
- 11.7% of studies are in outdoor, 75.3% in hospitals, and 13% in community public indoors.
- Average positivity rate was larger in hospital compared to outdoors and public indoor sites.
- Contamination of surfaces was more frequent than air but with a lower positivity rate.
- SARS-CoV-2 RNA concentrations in air follows outdoors < public indoors < hospitals.

ABSTRACT

Airborne transmission of SARS-CoV-2 has been object of debate in the scientific community since the beginning of COVID-19 pandemic. This mechanism of transmission could arise from virus-laden aerosol released by infected individuals and it is influenced by several factors. Among these, the concentration and size distribution of virus-laden particles play an important role. The knowledge regarding aerosol transmission increases as new evidence is collected in different studies, even if it is not yet available a standard protocol regarding air sampling and analysis, which can create difficulties in the interpretation and application of results. This work reports a systematic review of current knowledge gained by 73 published papers on experimental determination of SARS-CoV-2 RNA in air comparing different environments: outdoors, indoor hospitals and healthcare settings, and public community indoors. Selected papers furnished 77 datasets: outdoor studies (9/77, 11.7%) and indoor studies (68/77, 88.3%). The indoor datasets in hospitals were the vast majority (58/68, 85.3%), and the remaining (10/68, 14.7%) were classified as community indoors. The fraction of studies having positive samples, as well as positivity rates (i.e. ratios between positive and total samples) are significantly larger in hospitals compared to the other typologies of sites. Contamination of surfaces was more frequent (in indoor datasets) compared to contamination.
of air samples; however, the average positivity rate was lower compared to that of air. Concentrations of SARS-CoV-2 RNA in air were highly variable and, on average, lower in outdoors compared to indoors. Among indoors, concentrations in community indoors appear to be lower than those in hospitals and healthcare settings.

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1. Introduction

Since the beginning of the global pandemic in 2020, several studies raised a scientific debate on the possible role of airborne transmission of the disease in the spread of COVID-19 (Contini and Costabile, 2020; Domingo et al., 2020; Ishmatov, 2021; Klompas et al., 2020; Morawska and Cao, 2020; Prather et al., 2020; Ram et al., 2021). This mechanism of transmission could arise from virus-laden coarse and fine aerosols emitted by infected individuals during cough, sneezes, respiration, speaking, singing, and shouting. These could remain suspended in air, following different dynamics in indoor and outdoor environments, and being potentially inhaled by other susceptible individuals (Allen and Marr, 2020; Asadi et al., 2020; Belosi et al., 2021; Borouiba, 2020; Tang et al., 2021).

Several parameters are important to determine risks of airborne transmission: concentration and size distribution of virus-laden particles; fraction of infectious (viable) virus in air; minimum dose necessary to transmit infection to a susceptible individual. The first two parameters are depending on meteorological conditions (with differences between indoor and outdoor), on the dynamics of air currents and on physical-chemical properties of droplets (Niazi et al., 2021a, 2021b; Ratnesar-Shumate et al., 2020); the third parameter is influenced by specific vulnerabilities of susceptible individuals and vary strongly (Buonanno et al., 2020). Knowledge regarding airborne transmission of SARS-CoV-2 is continuously evolving as new evidence accumulates. This knowledge would directly impact on policy decisions regarding adequate mitigation measures to be implemented for efficient reduction of COVID-19 spread (Morawska and Cao, 2020; Morawska and Milton, 2020).

Mini-reviews about air detection methods for coronavirus in general and specific for SARS-CoV-2 have been published discussing problems and controversies, showing that more studies are necessary to find the method with best performance for SARS-CoV-2 detection in air and that it would be useful to develop a standard procedure for air sampling and analysis (Borges et al., 2021; Pena et al., 2021; Rahmani et al., 2020; Ratnesar-Shumate et al., 2021; Robotto et al., 2021; Yun et al., 2020). Other reviews (Heneghan et al., 2021; Tang et al., 2020) focused on the role of airborne transmission concluding that SARS-CoV-2 genetic material (RNA) is observed intermittently in air in different indoors environments and that the lack of detailed information on recoverable virus cultures prevents solid conclusions on the weight of airborne transmission. Finally, some review papers are focused on results, available during the first wave of pandemic during 2020, relative to detection of SARS-CoV-2 genetic material in air in hospital settings and working places finding that about 6% of air and surface samples in hospital were positive, although the data is very limited for non-healthcare settings and that only a few of the studies report quantification of concentrations in air (Anand et al., 2021; Birgand et al., 2020; Cherrie et al., 2021).

This work is aimed at presenting a systematic review of current knowledge, from the beginning of pandemic and until 31/08/2021, regarding identification/quantiﬁcation of SARS-CoV-2 RNA in airborne samples comparing different sites: outdoor sites, indoors in hospitals and healthcare settings, and community indoor locations. The analysis will also investigate positivity rates and concentrations comparing the different environments and the current methodologies used for samples collection and analysis. Conclusions about risks of airborne transmission and efﬁciency of mitigation measures are included, together with a discussion of the aspects that need further studies.

2. Methodology

The records used in this review include paper published since the start of COVID-19 pandemic until 31/08/2021. Published papers were selected from the Web of Science, Scopus, and Google Scholar databases using two search strings: “covid-19 and airborne” and “SARS-CoV-2 and airborne transmission”. This gave to 945 entries plus an additional 15 entries that were obtained from different sources, mainly speciﬁc searches and reading of the authors. Fig. 1 reports the ﬂowchart summarising the identiﬁcation and selection of the records using the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement (Moher et al., 2009; Page et al., 2021). After the exclusion of 305 duplicate records, the entries were screened to select records complying with the goal of this review, i.e. studies containing analysis of the presence/concentration of SARS-CoV-2 genetic material through measurements in both indoor and outdoor environments. Successively, the papers were assessed to select cases based on active air stationary sampling (i.e. excluding passive sampling and samples collected with personal samplers), irrespective of the approach/equipment used, and in which the identiﬁcation/quantiﬁcation of SARS-CoV-2 RNA traces was done by polymerase chain reaction (PCR). In this step only original studies reporting measurement results were maintained, i.e. review papers and commentaries were excluded. This left a ﬁnal number of 73 papers included in this review and six of them were preprints.

Fig. 2 shows the geographical distribution of the 73 studies that were performed in 22 different Countries. The majority of the studies (26.0%) were done in China, in Europe (24.6%), and in North America (13.7%).
Limited or no information is available from South America, Africa, India, and Oceania.

The selected papers furnished 77 datasets because three papers (Hu et al., 2020; Liu et al., 2020; Passos et al., 2021) included outdoor and indoor datasets, and one paper (Habibi et al., 2021) included datasets collected in different typologies of indoor environments. The datasets were separated into two categories: outdoor studies (9/77, 11.7%) and indoor studies (68/77, 88.3%). The indoor datasets were additionally separated into two groups: datasets dealing with measurements in hospitals, health care structures, and quarantine areas, that were the vast majority (58/68, 85.3%), and studies dealing with measurements taken at indoor community sites (10/68, 14.7%) such as commercial centres and markets, schools and universities, pharmacies, hair salon, stations, and vehicles of public transport. In the latter category it has been included a study on a mink farm (de Rooij et al., 2021a), in a meat processing plant (de Rooij et al., 2021b), in the Diamond Princess cruise ship (Yamagishi et al., 2020), and in two houses of a residential building in China (at Guangzhou) by Xie et al. (2020).

