

How efficiently can HEPA purifiers remove priority fine and ultrafine particles from indoor air?

Scott D. Lowther^{1,2}, Wei Deng², Zheng Fang², Douglas Booker³, J. Duncan Whyatt¹, Oliver Wild¹, Xinming Wang², Kevin C. Jones¹

Corresponding authors: Kevin C. Jones k.c.jones@lancaster.ac.uk and Xinming Wang wangxm@gig.ac.cn

1 Lancaster Environment Centre, Lancaster University, Lancaster LA1 4YQ, United Kingdom

2 State Key Laboratory of Organic Geochemistry, Guangzhou Institute of Geochemistry, Chinese Academy of Sciences, 511 Kehua Rd, Tianhe, Guangzhou 510640, China

3 NAQTS, Lancaster Environment Centre, Lancaster University, Lancaster LA1 4YQ, United Kingdom

Highlights

1. It is currently unclear how particles of different sizes are removed by air purifiers
2. Three popular models were tested in China's largest indoor smog chamber
3. Particles <100nm were removed efficiently
4. 200-250nm particles were least efficiently removed
5. Ambient air particles were removed at a similar rate to standard particle types

Abstract

More than 1 million premature deaths in Asia annually are estimated to be associated with indoor air quality. HEPA (high-efficiency particulate air) filter air purifiers (APs) are widely used in urban Chinese residences by the growing middle class, as public awareness of air pollution increases. Currently, understanding of how particle size affects particle removal is inconsistent, and the rate at which different particle types are removed remains largely unknown. Therefore, this investigation aimed to determine the relationship between particle size and the removal efficiency of particles, and how efficiently ambient air is filtered compared to standard particle types which are typically used for such tests (tobacco smoke, dust and pollen). Three of the most popular AP models in China were tested in China's largest indoor controlled chamber laboratory and the removal efficiencies of particles in the 18-514nm range were identified. Each AP had a distinct profile of removal efficiency against particle size, but the three APs shared similarities in performance, with removal efficiency consistently lowest at 200-250nm. This size fraction is important in an exposure context as these particles are abundant in ambient air in mega-cities, can penetrate through building shells effectively, remain airborne for long periods of time and can penetrate the deepest areas of the lungs. Ambient air particles were removed at a similar rate to test particles; this confirms that the Association of Home Appliance Manufacturers' (AHAM) standards are a suitable proxy for "real world" performance.

Keywords: HEPA, Air Purifiers, Air Filtration, Particulate Matter, Ambient PM

35 1. Introduction

36 An estimated 4.2 million premature deaths globally were attributed to indoor air pollution in 2016, compared
37 to 3.8 million from outdoor air pollution (WHO, 2018). It is estimated that 90% of people breathe air that
38 does not comply with the World Health Organization Air Quality Guidelines (WHO, 2016). Poor indoor air
39 quality is estimated to be the 9th largest global burden of disease risk (Forouzanfar et al., 2015). The
40 Institute for Health Metrics and Evaluation (2017) attributed 2.6 million premature deaths to indoor air
41 pollution in 2016; Roser and Ritchie (2018) partitioned this estimate by continent with Asia, Africa, Europe
42 and the Americas contributing 74%, 23%, 1% and 2% respectively, demonstrating the significance of
43 premature deaths in Asian countries. On average, modern populations spend more than 80% of their time
44 indoors (Duan et al., 2015; Klepeis et al., 2001), with the indoor environment contributing 19-76% of an
45 individual's ultrafine particle (UFP) exposure (Morawska et al., 2013).

46 Particulate matter (PM) is defined as the total of all solid and liquid particles suspended in air and is a major
47 determinant of indoor air quality (IAQ) (Lowther et al., 2019). PM is strongly associated with negative
48 health outcomes including strokes, heart failure, asthma and lung cancer (Lim et al., 2012). Size is an
49 important property of PM with regard to its potential health effects. Therefore, PM is commonly categorized
50 based on its aerodynamic diameter into the commonly regulated standards of $<10\mu\text{m}$ (PM_{10}), $<2.5\mu\text{m}$
51 ($\text{PM}_{2.5}$), and $<100\text{nm}$ (UFPs). Smaller particle size fractions are able to penetrate further into the respiratory
52 tract and are thought to have a higher toxicity per unit mass due to a larger surface area to mass ratio
53 (Harrison and Yin, 2000; HEI Review Panel, 2013).

54 In China, more than 1 million premature deaths were attributed to long-term exposure to $\text{PM}_{2.5}$ in 2016
55 (Health Effects Institute, 2018). In 2017, the average annual ambient $\text{PM}_{2.5}$ concentration across 338
56 Chinese cities was $44\ \mu\text{g}/\text{m}^3$, with 73% of these cities failing to meet the national air quality standard of
57 $35\ \mu\text{g}/\text{m}^3$ (Ministry of Ecology and Environment the People's Republic of China, 2018). Furthermore, in
58 China it is estimated that 66-87% of total exposure to $\text{PM}_{2.5}$ of outdoor origin occurs within indoor
59 environments (Xiang et al., 2019). It should, however, be noted that although PM levels in China are severe,

60 rapid reductions in PM concentrations are being observed. For example, the average PM_{2.5} concentration in
61 Beijing dropped from 90 µg/m³ in 2013 to 58 µg/m³ in 2017 ([Ministry of Ecology and Environment of the](#)
62 [People's Republic of China, 2018](#)).

63 The two fundamental sources of PM in indoor environments are: (i) PM generated by indoor sources and
64 activities and (ii) PM generated by outdoor (ambient) sources penetrating indoors. Important indoor PM
65 sources in China include solid fuel use, cooking, smoking and incense burning ([Apte and Salvi, 2016](#); [Tse](#)
66 [et al., 2011](#)). Solid fuel use is especially dangerous in China from a health perspective ([Zhang and Smith,](#)
67 [2007](#)), with solid fuel combustion generating high levels of PM with substantial concentrations of carbon,
68 iron, lead, cadmium and silica ([Apte and Salvi, 2016](#)). However, in the absence of major indoor sources,
69 outdoor to indoor air exchange is the most significant source of PM indoors. In a study of 41 Beijing
70 residences, a strong correlation ($R \geq 0.90$) was found between ambient and indoor PM_{2.5}, with ambient levels
71 accounting for $\geq 84\%$ of the variance of indoor levels ([Huang et al., 2015](#)). In a summary of 77 studies
72 involving over 4000 homes, indoor/ambient PM_{2.5} ratios were found to vary substantially, from 0.5-3.5
73 ([Chen and Zhao, 2011](#)). Additionally, buildings in China are often ineffective in preventing ambient fine
74 particles from entering indoor environments, with standards for air tightness of residential buildings being
75 less restrictive than in the United Kingdom or United States ([Hu et al., 2018](#)). Given that ambient air strongly
76 influences indoor air in China, the composition and properties of the air are likely to be very similar, in
77 contrast to conditions where indoor sources dominate. Therefore, this study focuses on ambient particles
78 that contribute significantly to indoor environments, to estimate the performance of HEPA type air purifiers
79 in real world indoor environments.

