# Supplemental Material for:

# <sup>2</sup> Combining low-cost, surface-based aerosol monitors with

## <sup>3</sup> size-resolved satellite data for air quality applications

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# <sup>12</sup> S1. Dataset Details of the Nairobi Example

13 This section provides details of the MISR and MODIS satellite and OPC suborbital data collected for the

14 Nairobi example, as well as the GEOS-Chem model input used to assess aerosol vertical distribution and

15 composition.

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## 17 S1.1. OPC-N2 deployment in Nairobi

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19 On May 1, 2016, a team at the United Nations Environment Program (UNEP) deployed six low-cost PM monitors (Alphasense OPC-N2 models) in schools around Nairobi (deSouza, 2017). This was the first 20 relatively long-term, continuous, in situ air quality monitoring project at multiple points around the city of 21 multiple pollutants: NO2, NO, SO2, and PM. Ten months of PM data from five of the six Alphasense 22 OPC-N2s deployed in Nairobi from May 2016 till March 2017 are used in this paper. In addition to PM 23 estimates, these OPCs report the raw particle counts in 16 bins based on particle diameter, ranging from 24 25 0.38 µm to 17.5 µm (Table S1), The sampling flow rate logged by each OPC-N2 at each site varied between 3-8 ml/s, and the data integration period was constant at 5 seconds per sample for this 26 deployment. 27

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29 Each Alphasense OPC-N2 costs  $\sim$  US \$450 (much less than the tens of thousands of dollars that reference air quality monitors cost). The monitors were deployed at Kibera Girls Soccer Academy (KGSA), located 30 in the Kibera informal settlement near the railway; St Scholastica, situated ~20 meters away from the 31 32 notoriously congested Thika highway, where we expected to capture pollution from vehicles; All Saints 33 Cathedral School, close to Mbagathi road and several small shops and industries; UN Environment 34 headquarters, located off UN Avenue in the relatively green area of Gigiri, and Alliance Girls School in 35 Kikuyu, considered an urban background site (deSouza et al., 2017). A sixth monitor was deployed at the 36 Viwandani Community Center in an informal settlement in the industrial area. However, due to an extended power outage, it did not log data after July 2016, so we do not use the data from this monitor in 37 the current study. Note that some of the other monitors also experienced power-outages over shorter 38 periods. The choice of these sites (Figure S1) was intended to allow us to measure the whole gamut of 39 pollutant signatures from different sources in Nairobi. The monitors were mounted on walls at 40 approximately 2 meters above the ground, close to adult breathing height, but out of reach of children, 41 and protected from the rain with plastic shields (deSouza et al., 2017). 42

45 Figure S1: Image of the five deployment sites indicated by white stars. For scale, the distance between UNEP and 46 St Scholastica is 5 km as the crow flies (© Google Maps)

47

48 Table S1: Minimum, maximum and center diameters for the 16 bins that the Alphasense OPC-N2 measures particle 49 counts.

Bin number	Minimum Diameter (µm)	Maximum Diameter (µm)	Bin Center Diameter (µm)
1	0.38	0.54	0.46
2	0.54	0.78	0.66
3	0.78	1	0.89
4	1	1.3	1.15
5	1.3	1.6	1.45
6	1.6	2.1	1.85
7	2.1	3	2.55
8	3	4	3.5
9	4	5	4.5
10	5	6.5	5.75
11	6.5	8	7.25
12	8	10	9
13	10	12	11
14	12	14	13
15	14	16	15
16	16	17.5	16.75

• United Nations ♀ St. Scholastica 0 Alliance Girls **◊** Kibera ♀ All Saints

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Particle hygroscopicity is a consideration, as the OPCs operate at ambient relative humidity (RH), 53 whereas PM<sub>25</sub> is assessed as dry mass. The Alphasense OPC-N2 sizing does depend on RH and aerosol 54 hygroscopicity (Crilley et al., 2018; Hagan et al., 2019). However, the ambient average RH for the current 55 study was 34% (Table S2); RH this low is unlikely to affect the OPC sizing. Further, previous studies 56 Gatari et al., (2005, 2009) found that organic carbon and black carbon dominate the urban air pollution 57 particles in Nairobi, which are sourced from various types of combustion. The higher the amount of 58 organic content, the less hygroscopic they tend to be (e.g., Samset et al., 2018). This means that the 59 particles present are likely to have low hygroscopicity. 60

