1	Particulate matter from re-suspended mineral dust and emergency cause-
2	specific respiratory hospitalizations in Hong Kong
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4	Vivian C. PUN: Jockey Club School of Public Health and Primary Medicine, Chinese University
5	of Hong Kong, Hong Kong, Prince of Wales Hospital, Shatin, New Territory, Hong Kong;
6	Department of Health Sciences, Northeastern University, Boston Massachusetts, 02115 USA.
7	<u>c.pun@neu.edu</u>
8	Linwei TIAN: School of Public Health, University of Hong Kong, Hong Kong. linweit@hku.edu
9	Kin-fai HO: Jockey Club School of Public Health and Primary Medicine, Chinese University of
10	Hong Kong, Hong Kong, Prince of Wales Hospital, Shatin, New Territory, Hong Kong.
11	kfho@cuhk.edu.hk
12	
13	
14	*Corresponding author
15	
16	Vivian C. Pun
17	Northeastern University
18	Department of Health Sciences
19	Boston, MA 02115
20	Phone: 617-373-7756
21	Email: c.pun@neu.edu
22	
23	

24 ABSTRACT

25 While contribution from non-exhaust particulate matter (PM) emissions towards traffic-related emissions is 26 increasing, few epidemiologic evidence of their health impact is available. We examined the association of short-27 term exposure to PM₁₀ apportioned to re-suspended mineral dust with emergency hospitalizations for three major 28 respiratory causes in Hong Kong between 2000 and 2008. Time-series regression model was constructed to examine 29 association of PM₁₀ from re-suspended mineral dust with emergency hospitalizations for upper respiratory infection 30 (URI), chronic obstructive pulmonary disease (COPD) and asthma at exposure lag 0 to 5 days, adjusting for time 31 trends, seasonality, temperature and relative humidity. An interquartile range (6.8 μ g/m³) increment in re-suspended 32 mineral dust on previous day was associated with 0.66% (95% CI: 0.12, 0.98) increase in total respiratory 33 hospitalizations, and 1.01% (95% CI: 0.14, 1.88) increase in URI hospitalizations. A significant 0.66%-0.80% 34 increases in risk of COPD hospitalizations were found after exposure to re-suspended mineral dust at lag 3 or later. Exposure to mineral dust at lag 4 was linked to 1.71% increase (95% CI: 0.14, 2.22) in asthma hospitalizations. 35 Associations from single-pollutant models remained significant in multi-pollutant models, which additionally 36 37 adjusted for PM₁₀ contributing from vehicle exhaust, regional combustion, residual oil, re-suspended mineral dust, fresh sea salt, aged sea salt, secondary nitrate and secondary sulfate, or gaseous pollutants (i.e., nitrogen dioxide, 38 39 sulfur dioxide, or ozone), respectively. Our findings provide insight into the biological mechanism by which non-40 exhaust pollution may be associated with risk of adverse respiratory outcomes, and also stress the needs for 41 strategies to reduce emission and re-suspension of mineral dust. More research is warranted to assess the health 42 effects of different non-exhaust PM emissions under various roadway conditions and vehicle fleets. 43

Keywords: re-suspended mineral dust, respiratory hospitalization, upper respiratory, COPD, asthma, time-series
 analysis

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46 1. INTRODUCTION

47 Road traffic emissions are a major source of urban air pollution (Charron et al., 2007). They contribute to particulate 48 matter (PM) pollution via two ways: primary exhaust emissions from the tailpipe of motor vehicles, and non-exhaust 49 emissions from brake/tyre wear, road surface abrasion (road wear), or re-suspension of deposited material (e.g., 50 mineral dust), which may be generated from a wide variety of sources (e.g., construction) near the roadside, due to 51 traffic-induced turbulence (Grigoratos and Martini, 2014; HEI, 2010; Pant and Harrison, 2013; Thorpe and Harrison, 52 2008). Non-exhaust emission of PM becomes increasingly important, particularly in developed countries in recent 53 decades. It is because contribution from primary exhaust emissions to total traffic PM pollution have reduced 54 considerably, due to the introduction of stringent standards for exhaust emission from diesel and gasoline vehicles, 55 technological upgrades of engine and exhaust systems, and cleaner fuels (HEI, 2010; Pant and Harrison, 2013; 56 Thorpe and Harrison, 2008). Thus, it is estimated that 80-90% of the total traffic PM emissions will come from nonexhaust emission by the end of the decade (Rexeis and Hausberger, 2009). 57 58 59 As the contribution from non-exhaust PM emissions increases, so should our understanding of the potential toxicity and health impact of non-exhaust-related particles. A few toxicological studies reported possible link between brake 60 61 wear particles and pro-inflammatory responses in lung cells in vitro (Gasser et al., 2009; Mazzarella et al., 2007), 62 and that studded tyre wear particles may have associated with considerable pro-inflammatory potential and profound 63 effects on macrophages in vitro and animal models (Gustafsson et al., 2008; Karlsson et al., 2011; Lindbom et al., 2006). Yet, no epidemiologic studies have directly linked brake/tyre wear PM with adverse health effects on humans. 64 65 In contrast, several epidemiologic studies have investigated the potential association between fine particles ($PM_{2.5}$)