The different studies selected were analysed to extract the main information and results regarding: the methodology used for air sampling; the methodology used for RNA detection; the number of positive samples found in air and the eventual other results found on environmental samples (surface swabs) collected; the concentrations found, when reported, in air samples.

Fig. 1. Flowchart of the identification, screen, and assessment of the records included in this review according to PRISMA statement (Page et al., 2021).

Fig. 2. Geographical distribution of the 73 studies included in this review.
3. Results

The first analysis was made to estimate the number of datasets that reports negative (i.e. SARS-CoV-2 not detected in any of the air samples) and positive (at least one sample found positive to SARS-CoV-2). Results show (Fig. 3) that the majority of the datasets refers to hospitals and care settings where the datasets reporting positive samples were 35 out of 58 (60.3%). Significantly lower percentages were observed for datasets in outdoors (3/9, 33.3%) and in indoor community environments (3/10, 30%) even if statistics is more limited for these last two categories.

3.1. Results in outdoor environments

The results of the papers dealing with measurements of airborne SARS-CoV-2 RNA traces in outdoor sites are reported in Table 1. Table 1 reports a summary of the methodology used and the results found in the different papers that include outdoor measurement in public areas. Results are available for different areas in Europe (Italy, Germany, and Spain), China, Turkey, and Brazil. All studies are based on samples collected in different periods of the first wave of pandemic. In three cases, the study includes both outdoor and indoor measurements (Hu et al., 2020; Liu et al., 2020; Passos et al., 2021). Five studies out of the eight available (62.5%) reported negative results with SARS-CoV-2 concentrations lower than the detection limit (LOD). Of the three studies reporting positive samples, one (Setti et al., 2020) does not report concentrations, the other two (Liu et al., 2020; Kayalar et al., 2021) quantifies concentrations of viral copies in air. Liu et al. (2020) in Wuhan (China) collected samples mainly in indoor environments (different locations hospitals) but eight of them were collected outdoor in public spaces. Authors concluded that these samples presented undetectable or very low concentrations (<3 copies m\(^{-3}\)) with the exclusion of the crowded sites in which traces of SARS-CoV-2 were observed up to 11 copies m\(^{-3}\). These concentrations appear to be lower than those found in indoor in the same study (i.e. 1–42 copies m\(^{-3}\) with a positivity rate of 67%). Kayalar et al. (2021) report 3% of samples (2 out of 68) positive in urban or urban background outdoor sites and the rate of positivity increases to 15% at hospital gardens, outdoor sites near SARS-CoV-2 sources, even if concentrations are not significantly different. The average positivity rate among the three datasets, i.e. the ratio between positive and total samples collected summed in the three datasets, was 17.6%. Concentrations were estimated only in two of the three datasets in which positive outdoor samples were identified. The concentration range considering all the data of the two datasets were 0.1–23 copies m\(^{-3}\) and the average value of all quantified samples was 7.9 copies m\(^{-3}\) (median 7.2 copies m\(^{-3}\)).

de Rooij et al. (2021a) investigated airborne SARS-CoV-2 concentrations indoor in mink farms in Netherlands collecting samples of total suspended particles as well as inhalable dust mainly in indoor. Some samples were also collected outdoors finding either negative or low concentrations cases at 1.5 m from the open entrance of the farm and negative samples at 20 m distance. This suggests that in outdoor conditions, virus-laden particles are quickly transported and dispersed by wind lowering concentrations when distance from the source increases. Hu et al. (2020) also collected outdoor samples at 10 m from the doors of in-patients and out-patients buildings of a hospital in Wuhan (China) finding 3 positive samples out of 20 (15%) while they did not detect SARS-CoV-2 in residential and open public areas. Stern et al. (2021b) collected outdoor air samples at the gates of COVID-19 hospitals in Kuwait finding 5 positive samples out of 33 (15%) with concentrations in the range 3–17 copies m\(^{-3}\). Habibi et al. (2021) studied indoor locations in Kuwait but also collected two outdoor samples in residential areas for control and both resulted negative.

The results discussed give the indication that, in outdoor conditions excluding crowded areas or zones located near the sources (close contact), virus-laden particles are quickly transported and dispersed by winds resulting in low or undetectable concentrations. In these conditions, airborne transmission of COVID-19 disease is very unlikely or negligible. However, the analysis reported reinforce the importance of avoiding crowds and large gathering of people, as well as to maintain social distancing from the sources to minimise the risks of airborne transmission in outdoor. This aspect is also supported by the simulations in outdoor for Milano and Bergamo areas, the epicentre of the COVID-19 outbreaks in Italy in winter 2020, reported in Belosi et al. (2021). The work of Belosi et al. (2021) focuses on these areas in Lombardia region (northern Italy) located in the “Po Valley” that is an atmospheric pollution hot-spot due to the local meteorological and micrometeorological conditions that favour shallow boundary layer, stable atmospheric conditions, and limited ventilation, especially during winter. Simulation of Belosi et al. (2021) showed low average concentrations (<0.6 copies m\(^{-3}\)) even in the grim hypothesis of a 10% of population currently infected and in the worst-case scenario for pollution dispersal.

It must be mentioned that different sampling approaches have been used, as well as different extraction procedures and the LODs are not clearly reported in all studies so that it could be difficult to interpret what a negative result really means. In addition, the recovery, i.e. the efficiency with which SARS-CoV-2 RNA is extracted and identified by PCR in collected samples, is not often reported. Moreover, it is analysed with specific tests only in a few studies and this also could influence results found and comparability of different studies. It was not observed a direct correlation between positivity rate and sampling volumes or sampling supports used. Positive samples were obtained from samples on different typologies of filters (Kayalar et al., 2021; Setti et al., 2020) as well as on gelatine substrates (Liu et al., 2020). On the other hand, datasets with all negative samples were obtained from studies with sampling on filters (Chirizzi et al., 2021; Linillos-Pradillo et al., 2021; Passos et al., 2021; Civato et al., 2021), using cyclone for sampling in centrifugal tubes (Dunker et al., 2021), as well as using high volume centrifugal sampling on phosphate buffered saline (PBS) (Hu et al., 2020).