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Figure S2 shows the derived  $PM_{2.5}$  time series plots (in  $\mu g/m^3$ ) for each Nairobi site from May 1 2016 to March 2 2017. PM data are recorded every minute. (The actual sampling time of the OPC-N2 is 5 seconds, after a 20 second warm-up period.) Some PM spikes registered are as high as 1000  $\mu g/m^3$ . In order to smooth the time-series and suppress outliers, that are likely due to instrument noise, we averaged

- 66 the OPC measurements over 30-minute time intervals.
- 67

68 Co-locating the OPC with a reference monitor to obtain high-quality PM data would be required to 69 calibrate the raw OPC measurements and distinguish the signal from noise directly. However, this would

70 be costly and possibly time-consuming (Castell et al., 2017; Rai et al., 2017). Due to limited resources,

71 we were unable to co-locate our low-cost sensors with a reference monitor in Nairobi. As such, we rely

72 primarily upon the more robust raw particle counts per size bin reported by the monitors, rather than the

73 reported PM<sub>2.5</sub>.

74

75 Table S2: 30 minute-average dN/dln(D) (Units: #/ml) for Bins 2-7 from OPC surface measurements, coincident

76 with the 10 favorable MISR retrievals, along with the daily-averaged temperature and relative-humidity. PM values 77 are given in  $(\mu g/m^3)$ .

78

date	orbit	site	temperature	relative humidity	PM <sub>1</sub>	PM <sub>2.5</sub>	PM <sub>10</sub>	bin2	bin3	bin4	bin5	bin6	bin7
2016-08-02	88423	Alliance	28.7	35.1	6	9	26	8.2	5.0	2.1	1.2	1.1	0.6
2016-08-02	88423	UNEP	25.7	40.4	14	20	56	21.3	12.7	4.4	2.1	1.8	1.3
2016-10-14	89486	UNEP	28.9	35.8	11	19	71	18.4	14.9	6.0	3.7	3.0	2.2
2016-12-17	90418	UNEP	31.7	24.8	5	8	31	7.0	5.7	2.7	1.8	1.5	1.3
2016-08-02	88423	Scholastica	23.8	45.6	24	36	80	39.0	27.7	8.8	4.2	3.2	2.4
2016-10-14	89486	Scholastica	28.2	40.6	10	17	47	16.9	14.9	6.3	3.6	3.0	2.3
2016-08-02	88423	All Saints	26.7	39.9	6	9	25	9.2	6.6	2.8	1.3	1.2	0.8
2016-10-14	89486	All Saints	30.6	33.4	9	16	59	13.6	13.0	5.3	3.3	3.0	2.1
2016-08-02	88423	KGSA	27.3	41.4	11	18	71	18.8	13.7	4.8	2.4	2.4	2.0
2016-10-14	89486	KGSA	26.5	42.8	10	18	61	15.3	18.2	8.6	4.3	3.5	2.6



**Figure S2:**  $PM_{2.5}$  (in  $\mu g/m^3$ ) time-series plots for each of the five Nairobi deployment sites, from May 1, 2016 to March 2, 2017. There are gaps in the Alliance and UNEP plots because the monitors routinely lost electricity and

84 thus did not make measurements during these periods.

86 S1.2. Satellite Data

## 87 S1.2.1. MISR Observations Over the Nairobi Sites

MISR Level 1B2 radiance data covering Nairobi from May 1 2016 to March 2 2017 were downloaded from the Langley ASDC<sup>1</sup> for retrieval regions directly above each air quality monitoring site, and analyzed using the MISR Research Algorithm (RA; (Limbacher and Kahn, 2014; 2019) with the standard set of 74 mixtures. Each mixture is comprised of up to three of eight aerosol components (Kahn et al., 2010; Table S3). Separate MISR RA AOD retrievals, reported in the MISR green band (centered at 558 nm wavelength), were obtained for all eight MISR aerosol components (Table S3) for pixels within a radius of 1.6 km from each of the five Nairobi surface monitoring sites.

- 96 Over the course of the study period, we had only 28 successful MISR retrievals, during eight unique days.
- 97 This is the main motivation for combining MISR retrievals with those of MODIS, as described below.
- 98 The relatively low AOD over Nairobi during the study period makes estimating aerosol-type uncertainty
- 99 difficult. We restricted the MISR retrievals used in this study to the 10 that have total MISR AOD  $\ge 0.15$

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<sup>&</sup>lt;sup>1</sup> https://eosweb.larc.nasa.gov/project/misr/mil2asae\_table (Last accessed on August 12, 2019)

100 (over three unique days), i.e., those meeting the good-quality MISR aerosol property criterion (Kahn et al., 2010).

