What do we know about indoor air quality of nurseries? A review of the literature

Shuo Zhang, D Mumovic, Samuel Stamp, Katherine Curran and Elizabeth Cooper

Abstract
Considering the alarming rise in the rate of asthma and respiratory diseases among school children, it is of great importance to investigate all probable causes. Outside of the home, children spend most of their time in school. Many studies have researched the indoor environmental quality of primary and secondary school buildings to determine the exposure of school children to indoor air pollution. However, studies of very young children in nurseries are scarce. Unlike at elementary schools or universities, children in nurseries are more vulnerable due to their physiology, inability to articulate discomfort and to adapt their behaviour to avoid exposures. This article reviews current studies on the indoor environment in nurseries. It summarizes air pollution levels and related environmental and behavioural factors in nurseries that have been reported in the literature. Additionally, exposure to indoor air pollution and related potential health outcomes are examined. This review concludes that indoor air pollution in nurseries often exceeds current guidelines, and designers and policymakers should be made aware of the impact on the health and wellbeing of children in nurseries. Proper interventions and guidelines should be considered to create a healthy indoor environment for nursery children.

Practical application: Previous IAQ assessments have mainly focused on indoor temperatures and CO2 levels. Data on comprehensive monitoring (including PMs, NO2, O3 and other pollutants) of indoor air quality of nurseries are scarce. Particularly in the UK, studies about indoor air quality in nurseries have not been founded. This paper categorized relevant articles according to the focus of the study, to provide evidence to a better understanding of current indoor air quality in nursery environments.

Keywords
IAQ, carbon dioxide, nitrogen dioxide, ozone, particulate matter, volatile organic compounds, allergens, bacteria, fungi, nursery, health impacts

Received 6 October 2020; Revised 22 March 2021; Accepted 25 March 2021

Introduction
Outdoor air pollution is recognized as a severe problem worldwide. Epidemiological evidence indicates that air pollutants contribute to increasing mortality rates and hospital
admissions. In recent decades, researchers have paid more attention to indoor air quality because people spend about 90% of their time indoors leaving them at risk for higher periods of exposure. However, monitoring and analysis of indoor environments can be challenging due to several factors, including: building characteristics, occupant behaviours, and pollutants from outdoor sources. Building characteristics include conditions such as ventilation rates, envelope ‘leakiness’, and age of the structure. With such a wide variety of building types, establishing rigorous and repeatable protocols for monitoring pollutants is difficult. Occupant behaviour has nearly as many varieties as there are occupants, and includes indoor activities such as cooking, as well as, patterns of use (e.g. time spent at home). Sources of pollutants also vary widely by location and building use. For example, buildings located along a busy urban road will have a different profile of pollutants than a suburban school in a greenfield. Due to the myriad building and occupancy configurations, generalizability is limited, and analysis is a challenge. However, studies have shown that indoor air can be more polluted than outdoor air, and poor indoor air quality has been linked to negative health outcomes.

Children, especially those under six years old, are more vulnerable than adults to environmental pollutants because their immune and respiratory systems are not fully developed; children have a larger surface area to volume ratio; and a faster rate of respiration. Furthermore, children spend more time in the indoors, averaging 7–11 h per weekday in classrooms alone. It has been shown that indoor air pollutants have the potential to damage children’s central nervous system. Also, exposures to air pollutants before the age of one-year may contribute to the development of childhood asthma. The evidence gathered for these observed outcomes has been primarily focussed on offices and residential buildings. The research on schools tends to be on primary and secondary schools, and there is a substantial and important lack of data and guidance on the indoor air quality of nurseries.

In this review we identified 33 studies (Table 1) that focussed on indoor air quality in nursery settings. These studies found that many children are in nurseries that have poor indoor environmental quality. Most of the studies only focussed on one or two specific aspects of indoor air quality (e.g. particulate matter, carbon dioxide, allergens), and only a few of them attempted to give an overall analysis of the indoor environmental quality in nurseries.

In the future, buildings should be designed or retrofitted with a comprehensive approach that integrates physical characteristics, occupant behaviour patterns, and avoidance of harmful microbial and chemical exposures in their design and operation. The aims of this paper are: first, explore the perception of thermal comfort of nursery children; second, describe the current ventilation strategies in nurseries; third, identify the type and scale of exposure of children to pollutants in nursery environments. This review aims to provide scientific evidence to guide policymakers, design professionals and researchers to a better understanding of current indoor environmental quality in nursery settings.

**Research overview**

Studies focusing on indoor air quality in nursery environments were conducted in Europe, including Portugal, France and Poland, as well as in South Korea, Singapore and Canada, only a few studies were available from developing countries. As shown in Table 1, many studies focussed solely on a single parameter of indoor air quality. Temperature, humidity, CO₂, particulate matter (PM), and volatile organic compounds (VOCs) were the parameters most often studied. The number of nurseries monitored varied from 1 to 310, and classroom sizes also varied between studies. Most of the investigated classrooms were naturally ventilated, and the age of children was mainly older than three years. For studies conducted in a location with a varied climate, few included any observed seasonal differences. Most studies
Table 1. A summary of papers reviewed with study location, number of nurseries monitored, primary ventilation method employed at the site, number and location of monitoring, age of the children in the room, the height of the sensor above finished floor, and the indoor environmental factor measured.

| Research                | Country    | Number of nurseries monitored | Building ventilation type | Sample number | Age of children | Height of sensor | Measured elements                                                                 |
|-------------------------|------------|-------------------------------|---------------------------|---------------|----------------|-----------------|----------------------------------------------------------------------------------|
| Zuraimi and Tham11      | Singapore  | 104                           | Naturally Ventilated      | Two indoor classrooms and an outdoor sampling point for each nursery | 0.5–0.7 m     | T, RH (relative humidity), CO₂, CO, PM₁₀, PM₂.₅, O₃, bacteria, fungi            |
| Yoon et al.30           | South Korea| 17 (13 urban 4 rural)         | Air conditioned           | One indoor and one outdoor for 71 classrooms                          | 1 m           | CO₂, CO, VOCs, Formaldehyde                                                  |
| Mendes et al.18         | Portugal   | 45                            |                           | 52 classrooms                                                        | 0.6–1.5 m     | T, RH, CO₂, CO, PM₁₀, VOCs, formaldehyde, bacteria, fungi, allergens           |
| Cano et al.49           | Portugal   | 19                            |                           | 125 classrooms                                                       | 3 months up to 5–6 years | 0.5–0.7 m | T, RH, CO₂, CO, PM₁₀, VOCs, formaldehyde, bacteria, fungi                      |
| Branco et al.66         | Portugal   | 4 (urban)                     |                           | 14 (includes 4 lunchrooms)                                           | >1–5          | 0.5–0.7 m | T, RH, CO₂                                                                     |
| Branco et al.66         | Portugal   | 4 (URBAN)                     |                           | 14 (includes 4 lunchrooms)                                           | >1–5          | 0.5–0.7 m | CO, NO₂, VOCs, formaldehyde, O₃, PM₁₀, PM₂.₅, PM₁, TSP (total suspended particles) |
| Branco et al.47         | Portugal   | 3 (urban)                     |                           | 11 (includes 3 lunchrooms)                                           | >1–5          |                                |                                                                                   |
| Nunes et al.44          | Portugal   | 4 (3 rural 1 urban)           | 10 natural ventilation 2 forced ventilation | 12 (includes 4 lunchrooms)                                           | >1–5          | 0.5–0.7 m | PM₂.₅, PM₁₀, PM₁, TSP, CO₂, NO₂, VOCs, formaldehyde, O₃,                  |
| Nunes et al.34          | Portugal   | 4 (3 rural 1 urban)           | 10 natural ventilation 2 forced ventilation | 12 (includes 4 lunchrooms)                                           | >1–5          |                                |                                                                                   |
| Gładyszewska-Fiedoruk124 (2013) | Poland | 3                             | Natural ventilation      | 31 (mixed spots, indoor and outdoor)                                 | 1–1.1 m       | T, RH, CO₂                                                                     |
| Mainka et al.4850       | Poland     | 2                             | Natural ventilation      | 4 indoor and 2 outdoor spots                                          | 3–6 years     | 0.8–1 m | CO₂, PM₂.₅, PM₁₀, PM₁, TSP, VOCs, bacteria, fungi                          |

(continued)
Table 1. Continued.

| Research                      | Country       | Number of nurseries monitored | Building ventilation type | Sample number | Age of children | Height of sensor | Measured elements |
|-------------------------------|---------------|------------------------------|--------------------------|---------------|-----------------|------------------|-------------------|
| Mainka and Zajusz-Zubek       | Poland        | 4 (2 rural 2 urban)          | Natural ventilation     | 8 playgrounds and inside 8 classrooms | 3–6 years       | 0.8–1 m          | CO₂, PM₁₀, PM₁₀₀, PM₁, TSP |
| Bragoszewska et al. 2016      | Poland        | 1                            | Natural ventilation     | 2 classrooms  | 3–6 years       |                  | Bacteria          |
| Mainka et al. 33              | Poland        | 1 rural                      | Natural ventilation     | 2 indoor and 1 outdoor spot | 3–6 years       | 0.8–1 m          | CO₂, PM₁₀, PM₁₀₀, PM₁, TSP, VOCs, bacteria, fungi |
| Kamaruzzaman and Razak        | Malaysian     | 2 (1 rural 1 urban)          | Natural ventilation     | 2             |                 |                  | T, RH, CO₂, CO, NO₂, VOCs, formaldehyde |
| Roda et al. 13                | France        | 28                           |                          |               |                 |                  | T, RH, CO₂, CO, NO₂, VOCs, aldehydes, fungi, allergens |
| Canha et al. 58               | France        | 17 (7 nursery 10 elementary) | Naturally Ventilated 73% Mechanical Ventilated 27% | 51 classrooms | 3–10 years      |                  | T, RH, CO₂, PM₁₀, PN, VOCs, formaldehyde |
| Ramalho et al. 76             | France        | 567 dwellings and 310 nurseries, kindergartens and elementary schools | Naturally Ventilated 73% Mechanical Ventilated 27% | 51 classrooms | 3–5 years 6–10 years | CO₂, PM₁₀, PM₁₀₀, formaldehyde |
| Michelot et al. 2013          | France        | 160 schools and day-care centres |                     |               |                 |                  | Formaldehyde |
| Daneault et al. 9             | Canada        | 91                           | Naturally Ventilated 19% Mechanical Ventilated 81% | 1672          | 18–60 months    |                  | T, RH, CO₂ |
| St-Jean et al. 15             | Canada        | 21                           | Naturally Ventilated 19% Mechanical Ventilated 81% | 1672          | 18–60 months    |                  | T, RH, CO₂, VOCs, formaldehyde |
| Andrade et al. 84             | France        | 30                           | Naturally Ventilated 19% Mechanical Ventilated 81% | 1672          | 18–60 months    |                  | Allergens         |
| Vaupotic 107                  | Slovenia      | 10                           | Naturally Ventilated 19% Mechanical Ventilated 81% | 1672          | 18–60 months    |                  | Allergens         |
| Rullo et al. 79               | Brazil        | 60 (15 elementary schools)   |                          |               |                 |                  |                   |