66 from crustal soil or re-suspended mineral dust and human health using source apportionment modeling (Andersen et

67 al., 2007; Bell et al., 2014; Halonen et al., 2009; Lall et al., 2011; Sarnat et al., 2008; Schreuder et al., 2006). Though

68 these studies show suggestive association of re-suspended mineral dust with adverse respiratory outcomes,

69 heterogeneity in the findings remains. This may be explained by the application of PM_{2.5} rather than coarse particles

- 70 (PM_{2.5-10}) that are predominantly released by much of the non-exhaust processes (e.g., mechanical abrasion and
- 71 corrosion). Other possible reasons include variation in the chemical composition depending on factors including
- 72 brake and tire manufacturers, vehicle type and movement, street maintenance, season and meteorological parameters
- 73 (Pant and Harrison, 2013; Stanek et al., 2011), as well as difference in the underlying study populations.

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- 75 In a recent study conducted by our research group, we found positive association between risks of emergency 76 hospitalizations for total respiratory causes and several PM₁₀ apportioned factors in Hong Kong (Pun et al., 2015). 77 Among the observations, the statistically significant association with PM_{10} factor whose chemical profile was most 78 reflective of that of re-suspended mineral dust was intriguing, given that this was the first Asian study to report such 79 association, and the only other study that examined such relationship reported no significant association in 80 Copenhagen (Andersen et al., 2007). The current study built upon this observation by assessing the association of 81 PM_{10} apportioned to re-suspended mineral dust with emergency hospitalization for three major respiratory causes 82 respectively. Parallel epidemiologic evaluation was also performed for aluminum, calcium and iron, the chemical 83 tracers that were largely apportioned to the re-suspended mineral dust factor, as well as evaluation for other PM_{10} 84 sources for comparison. Findings from this study may provide a useful perspective on and further understanding of non-exhaust particles and their associated health impact, especially in Asia where studies of such kind is scarce. 85
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87 **2. METHODS**

88 2.1. Hospitalization Data

89 We have previously compiled daily count data of emergency hospitalization into twenty-six publicly funded, 90 regulated by the Hong Kong Hospital Authority, hospitals across the territory for the period of 2001 to 2008 (Figure 91 S1; Pun et al., 2015, 2014a). In this study, we used the 9th revision of the International Classification of Diseases 92 (ICD-9) code to extract hospitalizations for three major respiratory causes: chronic obstructive pulmonary disease (COPD; ICD-9: 491,492 and 496), asthma (ICD-9: 493). ICD-9 classification for acute upper respiratory infection 93 (URI, ICD-9: 464-466) was only available from 2001 to 2007; thus analysis for URI was restricted to 2001-2007, 94 95 whereas analyses for all other respiratory causes focused on 2001-2008. Since prior work has shown influenza outbreaks to be a confounder of the association between PM and respiratory hospital admissions (Ren et al., 2006), 96 97 influenza hospitalizations (ICD-9: 487) were excluded from the total count of respiratory hospitalizations, and were 98 subsequently adjusted in the regression analysis.