102



All Saints in January 2017









**Figure S3:** Examination of monthly particle-type variability for three cases where MISR obtained two retrievals at a given site in the same month. The fraction of the total near-surface AOD ( $AOD_{N-S}$ ) ascribed to different MISR aerosol components is plotted for each case. Note that Components 2, 8, and 14 are aggregated, as discussed in Section 2.2.1 in the main text.

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109 Unfortunately, we did not have multiple MISR observations with  $AOD \ge 0.15$  corresponding to a specific 110 site in the same month. Therefore, we used the available MISR component AOD values obtained for a

111 given site to represent the 'monthly effective' particle size distribution for aerosols over that site. We

112 excluded months when there were no MISR retrievals for a site. As such, we assume the fractional



113 contributions of the near-surface MISR aerosol components to the total AOD remain nearly unchanged in 114 the study region for a given month, even if the total AOD varies. We tested this assumption to some 115 extent using all 28 MISR retrievals, as we do have several instances of two observations in a month over 116 a specific site if we include the lower-AOD cases. Figure S3 shows the fractional AOD values for 117 different MISR aerosol components where two observations over a specific site were made in the same 118 month. It can be seen that the fractions are very similar, which lends some credence to our particle 119 property assumption.

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**Table S3**: Aerosol components assumed in MISR Research Algorithm for this study. This is the same component set as is used in the MISR Standard Algorithm, versions 22 and 23. For each aerosol component, Rc is the characteristic radius,  $\sigma$  is the distribution width of the lognormal size distribution, Re is the effective radius, and R1 and R2 are the minimum and maximum radii of the respective size distributions. The Spectral extinction coefficient

125 represents the optical scattering + absorption of each aerosol component.

126

MISR comp onent	Component name	Particle size/Shape category	Rc µm	Re µm	σ	R1 μm	R2 μm	Spectral extinction coefficient µm <sup>2</sup>	
1	Spherical_nonabsorbing_ 0.06	Small spherical	0.03	0.06	1.65	0.001	0.4	0.00039594	
2	Spherical_nonabsorbing_ 0.12	Small spherical	0.06	0.12	1.7	0.001	0.75	0.013401464	
3	Spherical_nonabsorbinb_ 0.26	Medium spherical	0.12	0.26	1.75	0.01	1.5	0.18231007	
6	Spherical_nonabsorbing_ 2.8	Large spherical	0.5	2.8	1.9	0.1	50	16.189339	
8	Spherical_absorbing_0.1 2_ssa_green_0.9	Small spherical, moderately absorbing	0.06	0.12	1.7	0.001	0.75	0.014069699	
14	Spherical_absorbing_0.1 2_ssa_green_0.8	Small spherical, strongly absorbing	0.06	0.12	1.7	0.001	0.75	0.014874	
19	Grains_mode1_h1 (dust)	Medium dust	0.5	0.21	1.5	0.1	1	3.168676	
21	Grains_mode2_h1 (dust)	Coarse dust	1	3.32	2	0.1	6	15.51	

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Figure S4 shows the MISR columnar AOD for the 10 cases where the average total AOD  $\geq$  0.15. Small, 129 130 spherical, light-absorbing (components: 8 and 14) and non-absorbing (component 2) particles are characteristic of organic aerosols and sulfate/nitrate, respectively (Kahn and Gaitley, 2015), and are 131 typical of urban pollution particles. The component AOD values for these particle types are relatively 132 high for all coincident observations, consistent with our expectation that local pollution dominates the 133 aerosol loading in the Nairobi region (see Section S1.3 on the GEOS-Chem model, below). Further, we 134 know that the primary aerosol source in Nairobi during the study period is local pollution, so, lacking 135 other constraints, assuming the particle type is constant on a monthly basis seems plausible. The AOD of 136 component 6, a medium-large spherical particle often retrieved for dust, but also representing a generic 137

138 large, spherical non-absorbing particle (Kahn and Gaitley, 2015), is moderately high for all coincident 139 observations, especially for the five cases with the lowest overall AOD. This could be a transported 140 aerosol, representing a background component, less likely to be concentrated near the surface. The AOD 141 for component 21 is 0, implying this component was not present at a detectable level in any of the MISR 142 retrievals considered. In light of limited MISR aerosol-type retrievals, the assumption of relatively 143 constant aerosol-type fractions remains a limitation of the current study. Future experiments, performed 144 farther away from the equator, will have more frequent MISR observations, and likely higher numbers of 145 successful retrievals.