80 High-efficiency particulate air (HEPA) filters are an effective technology for improving IAQ when removing
81 PM is the priority ([Zhang et al., 2011](#)). To be defined as such, a HEPA filter must be able to remove 99.97%
82 of particles greater than or equal to 0.3 µm. In a HEPA Air Purifier (AP), air is forced through the HEPA
83 filter and particles are physically captured. The four key mechanisms through which particles are captured
84 are diffusion, interception, inertial impaction and sieving. Diffusion causes the smallest particles to be

85 removed, whereas interception, inertial impaction and sieving processes are more effective at removing
86 the largest particles (Yang, 2012). This means that particles of an intermediate size (100-400nm) are the
87 least efficiently removed (Kowalski et al., 1999). Particle size, charge and shape are the controlling factors
88 determining how effectively particles are removed by the HEPA medium. Studies have shown that HEPA
89 filters can reduce particulate mass and particle number concentrations by >50% (Batterman et al., 2012,
90 2005; Kelly and Fussell, 2019; Ward et al., 2017; Wheeler et al., 2014). There is also limited evidence to
91 suggest that these reductions lead to improvements in cardio-respiratory health (Fisk, 2013; Morishita et
92 al., 2015). Collectively, studies have reported that use of indoor APs may be associated with reductions in
93 blood pressure, oxidative stress, systematic inflammation and improved lung function (Kelly and Fussell,
94 2019). Health benefits are most consistently observed in Asian mega-city homes, likely due to higher
95 baseline indoor concentrations and therefore more significant absolute reductions (Kelly and Fussell, 2019).

96 The Chinese AP market stood at \$ 2 billion in 2017 and is predicted to surpass \$ 4.3 billion in 2023 (BIS
97 Research, 2018). HEPA AP technology held ~40% of market share in 2016 and is the fastest growing
98 segment of the market (BIS Research, 2018). This growth in the market can be attributed to the growing
99 Chinese middle class and improved awareness of IAQ, with APs mainly used by more affluent members of
100 Chinese society.

101 The Association of Home Appliance Manufacturers (AHAM) is the main body which verifies the performance
102 of HEPA APs and although they are based within the United States, they produce certified ratings for AP
103 brands all over the world. They measure the filtering efficiency of HEPA APs using the Clean Air Delivery
104 Rate (CADR) metric - the flow rate of particle-free air output in cubic feet per minute (ft^3/min ; note: 1
105 $\text{ft}^3/\text{min} = 0.028 \text{ m}^3/\text{min}$). AHAM test the CADR of HEPA APs for three particle types, tobacco smoke (0.09-
106 $1 \mu\text{m}$), household dust (0.5-3 μm) and pollen (5-11 μm) (AHAM, 2002). However, within a laboratory
107 context it is currently unknown how efficient HEPA APs are in removing "real world" particles, i.e. those
108 found in ambient air. Therefore, it is valuable to investigate how well ambient air particles are removed in

109 comparison to AHAM standard particle types, to see whether the selected particle types are representative
110 of real-world performance.

111 Combustion-generated particles can penetrate National Institute for Occupational Safety and Health
112 (NIOSH) N95 filtering face-piece respirators more efficiently than standard sodium chloride particles (Gao
113 et al., 2015). Peck et al. (2016) investigated whether this applied to HEPA APs, concluding that diesel
114 combustion particles were removed more efficiently than both NaCl and AHAM test particles, with lowest
115 and highest removal efficiencies at 42-100 nm and 100-700 nm respectively. For standard particle types
116 Sultan et al. (2011) and Waring et al. (2008) both observed erratic CADR performance below ~40 nm
117 (potentially due to instrument sensitivity), and consistent performance above 40 nm. Mølgaard et al. (2014)
118 tested two HEPA APs between 12-660 nm; one performed consistently with increasing size whilst the other
119 experienced a peak in removal efficiency at ~200 nm. Furthermore, Lee et al. (2015) found the lowest
120 filtration efficiencies for three APs to fall within the UFP size range. The findings of these studies contradict
121 our current understanding of the filtration efficiency of HEPA filters - a minimum efficiency of around 200-
122 300 nm varying from filter to filter (Kowalski et al., 1999). However, it is worth noting that a filter may not
123 perform as efficiently within an AP as it does in laboratory tests, given processes like filter bypassing (a
124 result of AP design) and short circuiting of filtered air (Shaughnessy and Sextro, 2006). Therefore, it is
125 currently unclear how effectively “real world” particles of different sizes are removed by commonly available
126 household APs, and why some measurements of performance do not align with the current understanding
127 of the removal processes of HEPA APs. This paper aims to resolve these uncertainties.

128 Using the Guangzhou Institute of Geochemistry’s state of the art chamber laboratory, the largest indoor
129 chamber in China (Wang et al., 2014), this investigation aimed to determine: (a) which particle sizes from
130 ambient air are most and least efficiently removed by APs and explain how this might be important in a
131 real-world context; and (b) whether ambient air particles are removed more or less efficiently than AHAMs
132 standard particle types (tobacco smoke, dust and pollen) and whether AHAM should therefore consider
133 adjusting their CADR measurements accordingly.