(continued)
| Research              | Country  | Number of nurseries monitored | Building ventilation type | Sample number | Age of children | Height of sensor | Measured elements                  |
|-----------------------|----------|-------------------------------|---------------------------|---------------|-----------------|-----------------|-----------------------------------|
| Ferng and Lee         | US       | 26 (10 day-care home)         | T, RH, CO₂                | 1 m           | Bacteria, fungi  |                 |                                   |
| Kim and Kim           | South Korea | 40 (10 childcare centres)    | CO₂, CO, PM₁₀, VOCs, formaldehyde | 1.5 m        |                 |                 |                                   |
| Yang et al.           | South Korea | 55 (5 kindergartens)         | CO₂, CO, PM₁₀, VOCs, formaldehyde | 1.5 m        |                 |                 |                                   |
| Alves et al.          | Portugal | 1 kindergarten 8 elementary schools | Natural ventilation      | 9             | 1.2 m           | T, RH, CO₂, CO, PM₂.₅, VOCs, formaldehyde |                                   |
| Madureira et al.      | Portugal | 9 (68 homes, 20 primary schools and 22 elderly care centres) | Naturally Ventilated Mechanical Ventilated 1 | 50           | 0.7–1.5 m       | Bacteria, fungi  |                                   |
| Branco et al.         | Portugal | 5 nursery schools; 12 pre-schoolers; 8 primary schools | Under 3 3–5 years | 1.5 ± 0.5 m    | Radon           |                 |                                   |
| Oliveira et al.       | Portugal | 1 138 samples (70 indoor and 68 outdoor ones) | 3–5 years | 1.5 m         | CO₂, CO, PM₂.₅, PM₁₀, VOCs, O₃ |                 |                                   |
| Kalimeri et al.       | Greece   | 1 kindergarten 2 primary schools | Natural ventilation       | 3 classrooms 1 outdoor | 1–1.5 m       | T, RH, CO₂, CO, PM₂.₅, PM₁₀, NO₂, VOCs, formaldehyde, O₃, radon |                                   |
| Villanueva et al.     | Spain    | 18 (and 18 primary schools combined) | NO₂, VOCs               | 1.5–2 m       |                 |                 |                                   |
only measured for one day during occupied hours. Few studies measured for more than one day, and for those that did the measurement periods lasted from two to nine days. Sensors were generally placed at a height of 0.5–1.5 m, in the breathing zone for children less than six years old.

The measuring methods used in the studies can be found in the appendix. Most studies employed active sampling methods to measure different indoor air quality (IAQ) parameters, with some studies using passive sampling methods for gas pollutants (VOCs, NO₂, etc.). However, there are several limitations in the IAQ monitoring methods used. For instance, many studies had insufficient measuring periods, or the measuring time/date was inconsistent between monitored nurseries. Some studies did not include outdoor pollutant levels, or if they did, used publicly available monitoring data as their outdoor reference. No study used both active and passive sampling methods for gas pollutants, and any room height difference of indoor PM levels was not determined.

This review categorizes IAQ into two themes; environmental conditions and IAQ control solutions. Environmental conditions include thermal comfort, ventilation rate and CO₂ level and indoor air pollutants (e.g. PM, chemical concentrations). Research outcomes within the same theme were analysed to get a general understanding of the indoor air quality conditions in nurseries.

Environmental conditions

Thermal comfort. According to ASHRAE (American Society of Heating, Refrigeration and Air Conditioning Engineers) Standard 55-2017, the recommended indoor temperature range is around 19–27°C, and the recommended indoor relative humidity is between 30 and 60%. Some studies measured the indoor air temperature and relative humidity in nurseries and compared the outcomes with current guidelines. Temperature (Figure 1) and relative humidity (Figure 2) in most nurseries fell within the comfort range. A notable exception to the adherence to temperature and relative humidity guidelines is one study that measured four nurseries in Portugal. The conditions there may have been due to a poorly constructed or ageing building (e.g. insufficient

![Figure 1. Summary of reported indoor temperature means and ranges.](image-url)
Zhang et al. (2012), Canada
Roda et al. (2011)a, France
Roda et al. (2011)b, France
Kamaruzzaman and Razak (2011), Malaysian
Branco et al. (2015), Portugal
Zuraimi and Tham (2008), Singapore
St-Jean et al. (2012), Canada

Figure 2. Summary of reported indoor relative humidity (RH) means and ranges.

thermal insulation and water intrusion), and the inappropriate use of heaters or air conditioning systems. Another study investigated 26 nurseries in the western United States. They reported that 42% of the monitored nurseries were outside of the temperature comfort zone (34.6% lower and 7.7% higher), and during naptime, 26.1% of the nurseries have a higher relative humidity than the comfort zone. In Poland, a study found higher than recommended indoor temperatures, ranging from 24.0 to 29.6°C during the daytime. However, comparisons between studies are tricky because the measuring periods, climate, countries and building characteristics were different.

In addition to collecting temperature and relative humidity data from nurseries, researchers assessed the thermal comfort of the children there. Children have higher metabolic rates than adults, and when they are unsatisfied with the thermal conditions, they do not necessarily behave like adults to adapt to the environment (e.g. take off/add clothes, open/close windows). One study focussed on the thermal comfort in nurseries in winter and spring. They reported a predicted mean vote (PMV) between “neutral” (0) and “slightly cool” (≤−1), on the thermal sensation scale of −2 to 2. In Korea, a study reported that children prefer lower temperatures (about 3°C lower) than adults and girls prefer temperatures about 1°C lower than boys of nursery age.

All in all, temperature and humidity are important elements in the studies of indoor environmental quality. It has been well demonstrated that temperature and humidity have a strong influence on the perception of indoor air quality and on the volatilisation of chemicals used indoors. More studies focussed on overheating as a problem, due to global climate change. High indoor temperatures can have many adverse impacts on human health, causing problems such as heatstroke and aggravating chronic conditions like cardiovascular and respiratory diseases. Low indoor relative humidity can cause problems such as dry eyes, nose, ears and throat, and high indoor relative humidity is associated with dust mites and fungal moulds.
Ventilation rate and CO₂ level. The indoor concentration of CO₂ can be used as an indicator of ventilation rate and indoor air quality. However, it is a poor indicator of outdoor-associated pollutants (e.g. traffic-related pollutants and fungi species). As mentioned by BB101 (2018), several factors can affect the concentration of CO₂ in indoor environments, including the ventilation rate, occupant density, activity level of occupants, and the occupied time. ASHRAE Standard 62.1-2016 recommends that indoor CO₂ concentrations should not exceed 700 ppm above the outdoor concentration (typically around 400 ppm), and when mechanical ventilation is used, indoor CO₂ concentrations in schools should be maintained at/or below 1000 ppm.

Studies commonly report that CO₂ concentrations in nurseries are high. Published results from several studies found 75% (out of 6 schools), 89.3% (out of 28 samples) and 90% (out of 91 schools) of measured indoor CO₂ concentrations exceeded 1000 ppm. Across numerous studies measured indoor CO₂ concentrations ranged from 377 to 2750 ppm. However, as monitoring methods (e.g. monitoring periods) used in the studies was different, comparing the results is difficult. This range provides a snapshot of the CO₂ concentration published in the current research. As shown in Figure 3, indoor CO₂ concentrations are relatively high in most studies. However, low CO₂ concentrations (below 1000 ppm) do not guarantee acceptable indoor air quality. As one study done in South Korea reported, 41% of rural schools exceeded the South Korean IAQ standard for TVOC concentrations (400 μg/m³), even though the average CO₂ concentration was 607.8 ppm in these same nurseries.

It is worth noting, studies reported that classrooms of younger children tend to have a higher CO₂ concentration than classrooms of older children. Also, higher CO₂ concentrations

![Figure 3. Summary of reported indoor CO₂ concentration means and ranges.](image-url)
occur during nap-time, with about 104 ppm higher than non-nap-times.\textsuperscript{17,18,33} Urban nurseries tend to have a higher concentration than rural nurseries,\textsuperscript{30,34} and publicly managed nurseries have poorer indoor air quality than privately managed nurseries.\textsuperscript{14}

The effect of inadequate ventilation on human health and performance includes; respiratory illnesses, allergies and asthma, sick building syndrome symptoms (SBS), reductions in performance and productivity and perceived air quality.\textsuperscript{35} Previous meta-analyses have reported that low ventilation rates might have adverse effects on the health of school children.\textsuperscript{36} Sundell et al.\textsuperscript{37} and Smedje et al.\textsuperscript{38} reported that increasing the outdoor air flow rate from 1.3 to 12.8 l/s-p (corresponding to a decreased mean indoor CO\textsubscript{2} concentration of 1050–780 ppm), reduced asthma symptoms in pupils from 11.1\% to 3.4\% over a two-year period. In addition to health outcomes, a study in primary and secondary schools reported that a 1000 ppm increase in indoor CO\textsubscript{2} related to a 10–20\% increase in student absenteeism.\textsuperscript{39}

**Particulate matter.** Particulate matter is a leading cause of death and disability worldwide,\textsuperscript{40} and the negative impact on health is especially consequential for children.\textsuperscript{40,41} In general, the smaller the particle size, the more deeply it penetrates and deposits within the respiratory system, posing a greater threat to human health. Studies have shown that large-scale international or national interventions, as well as personal prevention approaches, might help to reduce particulate matter and improve indoor air quality.\textsuperscript{42}

Measured indoor particulate matter levels are often higher than those reported outdoors. Indoor PM concentrations are strongly influenced by outdoor sources (mainly from traffic emissions), and urban nurseries tend to have higher PM levels than rural nurseries.\textsuperscript{43,44} There are also indoor determinants that strongly influence the PM level. In indoor environments, particulates can be generated from human-related activities like cooking, activities of children (playing/walking), cleaning activities, office equipment (e.g. printers), and from construction-related activities like renovation and reconstruction.\textsuperscript{11,29} Studies also find that higher indoor PM levels are associated with high occupant density and PM\textsubscript{10} concentrations are more sensitive to occupancy than PM\textsubscript{2.5}.\textsuperscript{45–47} However, a small number of children in the classroom is enough to increase PM concentrations.\textsuperscript{44} It is worth noting that indoor PM levels are higher in the classrooms of older children, due to the high level of activity of older children.\textsuperscript{47,48}