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148 **Figure S4:** AOD for each MISR aerosol component retrieved from the Research Algorithm for the 10 coincident 149 observations for which the total  $AOD_{558} \ge 0.15$ .

150 It takes about seven minutes for all 9 MISR cameras to observe a given location on the surface. We 151 averaged the OPC PM25 readings over 30 minutes, centered on the MISR overpass time. We matched 152 these OPC values with the corresponding MISR RA AOD values for each component group, using a 153 universe of 74 mixtures, within a radial distance of 1.6 km from each ground site. Each of the five sites 154 has up to  $\sim$ 7 MISR RA coincidences within this radius. The MISR retrievals yielded a spatial distribution 155 156 of spherical, light-absorbing and non-absorbing particles, with an effective radius of about 0.12  $\mu$ m everywhere. Note that the size distribution for the 0.12  $\mu$ m particles extends to 0.75  $\mu$ m (Table S3); we 157 158 rely on the tail of this distribution for comparisons with the OPCs, due to limited OPC sensitivity to smaller sizes. 159

## 160 S1.2.2. MAIAC/MODIS over Nairobi Sites

161 During the study period, the Terra satellite passed over Nairobi between  $\sim 10:30$  am - 11:30 am local time. 162 Aqua overpasses occurred approximately three hours after each Terra overpass. MAIAC AOD at 550 nm 163 was extracted for the study period from the Land Processes Distributed Active Archive Center (LP 164 DAAC).<sup>2</sup> Although the MAIAC AOD at 550 nm is slightly more error-prone than that at 470 nm, we 165 chose this AOD so it would be at essentially the same wavelength as the MISR-reported AOD. We 166 selected only MAIAC AOD values where the cloud mask was 'clear' and the adjacency mask indicated

<sup>&</sup>lt;sup>2</sup> https://lpdaac.usgs.gov/products/mcd19a2v006/ (Last accessed on June 12, 2019)

that the adjacent pixel cloud masks were also clear. This led to a reduction of  $\sim 70\%$  in the number of 167

- successful AOD retrievals, because Nairobi is at relatively high elevation and is often cloudy. 168
- 169

170 Hu et al. (2014) filled in the gaps of Terra AOD retrievals for different overpasses, when a successful 171 Aqua AOD retrieval was obtained, and vice versa, using a simple linear regression. However, predicted 172 AOD values inevitably contain additional uncertainties. Thus, we only used Terra AOD in this study. An

additional advantage of working exclusively with Terra AOD is that MISR is also aboard Terra, so their 173

observations are temporally coincident. 174

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176 As with MISR, we matched the PM<sub>25</sub> reading of each OPC site with the corresponding MAIAC AOD 177 values from the MODIS grid cells within a radial distance of 1.6 km, at Terra overpass time. We tested the

178 robustness of our results to using radial distances of 1 km and 0.5 km. There were up to nine MAIAC grid

179 cells linked with each site. As the KGSA and All Saints sites are 2.63 km apart, some of nine the grid

- 180 cells associated with the two sites overlap.
- 181

182 The mean AOD of the 1712 successful Terra MAIAC retrievals is 0.095 (min=0.014, max=0.301, 183 sd=0.053). The successful retrievals occur over 66 unique days. A total of 304 successful Terra MAIAC 184 AOD retrievals were acquired, over 20 unique days, during months when favorable MISR retrievals 185 (MISR AOD  $\geq 0.15$ ) were also obtained. Of these, 79 retrievals were made over UNEP, 48 were made 186 over St Scholastica, 78 over KGSA, 87 over All Saints and 12 over Alliance. The mean successful Terra MAIAC AOD retrievals over these days is 0.126 (min=0.038, max=0.268, sd=0.048). We separately 187 188 considered cases where the total MAIAC AOD  $\geq 0.15$  (85 of the 304 retrievals), in order to ensure that we were considering days when the surface aerosol dominated in the column. 189

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#### S1.3. The GEOS-Chem Model 191

The GEOS-Chem model results are for 2012, the latest available, following two months of spin-up for 192 193 chemical initialization, whereas the observations used in this paper are from 2016. Therefore, the model 194 will underestimate any pollution sources that track population growth from 2012 to 2016. Population growth rates in Nairobi between 2012 and 2016 were ~8%; this is well within the GEOS-Chem model 195 196 uncertainties, and given other uncertainties in the Nairobi example as well, we accept the 2012 estimates 197 in the current analysis.