134 2. Methodology

135 Selection of air purifiers

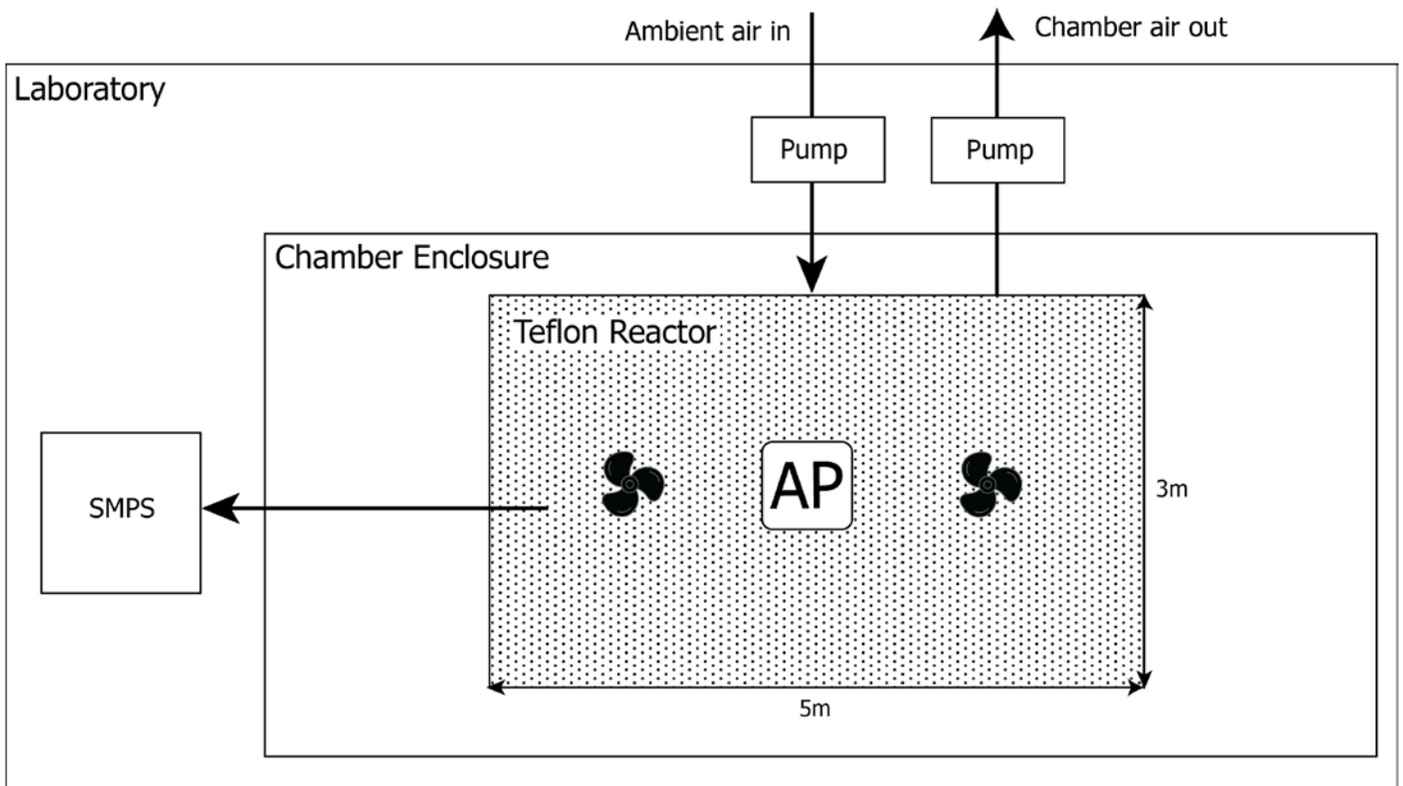
136 For this investigation, three HEPA APs were selected to represent the small (CADR 100-200), medium
 137 (CADR 200-300) and large (CADR >300) AP sizes (Table 1). All three APs were purchased on the Chinese
 138 market and were certified by AHAM. They were selected as popular models that represent different filter,
 139 AP design types and sizes. AHAM certification provides a means of allowing performance comparisons to
 140 be made between models for the removal of different particle types and their associated size fractions. The
 141 reason that tobacco smoke (90-1000 nm), dust (500-3000 nm) and pollen (5000-11000 nm) CADRs for the
 142 same AP are different is due to differential removal based on their respective particle sizes.

	CADR (ft ³ /min)		
	100-200	200-300	300<
Model	Blueair 203	Midea KJ400G-E33	Philips AC6608
Referred to as	AP(Small)	AP(Medium)	AP(Large)
AP Type	Compact	Tower	Cube
Filter type	Single Filter	Circular Filter	Dual Filter
Tobacco Smoke CADR (ft ³ /min)	155	226	369
Dust CADR (ft ³ /min)	155	229	389
Pollen CADR (ft ³ /min)	155	236	451
Purchase Cost RMB (USD)	2000 (200)	1700 (250)	4000 (700)
Filter Replacements RMB (USD)	200 (50)	600 (90)	600 (90)
Recommended Room Size (sq ft)	240	350	572
*RMB costs represent the cost on the Chinese market, USD represents price on the US market			
* 1 ft ³ /min = 28.3 litres/min			

143 **Table 1.** Summary of selected HEPA APs (AHAM, 2018).

144 Experimental setup

145 The atmospheric chamber laboratory at the Guangzhou Institute of Geochemistry, Chinese Academy of
146 Science was used for this investigation. The properties of this chamber are described in detail by Wang et
147 al. (2014). It consists of a 30m³ fluorinated ethylene propylene Teflon film reactor (hereon referred to as
148 a Teflon reactor) housed within a temperature-controlled enclosure (hereon referred to as the chamber
149 enclosure). The Teflon reactor can be filled and vented using pumps with a flow rate in excess of 1m³/min,
150 meaning that it may be filled entirely within 30 minutes. A blower motor from a high-volume air sampler
151 using a tube with a 6 cm bore was used to minimize particle losses. Figure 1 illustrates the layout of the
152 chamber laboratory.



153 **Figure 1.** Layout of the chamber laboratory, chamber enclosure and Teflon reactor at the Guangzhou
154 Institute of Geochemistry.

155 During the experiments, the Teflon reactor was filled entirely with ambient air from outside the laboratory.
156 It is important to understand the composition of this ambient air. Liu et al. (2014) have previously reported
157 that on the Guangzhou Institute of Geochemistry site, carbonaceous aerosols (which contribute a large
158 fraction of PM_{2.5}) could be attributed to fossil fuel (46%), non-fossil fuel (51%) and biomass burning (3%).

159 In a larger study of the city, in the dry season, when this investigation was conducted, ambient PM was
160 largely composed of emissions from vehicular (21%), industrial (20%), residential (4%), power generation
161 (2%) and other unknown sources (53%) (Cui et al., 2015). In 2017 Guangzhou had an annual average
162 PM_{2.5} concentration of 35 µg/m³ (Ministry of Ecology and Environment the People's Republic of China, 2018),
163 with PM_{2.5} in the dry season of 2013 composed of secondary organic aerosol (23%), primary organic aerosol
164 (14%), sulphate (14%), nitrate (11%), ammonium (7%), elemental carbon (4%) and an unidentified
165 fraction (28%) (Cui et al., 2015). The atmospheric chamber laboratory is located ~250m from an 8-lane
166 highway, and therefore UFPs will likely be of vehicular origin. Air was sampled at a height of 1m, directly
167 outside the atmospheric chamber laboratory.