**PM\textsubscript{10}.** The reported PM\textsubscript{10} levels in indoor nursery environments ranged from 6.8 to 216 µg/m\textsuperscript{3}.\textsuperscript{29,31,46,49–52} As shown in Figure 4, almost all the studies report indoor PM\textsubscript{10} levels that exceed the 50 µg/m\textsuperscript{3} 24-hour mean guidelines recommended by WHO.\textsuperscript{53} A study from Cano et al.\textsuperscript{49} reported that floor covering material might be a crucial element that influences the indoor level of PM\textsubscript{10}. Among the hard surface flooring materials (e.g. wood, tile/stone or PVC), wooden floors are more likely to become the source of PM\textsubscript{10}. The authors speculated that this might be due to the difficulty of adequately cleaning the joints in wooden floors. A meta-analysis reported that with an increase of 10 µg/m\textsuperscript{3} of PM\textsubscript{10}, there was an increase of 2.8\% in asthma symptoms and 1.2\% in cough.\textsuperscript{54} Exposure to air pollutants such as PM\textsubscript{10} was associated with illness-related absenteeism. In a three-year study, the estimated relative illness-related absenteeism risks were 1.06 (95\% confidence interval, 1.04–1.09) per 42.1 µg/m\textsuperscript{3} increase in PM\textsubscript{10}.\textsuperscript{55}

**PM\textsubscript{2.5}.** The reported PM\textsubscript{2.5} levels in nursery environments ranged from 3.2 to 177.2 µg/m\textsuperscript{3}.\textsuperscript{11,43,46,50,51,56–58} As shown in Figure 5, almost all the studies reported substantially higher indoor PM\textsubscript{2.5} levels than the 25 µg/m\textsuperscript{3} 24-h mean guideline recommended by WHO.\textsuperscript{53} One study found that children exposed to an excess level of PM\textsubscript{2.5} have a greater risk of
respiratory symptoms and reduced lung function. An epidemiological study reported that a 10 μg/m³ increase in PM$_{2.5}$ was correlated with a 15% rise in hospital admissions for asthma. Ultra-fine particulates. Limited data is available on the concentration of ultrafine particles (particulate matter of nanoscale size; less than 0.1 μm or 100 nm in diameter) in nurseries. The main
elements influencing UFP level are summarised as: children’s activities during classes (e.g. painting and other arts and crafts activities), combustion sources (e.g. candles on a birthday cake), and classroom cleaning (e.g. dusting and wood polishing).61

In Portugal, a study investigated three nurseries and reported a mean concentration of $1.82 \times 10^4$ particle/cm$^3$ and $1.32 \times 10^4$ particle/cm$^3$ in urban nurseries, and $1.15 \times 10^4$ particle/cm$^3$ in a rural nursery. They concluded that canteens have the highest UFP level, likely because they were directly connected to the kitchen with a gas stove. Also, the UFP levels in playrooms were about two times higher than in classrooms. It’s worth noting that during two activities (candles burning on a birthday cake and clay grinding), the concentrations were 13 times higher than the estimated mean value.62 Due to the small size of UFPs, they can penetrate biological membranes and pass into the systemic circulation, and eventually get into organ systems including the brain and nervous system. Studies about independent health effects of UFP are scarce. A review study identified 85 studies and reported that there were inflammatory and cardiovascular changes associated with short-term UFP exposure.63

**Nitrogen dioxide (NO$_2$).** Compared to other indoor environments, nurseries tend to have higher indoor NO$_2$ levels. Indoor levels are strongly influenced by outdoor levels associated with road traffic. For convenience, nurseries are often located on the ground floor and close to main roads making them vulnerable to this pollutant.13,64 Studies on indoor NO$_2$ concentrations in nursery environments are scarce. Reported indoor NO$_2$ levels in nurseries (Figure 6) ranged from undetectable to 30.2 µg/m$^3$, which does not exceed the annual mean value of 40 µg/m$^3$ recommended by WHO.12,13,64,65 However, one study in Portugal found the mean NO$_2$ concentrations in 10 urban and 5 rural classrooms ranged from undetectable to 136 µg/m$^3$ and 16.67–125.17 µg/m$^3$, respectively. The classroom with the highest NO$_2$ concentration was the one located on the ground floor with windows in the front (roadside) façade of the building.34,66

It is worth noting that one study reported higher NO$_2$ levels in classrooms with more students.64 However, in another study, a classroom was measured both fully occupied and partially occupied for NO$_2$ concentrations, and the outcomes were 16.67 and 41.18 µg/m$^3$, respectively. Indicating lower NO$_2$ levels with more students.34 The relationship between indoor NO$_2$ levels and occupant density warrants further exploration.

The health impact of NO$_2$ is primarily on the respiratory system, increasing the risk of lung infection and causing problems such as wheezing, coughing, colds, flu and bronchitis. A meta-analysis found that with an increase of 10 µg/m$^3$ of NO$_2$, there was an increase in asthma symptoms of 3.1%.54 However, compared with other pollutants, the adverse impact of NO$_2$ on health may have a longer lag period, which contributes to the difficulty in studying the relationship between NO$_2$ exposure and health outcomes.67

**Ozone (O$_3$).** Indoor O$_3$ concentrations are mainly influenced by outdoor air. In most circumstances, indoor O$_3$ levels are significantly lower than outdoor levels, because few indoor sources (e.g. printers, electronic air cleaners) are found in nurseries, and O$_3$ is highly reactive.68 One study investigated 10 classrooms in four urban nurseries, the mean O$_3$ concentrations in classrooms ranged from 9 to 24 µg/m$^3$.66 In Singapore, a study focused on the difference between air-conditioned and naturally ventilated nursery classrooms. Naturally ventilated classrooms had a mean O$_3$ concentration of 71.0 µg/m$^3$, which was significantly higher than air-conditioned classrooms with a mean concentration of 31.5 µg/m$^3$.11 A study from Portugal reported a mean O$_3$ concentration of 119 µg/m$^3$ which exceeds the 100 µg/m$^3$ (over an 8-hour period) recommended by WHO.65 The outdoor mean O$_3$ concentration was 188 µg/m$^3$. The authors did not provide a reason for the high concentrations,
but reported that the studied nursery is situated on moderately trafficked streets. A list of studies and their reported findings on O₃ concentration can be found in Figure 7. One study found that outdoor O₃ concentration and total area cleaned are important elements that influence indoor O₃ concentration. During the cleaning process, O₃ and terpene (a constituent of some cleaning products) react to reduce indoor O₃ concentrations.

Exposure to O₃ is associated with various respiratory symptoms including coughing, wheezing, dyspnoea, and other symptoms such as nausea and headache. In Mexico, a study reported that when nursery children were exposed for two consecutive days to relatively high O₃ levels (>0.13 ppm, or 259.4 µg/m³), there was a 20% increased risk of respiratory illness. Another study focused on elementary schools quantifies that further, with an increase of 20 ppb of O₃, there is an 82.9% increase in upper respiratory illnesses, 173.9% for lower respiratory illness with wet cough, and 62.9% for illness-related absence. A separate study, with similar outcomes, estimated that relative risks of illness-related absenteeism for O₃ were 1.08 (95% CI, 1.06–1.11) per 15.94 ppb.

Carbon monoxide (CO). Studies report that CO found indoors is mainly from outdoor sources, and generally traffic related. As a result, nurseries located in urban areas, and in naturally ventilated buildings, tend to have a higher indoor concentration. However, there are still indoor sources that should be considered such as, heating systems, wood-burning stoves, fireplaces, water heaters, clothes-dryers, and stoves.

Studies about CO levels in nursery environments reported an average concentration range from 4.2 to 2786.0 µg/m³. In Greece, a study investigated two primary schools and one kindergarten in their research. They report that, during winter, one room with
kitchen facilities in the kindergarten had an extremely high CO concentration of 4.2 ppm (approximately 4900.0 mg/m³).68 Another study mentioned that schools constructed within one year had significantly higher CO concentrations. As the main heating systems within the schools were electric, higher CO concentrations might be caused by the introduction of outdoor pollutants through open windows during the summer.29

The health effects of breathing CO include headache, dizziness, vomiting, and nausea. If levels are high enough, people can lose consciousness or die. The CO concentrations in the reviewed studies do not exceed the 6.1 ppm for 24-h exposure (approximately 7015.0 mg/m³) established by WHO guidelines.65 However, it should be noted that a study on elementary schools concluded when CO levels increased by 1.0 ppm, absenteeism increased by 3.79%.72

**Volatile organic compounds (VOCs).** In studies on the indoor air quality of nurseries, total VOCs (TVOCs) is used to report the indoor organic chemical compounds level. As shown in Figure 8, reported indoor TVOCs ranged from nondetectable to 6440 mg/m³ (with a mean concentration that ranged from 114 to 642.11 mg/m³). Some high TVOC peaks are included in this range, but further studies that may explain those high peaks have not been conducted.10,18,29,30,49,51

Some detailed research on indoor VOCs in nursery environments have been conducted, with BTEX (benzene, toluene, ethylbenzene and xylenes) the most commonly reported compounds. Reported mean concentration of benzene, toluene, ethylbenzene, m,p-xylenes and o-xylene ranged from 1.4 to 2.93 µg/m³; 2.2 to 7.9 µg/m³; 0.6 to 2.2 µg/m³; 1.6 to 5 µg/m³ and 1.3 to 1.6 µg/m³, respectively.13,30,33,48,58,73 As mentioned by BB101 (2018), 25 trichloroethylene, tetrachloroethylene, naphthalene and d-Limonene are also important chemicals in indoor environments. However, there is limited information about indoor concentrations of those pollutants in nursery environments. One study investigated 7 nurseries and 10 elementary schools in France and reported a mean trichloroethylene concentration of 2.3 µg/m³ with a range of 0–28.3 µg/m³ and mean tetrachloroethylene concentration of 1.1 µg/m³ with a range of 0–11.5 µg/m³.58 Two studies reported a naphthalene concentration that ranged from 0.3 to 3.1 µg/m³.15,48 Studies about the indoor d-Limonene level in nurseries were not found.

![Figure 7. Summary of study reported indoor O₃ concentration means and ranges.](image-url)
Due to the relative complexity of individuals’ susceptibilities to TVOCs, only indicators of sensory effects are reported. The complex mixture of chemicals in TVOCs can cause eye, nose and throat irritation, shortness of breath, headaches, fatigue, nausea, dizziness and skin problems. Higher concentrations may cause irritation of the lungs, as well as damage to the liver, kidney, or central nervous system. An increased cumulative incidence of lower respiratory symptoms was associated with a 2 μg/m³ change in process-related compounds (OR = 1.08).

As products containing formaldehyde such as plywood, particleboard, carpets, and foam insulation are frequently used in indoors, many studies focus on the indoor level of this compound. As shown in Figure 9, mean indoor formaldehyde levels reported are relatively low. However, high concentration peaks were reported in many studies. One study reported those peaks corresponded with poor ventilation and the activities of cleaning and moving furniture (i.e. scraping the floor). Most studies report high formaldehyde during the hot/non-heating season. It is worth noting that there was a strong correlation between benzene and CO with formaldehyde, which might suggest they are from common sources.