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100 2.2. Exposure Assessment

101 We employed the positive matrix factorization (PMF) 5.0 with the inclusion of nineteen PM_{10} chemical species, 102 characterized from filter samples collected from six monitoring stations between 1 January 2001 and 31 December 103 2008, to identify physically realistic sources. Detailed analytical methods and profiles for source-apportioned PM_{10} 104 have been published elsewhere (Pun et al., 2014b, 2015; Z. B. Yuan et al., 2013), and a summary can be found in the 105 eAppendix (Supplementary). Briefly, eight apportioned factors, reflective of chemical profiles for vehicle exhaust, 106 regional combustion, residual oil, re-suspended mineral dust, fresh sea salt, aged sea salt, secondary nitrate and 107 secondary sulfate, were obtained (Figure S2). Some factors have distinctive chemical profile corresponding to a 108 specific emission source, such as vehicle exhaust emission that explained 80% of the variation in elemental carbon. 109 Two factors are identified as composite of two or more sources that cannot be reasonably separated in the PMF 110 model due to similarity in the source profile. Regional combustion emission was associated with large variation for a 111 mixture of combustible constituents, arising from wood/biomass burning (e.g., potassium ion) and coal combustion (e.g., arsenic, cadmium, lead and zinc) in power plants and industrial facilities in the adjacent Pearl River Delta 112 region (Z. Yuan et al., 2013). In this study, we primarily focused on PM_{10} from re-suspended mineral dust, a 113 114 composite characterized by high loadings of aluminum, calcium and iron. This factor refers to material that is derived from a combination of exposed soil and dust from traffic, unpaved roads and construction activities from 115 116 within Hong Kong and nearby Pearl River Delta region (Yuan et al., 2006). It might also take into account, to some 117 degree, the particles emitted directly from brake/tyre/road wear and abrasion and then re-suspended, as evident by 118 the present of aluminum and iron that appear to be ubiquitous in brake linings, tyres and car paint. However, further 119 separation of brake/tire wear emission, road surface wear emission and re-suspension emissions was not attainable 120 in the PMF model.

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Daily mean temperature and relative humidity were also obtained from the Hong Kong Observatory for the same
study period. Daily mean concentrations of nitrogen dioxide (NO₂) and sulfur dioxide (SO₂), as well as 8-hour mean
(10:00 AM-6:00 PM) concentration of ozone (O₃) were also calculated from hourly air pollutant data.

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126 **2.3.** Statistical analysis

127 Given fairly high temporal correlations of PM_{10} constituents and sources across the six monitoring stations (r >

128 0.70), we used *a priori* method of centering and averaging to remove the station-specific influence on the resulting

PMF-resolved PM_{10} source contributions (Pun et al., 2014a, 2014b; Tian et al., 2013). The resultant time series contained nonmissing territory-wide mean source-apportioned PM_{10} for five consecutive days a week over the entire study period. No notable difference was observed between days with nonmissing sampling data and those with missing data in terms of cause-specific respiratory hospitalizations, apportioned sources and meteorological factors; thus missing data were not imputed. Previous analyses have shown that risk estimates of PM_{10} constituents and sources were insensitive to either regression models in which no data imputation was used or models in which imputation of missing sampling days was applied (Pun et al., 2014a, 2014b).

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137 We used generalized additive model with log link, Poisson-distributed errors and autoregressive terms to estimate 138 the associations of PM₁₀ from re-suspended mineral dust and hospital admissions for total respiratory causes, COPD, 139 URI and asthma, respectively (Hastie and Tibshirani, 1990). Each model adjusted for time-varying confounders, with smoothing splines with 8 degrees of freedom (df) per year for time trends and seasonality, 6 df for current day 140 temperature and previous 3-day moving average, and 3 df for current day relative humidity and previous 3-day 141 142 moving average selected *a priori* to minimize problems associated with multiple testing and core model selection strategies (Bell et al., 2009). Dummy variables for day of week, public holidays and influenza epidemics were also 143 144 controlled for. We estimated hospitalization risks associated with levels of pollution on the same day (lag_0) and up to 145 5 days (lag₅) prior to hospitalization, while controlling for time trend, seasonality, meteorological conditions, 146 calendar effects and influenza epidemic. Parallel epidemiological evaluation for aluminum, calcium and iron, the 147 chemical tracers of re-suspended mineral dust, was also conducted. We further tested the robustness of the association in multi-pollutant models, where we adjusted for all eight PM₁₀ sources simultaneously. Although not 148 149 the primary focus of the study, the associations of cause-specific respiratory hospitalizations with the remaining PM₁₀ sources were also explored in the multi-pollutant models to provide comparison with that of re-suspended 150 151 mineral dust. In addition, we constructed two-pollutant models adjusting for NO₂, SO₂ or O₃ respectively. In an 152 exploratory analysis, the average $PM_{2,5}$ -to- PM_{10} ratios for calcium and iron were calculated using the available daily 153 constituent concentrations in PM_{2.5} and PM₁₀ measured in Tsuen Wan air monitoring station, which is located close 154 to the geographic center of Hong Kong and is likely to be more representative of Hong Kong's air quality in general, 155 between November 2004 and November 2005. Ratio for aluminum was not computed due to unstable PM2.5 156 aluminum measurements. All estimates were reported as the percent increase [(relative risk-1)×100%] in daily