168 Before each test the Teflon reactor was evacuated, and ambient air was drawn in from directly outside the
169 laboratory. Two Teflon coated fans located within the reactor gently mixed the air during filling and
170 throughout the duration of each experiment. The Teflon reactor was not a fixed volume or shape like a
171 stainless-steel chamber, and so there was some variation in the reactor volume between experiments. This
172 is addressed in more detail later. Given that the air was purged entirely from the reactor before it was
173 refilled, no additional cleaning was required between test runs. A TSI SMPS (Scanning Mobility Particle
174 Sizer) consisting of a Differential Mobility Analyzer (DMA - classifier model 3080) and Condensation Particle
175 Counter (CPC - model 3775) was used to measure the total particle number concentration (PNC) and
176 particle size distribution (PSD) between 18-514nm in 94 size bins, with a full scan completed once every
177 minute. Once the Teflon reactor was filled with ambient air, the AP was started. Experiments were repeated
178 a minimum of four times for each AP, at each of three fan speeds (low, medium and high). A new HEPA
179 air filter was used for each AP for the duration of the repeats, therefore, filter loading had a minimal effect
180 on performance given that a single filter was used for no more than 24 hours in total (filters are rated for
181 roughly ~1-2 years of regular use).

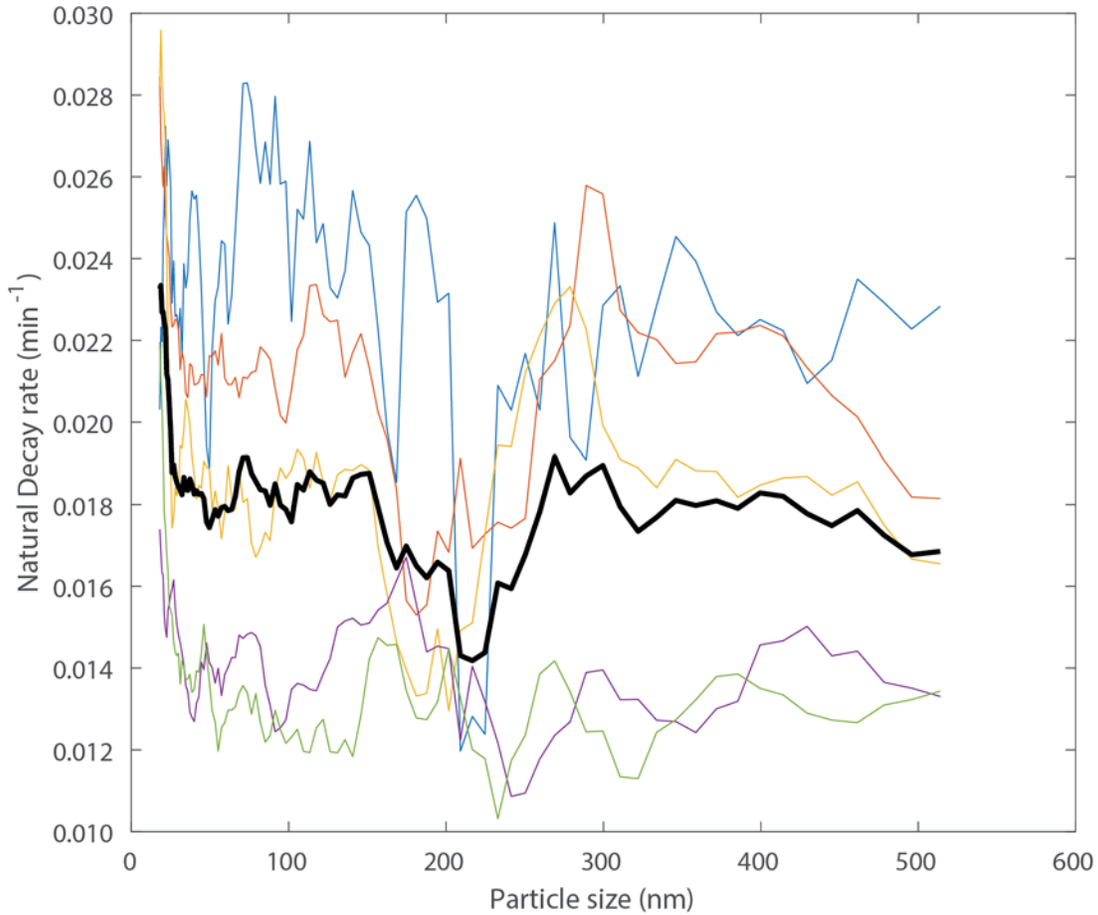
182 Clean air delivery rate

183 CADR was calculated using the equation $CADR = V(Dm - Dn)$ where V is the volume of Teflon reactor in ft^3 ,
184 Dm is the particle number decay rate when the AP is active and Dn is the natural particle number decay
185 rate in the reactor when the AP is inactive (AHAM, 2002). Dm and Dn are first-order loss rates (min^{-1}), the
186 decay constants of an exponential decay in particulate number concentrations, as measured by the SMPS.
187 Initial total particle number concentrations within the reactor varied between $8 \times 10^4 - 2 \times 10^5 \text{ \#/cm}^3$,
188 depending on the ambient conditions at the time.

189 If a decay series met either of the following criteria, then it was excluded; (a) if the decay series contained
190 less than 9 points, meaning that the minimum test duration was less than 9 minutes (AHAM, 2002), and
191 (b) if greater than 30% of values in the decay series exceeded their previous values.

192 Criteria (a) was responsible for identifying failed decay series on the largest AP at the highest fan speed,
193 with the AP cleaning the reactor too quickly (<10 minutes), making calculations of decay rate and therefore
194 CADR unreliable. Criteria (b) was mainly used to identify failures on the smallest AP at the lowest fan speed,
195 with some decay series being difficult to identify amongst variability caused by mixing. Failure to meet
196 these criteria is illustrated in Table 2.

197 Calculating natural decay rate was essential to determine how much particle removal was due to the AP
198 and how much was due to other processes including agglomeration, wall loss and deposition. Natural Decay
199 rates in the Teflon reactor were calculated with an AP present, but not actively running, using the SMPS
200 for each of the 94 size bins from 18-514nm. In this way, both the measured decay rate and natural decay
201 rate were specific to the particle size. The decay rates were measured five times within a single day, see
202 Figure 2.