Allergens. Studies have demonstrated that indoor allergens are common in nurseries, and allergens can be different due to different geographic, climatic, and cultural factors. Most studies in nursery environments reported a low concentration of allergens which did not exceed recommended levels, however low levels of exposure still have a potential to cause allergic reactions.

Based on current studies, cat (Fel d 1) and dog (Can f 1) allergens were the dominant allergens found in nurseries. Measured cat allergen (Fel d 1) ranged from undetectable to 1.48 μg/g. Measured dog allergen (Can f 1) ranged from undetectable to 3.3 μg/g.

Dust mite (Der f 1 and Der p 1) and cockroach (Bla g 1 and Bla g 2) allergens were also detected in some studies. Dust mite allergens (Der f 1 and Der p 1) ranged from 0.13 to 5.40 μg/g and 0.05 to 21.8 μg/g. In Brazil, a study reported that dust mite allergens were greater than 2 μg/g in 67% of samples collected from day-care centres and preschools, and the highest levels were seen in a preschool bed with a mean Der 1 (Der p 1 + Der f 1) concentration of 6.3 μg/g. Cockroach allergen levels were comparatively low or undetectable in other studies.

The common reservoirs for allergens were carpeting, upholstered furnishings, and
clothing. Animals were not allowed in almost all nurseries, so the indoor allergens were mainly from the hair and clothing of children or nursery staff with a pet at home. It should be noted that nursery children are more likely to play on the floor than school children and therefore may be exposed to a higher allergen level. Cleaning was beneficial at reducing indoor allergens as reported by Smedje et al. Studies have associated wheezing, daytime breathlessness, sensitization and asthma with indoor allergens. A cased-controlled study reported that mite allergens above 10 \( \mu g/g \) of dust was positively associated with wheezing and breathlessness, and that cat allergens above 8 \( \mu g/g \) of dust in home environments was a risk factor for coughing at night.

**Fungi species.** An increase in fungal levels in the indoor environment is associated with mould/water damage in the building structure. Exposure to fungi can cause adverse human health effects from three aspects: immune response, infection by the organism, or toxic-irritant effects from by-products of mould (mycotoxins, MVOCs etc.). The symptoms caused by indoor fungi include respiratory complaints, eye symptoms, and mucous membrane irritation. However, little is known about the relationship between inhalation and response, and there are no unified sampling or analytical methods for mould exposure. Kim et al. reported mean MVOCs concentrations of 423 ng/m\(^3\) in eight primary schools, they also mentioned that nocturnal breathlessness and doctor diagnosed asthma were associated with higher indoor concentrations of total MVOC.

Studies in South Korea and Portugal reported that nurseries tend to have higher fungi concentration compared to homes, hospitals, postpartum nurse centres, primary schools and elderly care centres. The fungi found in the indoor environment were mainly from outdoor sources. *Penicillium* and *Cladosporium* were two main fungi genera found in indoor environments. Studies investigating total indoor fungi concentrations reported results ranging from 69.2 to 707 CFU/m\(^3\), with
a higher concentration in summer.13,18,32,33,48,49,92,94

In tropical countries, the indoor fungi concentrations in nurseries tended to be much higher. In Singapore, a study reported that the total fungi concentration was 1424.2 CFU/m³ on dry days and 2930.5 CFU/m³ on rainy days.95 They also reported that most of the indoor airborne culturable fungi had a size range between 1.1 and 3.3 μm. However, another study reported a dominant size range of 3.3–4.7 μm.92 Further studies are needed to explain this inconsistency. In addition to outdoor air (the main determinant), occupant density, cleaning, pets, plants, plumbing systems, heating, ventilation, air-conditioning systems, mold and dust resuspension all had an impact on the fungi concentrations indoors.11,28

Bacterial concentrations. Studies about indoor airborne bacteria of nurseries mainly focused on the total bacterial concentrations. It is difficult to determine if indoor bacteria have a specific influence on health, because of a lack of speciation information. However, long exposure time in an environment with high levels of bacteria was shown to have adverse health effects.36 It is worth noting that nurseries tended to have higher bacterial concentrations compared with other indoor environments that were tested. These high levels may be due to higher occupant densities, activities of children, and poor ventilation. A study done in Portugal investigated four environments including homes, child day-care centres, primary schools and elderly care centres, they reported the highest bacterial concentration with a median of 3870 CFU/m³ in 50 classrooms of nine child day-care centres, and found that children have at least two times the dose rates of bacteria than older people.32

Most studies reported significantly higher indoor total bacterial concentration than outdoor, with results that ranged from 1596 to 4630 CFU/m³.18,33,48,49,94 Based on these higher indoor concentrations, the main airborne-bacteria sources are likely from indoors. Human oral and respiratory droplets emitted during coughing, sneezing, talking, breathing, and skin shedding are likely sources.94 The reported bacteria concentrations were much lower in some locations. Researchers in South Korea studied 43 child care facilities and the mean total suspended bacteria was 418 CFU/m³.31 Another study in South Korea reported that the mean concentrations of total and respirable airborne bacteria were 931 and 358 CFU/m³ in childcare centres.92

In addition to studies about total bacteria concentration, a few studies focussed on determining the size distribution and the genera of indoor bacteria in nursery environments. One study reported that Staphylococcus spp., Micrococcus spp., Corynebacterium spp., and Bacillus spp. (mainly gram-positive bacteria), were dominant genera and accounted for over 95% of the total airborne bacteria.92 In Poland, a study investigated one urban nursery and identified Micrococcus spp. (a gram-positive bacteria) as the dominant indoor bacteria. They also analysed the size distribution of bacterial aerosols and concluded that small particles (<4.7 μm) contributed up to 85% of the total bacterial aerosols in indoor air.

Radon. Radon is a naturally occurring radioactive gas produced from the decay of uranium in soil and rocks. It can penetrate buildings through cracks in the foundation. When inhaled, it remains in the lungs and continuously releases ionizing radiation that damages tissue.96 A review paper reported that Radon is a human lung carcinogen, and is the second leading cause (after tobacco smoke) of death from lung cancer.97

Most studies that looked at the radon levels in nursery environments reported acceptable average indoor values, within the 100 Bq/m³ recommended by WHO.98–101 Some studies reported ranges of 100–300 Bq/m³ recommended by ICRP (The International Commission on Radiation Protection).102–106

However, in some high radon areas, studies reported relatively high levels. In Slovenia, a
study investigated 10 high radon level kindergartens. The average indoor air radon concentration ranged from 264 to 1700 Bq/m³. Studies have reported high radon levels in other countries (Italy; Slovenia; Bulgaria), with results ranging as following: 50–1047 Bq/m³; 145 to 794 Bq/m³; 104–1761 Bq/m³. Although in many places, radon is not a frequent contamination, in high-risk locations, radon should be taken into consideration in the indoor environments of nurseries.

**IAQ control solutions**

There are three main approaches to improving indoor air quality: (1) eliminate sources of indoor air pollution, (2) improve ventilation, and (3) air cleaning. Many studies attempt to track the sources of indoor air pollutants to suggest means of reducing the levels, and ventilation is recognized as an efficient method of diluting pollutants from indoor sources. Nature ventilation and HVAC system could be used to increase the air exchange rate. However, these approaches are not always feasible, leaving air cleaning as an additional strategy to improve indoor air quality.

Air purifiers are designed to remove pollutants of different types. Air cleaning can be integrated into the ventilation system to serve multiple spaces within a building, or it can be a portable or fixed (wall, window, or ceiling) device installed in one room or area. Current air cleaning technologies were reviewed by Luengas et al. and Kelly et al. A summary of these technologies, and their pros and cons, can be found in Table 2. The pollutants targeted in air cleaning are PM, VOCs and bioaerosols. Studies report efficient removal rates in many circumstances. However, there are still limitations for each technology, including unwanted and potentially harmful by-products such as O₃ and NOₓ. The literature review by Luengas et al. investigated various types of indoor air treatment and reported that “mechanical filtration is a simple and extensively used technique for removing suspended particles from indoor air”.

The use of air cleaner interventions to reduce particulate pollutants at homes has been demonstrated to be effective in improving indoor air quality. In the U.S., a study found that an air cleaner intervention (using HEPA filtration) in homes substantially decreased the indoor PM₂.₅ levels, from 38 to 24 μg/m³ over a 12 month period. However, research on school and nursery environments are scarce, information about those few studies can be found in Table 3. One study selected 18 classrooms (nine control, nine intervention) in three urban elementary schools, they reported that the PM₂.₅ levels in the intervention classrooms with HEPA filters were substantially reduced compared with control classrooms, with mean PM₂.₅ concentrations reduced from 6.2 μg/m³ to 2.4 μg/m³. In South Korea, PM₁₀, PM₂.₅, airborne bacteria, and fungi were measured for five days in ten nurseries before the use of an air purifier system. They then took the same measurements with the system operating. The researchers reported that all pollutants were substantially reduced over the three weeks of air cleaner use. Concentrations dropped from 39.9 μg/m³ to 5.6 μg/m³ for PM₂.₅, and from 81.3 μg/m³ to 15.0 μg/m³ for PM₁₀. The bio-aerosol concentrations decreased from 794.1 CFU/m³ to 304.4 CFU/m³ and from 94.4 CFU/m³ to 42.5 CFU/m³ for airborne bacteria and fungi, respectively. Another pilot study tested the efficiency of HEPA filtration in four nurseries, they selected two classrooms (one with an air cleaner and one without) in each nursery building. The measured PM₂.₅ concentrations were, nursery A: 33.0 μg/m³ and 20.9 μg/m³; nursery B: 13.3 μg/m³ and 7.3 μg/m³; nursery C: 17.8 μg/m³ and 8.4 μg/m³; nursery D: 17.1 μg/m³ and 13.0 μg/m³. Outdoor PM₂.₅ concentrations were 35.5 μg/m³, 18.6 μg/m³, 26.9 μg/m³ and 21.9 μg/m³, respectively. Although air cleaning appears to be a good way to remove indoor air pollutants, the links between it and health improvements need further development.
Table 2. Air cleaning technology, targeted pollutant, and assessment.