198

199 The GEOS-Chem model results indicate that the dominant aerosol components over Nairobi are organic aerosols (OA), typically produced from biofuel use (Figure S5). The model also shows negligible 200 201 sulfate-nitrate-ammonium that is typical in most rapidly developing countries. The Nairobi low-cost 202 sensor sites coincide with two GEOS-Chem grid cells. The UNEP site is in one cell, and the other sites 203 are all in the other. The GEOS-Chem grid cells are of a different size compared with MISR grid cells, and we used the GEOS-Chem results for the non-UNEP sites only, as this cell has a maximum areal overlap 204 with the corresponding MISR cell. 205

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The fraction of AOD in the first layer of the GEOS-Chem model, as a proportion of the total-column 207 208 AOD, provides the AOD vertical scaling factor (Figure S5). This factor is used to calculate the fractional AOD at the surface of Earth, to connect the near-surface aerosols mass results with the AOD from satellite 209 210 data. The vertical scaling factors from GEOS-Chem, were  $\sim 0.4$  for anthropogenic aerosol and 0.07 for 211 dust. We attempted to validate the GEOS-Chem-derived vertical scaling factor by comparing with 212 statistical data from Cloud-Aerosol Lidar and Infrared Pathfinder (CALIPSO) L2 aerosol extinction 213 coefficient retrievals for the same season over the Nairobi region. Unfortunately, the narrow CALIPSO 214 swath ( $\sim 100$  meters) passes no closer than  $\sim 40$  km from the urban center of Nairobi, making the AOD 215 vertical scaling factor derived from the latter difficult to compare with that from GEOS-Chem. We

216 nevertheless calculated the monthly-averaged vertical scaling from CALIPSO, by dividing the sum of the

217 aerosol extinction coefficient at 532 nm for the near-surface aerosol layer by the sum of all the aerosol 218 extinction coefficients for the entire column. For cells within  $\sim$ 100 km from the ground-based monitoring 219 sites, the scaling factor varied between 0.1 and 0.2 over the months in 2016 for which we had MISR 220 retrievals > 0.15. This is less than that from the GEOS-Chem grid cells, most likely because over rural 221 areas, a larger fraction of aerosol in the column is transported, whereas the dominant aerosol in the urban 222 center is locally sourced and concentrated nearer the surface.

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We also attempted to evaluate this vertical scaling layer using the Cloud-Aerosol Transport System 224 (CATS) instrument (on board the International Space Station) L2 particle backscatter coefficient at 1064 225 nm wavelength over the Nairobi region. The monthly-averaged vertical scaling from CATS reported for 226 cells within  $\sim 50$  km from the ground-based monitoring sites, calculated by dividing the sum of the 227 particle backscatter coefficient at 1064 nm for the near-surface aerosol layer by the sum of all the particle 228 backscatter coefficients for the entire column, varies between 0.01 and 0.2 over the months in 2016 for 229 which we had MISR retrievals > 0.15. Unfortunately, the L2 CATS product does not contain information 230 about the backscattering coefficient at or near 550 nm wavelength. The 1064 nm measurements are much 231 232 less sensitive to the small sized urban particles and more likely weighted toward any transported soil or 233 dust particles, making it difficult to compare the vertical scaling parameter from CATS with that from GEOS-Chem. However, GEOS-Chem has been validated at the regional scale (Marais et al., 2019), so we 234 use the available vertical scaling parameters from the model. 235

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GEOS-Chem vertical aerosol mass concentrations [µg m-3]

Figure S5: The vertical distribution of aerosol optical depth obtained from the GEOS-Chem model for selected
 months, for the model pixel above the UNEP site. Note: OA: Organic aerosols, BC: Black Carbon

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242 The GEOS-Chem monthly scaling factors account for seasonal variation in the aerosol vertical distribution (Figure S5). However, as all of the ground-based sites fall within one GEOS-Chem grid cell, 243 244 it does not account for any variation between sites in the local AOD- $PM_{25}$  relationships. In the ensuing analysis, we allowed for site-specific factors in our regression analysis to compare these relationships 245 over the study area. For future deployments, local, direct measurements of vertical extinction or at least 246 247 backscatter from lidar would reduce our reliance solely on model-simulated scaling coefficients. Also, having higher-spatial-resolution modeling would allow us to account for any variability on scales smaller 248 than the  $\sim 200$  km resolution of the GEOS-Chem simulations. One barrier to running the model at higher 249 spatial resolution is the uncertainty in current emissions inventories, especially over much of the 250 developing world. 251