204 **Figure 2.** Natural decay rate (min^{-1}) in Teflon reactor without use of APs, $n=5$. Solid black line
 205 represents the average natural decay rate (min^{-1}).

206 Given that the Teflon reactor was inflated using ambient air, the reactor was not a fixed volume at the start
 207 of every experiment. The minimum and maximum volumes for the reactor at the fixed roof height were
 208 therefore calculated using the trace gas injection method (Mazzeo, 2012). The minimum and maximum
 209 volumes were 24.5 m^3 and 27.2 m^3 respectively, however, the reactor was inflated to an intermediate
 210 volume between the minimum and maximum volume. A volume of 25.9 m^3 (the midpoint between
 211 maximum and minimum volumes) was therefore used in the calculations.

212 In comparable chamber studies, high concentrations of tobacco smoke, vehicle exhaust, sodium chloride
213 or pollen were released into the chamber and mixed ([Mølgaard et al., 2014](#); [Peck et al., 2016](#); [Sultan et](#)
214 [al., 2011](#); [Waring et al., 2008](#)). In this experiment, the reactor was filled with much lower particle
215 concentrations in ambient air. The challenges associated with this investigation were likely larger than that
216 of comparable chamber studies; given the nature of using ambient air, which varies temporally in
217 composition, humidity and temperature. Furthermore, given that the reactor needed to be inflated to an
218 approximate size, this limited the ability to use a fixed volume of air. However, measuring ambient air (with
219 complex compositions) under real world conditions is likely more indicative of real-world performance than
220 laboratory tests utilizing standardized particle types.

221 3. Results and discussion

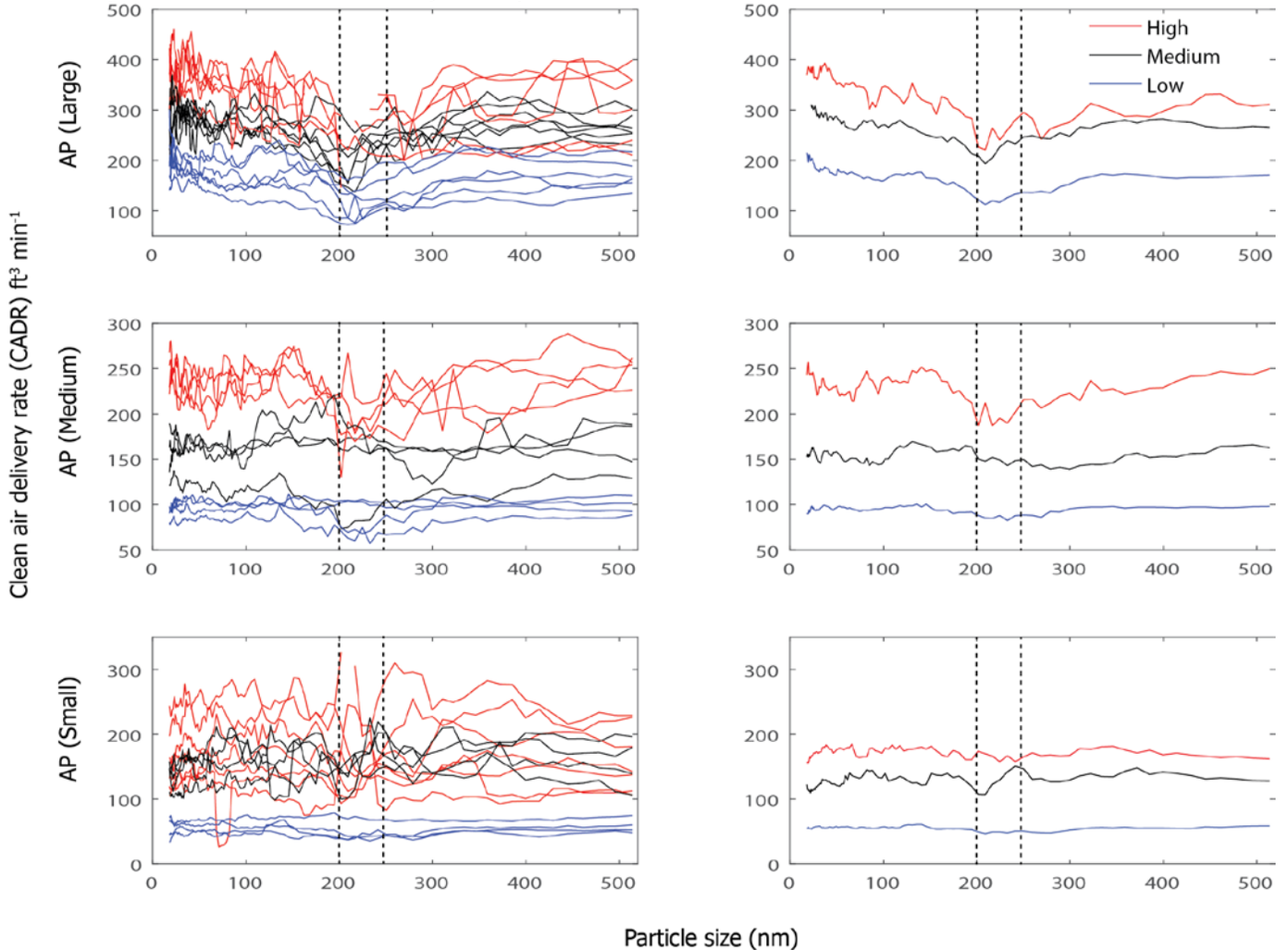
222 Air purifier performance statistics

Air Purifier	Fan Speed	CADR (ft ³ min ⁻¹) Statistics								Electrical Power Draw (W)	Energy efficiency (CADR/W)	Noise (dB)	Noise rating (CADR/dB)
		Min	Max	Mean (s.d.)	Coefficient of Variation (%)	Median	N	R	S				
AP(Large)	High	130	440	316 (58)	18	330	6	564	64	60.5	5.2	51.4	6.1
	Medium	123	346	251 (35)	14	251	6	564	7	35.0	7.2	44.2	5.7
	Low	58	279	151 (36)	24	152	6	564	1	20.0	7.6	34.5	4.4
AP(Medium)	High	130	288	230 (23)	10	232	4	376	1	36.3	6.3	49.9	4.6
	Medium	75	221	154 (25)	17	161	4	376	1	16.1	9.6	40.2	3.8
	Low	57	112	95 (10)	11	98	4	376	2	6.9	13.8	N/A	N/A
AP(Small)	High	25.9	327	172 (52)	31	160	7	658	4	61.5	2.8	47.0	3.7
	Medium	98	308	155 (29)	19	156	5	470	1	45.2	3.4	37.3	4.2
	Low	34	79	55 (10)	18	56	4	376	23	16.6	3.3	34.5	2.3

Table 2. Air Purifier Statistics, R is the number of repeats, N is the number of decay series measured (R multiplied by the number of size bins = 94), S is the number of runs that failed to meet the selection criteria. The CADR values presented were averaged over the 94 size bins from 18-514 nm.