| Technology                  | Targeted pollutants                  | Pros                                                                 | Cons                                                                 |
|-----------------------------|---------------------------------------|----------------------------------------------------------------------|----------------------------------------------------------------------|
| Mechanical filtration       | Suspended particles                   | Over 95% efficient in removing particles of all sizes (HEPA filter)  | Filters must be replaced to maintain removal efficiency. If not, there is risk of growing harmful microorganisms |
| Electronic filtration       | Suspended particles                   | 90 and 95 % efficient in removing particles from 0.3–6 μm (electrostatic filters and ion generators respectively) | High relative humidity will negatively affect removal efficiency; potential generation of hazardous by-products |
| Adsorption                  | VOCs, O₃, NO₂, SO₂, H₂S, bacteria and fungi | Over 90% removal efficiency for gaseous pollutants, bacteria and fungi | High relative humidity and pollutant load variations compromise the efficiency; airborne bacteria might thrive on carbon sorbents; waste pollutants might re-enter the air if the media is full |
| Ozonation                   | Microbial agents and odours           |                                                                      | The efficient removal rate of indoor air contaminants cannot be guaranteed (with a safe O₃ level of 50–100 ppb); produces potentially harmful secondary organic aerosols |
| UV photolysis               | Bioaerosols such as airborne viruses, bacteria, dust mites, animal dander and mould | Affordable and efficient                                              | Produces ozone and free radicals with harmful effects               |
| Photocatalytic oxidation    |                                       |                                                                      | Moderate performance; short lifetime of the catalyst; generates intermediates and harmful by-products |
| Cold plasma or non-thermal plasma (NTP) | PM, biological pollution and VOCs | Over 95% and 85–98% removal efficiency for bacterial and fungal species | Poor energy efficiency; formation of O₃, NOₓ and other hazardous organic by-products |
| Biofiltration               | VOCs and inorganic gases              | Cost-effective and eco-friendly                                       | Poor pollutant transfer from gas phase to biofilm; limitations in the case of poorly soluble or recalcitrant substances; the potential release of dust and microorganisms |
| Botanical purification      | VOCs                                  | A plant’s ability to take up VOCs is well documented in laboratory studies under controlled conditions | Further research on the full capacity of plants and their response in real indoor scenarios is needed |
| Study                | No. of nurseries | Targeted pollutants                        | Filtration technology | Reference classroom Outdoor levels | Study design                                                                                                           | Conclusion/effectiveness                                                                                                                                                                                                 |
|---------------------|------------------|--------------------------------------------|-----------------------|-----------------------------------|------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Rosén and Richardson | 2                | Particulate Matter (0.3–7 μm)              | Electrostatic Air No Cleaning (EAC system) | Taken into consideration          | Over three years, the EAC systems were turned on year 2, then turned off and left in place throughout year 3. Indoor particle counts were recorded for 24-h periods in one centre only. | Reductions of 78% for fine particles (PM$_{3.2}$) from outdoors. The very fine particles (PM$_{0.3-3}$) produced indoors were reduced by 45% compared with rooms without air cleaners, a substantial reduction from 428 to 232 particles/litre of air. |
| Oh et al.           | 10               | PM$_{2.5}$, PM$_{10}$, airborne bacteria and fungi | Air purifier (LA-NRI19SWF, Korea) | Not taken into consideration      | (1) PM$_{2.5}$, PM$_{10}$, airborne bacteria and fungi were measured with air purifier off; (2) air purifier was on for five days; (3) measured for twenty days. Concentrations compared before and after air purifier use. | The concentration went from 39.9 μg/m$^3$ to 5.6 μg/m$^3$ for PM$_{2.5}$ and from 81.3 μg/m$^3$ to 15.0 μg/m$^3$ for PM$_{10}$. The bioaerosol concentration went from 794.1 CFU/m$^3$ to 304.4 CFU/m$^3$, and from 94.4 CFU/m$^3$ to 42.5 CFU/m$^3$ for airborne bacteria and fungi, respectively. |
| Gayer et al.        | 4                | PM$_{2.5}$                                | HEPA filters          | Taken into consideration          | In each of the selected classrooms where routine work was performed the air purifier was turned on for 24 h, seven days a week, and in the other room the air purifier was turned off. Measurements of PM$_{2.5}$ concentration were taken over five days (three working days and weekend). | Two classrooms were measured (one with an air cleaner and one without) in each nursery, the measured mean PM$_{2.5}$ concentrations were 20.9 μg/m$^3$ and 33.0 μg/m$^3$; 13.3 μg/m$^3$ and 7.3 μg/m$^3$; 17.8 μg/m$^3$ and 8.4 μg/m$^3$; 17.1 μg/m$^3$ and 13.0 μg/m$^3$ for the classroom with and without an air cleaner respectively. |
During the COVID-19 pandemic, indoor air quality has become even more critical. Qian et al.\textsuperscript{116} investigates 318 outbreaks in China and reported that they all occurred in indoor environments. Respiratory droplets (generally $>5\, \mu m$) and aerosol droplets (generally $<5\, \mu m$) which can carry the SARS-CoV-2 virus (the causative agent for COVID-19) are the primary means of airborne transmission of COVID-19.\textsuperscript{117} Respiratory droplets deposit on the ground or surface rapidly, but aerosol droplets may remain suspended in indoor air for one or more hours. Knibbs et al.\textsuperscript{118} discussed that increased air exchange rates could decrease the risk of airborne disease transmission through dilution with outdoor air. In addition to, or in lieu of, increased ventilation, air filtration could also be used to help reduce the transmission risk by reducing the concentration of virus-laden droplets. One recent review paper reported that air purifiers with HEPA filters which have a high removal rate for indoor particles larger than 0.3\,\mu m, may be an effective method for reducing viral load in hospital environments.\textsuperscript{119} Previous experiments on SARS-CoV-1 (the causative agent of the SARS outbreak) demonstrated the efficacy of HEPA filters in the ‘capture and containment of diseases of similar particle size’.\textsuperscript{120} However, as no direct studies have been conducted to validate this assumption, more specific studies on the impact of using air purifiers on indoor viral load of SARS-CoV-2 are needed.\textsuperscript{117}

**Discussion and conclusions**

This report focuses on indoor air quality in nursery environments. The overall evidence indicates that the indoor air quality in nurseries is poor which warrants further attention and remediation. Poor indoor air quality might lead to some negative health outcomes which cannot be ignored. Key findings are as follows:

1. Regarding thermal conditions in nurseries, most reported temperature and relative humidity levels lie within the comfort range. However, both lower and higher temperatures occurred due to poor building facilities. HVAC systems should be properly operated and maintained, which may require appropriate training or additional facilities personnel. Also, it was reported that children prefer a lower temperature than adults, and that there is a difference between the preferences of boys and girls. The methods of collecting accurate feedback on the thermal comfort of nursery children, and guidelines based on the needs of nursery children, warrants further development.

2. Ventilation in nurseries appears inadequate based on CO$_2$ concentrations which commonly exceeded recommended standards (mean concentrations: 377–2750 ppm). The main reasons for the high CO$_2$ levels were overcrowding and poor ventilation of the classrooms. Higher ventilation rates, reducing occupant density, and additional mechanical ventilation are recommended. Additionally, sleep time and sleeping-only rooms should be of special consideration because higher CO$_2$ concentrations were often reported during naptime.

3. Particulate matter of both indoor and outdoor origin was reported in studies. PM levels in nurseries often exceeded current guidelines. Air cleaning systems may be useful in improving indoor air quality. High intensity activities of children, as well as activities that produce indoor particles (e.g. cooking, burning candles, clay grinding), should be especially noted.

4. Indoor NO$_2$, O$_3$ and CO levels were often influenced by outdoor levels, and although limited information was reported, some measurements exceeded the current guidelines. Urban nurseries, or nurseries adjacent to high traffic areas, should be aware of these pollutants. When ambient air quality is not ideal (e.g. during peak traffic periods), ventilation from outdoor air without adequate filtration may not be advisable.

5. High peaks in VOC concentration were reported in most studies, the mean concentrations however, were generally low. The effect
of short-term exposures to VOCs on children’s health needs further study. Construction materials, interior decoration, cleaning and office products should be carefully selected.

6. Bioaerosols like allergens, fungi species and bacteria were reported in some studies. Schools could be potential important sources of exposures to those contaminants. Low levels of exposure might still cause adverse health outcomes. Well defined thresholds for biological contaminants are needed.

7. In high-risk locations, indoor radon levels should be measured, and appropriate remediation actions taken if standards are exceeded.

8. Air purifiers may be a useful tool to help improve indoor air quality when source control and ventilation alone cannot achieve the necessary levels. The filtration technology should be carefully selected as some air cleaning technologies produce unwanted by-products. Currently, HEPA filters are suggested as one of the best options (especially for reducing PM levels). However, it should be noted that the costs of purchasing and maintaining air purifiers could exacerbate existing health inequalities.121

Studies reported in this review originate from different countries with different climates. Also, the methods used are different, for instance, monitoring devices, monitoring periods, and monitored parameters varied in different studies. It’s therefore difficult to directly compare results, and we can only gain a general understating about the current IAQ performance in nursery environments. More comprehensive studies with longer monitoring campaigns and more considered confounders are needed to help us further understand the issues.

Additionally, more research from developing countries, where approximately 70% of the world’s population lives, is needed. Studies have found that poor indoor air quality in homes in developing countries has a fundamental impact on health.122,123 Issues around access to childcare and early childhood education go beyond the scope of this review. However, it is noteworthy that most of the studies cited in this review were from developed countries, and yet the overall IAQ performance was unacceptable. The authors express concern, therefore, that the air quality in nurseries in developing countries may be even more precarious, especially where outdoor pollution is high and the structural fabric of buildings is poor.

The present review of the literature highlights the poor indoor air quality in nurseries and its potential effect on the health of children. When it comes to nurseries, designers should take ambient pollution levels, ventilation, filtration, decoration and construction materials, and occupant density into consideration to design for healthier indoor environments in the future.

Declaration of conflicting interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding
The author(s) disclosed receipt of the following financial support for the research, authorship, and/or publication of this article: EPSRC Impact Acceleration Award: Indoor Air Quality in London’s Nurseries.