3.2. Results of indoor measurements in hospitals, quarantine areas, and COVID-19 health care facilities

The summary of results obtained from the 58 datasets dealing with measurements of airborne SARS-CoV-2 RNA traces in hospitals and care facilities are reported in Tables 2 and 3. Table 2 deals with the 41 datasets that have negative (i.e. all air sampled tested negative for presence of SARS-CoV-2) or positive (i.e. at least one air sample tested positive at PCR analysis) without quantification of concentrations. Table 3 deals with the 17 datasets (29.3% of the total) that have quantification of concentrations in air samples in hospitals and health care facilities.
It is interesting to observe that several studies in this category of sites also provided measurements of surface collected (swabs) samples. The datasets having at least one positive air sample are 60.3% of the total, instead, for surface samples the datasets reporting positive samples are 89.5% (i.e. 34 out of 38). This means that, in this category of indoor environments, SARS-CoV-2 is found more frequently on surface rather than in air. There are 15 studies (Tables 2 and 3) in which air samples tested negative, but traces of SARS-CoV-2 were found on different surfaces; on the contrary only one paper (Jin et al., 2020) reports one positive sample in air and no positive samples on surfaces. Considering that surface stability of SARS-CoV-2 on different kind of surfaces could be of several hours, arriving up to a few days (van Doremalen et al., 2020; Marquès and Domingo, 2021), indirect transmission of the virus through contaminated surfaces (i.e. fomites), in these indoor environments, cannot be ruled out. This could happen by touching contaminated surfaces and objects followed by touching the mouth, nose, or eyes. However, it is extremely difficult, in real cases, to ascertain the mechanism of transmission and the exact role of transmission via fomites in the spread of COVID-19. Epidemiological investigations, or structured analysis (like randomized controlled trials), are not feasible to investigate the role of fomites transmission because fomite-mediated contagions are likely a rare event and it is difficult to decouple from other, more likely, transmission routes (Pitol and Julian, 2021). Different studies indicate that the risks of SARS-CoV-2 infection from contact with a fomite are estimated to be low (Marquès and Domingo, 2021; Pitol and Julian, 2021; Zhou et al., 2021b). In addition, these risks could be furtherly reduced by washing hands and with regular disinfection practices of indoor surfaces (Gonçalves et al., 2021; Marquès and Domingo, 2021).

The difference between air and surface samples results could originate from several factors. One factor is that surfaces could be contaminated also by direct contact from infected individuals and not only from respiratory aerosol. A second factor is that virus-laden particles that deposit on surfaces are large respiration droplets that could have a different viral load if the site of infection is the same or very close to the site of particle formation. The different datasets showing positive air samples and/or positive surface samples, were analysed to investigate the positivity rate as

| Reference                  | Sites                                    | Sampling                                                                 | Method                                  | Results                                                                 | Notes                               |
|---------------------------|------------------------------------------|--------------------------------------------------------------------------|-----------------------------------------|------------------------------------------------------------------------|-------------------------------------|
| Chirizzi et al. (2021)     | Two urban background sites: Veneto (Venice, North Italy) and Apulia (Lecce, South Italy); simultaneously studied, period 13/05/2020–27/05/2020. | At each site, 6 PM10 samples on quartz fibre filters (48 h at 38.3 L min⁻¹) and 24 multi-stage impactor samples (6d at 30 L min⁻¹), size range from D < 0.055 μm up to D > 18 μm, Volumes 110 m³ or 250 m³. | RT-PCR targeting E and RdRp genes. | 100% of samples negative with both methods RT-PCR and dd-PCR.           | LOD - PM10 0.8 copies m⁻³; LOD - impactor 0.4 copies m⁻³; Recovery 49%. |
| Dunker et al. (2021)       | University of Leipzig Medical centre. Samples collected between 11/03/2020 and 28/05/2020. | 7 weekly air samples and one 14 days sample. Sampling at 15 min⁻¹ with a cyclone trap directly into 1.5 mL micro centrifuge tube. | RT-PCR targeting E or N and RdRp genes. | 100% of samples negative.                                                | LOD and recovery not reported. Fresh pollen samples were also collected finding no presence of SARS-CoV-2. Recovery not reported. |
| Hu et al. (2020)           | Public areas in Wuhan.                   | 20 samples collected with a centrifugal sampler WA-400 at 400 L min⁻¹ in PBS. Volumes 12 m³. | qRT-PCR targeting ORF1ab gene.         | 100% of samples negative.                                                | LOD and recovery not reported.      |
| Kayalar et al. (2021) Turkey | Samples from 13 locations in 10 towns, period 13/05/2020–14/06/2020. Hospital garden (HG) sites; urban (U) and urban background (UB) sites. | 80 TSP, 19 PM10, 23 PM2.5–10, 33 PM2.5 samples with different samplers and filters (PTFE, quartz and glass fibre, polycarbonate. Volumes 7.2–360 m³. 48 size segregated (6 sizes) samples on glass fibre filters, volume 1422 m³. | RT-PCR targeting N1 and RdRp genes. | LOD and recovery not reported. Fresh pollen samples were also collected finding no presence of SARS-CoV-2. Recovery not reported. |
| Linillos-Pradillo et al. (2021) | Spain (district 09) university area in the period 04/05–22/05/2020. | 6 PM10, 6 PM2.5, and 6 PM1, simultaneous samples on quartz fibre filters at 30 m³ h⁻¹ for 17.5–24 h. Volumes 525–720 m³. | RT-PCR targeting N1 and N2 genes and control of human RNA P (RP) gene. | 100% of samples positive.                                                | LOD reported. Recovery not reported. |
| Liu et al. (2020)          | Different sites near hospital, community check point, department stores and supermarket and residential buildings. | TSP sampled on gelatine substrate at 5 L min⁻¹. Volumes 1.5–5 m³. | dd-PCR targeting Orf1ab and N genes. | 3/8 (37%) of samples positive collected near hospital and near the door of a busy department store. | Outdoor concentration non-detectable or very low (<3 copies m⁻³) but at crowded sites that arrived at 11 copies m⁻³. Recovery not reported. LOD not reported. Recovery >100% |
| Passos et al. (2021)       | Metropolitan area of Belo Horizonte. Period 25/05/2020–06/08/2020. | 2 PM2.5 and 7 PM10 samples in total at: car parking of a COVID-19 hospital, sidewalk near hospital, bus station. Quartz fibre filter sampled at 1130 L min⁻¹. Volumes 7–4500 m³. | RT-PCR targeting N1 and N2 genes. | 100% of samples negative.                                                | LOD reported. Recovery not reported. |
| Privato et al. (2021) Italy | 10 sites (urban-rural background, traffic, industrial), NE Italy (Padua province) period 24/02/2020–09/03/2020. | 25 PM10 and 19 PM1.5 samples were collected in total over the 10 sites, on quartz fibre filters (24 h at 38.3 L min⁻¹), Volume 55.2 m³. | RT-PCR targeting N and Orf1b-14nsp genes. | 100% of samples positive.                                                | LOD 1.2 copies m⁻³; Recovery not reported. |
| Setti et al. (2020)        | Industrial area of Bergamo over a continuous 3-week period 21/02/2020–13/03/2020. | 34 PM10 samples on quartz fibre filters (24 h at 38.3 L min⁻¹), Volume 55.2 m³. | RT-PCR targeting E, RdRp, and N genes. | 58.8% of samples (20/34) positive in 1 gene: 11.8% (3/24) for 2 genes; none for 3 genes. | Concentrations not reported. LOD not reported. Recovery not reported. |
Table 2
Summary of the methodology used and of the results found in different datasets focused on detecting SARS-CoV-2 genetic material (RNA), without quantification of concentrations in indoor sites in hospitals and healthcare settings in which were present COVID-19 patients.