223 Our results show that larger APs and higher fan speeds generate larger average CADRs than smaller APs and lower fan speeds, as expected. This
 224 aligned with electrical power draw, which also increased with increasing AP size and fan speed. The coefficients of variation measured over 18-
 225 514nm were comparable to those of [Waring et al. \(2008\)](#), who measured 16% and 14% for the two HEPA APs tested. Table 2 also shows that the
 226 APs were most noise and energy efficient when running on their lowest fan speeds. On lower fan speeds AP (Medium) was substantially more
 227 energy efficient than AP (Large) or AP (Small).

228 Air purifier removal efficiency with particle size



229 **Figure 3.** CADR as a function of particle size for three APs and for three fan speeds. Each line on the
 230 left plots represents a single AP decay series whilst the lines on the right plots show the average for each
 231 fan speed.

232 Figure 3 shows that AP (Large), (Medium) and (Small) are all effective at removing UFPs from ambient air.
 233 Each AP showed a distinctive removal profile which was consistent across the fan speeds, most likely
 234 attributed to the design of the HEPA filter and sealing. These profiles, although distinct, share some
 235 common themes. Generally, the APs performed least well between $\sim 200\text{-}250\text{nm}$, which is consistent with

236 the understanding of the removal processes of HEPA filters (Kowalski et al., 1999; Stafford and Ettinger,
237 1972). This can be seen more clearly in Figure 4.

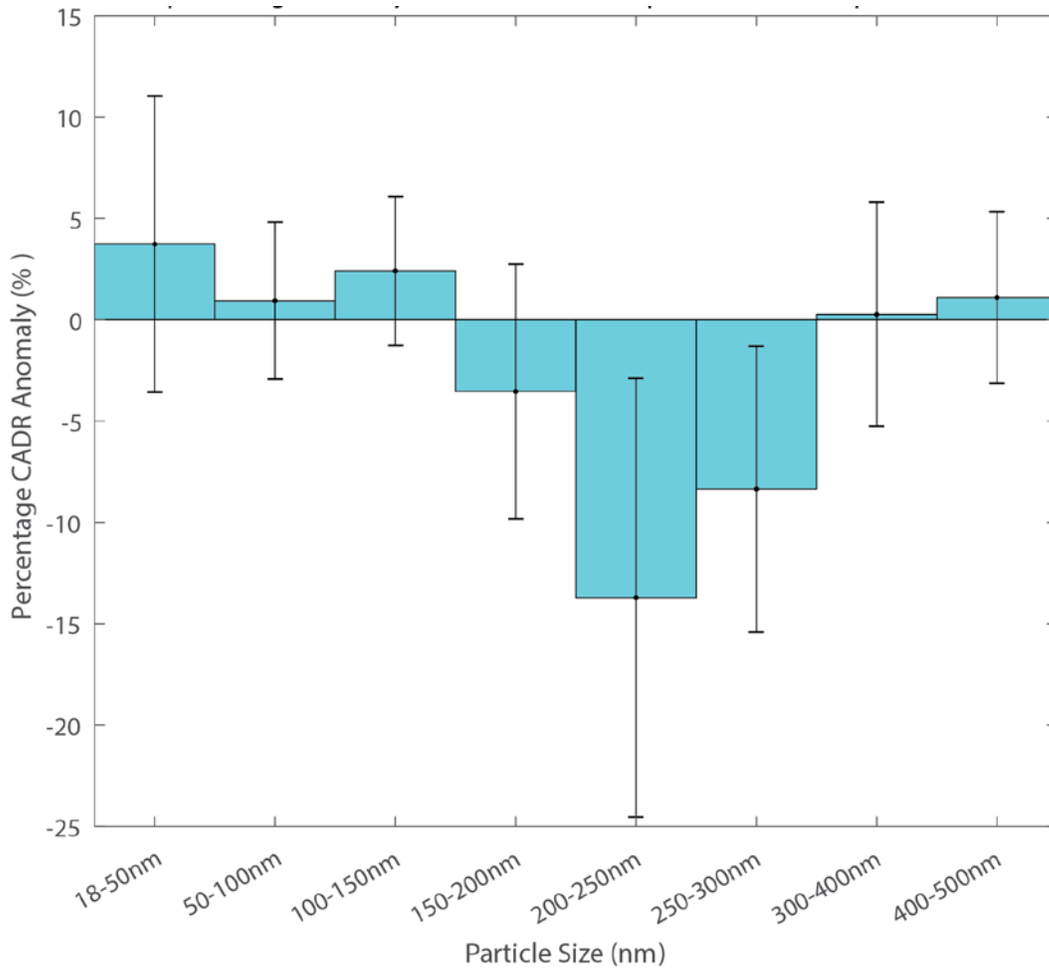


Figure 4. Percentage change in CADR relative to mean CADR for particle sizes between 18-500nm averaged over all tested air purifiers and fan speeds. Each value is the average value for the size bin. Percentage anomaly was calculated for the average of each of the APs for each given fan speed (n=9) and was divided into size bins. The standard deviation was calculated across the 9 arrays and is shown with the error bars indicating one standard deviation.

238 Figure 4 is consistent with the typical performance of a HEPA filter (minimum efficiency 200-300nm)
239 (Kowalski et al., 1999; Stafford and Ettinger, 1972), and aligns with the understanding that diffusion
240 primarily removes the smallest particles and that interception, inertial impaction and sieving primarily
241 remove the largest particles, with particles in the intermediate size range (~100-400nm) least efficiently
242 removed. However, this is contrary to the findings of Peck et al. (2016) who observed peak performance

243 between 100-700nm and [Sultan et al. \(2011\)](#), [Waring et al. \(2008\)](#) and [Lee et al. \(2015\)](#) who observed
244 lowest performance for particles <100nm and consistent performance above this. In [Sultan et al. \(2011\)](#)
245 and [Waring et al. \(2008\)](#), these unexpected performances were attributed to non-uniform mixing in the
246 chamber, with air flows short circuiting the APs and isolated flows forming due to particle size and flow
247 dynamics. However, our results, based on the use of a Teflon reactor, may be more reliable than those
248 generated in stainless-steel chambers, as our reactor was specifically designed to mix uniformly and reduce
249 particle deposition. In addition, given that the Teflon reactor is more rounded than a stainless-steel chamber,
250 this will promote mixing, reducing the likelihood of isolated flow-pathways forming. [Waring et al. \(2008\)](#)
251 also attributed lower performances for UFPs due to particles within the APs bypassing the filter medium.
252 Differences between our measurements and those of [Waring et al. \(2008\)](#) could be due to AP housing and
253 filters being designed to be sealed more tightly during the past 10 years, in order to force particles through
254 the filter medium. Alternatively, the subset of APs selected in this study could be especially well sealed;
255 this may be linked to the bias towards selecting APs that were popular on the Chinese market and were
256 therefore likely effective.