157 cause-specific emergency hospital admissions for an interquartile range (IQR) increment in pollutant concentrations.

158 The analyses were performed in the statistical environment R Software, version 3.1.2 (R Development Core Team,

159 Vienna).

160

161 **3. RESULTS**

162 Between 2001 and 2008, there were 659,963 emergency respiratory hospitalizations (~226 admissions per day) in

163 Hong Kong (Table 1). COPD hospitalizations accounted for 23.9% of total respiratory causes, followed by URI

164 (18.6%) and asthma (7.5%). The daily mean temperature and relative humidity were 23.5°C and 78.1% respectively.

165 The average daily PM_{10} concentration was 55.9 μ g/m³, of which 13% was from re-suspended mineral dust that was

almost comparable to the contribution from vehicle exhaust (15%; see Table S1 for descriptive statistics of all

apportioned PM₁₀ sources). The annual average contribution of mineral dust was relatively constant (p-trend =

168 0.053), compared to the rapid decreasing contribution of vehicle exhaust over the study period (p-trend = 0.015;

169 Figure S3 in the Supplementary). Refer to Table S1 in the Supplementary for Pearson's correlations between source-

- apportioned PM₁₀ and meteorological factors.
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172 PM_{10} from re-suspended mineral dust and its chemical tracers was positively associated with hospitalizations for total respiratory, COPD, URI and asthma at various lag exposures, respectively (Figure 1). An IQR of 6.8 µg/m³ 173 174 increment in PM_{10} contribution from mineral dust on previous day (lag 1) was statistically significantly associated 175 with 0.66% (95% CI: 0.12, 0.98) increase in total respiratory hospitalizations, and 1.01% (95% CI: 0.14, 1.88) increase in URI hospitalizations. Re-suspended mineral dust was also positively linked to increased risk of COPD 176 177 hospitalizations at all lags examined, and the associations were statistically significant at 3 or more days prior to 178 hospitalization (lag 3), corresponding to 0.66%-0.80% increases in risk. Risks of asthma hospitalizations were significantly increased (1.71%; 95% CI: 0.14, 2.22) with exposure to mineral dust at lag 4. 179

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181 Results from multi-pollutant models that further adjusted for potential confounding of other PM₁₀ sources were

182 similar to those from single-pollutant models (Table 2). The increased hospitalization risks of total respiratory

- 183 causes (0.63%; 95% CI: 0.22, 1.04), URI (0.93%; 95% CI: 0.01, 1.85) and COPD (0.77%; 95% CI: 0.17, 1.36) were
- 184 slightly attenuated relative to single-pollutant model estimates; the risks of asthma hospitalizations was strengthened

185 at lag 4 based on the multipollutant model (1.44%; 95% CI: 0.34, 2.55) compared with the single-pollutant model. In 186 contrast to re-suspended mineral dust that exhibited differential temporal lag pattern of associations by cause-187 specific respiratory causes, increment in PM₁₀ apportioned to vehicle exhaust, secondary nitrate, secondary sulfate at 188 earlier days prior to hospitalization (e.g., lag 3-5) were generally and positively associated with respiratory 189 hospitalizations regardless of causes, after adjusting for all other sources (Figure S4). Level of PM₁₀ from aged sea 190 salt was consistently linked to cause-specific respiratory hospitalizations at shorter lag (e.g., lag 0). However, no 191 discernible lag pattern was found for PM₁₀ apportioned to regional combustion, residual oil and fresh sea salt. 192 Additional sensitivity analysis showed that the patterns of associations were similar in two-pollutant models 193 adjusting for O₃, NO₂, or SO₂ (Figure S5 in the Supplementary). Figure 2 shows that tracer species of mineral dust 194 were most abundantly found in the coarser mode of PM_{10} , with average $PM_{2.5}$ -to- PM_{10} ratio ≤ 0.38 .