| Reference                  | Sites                                      | Sampling                                                                 | Method                  | Results for air samples                                      | Notes                                      |
|---------------------------|--------------------------------------------|-------------------------------------------------------------------------|-------------------------|-------------------------------------------------------------|---------------------------------------------|
| Ahn et al. (2020)         | Korea                                      | Negative pressure isolation rooms (12 air exchange h⁻¹) with confirmed COVID-19 patients. | RT-PCR targeting E and RdRp genes. | All of the 3 air samples tested negative.                  | LOD and recovery not reported.           |
| Cai et al. (2020)         | China                                      | General wards in Wuhan hospital. Sampling with a Bobcat sampler on electret sampler. | RT-PCR targeting ORF1ab gene. | All of the 15 air samples tested negative.                | LOD and recovery not reported.           |
| Cheng et al. (2020a)      | Hong Kong                                  | Hospital, room of Covid-19 patient. TSP collected with a SAS sampler at 180 L min⁻¹ at 10 cm from the face of patient with and without facemask. | RT-PCR targeting RdRp gene. | All of the 8 air samples tested negative.                  | LOD and recovery not reported.           |
| Cheng et al. (2020b)      | Hong Kong                                  | Airborne infection isolation rooms (AIIRs) with 12 air exchanges per h. TSP using MD8 portable sampler at 50 L min⁻¹ on gelatine. | RT-PCR targeting RdRp gene. | All of the 6 air samples tested negative.                 | LOD and recovery not reported.           |
| Dedeclimenti et al. (2020) | Italy                                      | Non-intensive care units in northern Italy. Sampling with a SKP pump on PTFE filters at 15 L min⁻¹. | RT-PCR, not clear the targets. | All of the 8 air samples tested negative, one outdoor control sample negative. | LOD and recovery not reported. 0.24 surface samples positive. |
| Döhlé et al. (2020)       | Germany                                    | 21 quarantine households in the Bonn area. Air samples with Coriolis cyclone sampler at 300 L min⁻¹ in 15 mL of 0.9% NaCl, with no close contact (< 2 m) of patients. | RT-PCR targeting RdRp gene. | All of the 15 air samples tested negative.                | LOD and recovery not reported. 4/119 surface samples positive. No infectious virus under cell cultures. |
| Dumont-Leblond et al. (2021) | Canada                                   | 31 rooms in 7 long-term care settings in major cities of Quebec. Sampling with IOM Multidust sampler on gelatine filters, at about 2 m from the residents, at 3 L min⁻¹. | RT-PCR targeting ORF1ab gene. | All of the 31 air samples tested negative.                 | LOD and recovery not reported. 20/62 surface samples positive. Cultures of virus from surface samples were negative. |
| Faridi et al. (2020)      | Iran                                       | Hospital ward in Tehran with confirmed COVID-19 patients. Sampling with impinger in 20 mL solution between 2 and 5 m from patients. | RT-PCR targeting E and RdRp genes. | All of the 10 air samples tested negative.                 | LOD and recovery not reported.           |
| Kim et al. (2020)         | Korea                                      | Four health care facilities with COVID-19 patients. TSP using MD8 portable sampler at 50 L min⁻¹ on gelatine. | rRT-PCR targeting E and RdRp genes. | All of the 52 air samples tested negative.                | LOD and recovery not reported. 89/320 surface samples positive. LOD and recovery not reported. |
| Lane et al. (2020a)       | USA                                        | Infection isolation room of a ventilated COVID-19 patient in Atlanta. Air sampling with 2 NIOSH 251 2 BCE-stage samplers for separation in three size fractions at 3.5 L min⁻¹. | RT-PCR targeting N and human RNase P genes. | All of the 28 air samples tested negative.                 | LOD and recovery not reported. LOD and recovery not reported. |
| Lane et al. (2020b)       | USA                                        | Hospital, different sites outside patients' rooms. Air sampling with 2 NIOSH 251 2 BCE-stage samplers for separation in three size fractions at 3.5 L min⁻¹. | rRT-PCR targeting N1, N2, N3 genes or N2, E, RNAase P. | All of the 528 air samples tested negative.                | LOD and recovery not reported. LOD and recovery not reported. 2/00 surface swab positive (facemasks of patients). |
| Li et al. (2020)          | China                                      | Different wards in a hospital for Covid-19 patients. Impinger sampler (BIO-Capturer-6) in a sampling buffer at 80 L min⁻¹. | RT-PCR not reported the targets. | All of the 135 air samples tested negative.                | LOD and recovery not reported. LOD and recovery not reported. |
| Masoumbeigi et al. (2020) | Iran                                       | Different wards of a Covid-hospital in Tehran. Samples taken with all-glass impinger (AGI) at 5–40 L min⁻¹. | rRT-PCR, not clear the targets. | All of the 31 air samples tested negative.                 | LOD and recovery not reported. LOD and recovery not reported. 4/141 surface samples positive. LOD and recovery not reported. 17/27 surface samples positive. LOD and recovery not reported. 25/1502 surface samples positive. |
| Nakamura et al. (2020)    | Japan                                      | Health care facility, different indoor locations. TSP using MD8 portable sampler at 50 L min⁻¹ on gelatine. | RT-PCR targeting N gene. | All of the 11 air samples tested negative.                 | LOD and recovery not reported. LOD and recovery not reported.  |
| Reference          | Sites                                      | Sampling                                                                 | Method                        | Results for air samples | Notes                                           |
|--------------------|--------------------------------------------|--------------------------------------------------------------------------|-------------------------------|-------------------------|------------------------------------------------|
| Wei et al. (2020b)| Six negative pressure non-ICU rooms in isolation ward in Chengdou. | Samples taken with microbiological sampler (CSC-V) on filter membranes at 100 L min⁻¹, Volume 1.5 m³. | RT-PCR targeting ORF1ab and N genes. | All of the 6 samples tested negative. | 0/55 samples from PPE positive. LOD and recovery not reported. |
| Kotwa et al. (2020)| 18 sites in four rooms of quarantine non-healthcare settings. | Coriolis cyclonic air sampler at 300 L min⁻¹ in viral transport media (VTM), Volume 9 m³. | RT-PCR targeting RdRp gene. | All of the 6 air samples tested negative. | 44/112 surface samples positive. LOD and recovery not reported. |
| Zhou et al. (2020b) | Fangcang shelter hospital in Wuhan. | Air sampling with Ningbo IGene Tec at 6 m³ h⁻¹ on gelatine filters in clean, buffer, and contaminated areas. Volume 1 m³. | RT-PCR targeting ORF1ab gene. | All of the 24 air samples tested negative. | 2/482 surface samples positive. LOD 100 copies ml⁻¹ (liquid phase). Recovery not reported. |
| Baboli et al. (2021) | Different sites in four hospitals in Wuhuan. | Two impingers (WA-15 and WA-400) at 15 and 400 L min⁻¹. Volumes 0.6 m³ and 16 m³. | RT-PCR targeting ORF1ab and N genes. | All of the 44 air samples tested negative. | LOD 100 copies ml⁻¹ (liquid phase). Recovery not reported. 4/318 surface samples positive. |
| Barbieri et al. (2021) | Different areas of a Covid-19 hospital in Ahvaz. | Glass impinger, SKC pump with PTFE filters, QuickTake30 kit at 4 L min⁻¹, 0.12 m³ volume, 1−3 m from patient beds. | RT-PCR targeting RdRp and N genes. | 5/51 air samples positive. | LOD and recovery not reported. |
| Ge et al. (2020) | Two hospitals and one quarantine facility. | MD8 sampler at 50 L min⁻¹ on gelatine filters. Volume 1 m³. | RT-qPCR targeting RdRp gene. | 1/5 air sample positive for all replicates. | LOD and recovery not reported. |
| Ben-Shmuel et al. (2020) | Single-occupant rooms at Duke University hospital. | 8 NIOSH BC251 samplers, 1−3.2 m from patients, at 3.5 L min⁻¹. Volume 0.84 m³. | RT-PCR targeting N gene. | 3/143 samples positive at 1.4−2.2 m from patients. | LOD and recovery not reported. 6/70 surface samples positive. Virus cultures negative. |
| Ding et al. (2021) | 4 three-bed isolation rooms of Covid-19 hospital in Nanjing and other indoor areas. | 4 bioaerosol samplers QuickTake-30, a MD8 sampler, an impinge WA-15, an ASE100 sampling at different flow-rates. | RT-PCR targeting RdRp and E genes. | 1/46 samples were positive. | LOD and recovery not reported. 7/107 surface samples positive. |
| Dubey et al. (2021) | Different wards of a COVID-19 dedicated hospital in Delhi. | Sampling at 1.5, 16.7, 27 L min⁻¹ on PVDF filters at 1−3 m distance from patients. Volumes 0.09, 1, 1.62 m³. | RT-PCR targeting RdRp and E genes. | 54/126 air samples positive. | LOD and recovery not reported. 14/18 surface samples positive. LOD and recovery not reported. 17/112 surface samples positive. |
| Ge et al. (2020) | 6 sites in three hospitals in Changsha, Changzhou, and Shaoyang. | NIOSH biosampler (BC251) at 3.5 L min⁻¹. Volume 0.1 m³. | qRT-PCR targeting NP gene. | 4/33 samples were positive. | 2/40 virus cultures positive. 42/89 surface samples positive. Virus cultures negative. LOD and recovery not reported. |
| Hemati et al. (2021) | Different wards of an hospital in Shahrekord. | Sampling with impinger (SKC) at 2 L min⁻¹. Volume 0.48 m³. | RT-PCR targeting RdRp and N genes | 6/45 samples were positive. | LOD and recovery not reported. |
| Kenarkoohi et al. (2020) | ICU and wards of Covid-19 hospital in Ilam province. | Liquid impinger (SKC) at 12 L min⁻¹. Volume 2.16 m³. | RT-PCR targeting ORF1ab, N genes. | 2/14 air samples were positive. | LOD 200 copies ml⁻¹ (liquid phase). Recovery not reported. |
| Kotwa et al. (2021) | Six acute care hospitals in Toronto. | 3 samplers at 3.5 L min⁻¹ on PFTE, Polycarbonate, gelatine filters. A NIOSH bioaerosol sampler. Volume 0.42 m³. | RT-qPCR targeting ORF1ab and E genes. | 3/146 surface samples positive. | LOD and recovery not reported. 125/474 surface samples positive. Virus cultures negative for air but positive on 6 of 36 surfaces. |
| Jin et al. (2020) | Isolation room and PPE dressing room of a hospital in Guiyang. | WA400 impinger at 400 L min⁻¹. Volume 6 m³. | qRT-PCR targeting ORF1ab, NP genes | 1/2 sample was positive. | LOD and recovery not reported. 0/5 surface samples positive. LOD and recovery not reported. 0/218 ICU samples (air and surfaces) positive. |
| Lei et al. (2020) | An ICU and an isolation ward of hospital in Guangzhou. | A two-stage cyclonic NIOSH sampler and an aerosol particle liquid concentrator (W-15, DingBlue) operating at 3.5 L min⁻¹. Volume 0.84 m³. | RT-PCR targeting ORF1 and N genes. | All samples collected at ICU were negative. 2 air samples positive in isolation ward. |... (continued on next page)
Table 2 (continued)