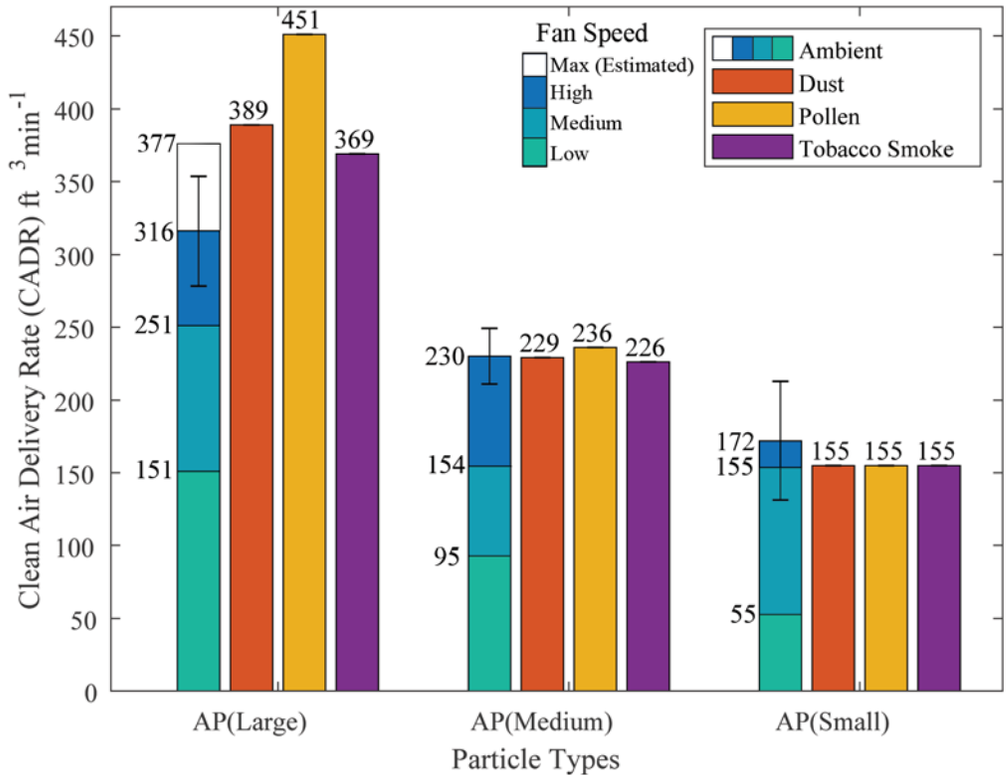
257 Given that the lowest removal efficiencies were observed within this 200-300nm range, it is worth
258 considering the real-world importance of this size fraction. Firstly, these particles are relevant in a health
259 context. Particles of <300nm can penetrate into the alveolar region of the lungs ([Heyder, 2004](#)) and pass
260 into the circulation system, with particles <200nm being found in the brain ([Maher et al., 2016](#)) and it is
261 thought that particles <240nm can cross the placental barrier, potentially impacting upon fetuses ([Wick et
262 al., 2010](#)). Secondly, because the removal properties of building shells are similar to those of a HEPA filter,
263 the particle size that most effectively penetrates cracks in building shells is ~200nm, similar to the 200-
264 250nm for our HEPA APs ([Hänninen et al., 2013](#); [Liu and Nazaroff, 2001](#)). Thirdly, the deposition velocity
265 (m/s), the rate at which particles are deposited onto surfaces, is also lowest at ~200 nm which is consistent
266 with the reactor deposition rates in this investigation ([Lai, 2002](#)). This means that particles within this size
267 range can effectively penetrate building shells and will have longer airborne residence times, due to lower
268 depositional velocities.

269 Particles within the 200-300nm size range are usually found at low concentrations in the atmosphere,
270 typically falling between the Aitken (10-50 nm) and Accumulation (50-1000 nm) particle modes, subject to
271 controls such as composition, humidity and turbulence. Irrespective, [Cai et al. \(2017\)](#) showed that there
272 are still a significant number of particles found within this size range in Guangzhou, in fact, a second
273 accumulation mode was observed with peak number concentrations within the 200-300nm range. Another
274 investigation across 60 Hong Kong residences concluded that particles <400nm contributed the most to
275 total particle mass ([Chao et al., 2002](#)). This is unusual, given that the smallest particles usually contribute
276 the least to total mass measurements. The large concentrations of these particles in megacities could be
277 attributed to secondary aerosols, vehicular and industrial emissions, which generate smaller sized particles
278 ([Zhang et al., 2018](#)).

279 In summary, within Asian mega-cities, particles within the 200-300nm range are abundant in ambient air,
280 can penetrate building shells effectively, can remain airborne for long periods, and are able to penetrate
281 the deepest and most sensitive regions of the body. This means that the population are more likely to be
282 exposed to particles of this size fraction than particles of other fractions in the indoor environment, which
283 may have important health consequences. It is therefore important to note that HEPA APs currently are
284 least efficient at removing this size fraction. It would be beneficial to design another filter media which
285 could remove these 200-300nm particles without dramatically changing the pressure gradient across the
286 filter medium.