195

196 4. DISCUSSION

Particles arising from re-suspension of mineral dust are primarily coarse particles generated by mechanical grinding 197 198 of tyre and brakes, and agricultural activities, in contrast to their combustion counterparts that are mostly fine particles (Peng et al., 2008). Currently, no study has examined the adverse health associated with PM2.5-10 from non-199 200 exhaust traffic sources, because quantitative measurements of composition or sources of coarse PM are lacking. 201 Instead, most epidemiologic studies used PM_{2.5} composition to identify non-exhaust traffic PM. While studies from 202 Connecticut and Massachusetts USA, Finland and Chile have reported that PM_{2.5} from soil and/or road dust are 203 predictors of respiratory emergency department visits/hospitalizations (Bell et al., 2014; Cakmak et al., 2009a; 204 Halonen et al., 2009), research from New York City, Atlanta, and Washington State USA and Korea found no 205 evidence of association (Heo et al., 2014; Lall et al., 2011; Sarnat et al., 2008; Schreuder et al., 2006). Such 206 conflicting findings might attest to the limitation that PM_{2.5} does not capture majority of the particles emitted by 207 non-exhaust traffic sources, and thus may not capture their true health impacts. 208

- 209 This study is one of the two existing research that apportioned PM_{10} composition to re-suspended mineral dust,
- along with other sources, and is the first to examine the dust association with cause-specific respiratory
- hospitalizations. The resultant exposure indicator, with an average $PM_{2.5}$ -to- PM_{10} ratio ≤ 0.38 , enabled us to capture
- a wider spectrum of health effects associated with particles in either fine or coarse fraction of PM_{10} , compared to the

213 use of PM2.5 composition alone. We associated an IQR increment in PM10 from re-suspended mineral dust on the 214 previous day with an increased risk of total respiratory hospitalization, after adjusting for covariates; this increase in 215 risk was mainly seen for URI hospitalizations. Statistically significantly positive associations were also observed for 216 COPD and asthma hospitalizations, though with pollutant exposure at 3 or more days prior to hospitalizations. The 217 patterns of associations were similar in models considering tracer species, multi-pollutant models adjusting for all PM₁₀ sources, and two-pollutant models adjusting for gaseous pollutants. Our observed positive association between 218 219 respiratory hospitalization and PM_{10} mineral dust is in contrast to findings reported by Andersen et al. (2007), which showed no significant association of PM10 crustal materials (comprised of road concrete, tire wear, igneous rock and 220 221 limestone) with respiratory and asthma hospital admissions among elderly and children in Copenhagen. The 222 heterogeneous findings may due to the difference in PM₁₀ composition and source profiles between cities, which 223 vary considerably depending on factors including brake and tire manufacturers, vehicle movement, street 224 maintenance, season and meteorological parameters. Other factors, such as longer study period, larger sample size in our study, as well as population susceptibility between cities, may also explain the difference. 225

226

227 Precise biological mechanisms of how PM influence pathogenesis of respiratory illnesses remain unknown; however, 228 several key factors (e.g., size fraction, chemical composition, surface area) must be taken into account in order to 229 better understand the toxicity and potential adverse health effects of PM. In this study, we found PM₁₀ mineral dust 230 or its tracer species, which were generally PM_{2.5-10}, to be associated with URI hospitalizations at shorter day 231 exposure lag, whereas association of hospitalizations for lower respiratory tract illnesses (i.e., COPD and asthma) was at longer lag (e.g., lag 3). Such observed differential temporal lag patterns of associations may be explained by 232 233 the different triggering mechanisms in the lung by the fine and coarse fraction of PM₁₀ soil/road dust (Centers for 234 Disease Control and Prevention et al., 2010). The differential temporal lag pattern of associations of respiratory 235 hospitalization by PM_{10} from vehicle exhaust and those from age sea salt may be also explained by particle size, 236 which has been shown to be important factor affecting particles deposition in the respiratory tract (Samet and 237 Dominici, 2000). Coarse particles, carrying more biological agents (e.g., endotoxins) than finer particles, deposit 238 primarily in the upper airway by impaction, thereby causing irritation and epithelial disruptions, inducing and/or 239 exacerbating upper respiratory illnesses with shorter delay (Ferguson et al., 2013; Schulz et al., 2000). On the other 240 hand, finer particles (e.g., PM_{2.5}) tend to travel deeper in airways and alveoli of the lung by impaction, sedimentation