| Reference         | Sites                                      | Sampling                                                                 | Method            | Results for air samples                                           | Notes                           |
|-------------------|--------------------------------------------|--------------------------------------------------------------------------|-------------------|-------------------------------------------------------------------|---------------------------------|
| López et al. (2021) | Mexico                                    | Two hospitals in Hermosillo (Sonora).                                    | Sampling on Millipore filters at 9.6 L min⁻¹, Volume 1.73 m³. | RT-PCR targeting E and RdRp genes.                                | Not clear the total number of air samples collected. positive. 9/182 air and surfaces at isolation ward positive. LOD and recovery not reported. |
| Ma et al. (2021)   | China                                     | Hospitals and quarantine hotels in Beijing.                              | Two impingers: WA-15 (15 L min⁻¹) and WA-400 (400 L min⁻¹). Volumes 0.6–16 m³. | RT-PCR targeting ORFlab, N genes.                                  | 1/26 sample were positive.       | LOD and recovery not reported. 13/242 surface samples positive. LOD and recovery not reported. 3/35 surface samples positive. 1/42 surface samples positive. |
| Morioka et al. (2020) | Japan                                    | Tertiary care hospital in Tokyo.                                          | Air samples collected on gelatine filters with a MD8 sampler at 50 L min⁻¹, Volume 1 m³. | RT-PCR targeting N gene.                                          | 0/4 air sample were positive.     | LOD and recovery not reported. |
| Mouchtouri et al. (2020) | Greece                                   | Three isolation wards and a long-term care facility.                      | Samples collected with a MD8 sampler on gelatine filters at 50 L min⁻¹, Volume 0.5 m³. | RT-PCR not clear the targeted genes.                              | 1/12 sample positive at 2.5 m from patient without mask. LOD and recovery not reported. 3/35 surface samples positive. LOD and recovery not reported. 9/37 surface samples positive. LOD and recovery not reported. 11/341 surface samples positive. |
| Nor et al. (2021)  | Malaysia                                   | 4 hospital wards hosting Covid-19 patients in Kuala Lumpur.                | PM2.5 samples collected with a Minovol sampler on glass microfiber filters at 5 L min⁻¹, Volume 14.4 m³. | RT-qPCR targeting N1 and N2 genes.                                | Positive samples were observed in 2/4 wards. LOD and recovery not reported. |
| Razzini et al. (2020) | Italy                                     | Covid-19 isolation ward of a hospital in Milan.                           | Samples collected with a MD8 sampler on gelatin filters at 50 L min⁻¹, Volume 2 m³. | RT-PCR not clear the targeted genes.                              | 2/5 samples were positive.         | LOD and recovery not reported. |
| Tan et al. (2020)  | China                                     | Covid-19 isolation wards and ICUs in Wuhuan.                             | Air samples collected at 5 L min⁻¹, Volume 0.3 m³ at <1 m from patients plus samples in clean areas. | RT-PCR targeting ORFlab gene.                                      | 1/12 sample positive taken during intubation. 0/17 in clean area were positive. LOD and recovery not reported. 11/341 surface samples positive. |