ORCID iDs
Shuo Zhang id https://orcid.org/0000-0002-3496-249X
Katherine Curran id https://orcid.org/0000-0001-7169-3359

References
1. Kampa M and Castanas E. Human health effects of air pollution. *Environ Pollut* 2008; 151: 362–367.
2. Madureira J, Alvim-Ferraz MCM, Rodrigues S, et al. Indoor air quality in schools and health symptoms among Portuguese teachers. *Hum Ecol Risk Assess* 2009; 15: 159–169.
3. Almeida SM, Canha N, Silva A, et al. Children exposure to atmospheric particles in indoor of Lisbon primary schools. *Atmos Environ* 2011; 45: 7594–7599.
4. Calderón-Garciduenas L, Torres-Jardón R, Kulesza RJ, et al. Air pollution and detrimental effects on children’s brain. The need for a multidisciplinary approach to the issue complexity and challenges. *Front Hum Neurosci* 2014; 8: 613.
5. Deng Q, Lu C, Ou C, et al. Effects of early life exposure to outdoor air pollution and indoor renovation on childhood asthma in China. Build Environ 2015; 93: 84–91.
6. Chatziidakou L, Mumovic D and Summerfield AJ. What do we know about indoor air quality in school classrooms? A critical review of the literature. Intell Build Int 2012; 4: 228–259.
7. Standard A. Standard 55–2017 thermal environmental conditions for human occupancy. Atlanta, GA: ASHRAE.
8. EPA (United States Environmental Protection Agency). A brief guide to mold, moisture, and your home. 2012. https://www.epa.gov/mold/briefguide-mold-moisture-and-your-home (2010, accessed 10 March 2021).
9. Daneault S, Beausoleil M and Messing K. Air quality during the winter in Quebec day-care centers. Am J Public Health 1992; 82: 432–434.
10. Ruotsalainen R, Jaakkola N and Jaakkola JJK. Ventilation and indoor air quality in Finnish daycare centers. Environ Int 1993; 19: 109–119.
11. Zuraimi MS and Tham KW. Indoor air quality and its determinants in tropical child care centers. Atmos Environ 2008; 42: 2225–2239.
12. Kamaruzzaman S and Razak R. Measuring indoor air quality performance in Malaysian government kindergarten. J Build Perform 2011; 2: 70–79.
13. Roda C, Barral S, Ravelomanantsoa H, et al. Assessment of indoor environment in Paris child day care centers. Environ Res 2011; 111: 1010–1017.
14. Branco PTBS, Alvim-Ferraz MCM, Martins FG, et al. Children’s exposure to indoor air in urban nurseries—part I: CO2 and comfort assessment. Environ Res 2015; 140: 1–9.
15. St-Jean M, St-Amand A, Gilbert NL, et al. Indoor air quality in Montréal area day-care centres, Canada. Environ Res 2012; 118: 1–7.
16. Zender - Swierzcz E and Telejko M. Indoor air quality in kindergartens in Poland. IOP Conf Ser: Mater Sci Eng 2019; 471: 092066.
17. Ferg S-F and Lee L-W. Indoor air quality assessment of daycare facilities with carbon dioxide, temperature, and humidity as indicators. J Environ Health 2002; 65(4): 14–22.
18. Mendes A, Aelenei D, Papoila AL, et al. Environmental and ventilation assessment in child day care centers in Porto: the ENVI RH project. J Toxicol Environ Health A 2014; 77: 931–943.
19. Yun H, Nam I, Kim J, et al. A field study of thermal comfort for kindergarten children in Korea: an assessment of existing models and preferences of children. Build Environ 2014; 75: 182–189.
20. Fang L, Clausen G and Fanger PO. Impact of temperature and humidity on perception. Indoor Air 1998; 8: 80–90.
21. Haghighat F and De Bellis L. Material emission rates: literature review, and the impact of indoor air temperature and relative humidity. Build Environ 1998; 33: 261–277.
22. Fang L, Wyon DP, Clausen G, et al. Impact of indoor air temperature and humidity in an office on perceived air quality, SBS symptoms and performance. Indoor Air 2004; 14: 74–81.
23. Taylor J, Davies M, Mavrogianni A, et al. The relative importance of input weather data for indoor overheating risk assessment in dwellings. Build Environ 2014; 76: 81–91.
24. Strachan DP and Sanders CH. Damp housing and childhood asthma; respiratory effects of indoor air temperature and relative humidity. J Epidemiol Community Health 1989; 43: 7–14.
25. Education and Skills Funding Agency. BB 101: Guidelines on ventilation, thermal comfort, and indoor air quality in schools. https://www.gov.uk/government/publications/building-bulletin-101-ventilation-for-school-buildings (2018, accessed 10 March 2021).
26. ASHRAE A. ASHRAE Standard 62.1-2016. Ventilation for acceptable indoor air quality. Atlanta, GA: ASHRAE.
27. Stankevica G and Lesinskis A. Indoor air quality and thermal comfort evaluation in Latvian daycare centers with carbon dioxide, temperature and humidity as indicators. J Civ Eng Archit 2017; 6: 633–638.
28. Rejc T, Kukec A, Bizjak M, et al. Microbiological and chemical quality of indoor air in kindergartens in Slovenia. Int J Environ Res Health 2019; 00: 1–14.
29. Yang W, Sohn J, Kim J, et al. Indoor air quality investigation according to age of the school buildings in Korea. J Environ Manage 2009; 90: 348–354.
30. Yoon C, Lee K and Park D. Indoor air quality differences between urban and rural preschools in Korea. Environ Sci Pollut Res Int 2011; 18: 333–345.
31. Kabir E, Kim KH, Sohn JR, et al. Indoor air quality assessment in child care and medical facilities in Korea. Environ Monit Assess 2012; 184: 6395–6409.
32. Madureira J, Paciência I, Rufo JC, et al. Assessment and determinants of airborne bacterial and fungal concentrations in different indoor environments: homes, child day-care centres, primary schools and elderly care centres. Atmos Environ 2015; 109: 139–146.
33. Mainka A, Zajusz-Zubek E, Kozielska B, et al. Investigation of air pollutants in rural nursery school—a case study. E3S Web Conf 2018; 28: 01022.
34. Nunes RAO, Branco PTBS, Alvim-Ferraz MCM, et al. Gaseous pollutants on rural and urban nursery schools in Northern Portugal. Environ Pollut 2016; 208: 2–15.
35. Seppäinen OA and Fisk WJ. Summary of human responses to ventilation. Indoor Air Suppl 2004; 14: 102–118.
36. Daisey JM, Angell WJ and Apte MG. Indoor air quality, ventilation and health symptoms in schools: an analysis of existing information. Indoor Air 2003; 13: 53–64.
37. Sundell J, Levin H, Nazaroff WW, et al. Ventilation rates and health: multidisciplinary review of the scientific literature. Indoor Air 2011; 21: 191–204.

38. Smedje G and Norbäck D. New ventilation systems at select schools in Sweden – effects on asthma and exposure. Arch Environ Health 2000; 55: 18–25.

39. Shendell DG. Associations between classroom CO2 concentrations and student attendance in Washington and Idaho. Indoor air, 14(5): 333–341.

40. Kim KH, Kabir E and Kabir S. A review on the human health impact of airborne particulate matter. Environ Int 2015; 74: 136–143.

41. McCormack MC, Breyssse PN, Matsui EC, Center for Childhood Asthma in the Urban Environment, et al. Indoor particulate matter increases asthma morbidity in children with non-atopic and atopic asthma. Ann Allergy Asthma Immunol 2011; 106: 308–315.

42. Kelly FJ and Fussell JC. Improving indoor air quality, health and performance within environments where people live, travel, learn and work. Atmos Environ 2019; 200: 90–109.

43. Oh HJ, Nam IS, Yun H, et al. Characterization of indoor air quality and efficiency of air purifier in childcare centers. Korea Build Environ 2014; 82: 203–214.

44. Nunes RAO, Branco PTBS, Alvim-Ferraz MCM, et al. Particulate matter in rural and urban nursery schools in Portugal. Environ Pollut 2015; 202: 7–16.

45. Alves C, Nunes T, Silva J, et al. Comfort parameters and particulate matter (PM10 and PM2.5) in school classrooms and outdoor air. Aerosol Air Qual Res 2013; 13: 1521–1535.

46. Rim D, Gall ET, Kim JB, et al. Particulate matter in urban nursery schools: a case study of Seoul, Korea during winter months. Build Environ 2017; 119: 1–10.

47. Branco PTBS, Alvim-Ferraz MCM, Martins FG, et al. Indoor air quality in urban nurseries at Porto city: particulate matter assessment. Atmos Environ 2014; 84: 133–143.

48. Mainka A, Bragoszewska E, Kozielska B, et al. Indoor air quality in urban nursery schools in Gliwice, Poland: analysis of the case study. Atmos Pollut Res 2015; 6: 1098–1104.

49. Cano M, Nogueira S, Papoila AL, et al. Indoor air quality in Portuguese children day care centers – ENVIRH Project. In: The Second International Conference on Building Energy and Environment (COBEE), Boulder, CO, USA, 1–4 August, 2012, pp.414–421.

50. Mainka A and Zajusz-Zubek E. Indoor air quality in urban and rural preschools in Upper Silesia, Poland: particulate matter and carbon dioxide. Int J Environ Res Public Health 2015; 12: 7697–7711.

51. Oliveira M, Slezakova K, Delerue-Matos C, et al. Assessment of air quality in preschool environments (3–5 years old children) with emphasis on elemental composition of PM10 and PM2.5. Environ Pollut 2016; 214: 430–439.
66. Branco PTBS, Nunes RAO, Alvim-Ferraz MCM, et al. Children’s exposure to indoor air in urban nurseries – part II: gaseous pollutants’ assessment. *Environ Res* 2015; 142: 662–670.

67. Mukala K, Pekkanen J, Tiittanen P, et al. Personally measured weekly exposure to NO2 and respiratory health among preschool children. *Eur Respir J* 1999; 13: 1411–1417.

68. Kalimeri KK, Saraga DE, Lazaridis VD, et al. Indoor air quality investigation of the school environment and estimated health risks: two-season measurements in primary schools in Kozani, Greece. *Atmos Pollut Res* 2016; 7: 1128–1142.

69. Carlisle AJ and Sharp NCC. Exercise and outdoor ambient air pollution. *Br J Sports Med* 2001; 35: 214–222.

70. Romieu I, Lugo MC, Velasco SR, et al. Air pollution and school absenteeism among children in Mexico city. *Am J Epidemiol* 1992; 136: 1524–1531.

71. Gilliland FD, Berhane K, Rappaport EB, et al. The effects of ambient air pollution on school absenteeism due to respiratory illnesses. *Epidemiology* 2001; 12: 43–54.

72. Chen L, Jennison BL, Yang W, et al. Elementary school absenteeism and air pollution. *Inhal Toxicol* 2000; 12: 997–1016.

73. Sofuoglu SC, Aslan G, Inal F, et al. An assessment of indoor air concentrations and health risks of volatile organic compounds in three primary schools. *Int J Hyg Environ Health* 2011; 214: 36–46.

74. McCloskey WW. American journal of therapeutics. *JAMA* 1996; 276: 504–505.

75. Gilbert NL, Guay M, Gauvin D, et al. Air change rate and concentration of formaldehyde in residential indoor air. *Atmos Environ* 2008; 42: 2424–2428.

76. Ramalho O, Wyart G, Mandin C, et al. Association of carbon dioxide with indoor air pollutants and exceedance of health guideline values. *Build Environ* 2015; 93: 115–124.

77. Salo PM, Sever ML and Zeldin DC. Indoor allergens in school and day care environments. *J Allergy Clin Immunol* 2009; 124(2): 185–192.

78. Grant T, Rule AM, Koehler K, et al. Sampling devices for indoor allergen exposure: pros and cons. *Curr Allergy Asthma Rep* 2019; 19: 9.