or diffusion, causing inflammation in the lower respiratory tract and lung tissues in a more delayed observable
fashion (Donaldson and Stone, 2003; Kreyling et al., 2006). Animal models have shown coarse particles, on a mass
basis, to have higher hydroxyl radical generating capacity, greater cytokine production of macrophages and bacterial
endotoxin content, and produce significantly more pulmonary inflammatory responses compared to fine particles
(Shi et al., 2003; Tong et al., 2010); whereas smaller particles may be more potent if taking into consideration of
particle number or surface area. Our findings of PM₁₀ mineral dust provide support for the importance of PM sizemediated impacts on upper and lower respiratory tract illnesses.

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249 In addition, the heterogeneous associations across cause-specific respiratory illnesses or PM₁₀ sources reported in 250 this study may reflect effects of different generation processes, and of the combination of chemical constituents 251 emitted from these sources (Kim et al., 2012). Toxicological studies have suggested that carbonaceous constituents, tracer for vehicle exhaust, may elevate levels of biomarkers for systemic inflammation and platelet activation 252 (Delfino et al., 2009), and exposure to Pb, present in coal combustion as well as construction material, enhanced the 253 254 production of oxidative stress, leading to inflammatory reactions (Saxena and Flora, 2004). Aluminum and iron, tracers for re-suspended mineral dust, may contribute to the formation of free radicals, induce oxidative processes 255 256 and up-regulate pro-inflammatory mediators in vitro, thereby resulting in pulmonary inflammation and respiratory 257 diseases (Garçon et al., 2000; Risom et al., 2005). There is also suggestive evidence that patterns of temporal lag structure of association varied by PM chemical constituents, as Kim et al. (2012) reported that delayed patterns of 258 increased relative risks for asthma hospitalizations began at later lags for sulfate and nitrate than for elemental and 259 260 organic carbon. Further work is needed to examine the biological mechanism in which PM constituents may affect 261 human health.

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As the relative non-exhaust contribution towards any traffic related emissions is becoming greatly significant and posing challenges for authority, findings from the current study stress the needs for strategies and measures for abatement of particle emission and re-suspension. While road/street sweeper alone may not significantly reduce PM₁₀ concentration (Amato et al., 2010; Keuken et al., 2010), other measures such as cleaning road and pavement with water, or applying dust suppressants/dust binders (e.g., aqueous solutions of magnesium/calcium chloride and calcium magnesium acetate) to maintain wet road surface have shown to be effective in certain conditions (Aldrin et al., 2008; Norman and Johansson, 2006). Amato et al. (Amato et al., 2014) concluded in a recent review that a
 combination of strategies aiming at minimizing the emission sources (e.g., improve wear properties of materials and
 traffic, better maintenance of road) and minimizing re-suspension (e.g., road cleaning, reduce traffic) would likely

272 provide optimal abatement of PM non-exhaust emissions from road traffic.

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274 Findings from our study should be interpreted with caution, as we cannot eliminate exposure measurement error. 275 Particles from re-suspended mineral dust tend to be more spatially heterogeneous, which might be subject to more 276 error and bias the estimates towards the null (Kioumourtzoglou et al., 2014b). Quantitative measurement of 277 composition or sources of coarse PM, or measurements of both PM2.5 and PM10 composition for the calculation of 278 PM_{2.5-10} composition for source apportionment are needed to determine clear differences in source contribution by 279 size fraction and validate our findings. Finally, we did not account for the uncertainty of the estimated source contributions directly in the health model. Any underestimation of the resulting inferences, however, is expected to 280 be small given the consistency in source identification across monitors and in effects obtained, and the agreement of 281 our findings with previous studies (Kioumourtzoglou et al., 2014a). 282

283

284 5. CONCLUSION

In summary, we found evidence of significantly positive association of total respiratory hospitalizations with exposure PM_{10} from re-suspended mineral dust, adjusting for covariates (e.g., vehicle exhaust). Exposure at early lag day was linked to URI hospitalization, whereas exposure at later lags was associated with COPD and asthma hospitalizations. These findings help prioritize research on the biologic mechanisms of PM effect on specific pulmonary conditions, and stress the needs for strategies to reduce emission and re-suspension of mineral dust.

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