function of the number of samples available in the dataset (Fig. 4). Results show a quite large variability for both kind of environmental samples, especially for cases with small number of samples, reflecting variabilities in source strength (i.e. number of infected people and their viral load), in geometries and ventilations of the different environments, and in the mitigation policies applied. The average positivity rate, considering all available datasets, was 22.8% (median 11.1%) for air samples and it was 12.4% (median 8.5%) for surface samples. This indicate that, even if contamination of surface samples is more frequent compared to air samples, the average positivity rate is lower in surface compared to air. It must be said that there are relevant differences in the total number of air and surface samples because, in the different datasets, the number of air samples is much lower than that of surface samples. This could create a bias in the evaluation of positivity rate because, for example, when only two samples are present the minimum positivity rate is 50% (i.e. one of the two positive). The red curves in Fig. 4a and b report the number of samples as function of the total number of samples in the dataset. If all air samples of Fig. 4a are considered together as if they was a single dataset, the positivity rate would be 13.3% against the value of 10% for the surface samples taken together. This suggest that positivity rates in air and surface samples could be more similar in datasets having comparable number of samples.

Fig. 4c shows the comparison of the frequency distributions of positivity rate for air and surface samples collected in hospital and care facilities. This shows that the larger positivity rate in air samples is due to two aspects: the absence of datasets with low positivity rate (i.e. < 2%) in air samples that are instead present in surface samples; a significantly larger number of air samples datasets having positivity rate in the range 30%–50% compared to surface samples.

The concentrations of SARS-CoV-2 RNA traces in air samples in these environments observed in the 17 datasets having quantification, span a quite large range: 0.1–94,000 copies m⁻³ with an average value of 3606 copies m⁻³. The median value of the samples collected in all available dataset was 540 copies m⁻³ (17.5–2890 copies m⁻³) was the inter-quartile range between 25th and 75th percentiles. The frequency distribution of measured concentrations is reported in Fig. 5. The histogram shows essentially three peaks: a large one at low concentrations <50 copies m⁻³, a second peak (less frequent) centred in the range 200–500 copies m⁻³, and a third peak at larger concentrations in the range 1000–10,000 copies m⁻³. These values indicate a large variability, as expected, mainly of the variability in the intensity of the sources (i.e. number of infected people indoor and their viral load), of the different sizes and ventilation conditions of the indoors environment studied. Dubey et al. (2021) collected, in a systematic way, air samples in hospital settings at two distances from patients: 1 m and 3 m. They observed that at 1 m from patient the positivity rate was 94.4% but it decreased at 22.2% at 3 m distance. This suggests that distance of sampling from infected individuals could also be an issue in comparison of different studies as well as in the risk of airborne transmission of contagion. As a consequence, the physical distancing among people is a measure likely efficient to be maintained for reducing risks of airborne transmission.

A direct correlation, among all the datasets studied, between sampling method (volumes and typologies of substrates used) and positivity rate was not observed similarly to the results discussed for outdoor datasets. However, Dubey et al. (2021) compared systematically air samples collected at different flow rates (for the same duration) giving three different sampling volumes: 0.09 m³, 1 m³, and 1.6 m³. Samples were collected at distances between 1 m and 3 m from patients in a hospital setting. Results show that positivity rate increases when sampling volume increase, going from 28.6% (samples of 0.09 m³) to 45.2% (samples of 1 m³) and reaching 54.8% for samples of 1.6 m³. This suggests that LOD and recovery could be an issue for estimates of positivity rate, especially if low sampling volumes are used. As a consequence, it would be necessary to include estimates of these parameters in the future studies reinforcing the necessity to develop a standard methodology for collection of air sampling devoted to detection/quantification of SARS-CoV-2 RNA. Likely other dedicated studies are necessary to define a protocol, however, the results analysed in this work seems to suggest that a standard operating procedure should include: an evaluation of recovery in the effective sampling conditions; an optimisation (also considering the recovery value) of the sampling substrate, sampling
### Table 3

Summary of the methodology used and of the results found in different datasets focused on detecting SARS-CoV-2 genetic material (RNA), with quantification of concentrations in indoor sites in hospitals and healthcare setting in which were present COVID-19 patients.