287

288 Air Purifier performance for differing particulate matter types



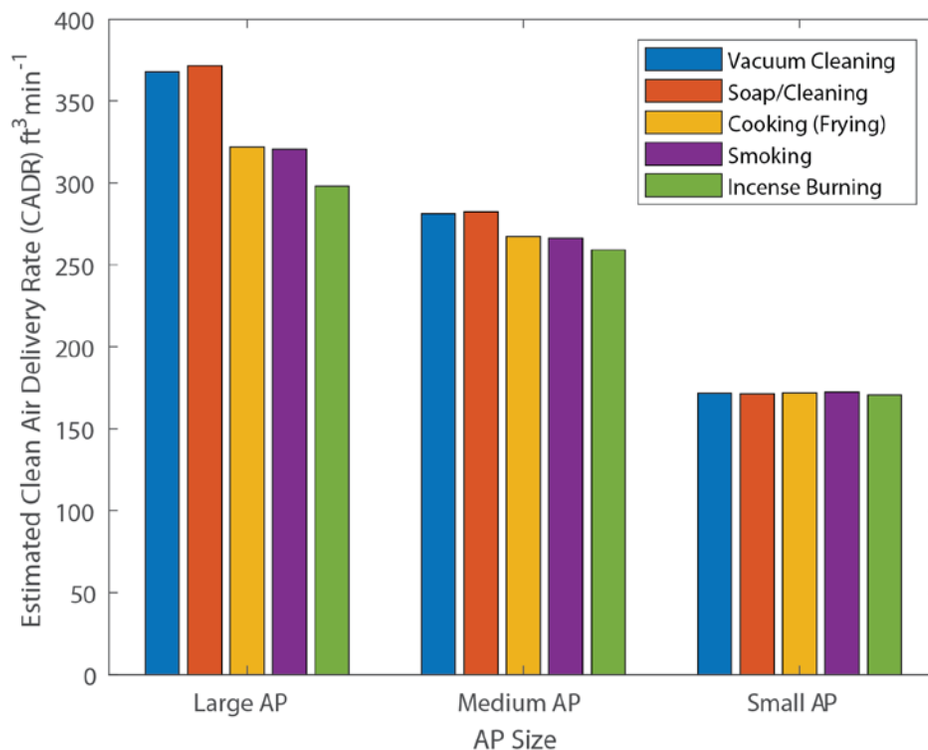
289 **Figure 5.** The CADR (ft³min⁻¹) for different particle types for three APs. Ambient measurements collected
 290 in this study were compared against AHAM dust, pollen and tobacco smoke CADRs for the same APs.
 291 Error bars represent 95% confidence intervals around the means. For AP (Large), performance on
 292 maximum fan speed is estimated based on energy consumption \approx fan rpm \approx CADR Performance. Given
 293 that APs are only tested by AHAM at max speed, this should be used for comparison with AHAM
 294 measurements.

295 In this investigation, ambient particles, despite representing a smaller size fraction (18-514nm) than
 296 tobacco smoke (90-1000nm), dust (500-3000nm) and pollen (5000-11000nm), were removed with similar
 297 (or greater) efficiency than AHAM's standard particle types, as seen in Figure 5. Therefore, the AHAM
 298 standards appear indicative of how efficiently ambient air particles are removed by APs, and hence seem
 299 an appropriate proxy for "real world" AP performance.

300 Our results support [Peck et al. \(2016\)](#) who found that particles generated by diesel combustion were
 301 removed at a greater rate than AHAMs "standard" particle types. This similarity could be due to the strong
 302 influence of vehicular emissions (~20%) in ambient air in Guangzhou. Given the size of the diesel
 303 combustion generated particles used by [Peck et al. \(2016\)](#), and the ambient particles used in this

304 experiment, we would expect them to be removed less efficiently than AHAM standard particle types. [Peck](#)
 305 [et al. \(2016\)](#) attributed this higher removal efficiency of diesel particles to differences in the measured size
 306 ranges between AHAM and those measured within their investigation. However, we hypothesize that this
 307 is likely due to smaller ambient and diesel particles having higher charge to mass ratios compared to
 308 tobacco, dust or pollen particles, which increases removal through the process of diffusion. As the filter
 309 media becomes more saturated with charged particles, this will more effectively remove particles with
 310 higher charge to mass ratios ([Hanley et al., 1994](#)).

311 Applying the relationship between particle size and AP removal efficiency as identified in Figure 5, we can
 312 estimate how efficiently different particle types commonly generated in indoor environments may be
 313 removed.



314 **Figure 6.** Estimated CADR values for different particle types for three APs running on high fan speed.
 315 The particle size distributions utilized to estimate CADR were adapted from [Vu et al., \(2017\)](#), assuming a
 316 log normal distribution of particle size generation. This estimation of CADR is based on particles being
 317 differentially removed based on particles size; it therefore does not account for other factors affecting
 318 removal, for example, particle shape, composition and electrostatic charges. Particle types are ordered in
 319 increasing mode particle size, with vacuum cleaning particles being the smallest and incense burning
 320 being the largest.

321

322 By applying PSDs for different particle types adapted from [Vu et al. \(2017\)](#), we can estimate CADR values
323 for different particle types for each of our APs, as shown by Figure 6. For the largest AP, a 20% difference
324 in CADR can be seen between the most and least efficiently removed particle types. The particle types with
325 the lowest CADR scores were those with high particle number concentrations in the 200-250 nm range,
326 where particles are least effectively removed. It is especially noteworthy that fry cooking, smoking and
327 incense particles are removed less efficiently, given that these are common practices in Chinese households
328 ([Apte and Salvi, 2016](#)).

329 Conclusions

330 Using the largest indoor smog chamber in China, this investigation aimed to determine (a) which particle
331 sizes from ambient air were most and least efficiently removed by APs and explain how this may be
332 important in a real world context and (b) whether ambient air particles were removed more or less
333 effectively than AHAMs standard particle types.

334 This investigation found that although UFPs were effectively removed by each of the APs, a reduced removal
335 efficiency was observed within the 200-250nm size range. This is important in a health context, with
336 particles within that size range being present in significant concentrations in mega-cities ([Cai et al., 2017](#)),
337 able to effectively penetrate the shells of buildings ([Hänninen et al., 2013](#); [Liu and Nazaroff, 2001](#)), remain
338 suspended ([Lai, 2002](#)), and penetrate into the deepest areas of the body ([Heyder, 2004](#); [Maher et al., 2016](#);
339 [Wick et al., 2010](#)). Furthermore, this investigation found that ambient air particles were removed at a
340 similar rate to AHAMs standard particle types, suggesting that these standards are representative of “real
341 world” performance.

342 Further investigations should try to identify technologies which may improve the removal of 200-250 nm
343 particles by HEPA filters without dramatically affecting the pressure drop. Additionally, it is necessary to
344 understand the degree to which other properties of particles, apart from size, affect their removal rates.

345 This could be used to further identify key particle types that may be important within a health context and
346 which are more difficult to remove through filtering. Furthermore, some aspects of HEPA AP use should be
347 explored, for example, how factors like AP placement, number of APs, rate of air exchange and mixing may
348 influence AP performance within a residential setting.

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