79. Rullo VEV, Rizzo MC, Karla LA, et al. Daycare centers and schools as sources of exposure to mites, cockroach, and endotoxin in the city of São Paulo, Brazil. *J Allergy Clin Immunol* 2002; 110: 582–588.

80. Instanes C, Hetland G, Berntsen S, et al. Allergens and endotoxin in settled dust from day-care centers and schools in Oslo, Norway. *Indoor Air* 2005; 15: 356–362.

81. Sooc O, Korkmaz M, Erdem N, et al. An important source for cat and house dust mite allergens: day-care centers. *Turkiye Klinikleri J Med Sci* 2012; 32: 750–758.

82. Cyprowski M, Buczyńska A and Szadkowska-Stańczyk I. Indoor allergens in settled dust from kindergartens in city of Łódź, Poland. *Int J Occup Med Environ Health* 2013; 26: 890–899.

83. Fernández-Caldas E, Codina R, Ledford DK, et al. House dust mite, cat, and cockroach allergen concentrations in daycare centers in Tampa, Florida. *Ann Allergy Asthma Immunol* 2001; 87: 196–200.

84. De Andrade AD, Charpin D, Birnbaum J, et al. Indoor allergen levels in day nurseries. *J Allergy Clin Immunol* 1995; 95: 1158–1163.

85. Kim JL, Elfman L, Mi Y, et al. Current asthma and respiratory symptoms among pupils in relation to dietary factors and allergens in the school environment. *Indoor Air* 2005; 15: 170–182.

86. Bardana EJ. Indoor pollution and its impact on respiratory health. *Am Allergy Asthma Immunol* 2001; 87: 33–40.

87. Gehring U, Heinrich J, Jacob B, Indoor Factors and Genetics in Asthma (INGA) Study Group, et al. Respiratory symptoms in relation to indoor exposure to mite and cat allergens and endotoxins. *Eur Respir J* 2001; 18: 555–563.

88. Bush RK, Portnoy JM, Saxon A, et al. The medical effects of mold exposure. *J Allergy Clin Immunol* 2006; 117: 326–333.

89. Portnoy JM, Kwak K, Dowling P, et al. Health effects of indoor fungi. *Ann Allergy Asthma Immunol* 2005; 94: 313–320.

90. Salonen H, Duchaine C, Mazaheri M, et al. Airborne viable fungi in school environments in different climatic regions – a review. *Atmos Environ* 2015; 104: 186–194.

91. Kim JL, Elfman L, Mi Y, et al. Indoor molds, bacteria, microbial volatile organic compounds and plasticizers in schools – associations with asthma and respiratory symptoms in pupils. *Indoor Air* 2007; 17: 153–163.

92. Kim KY and Kim CN. Airborne microbiological characteristics in public buildings of Korea. *Build Environ* 2007; 42: 2188–2196.

93. Aydogdu H and Asan A. Airborne fungi in child day care centers in Edirne city, Turkey. *Environ Monit Assess* 2008; 147: 423–444.

94. Carreiro-Martins P, Paposil AL, Caires I, et al. Effect of indoor air quality of day care centers in children with different predisposition for asthma. *Pediatr Allergy Immunol* 2016; 27: 299–306.

95. Zuraimi MS, Fang L, Tan TK, et al. Airborne fungi in low and high allergic prevalence child care centers. *Atmos Environ* 2009; 43: 2391–2400.

96. WHO. Handbook on indoor radon – a public health perspective. Geneva, Switzerland, WHO.

97. Al-Zoughool M and Krewski D. Health effects of radon: a review of the literature. *Int J Radiat Biol* 2009; 85: 57–69.
98. Planinić J, Vaupotić J and Kobal I. Indoor radon concentrations in kindergartens. J Environ Radioact 1993; 19: 167–171.

99. Al-Ghamdi SS, Al-Garawi MS, Al-Mosa TM, et al. Investigating indoor radon levels and influencing factors in primary schools of Zulfi City, Saudi Arabia. AIP Conf Proc 2011; 1370: 294–298.

100. Bem H, Bem EM, Krawczyk J, et al. Radon concentrations in kindergartens and schools in two cities: Kalisz and Ostrów Wielkopolski in Poland. J Radioanal Nucl Chem 2013; 295: 2229–2232.

101. Jönsson G, Theodörsson ÖHP and Karlsson S. Indoor and outdoor radon levels in Iceland. In: The NSFS XVII Conference, Copenhagen, Denmark, 24-27 August, 2015, pp.128–134. EDP Sciences.

102. Kullab M. Assessment of radon-222 concentrations in buildings, building materials, water and soil in Jordan. Appl Radiat Isot 2005; 62: 765–773.

103. Fojtíková I and Rovenská KN. Influence of energy-saving measures on the radon concentration in some kindergartens in the Czech Republic. Radiat Prot Dosimetry 2014; 160: 149–153.

104. Ivanova K, Stojanovska Z, Tsenova M, et al. Measurement of indoor radon concentration in kindergartens in Sofia, Bulgaria. Radiat Prot Dosimetry 2014; 162: 163–166.

105. Sousa SIV, Branco PTBS, Nunes RAO, et al. Radon levels in nurseries and primary schools in Bragança District – preliminary assessment. J Toxicol Environ Health A 2015; 78: 805–813.

106. Branco PTBS, Nunes RAO, Alvim-Ferraz MCM, et al. Children’s exposure to radon in nursery and primary schools. Int J Environ Res Public Health 2016; 13: 1–16.

107. Vaupotić J. Search for radon sources in buildings – kindergartens. J Environ Radioact 2002; 61: 365–372.

108. Trevisi R, Leonardi F, Simeoni C, et al. Indoor radon levels in schools of South-East Italy. J Environ Radioact 2012; 112: 160–164.

109. Vaupotić J, Bezek M, Kávási N, et al. Radon and thoron doses in kindergartens and elementary schools. Radiat Prot Dosimetry 2012; 152: 247–252.

110. Vuchkov D, Ivanova K, Stojanovska Z, et al. Radon measurement in schools and kindergartens (Kremikovtsi Municipality, Bulgaria). Rom Reports Phys 2013; 58(S): S328–S335.

111. Siegel JA. Primary and secondary consequences of indoor air cleaners. Indoor Air 2016; 26: 88–96.

112. Luengas A, Barona A, Horta C, et al. A review of indoor air treatment technologies. Rev Environ Sci Biotecnol 2015; 14: 499–522.

113. Eggleston PA, Butz A, Rand C, et al. Home environmental intervention in inner-city asthma: a randomized controlled clinical trial. Ann Allergy Asthma Immunol 2005; 95: 518–524.

114. Jhun I, Gaffin JM, Coull BA, et al. School environmental intervention to reduce particulate pollutant exposures for children with asthma. J Allergy Clin Immunol Pract 2017; 5: 154–159.e3.

115. Gayer A, Mucha D, Adamkiewicz L, et al. Children exposure to PM2.5 in kindergarten classrooms equipped with air purifiers – a pilot study. In: MATEC Web of Conferences 2018, Lwów, Ukraine, 7–8 November, 247, pp.1–7.

116. Qian H, Miao T, Liu L, et al. Indoor transmission of SARS-CoV-2. Indoor Air. Epub ahead of print 31 October 2020. DOI: 10.1111/ina.12766.

117. Christopherson DA, Yao WC, Lu M, et al. High-efficiency particulate air filters in the era of COVID-19: function and efficacy. Otolaryngol Head Neck Surg 2020; 163: 1153–1155.

118. Knibbs LD, Morawska L, Bell SC, et al. Room ventilation and the risk of airborne infection transmission in 3 health care settings within a large teaching hospital. Am J Infect Control 2011; 39: 866–872.

119. Mousavi ES, Kanazizadeh N, Martinello RA, et al. COVID-19 outbreak and hospital air quality: a systematic review of evidence on air filtration and recirculation. Environ Sci Technol. Epub ahead of print 11 September 2020. DOI: 10.1021/acs.est.0c03247.

120. Schentag JJ, Akers C, Campagna P, et al. SARS: clearing the air. In: Learning from SARS: preparing for the next disease outbreak: workshop summary. Washington, DC: National Academies Press, 2004.

121. Cheek E, Guercio V, Shrubsole C, et al. Portable air purification: review of impacts on indoor air quality and health. Sci Total Environ 2021; 766: 142585.

122. Bruce N, Perez-Padilla R and Albalak R. Indoor air pollution in developing countries: a major environmental and public health challenge TT – pollution atmosphérique à l’intérieur des locaux: un problème majeur pour l’environnement et la santé publique TT – contaminación del aire de locales ce. Bull World Health Org 2000; 78: 1078–1092.

123. Gall ET, Carter EM, Earnest CM, et al. Indoor air pollution in developing countries: research and implementation needs for improvements in global public health. Am J Public Health 2013; 103: e67–e72.

124. Gładyszewska-Fiedoruk K and Norback D. Correlations of air humidity and carbon dioxide concentration in the kindergarten. Energy Build 2013; 62: 45–50.

125. Brągoszewska E, Mainka A, Pastuszka JS. Bacterial aerosols in an urban nursery school in Gliwice, Poland: a case study. Aerobiologia (Bologna) 2016; 32: 469–480.

126. Michelot N, Marchand C, Ramalho O, et al. Monitoring indoor air quality in French schools and day-care centers. HVAC R Res 2013; 19: 1083–1089.