| Reference          | Sites                                                                 | Sampling                                                                 | Method                | Results for air samples | Notes                      |
|--------------------|----------------------------------------------------------------------|--------------------------------------------------------------------------|-----------------------|-------------------------|----------------------------|
| Chia et al. (2020) | Singapore: 3 AIIRs in the ICU and 27 AIIRs in the wards of a hospital. | 6 NIOSH BC251 bioaerosol samplers at 3.5 L min⁻¹. Volume 5.04 m³.        | RT-PCR targeting E, ORF1ab genes. | 4/10 air samples positive, D > 1 μm. Conc. 916–2000 copies m⁻³. 11/100 air samples positive. Conc. 10–514 copies m⁻³. | Recovery not reported. 56/245 surface samples positive. Recovery not reported. |
| Dumont-Leblond et al. (2020), Canada | Acute care hospital rooms in Quebec. | Two plastic IOM (SKC) samplers with gelatine or polycarbonate filters at 10 L min⁻¹. SASS 3100 dry sampler at 300 L min⁻¹. Volumes 2.4–10.8 m³. | RT-qPCR targeting ORF1b gene. | 1/12 air sample positive. Conc. 1857 copies m⁻³. | Recovery not reported. 4/202 surface samples positive. Recovery not reported. 23/161 surface positive in ICU. 2/134 surface positive in general ward. |
| Feng et al. (2021) | Hospital in Zhejiang. | NIOSH sampler at 3.5 L min⁻¹ at 0.2 m from the bed of patients (head position). Volume 0.105 m³. | RT-qPCR but not clear the gene target. | 4/81 positive in ICU. 0/38 positive in general ward. Conc. 520–3800 copies m⁻³. | Recovery not reported. |
| Guo et al. (2020) | Hospital: intensive care unit (ICU) and a general COVID-19 Ward in Wuhan. | Samples collected with SASS 2300 wetted cyclone sampler at 300 L min⁻¹ on viral transport medium. Volume 9 m³. | qRT-PCR targeting ORF1ab, N genes. | 9/16 air samples positive. Conc. 3200–42,000 copies m⁻³. | Recovery not reported. Virus cultures were negative. LOD 37.5 copies μL⁻¹. Recovery not reported. Virus cultures were negative. LOD 37.5 copies μL⁻¹ (liquid phase). Recovery not reported. One sample showed positive virus culture. Recovery not reported. |
| Habibi et al. (2021) | Three major hospitals in Kuwait dealing with Covid-19 patients. | Sampling at 30 L min⁻¹ in wash bottles with TRIzol (APB Bioscience). Volume 3.6 m³. | RT-qPCR targeting ORF1ab, N genes. | 5/13 air samples positive. Conc. 12–99 copies m⁻³. | Recovery not reported. |
| Hu et al. (2020) | Various sites in different health facilities of Wuhan. | Centrifugal sampler WA-400 at 400 L min⁻¹ in PBS. Volumes 12 m³. | qRT-PCR targeting ORF1ab gene. | 9/81 air samples positive. Conc. 1110–12,000 copies m⁻³. 1/2 air sample positive. Conc. 870 copies m⁻³. | Recovery not reported. Virus cultures were negative. LOD 37.5 copies μL⁻¹. Recovery not reported. Virus cultures were negative. LOD 37.5 copies μL⁻¹ (liquid phase). Recovery not reported. One sample showed positive virus culture. Recovery not reported. |
| Lednicky et al. (2020a) | Student health care centre in Florida for COVID-19 patients. | VIVAS sampler on PBS at 6.5 L min⁻¹. Volume 0.39 m³. | RT-PCR targeting N gene. | 4/4 air samples positive. Conc. 1600–94,000 copies m⁻³. | Recovery not reported. |
| Lednicky et al. (2020b) | A two patient room in a hospital in Florida with 6 air exh. h⁻¹. | VIVAS sampler and a BioSpot-VIVAS BSS300P on PBS at 8 L min⁻¹. Volume 1.44 m³. | RT-PCR targeting N gene. | 16/25 samples positive. Conc. 1–42 copies m⁻³. 4/55 samples positive with Coriolis sampler. 0/34 samples positive with MDB. Conc. 10–460 copies m⁻³. | Recovery not reported. 30/336 surface samples positive. Virus cultures were negative. |
| Liu et al. (2020) | Different sites in three hospitals of Wuhan. | TSP sampled on gelatine substrate at 5 L min⁻¹. 3 size-segregated samples. Volumes 1.5–5 m³. | dd-PCR targeting Orf1ab and N genes. | 6/12 positive samples in AIIRs conc. 179–2738 copies m⁻³. 1/9 positive sample in CIF, conc. 978 copies m⁻³. | Recovery not reported. |
| Moore et al. (2021) | 8 hospitals, different locations (11 AIIRs), 11 neutral pressure side rooms, six ICU/HUD, open cohorts and 12 non-ICU sites. | Coriolis sampler at 300 L min⁻¹ in PBS and a MDB sampler in gelatine filters at 50 L min⁻¹. Volumes 0.5–3 m³. <1 m from patients. | qRT-PCR targeting ORF1ab gene. | 6/12 positive samples in AIIRs conc. 179–2738 copies m⁻³. 1/9 positive sample in CIF, conc. 978 copies m⁻³. | Recovery not reported. 30/336 surface samples positive. Virus cultures were negative. |
| Ong et al. (2021) | ALLRs and a community isolation facility (CIF). | Samples using a BioSpot-VIVAS BSS300-P at 8 L min⁻¹ not clear sampling time. | qRT-PCR targeting E, ORF1ab genes. | 6/12 positive samples in AIIRs conc. 179–2738 copies m⁻³. 1/9 positive sample in CIF, conc. 978 copies m⁻³. | Recovery not reported. |
| Passos et al. (2021) | 2 hospitals in the area of Belo Horizonte. | Different low and high volume samplers on cellulose, quartz, and PTFE filters. Volumes 0.12–250 m³. | RT-PCR targeting N1 and N2 genes. | 3/33 samples positive. Conc. 0.14–0.33 copies m⁻³. 12/19 samples in rooms positive. Conc. 2420–8340 copies m⁻³. 14/24 hallway samples positive. Conc. 2080–8090 copies m⁻³. | Recovery not reported. LOD 5 copies μL⁻¹ (liquid phase). Recovery not reported. 60/74 surface positive. Cultivation of virus was tried but not confirmed. Recovery not reported. |
| Santarpia et al. (2020) | Rooms and hallways of quarantine and isolation care areas in Nebraska. | MD8 sampler on gelatine filters at 50 L min⁻¹. Volume 0.75 m³. | RT-PCR targeting E gene. | 8/90 air samples positive. | Recovery not reported. |
| Stern et al. | Different sites at a hospital in Boston. | Cascade impactor, 3 stages (<2.5 μm, 2.5–10 μm, RT-qPCR | | | |

(continued on next page)
flow, and sampling volume in order to achieve a LOD as low as 1–2 copies m$^{-3}$; low-temperature conservation of collected samples before the analysis and some results indicate that $-25^\circ$C or lower could be suitable (Conte et al., 2021).

It is important to mention that the detection of RNA traces of the SARS-CoV-2 virus does not imply infectivity of these airborne virus-laden particles. Some of the studies, summarized in Tables 2 and 3, tried to perform cultures of the SARS-CoV-2 in collected positive samples, both in air and on surfaces, to have a better insights of the eventual viability. Infectious virus particles were not observed in environmental (air and/or surface) samples collected and analysed in most of the available studies (Binder et al., 2020; Döhla et al., 2020; Dumont-Leblond et al., 2020, 2021; Hu et al., 2020; Moore et al., 2021). Santarpia et al. (2020) attempted virus culture but, given the low concentrations found in collected samples, cultivation of virus was not successful. Kotwia et al. (2021) investigated viability of positive samples (Ct < 34 on PCR analysis) and found that none of the positive air samples yielded viable virus, however, viable virus was observed in 6 out of 36 surface samples cultured. Virus cultures done in samples collected at a COVID-19 University health care facility in Florida were negative (Lednicky et al., 2020a) but resulted positive the cultures done in samples collected at a hospital in Florida with viable viral concentrations between 6000 and 74,000 TCID50 m$^{-3}$ (Median Tissue Culture Infectious Dose) (Lednicky et al., 2020b). Ben-Shmuel et al. (2020) used 97 positive air and surface samples to test viability of SARS-CoV-2 and none were found to contain infectious titres of the virus. These results suggest that cultural viable virus is present in a limited fraction of positive surface samples, however, there is need of additional studies before to reach a robust conclusion on this aspect.

3.3. Results of measurements in community indoor sites

The summary of results obtained from the 10 datasets dealing with measurements of airborne SARS-CoV-2 RNA traces in community indoor environments are reported in Tables 4 together with details of sampling methods used and results found. A fraction of 30% of available datasets includes at least one positive air sample; this fraction increases up to 67% when surface samples are considered. This means that contamination from SARS-CoV-2 RNA is found more frequently over surfaces rather than on air samples similarly to what has been observed for hospitals and healthcare settings. The positivity rate for air samples in community indoors ranges from 11.1% (de Rooij et al., 2021a) to 64.3% (Hadei et al., 2021). The positivity rate for surface samples ranges between 3% (de Rooij et al., 2021b) up to 42.2% (Moreno et al., 2021). Therefore, even if contamination is found more frequently on surfaces, the positivity rates are lower on surfaces compared to air samples, exactly the same results obtained also for hospitals and healthcare settings. In addition, the positivity rates found for air and surfaces in indoor community environments are similar to those observed in hospitals and healthcare settings.