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# 254 S2. Details of the Nairobi Example Regression Analysis

Because the satellite data are stable and self-consistent, we explore how well the PM<sub>25</sub> reported by the 255 OPCs track the variability in AOD<sub>N-S</sub> as estimated by MISR and by MAIAC. 256 257 We first ran a regression analysis to relate surface  $PM_{25}$  to the total satellite MISR AOD<sub>N-S</sub> using Equation 258 S1 for our 10 coincident readings, using the reported average PM2.5 from the OPC-N2 network acquired 259 over the 30 minutes around the Terra overpass time. 260 261  $PM_{25} = \gamma x$  (total near-surface AOD<sub>N-S</sub>) (with no intercept) (S1) 262 263 We obtained an adjusted R squared of 0.88, and a  $\gamma$  of 138.6 (95% Confidence Interval:107, 170.2). This 264 high correlation indicates that the vertical scaling from GEOS-Chem works well for our sites. Table 1 in 265 Section 4 of the main text shows the total near-surface scaling of the MISR AOD<sub>N-S</sub> and the 266 corresponding PM<sub>25</sub> (units: µg/m<sup>3</sup>) for the ten cases where there are MISR RA retrievals over individual 267 OPC sites in the Nairobi area, and the total-column mid-visible AOD exceeded 0.15. 268 269 270 We also scaled the MAIAC total-column AOD using the GEOS-Chem vertical distribution. We then ran simple linear regressions using the reported 30-minute-averaged PM<sub>2.5</sub> from the OPC-N2 network against 271 the MAIAC AOD<sub>N-S</sub> in Equation S1. We had a total of 1712 coincident measurements MAIAC AOD and 272 PM measurements over the course of this study. 273 274 We obtained an adjusted R squared of 0.66 and a  $\gamma$  of 170.7 (95% CI: 165.0, 176.5). Note, however, that 275 when we restrict the AODs considered in the regression to retrievals where the total MAIAC AOD  $\geq 0.15$ 276 (85 retrievals), i.e., when near-surface pollution likely dominates the total-column aerosol concentration, 277 we obtain an adjusted R squared of 0.80 and a  $\gamma$  of 138.6 (95% CI: 130.7, 146.5). 278 279 When we considered site-specific effects and added a site-specific interaction term in the regression 280 281 model i.e., using Equation S2.using all 1712 data points, we obtained an adjusted R squared of 0.68 and a  $\gamma_1$  of 183.2 (95% CI: 167.0, 199.4) for the Alliance site, 141.4 (95% CI: 130.0, 152.7) for All Saints, 282 194.8 (95% CI: 181.8, 207.8) for Kibera Girls Soccer Academy, 207.7 (95% CI: 195.8, 219.6) for St 283 Scholastica and 140.5 (95% CI: 128.8, 152.1) for UNEP. When we restricted the retrievals to those where 284 the total MAIAC AOD  $\geq 0.15$ , we obtained an adjusted R square of 0.85. 285 286  $PM_{2.5} = \gamma_1 x$  site-specific-factor x AOD (excluding intercept terms) 287 (S2)288 The lower R squared obtained with the MAIAC measurements using all 1712 measurements compared to 289 290 MISR is probably because we used only a subset of MISR retrievals for which AOD  $\ge 0.15$ , to ensure we were including only measurements for which the particle-type retrievals are likely of good quality. These 291 292 are also cases where the near-surface aerosol likely dominates, favoring assumptions in our application. When we include the low-AOD MAIAC retrievals, noise from the land surface can contribute 293 significantly to the satellite retrieval. In addition, small amounts of transported aerosol above the 294 near-surface layer can introduce larger uncertainties to the analysis when the AOD is low. When we 295 restrict retrievals to total AOD  $\geq =0.15$ , we obtain a much closer correlation between the surface PM and MAIAC AOD retrievals as well. 297 298



**Figure S6:** AOD for each near-surface MISR aerosol component obtained from the 304 successful MAIAC 302 retrievals.



<sup>307</sup> **Figure S7:** (Blue)  $PM_{2.5}$  values (in  $\mu g/m^3$ ) from the MAIAC Analysis 5 in Table 2 (Remember only 85 satellite

observations with the total MAIAC AOD  $\ge 0.15$  are considered in this analysis). The corresponding daily-averaged PM<sub>2.5</sub> from the ground-based OPC in units of  $\mu$ g/m<sup>3</sup> are shown in red. The correlation between the two estimates of PM is 0.47.

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