127. Rosén KG, Richardson G. Would removing indoor air particulates in children’s environments reduce rate of absenteeism - A hypothesis. Sci Total Environ 1999; 234: 87–93.
**Appendix 1. Sampling/measuring methods used in reviewed papers**

| Research            | Period                              | Method                                | Detailed method                                                                                                                                 |
|---------------------|-------------------------------------|---------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------|
| Zuraimi and Tham    | One day, 8 a.m.–5 p.m.              | Active sampling                       | HOBO H8 Family data loggers (Onset Corporation, Bourne, MA, USA) for T, RH, CO₂; T15v Langan CO Measurers (Langan Products, Inc., San Francisco, CA, USA) for CO; TSI Model 8520 DustTrak aerosol monitors (TSI Incorporated, St. Paul, MN, USA) for PM₂.₅; UV-absorbance ozone analysers (2B Technologies Inc., Golden, CO, USA) for O₃; Single stage Andersen impactor (Graseby–Andersen, Atlantis, GA, USA) for bacteria and fungi. |
| Yoon et al.         | One day, 6–8 h except for TVOCs, which were sampled for 60–100 min | Active sampling and passive sampling  | Direct-reading instrument (IAQ Calc 8762 meter, TSI, USA) for CO₂ and CO; Thermal desorption tube (Tenax TA SS, Supelco, USA) for TVOCs; 2,4-dinitrophenylhydrazine-coated silica tube (300/150 mg, 226-119, SKC, USA) for formaldehyde. |
| Mendes et al.       | Measured twice, during spring (between March and May 2011) and winter (between November 2011 and February 2012) seasons. Starting at 9 a.m. and continuing for at least 4 h during normal activities | Active sampling                       | INNOVA transducers MM0034 for T; INNOVA transducers MM0037 for RH; Photoacoustic multi-gas monitor (model 1312, INNOVA Air Tech Instruments, Ballerup, Denmark) for CO₂ and CO; Poly-tetrafluoroethylene (PTFE) filters on SKC Personal Environmental Monitors (PEM) for PM₁₀; Stainless-steel sampling tube containing Tenax TA using a personal air sampling pump (SKC Pocket pump) for TVOC; NIOSH 3500 method (chromatropic acid) for formaldehyde; MAS-100 (Merck Millipore, Billerica, MA) for bacteria and fungi; Duststream Collector (Indoor Biotechnologies, Cardiff, Wales) for allergens. |
| Research          | Period                                                                 | Method          | Detailed method                                                                                                                                                                                                 |
|-------------------|-------------------------------------------------------------------------|-----------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Cano et al.⁴⁹     | Routine school activities (from 10 AM to 5 PM) Continuous two-month period (March–April) | Active sampling | INNOVA 1221 Thermal Comfort Data Logger for T and RH; Photoacoustic Multi-gas Monitor Type 1312, INNOVA for CO₂ and CO; Active sampling on pre-weighted PTFE filters mounted on PM₁₀ collectors (PEM, SKC), using GilAir 5 personal pumps for PM₁₀; TENAX Tubes using SKC personal pumps for TVOC; Personal pumps GilAir 5 for formaldehyde; Microbiological Air Sampler MAS-100 (Merck) for bacteria and fungi. |
| Branco et al.⁶⁶   | February–November from two to nine days not simultaneously in each studied room (whole day) | Active sampling | Haz-Scanner IEMS Indoor Environmental Monitoring Station (SKC Inc., USA) for T, RH and CO₂                                                                                                                |
| Branco et al.⁶⁶   | February–November from two to nine days not simultaneously in each studied room (whole day) | Active sampling | Haz-Scanner IEMS Indoor Environmental Monitoring Station (SKC Inc., USA) for CO, NO₂, TVOC, formaldehyde and O₃.                                                                                     |
| Branco et al.⁴⁷   | February–June from two to nine days (whole day include weekend)         | Active sampling | TSI DustTrak DRX 8534 particle monitor for PM₁₀, PM₂,5, PM₁ and PMTotal.                                                                                                                                   |
| Nunes et al.⁴⁴    | April–June from one to six days (whole day include weekend)             | Active sampling | TSI DustTrak™ DRX 8534 Aerosol Monitor (Tsi, USA) for PM₁₀, PM₂,5, PM₁ and TSP.                                                                                                                           |
| Nunes et al.³⁴    | April–June from one to six days (whole day include weekend)             | Active sampling | Haz-Scanner IEMS Indoor Environmental Monitoring Station (SKC Inc., USA) for CO₂, CO, NO₂, TVOC, formaldehyde and O₃.                                                                                   |
| Gladyszewska-Fiedoruk¹²⁴ | Before the children’s and the staff’s arrival, as well as after all the classes had finished, at the same time of the day | Active sampling | Testo 4354 for T, RH and CO₂.                                                                                                                                                                                 |
| Mainka et al.⁴⁸,⁵⁰ | From 9 to 20 December 2013 (SU-1) and from 7 to 17 January 2014 (PU-2)  | Active sampling | Automatic portable monitors (model 77,535 Az Instruments International Ltd., Hong Kong) for CO₂; PN-EN12341 standard (2014) for PM₁, PM₂,5, PM₁₀ and TSP; Actively sampled according to the US EPA TO-17 method for VOCs; Six-stage Andersen cascade impactor for bacteria and fungi. |
| Research                  | Period                                  | Method                      | Detailed method                                                                 |
|---------------------------|-----------------------------------------|-----------------------------|---------------------------------------------------------------------------------|
| Mainka and Zajusz-Zubek   | Monday–Friday, 7:30–15:30                | Active sampling             | Automatic portable monitors (model 77,535 Az Instruments International Ltd., Hong Kong) for CO₂; PN-EN12341 standard (2014) for PM₁, PM₂.₅, PM₁₀ and TSP. |
| Bragoszewska et al.        |                                         | Active sampling             | Six-stage Andersen cascade impactor for bacteria.                                |
| Mainka et al.             |                                         |                             | Automatic portable monitors (model 77,535 Az Instruments International Ltd., Hong Kong) for CO₂; PN-EN12341 standard (2014) for PM₁, PM₂.₅, PM₁₀ and TSP; Stainless steel tube samplers containing Tenax GR according to the US EPA TO-17 method for VOCs; Six-stage Andersen cascade impactor for bacteria and fungi. |
| Kamaruzzaman and Razak    | Three days, 7:00–15:00                   | Active sampling             | MultiRAE meter for T, RH and CO₂                                                  |
| Roda et al.               | Cold season: October–March, hot season: April–September, Installed on Monday morning and collected on Friday afternoon (passive sampling) | Active sampling and passive sampling | Q-Traks IAQ monitor (TSI Incorporated, St. Paul, Minnesota, USA) for T, RH and CO₂; Passive samplers, Radiello (Fondazione Salvatore Mauger – IRCCS, Italy) for aldehydes, VOCs, and NO₂; Single-stage multi-holed impactor (Air IdealTM, bioMérieux sa, France) for fungi; Mitest dust collector (Indoor Biotechnologies, Charlottesville, Virginia, USA) for allergens. |
| Canha et al.              | Heating season (from 11 January 2010 to 2 April 2010); non-heating season (from 26 April 2010 to 25 June 2010), From Monday to Friday 8:00 to 17:00 | Active sampling and passive sampling | Q-Trak Plus IAQ monitor 8552 (TSI Incorporated, Shoreview, MN, USA) for T, RH and CO₂; Gravimetric MicroVol samplers (Europa Environmental, Twynning, England) for PM₂.₅; Passive samplers (Radiello, Fondazione Salvatore Mauger, Padova, Italy) for VOCs and formaldehyde. |
| Ramalho et al.            | 4.5 days from Monday morning to Friday evening during both heating and non-heating seasons | Active sampling and passive sampling | Non-dispersive infrared sensor (Q-Trak IAQ model 8550, TSI Inc.) for CO₂; PEMS impactor at 1.8L/min with Chempass model 3,40,02,100 Minipartisol air sampler (Rupprech & Patachnick Co, Inc.) for PM₁₀ and PM₂.₅; Radial passive sampler with 2,4-DNPH coated Florisil® (Radiello® cartridge code 165) for formaldehyde. |
| Michelot et al. 2013      | Monday morning to Friday afternoon, i.e., for 4.5 days, both heating and non-heating seasons | Passive sampling            | Radial passive sampler with 2,4-DNPH coated Florisil® (Radiello® cartridge code 165) for formaldehyde. |

(continued)
Continued.

| Research                         | Period                                          | Method                   | Detailed method                                                                                                                                 |
|----------------------------------|-------------------------------------------------|--------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------|
| Daneaultet al.                  | Active sampling                                 |                          | Sling psychrometer (Bacharach Instruments) for T, RH and CO₂                                                                                   |
| St-Jean et al.                   | January 14–February 22, 2008; from Monday or Tuesday morning to Thursday or Friday afternoon, the passive devices were deployed for at least 6 h in one of the monitored rooms and removed the same day (Thursday or Friday) | Active sampling and passive sampling | YES-206LH Falcon monitor (Yes Environment Technologies Inc., Delta, BC, Canada) for T, RH and CO₂; passively using 6.0 L cleaned and evacuated Summa canisters for VOCs; SKC UMEEx 100 Passive Samplers (SKC Inc. Eighty-Four, PA, USA) for formaldehyde |
| Andrade et al.                   | Samples for allergens were collected according to the recommendations of the First International Workshop on Dust Allergens and Asthma |                          |                                                                                                                                               |
| Vaupotic                       | Two months                                       |                          | Instantaneous: alpha scintillation cells, average: Karlsruhe etched track detectors for Radon mAb-based ELISAs for allergens                   |
| Rullo et al.                    | Same day 7:30–3:30                               | Active sampling          | TSI Q-Trak IAQ Monitor Model 8550 for T, RH and CO₂                                                                                           |
| Ferng and Lee                   | 11:30 am–1:00 pm once in spring (March–May) and autumn (September–November), respectively, in 2004. | Active sampling          | Six-stage viable particulate cascade impactor (Model 10-800, Andersen Inc, USA) for bacteria and fungi                                          |
| Kim and Kim                    |                                                |                          |                                                                                                                                               |
| Yang et al.                     | Active sampling and passive sampling             |                          | Non-dispersive infrared (NDIR) analyzer (TSI, model 8762) for CO₂ and CO; Pall flex membrane filter (47 mm, Gelman Science) for PM_{10}; Tenax-TA tubes for TVOC; Supelco LPDNPH S10 for formaldehyde |
| Alves et al.                    | 9–29 January 2012                                | Active sampling          | Indoor Air IQ-610 Quality Probe (Gray Wolf® monitor) for T, RH, CO₂, CO and TVOC; Echo TCR Tecora samplers for PM_{2.5}; TSI Model 8533 DustTrak DRX; AIRE-AIDE model AA-3500 and Topas monitor from Turnkey Instruments (each for one classroom) for PM_{10} |

(continued)
| Research          | Period                                    | Method                  | Detailed method                                                                 |
|-------------------|-------------------------------------------|-------------------------|--------------------------------------------------------------------------------|
| Madureira et al.  | Short-term samplings were performed from two to nine consecutive days in each room (not simultaneously) in nursery schools | Active sampling         | Merck Air Sampler MAS-100 for bacteria and fungi Radim 5B radon monitor for Radon |
| Branco et al.     | 24h, seven weeks of May–July 2013         | Active sampling         | Model TG 502; GrayWolf Sensing Solutions, Shelton, USA for CO₂, CO, TVOC and O₃; Polytetrafluoroethylene membrane disks (Ø47 mm, SKC Ltd., UK) for PM₁₀ and PM₂.₅ |
| Oliveira et al.   | In the heating and non-heating season, respectively, Monday morning to Friday noon | Active sampling and passive sampling | HOBO data loggers for T and RH; Telair 7001 for CO₂; aeroQUAL CO sensors for CO; Derenda LVS3.1/ PMS3.1-15 for PM₂.₅; Grimm 1.108 for PM₁₀; Radiello passive samplers for NO₂, VOCs, formaldehyde and O₃; Gammadata RAPIDOS sampler for Radon |
| Kalimeri et al.   | February–April, during the year 2013       | Passive sampling        | Radiello® passive samplers (Fondazione Salvatore Maugeri, Padova, Italy) for VOCs, carbonyl compounds and NO₂ |