Concentrations were quantified in 2 out of the 3 datasets having positive air samples and showed different values due to the same causes already mentioned for hospitals and healthcare setting. An average concentration of 14.5 copies m$^{-3}$ was found by Moreno et al. (2021) and an average of 3700 copies m$^{-3}$ was reported by de Rooij et al. (2021a). The average of all available samples was about 1857 copies m$^{-3}$. This suggests that the expected average concentrations in indoor community environments are larger than those typically found in outdoor and lower than those found in hospitals and healthcare settings even if the number of datasets in indoor community environments is quite limited and further studies are needed to confirm this conclusion.

The analysis done in public transport in Barcelona (Moreno et al., 2021) showed that positivity rate and concentrations in air samples are larger for subways compared to surface busses; instead, positivity rates for surface samples are comparable. The results found in a bus in central Italy (Di Carlo et al., 2020) during the first wave of pandemic showed all negative air and surface samples interpreted as a possible consequence of strict measures against COVID-19 spread applied. Hadei et al. (2021) performed logistic regressions between positivity rates and other parameters such as the number of people present during sampling, the percentage of use of face masks, outdoor temperature, and volumes of indoor sampling sites finding no statistically significant correlations.

The results of positivity rates and concentrations found in the datasets collected in community indoor environments, even if they are limited in numbers, suggest that there could be an intermediate risk of airborne transmissions between outdoor and hospital and health care. In addition, most of the study suggests that volumes and ventilations of indoor environments and the use of face masks are factors strongly influencing the presence of and concentrations of SARS-CoV-2 RNA in these environments and, consequently, influencing the risks of airborne transmission in general public.

4. Conclusions

This work reviewed 73 papers dealing with identification/quantification of SARS-CoV-2 genetic material in air by means of active sampling and PCR detection, published since the start of pandemic and until 31/08/2021. Selected papers furnished 77 datasets in different environments. Only 9 datasets (11.8% of the total) were obtained in...
outdoor sites and all of them were collected during the first wave of pandemic in 2020. The remaining datasets were collected in hospitals and healthcare settings (75.1% of the total) and in community indoor environments (13.1% of the total).

The fraction of datasets having at least one air sample positive to SARS-CoV-2 was 33.3% in outdoor, comparable to the 30% observed in community indoors and significantly lower than the value of 60.3% observed in hospitals and healthcare settings. The average positivity rate of air samples (i.e. the ratio between positive and total samples) was lower in outdoor (17.6%) compared to indoor (23.7%) sites. Among the indoor sites, the positivity rates in hospitals and healthcare settings varied in an interval comparable with that of indoor community sites. Several of indoor studies also included datasets of surface samples (swabs collected on different indoor surfaces). The fraction of datasets having at least one positive surface sample were larger than the fractions observed for air samples in both typologies of sites (i.e. hospitals and community indoors). Despite the fact that contaminated surfaces were found more frequently than contamination in air samples, the positivity rate was lower on indoor surfaces (12.4% in hospitals and 9.7% in community indoors) compared to that of indoor air (23.7%), even if these numbers are calculated using a significantly different number of samples. This suggests that indirect SARS-CoV-2 transmission (i.e. via fomites) could not be ruled out even if its role in the spread of contagion is not actually clear and quantified. However, measures such as increased frequency of disinfection of indoor surfaces and washing hands could be useful to reduce these risks and should be maintained during the next phases of pandemic.

The concentrations observed in outdoor are relatively low, compared to those observed in indoor sites, and positive samples are mainly observed near sources (i.e. crowds or hospital settings). This suggest that risks of airborne transmission in outdoor sites are relatively limited providing that physical distance is maintained avoiding crowds and vicinity to potentially large sources. Concentration in hospital and healthcare setting shows a large variability with average values in the different studies covering five orders of magnitude. This suggests a larger level of risk compared to outdoor and a strong influence of source (i.e. number of infected individuals) but also of volumes of rooms, distances from patients, and ventilation rates. A limited number of studies report concentrations for indoor community environments, that are likely the sites that needs to be further investigated in the next future, however, the concentration in these sites seems to be, on average, larger than in outdoors and lower than in hospital and healthcare settings. The studies that tried to culture SARS-CoV-2 from positive samples (air and surface samples) are still relatively limited; it seems to suggest that in the majority of cases airborne and surface virions are likely not viable, however, it would be advisable to have additional studies on this aspect.

There is not yet a standard methodology of sampling, conservation, and analysis of collected samples; however, datasets with positive samples were obtained from samples on different typologies of filters, on gelatine substrates as well as datasets with all negative samples. A direct correlation, among all the datasets studied, between sampling method (volumes and typologies of substrates used) and positivity rate was not thereby observed neither in outdoor nor in indoor sites. However, Dubey et al. (2021) show that positivity rate increases when sampling volume increase. This suggests that LOD and recovery could be an issue for estimates of positivity rate and it would be necessary to include estimates of these parameters in the future studies. In addition, this
work reinforces the usefulness of developing a standard methodology for collection of air samples devoted to detection/quantification of SARS-CoV-2 RNA. The results of positivity rates and concentrations found in community indoor environments, even if the number of datasets is limited, suggest that there could be intermediate risks of airborne transmissions between outdoor and hospital and healthcare settings. The risk appears strongly related to volumes and ventilations of indoor environments and the use of facemasks, both factors influencing the presence/concentrations of SARS-CoV-2 RNA. Therefore, also these mitigation measures should be maintained in the next phase of pandemic for indoor environments.

Despite the uncertainty on the viral load in positive subjects (which depends not only from subject to subject, but also from the lung region where the infection is taking place and its time course) and the difficulties due to the different sampling methods used, this review allowed a statistical evaluation of the prevalence of the SARS-CoV-2 in air, of the positivity rates, and of detected concentrations, mediated on the different contexts: outdoor, indoor (hospitals and similar) and indoor (living environments). These quantifications (including ranges of variability) represent an important input for epidemiological models (both outdoor and indoor) to estimate the risk of virus transmission, providing the scientific community with summarized data useful for risk assessment and not just for behavioral indications.

CRediT authorship contribution statement

D. Contini, A. Gambaro, F. Belosi, G. La Saldana conceptualized the study design; E. Barbaro, E. Gregoris, M. Feltracco, M. Conte, A. Dinoi, S. Trabucco collected published material and collaborated for statistical analysis; D. Chirizzi, G. Ciccarese, G. and G. La Bella collaborated to investigate protocols used in the different datasets. All authors collaborated to interpretation of results, wrote, read, commented, and approved the final manuscript.
Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have affected the work reported in this paper.

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