



Strong evidence for the continued contribution of lead deposited during the 20th century to the atmospheric environment in London of today

Eléonore Resongles^{a,b,1} , Volker Dietze^c, David C. Green^d, Roy M. Harrison^{e,f,g} , Raquel Ochoa-Gonzalez^{a,h} , Anja H. Tremper^d, and Dominik J. Weiss^{a,i,1}

^aDepartment of Earth Science and Engineering, Imperial College London, SW7AZ London, United Kingdom; ^bHydroSciences Montpellier, University of Montpellier, CNRS (Centre national de la recherche scientifique), IRD (Institut de Recherche pour le Développement), 34090 Montpellier, France; ^cAir Quality Department, Research Center Human Biometeorology, German Meteorological Service, 79104 Freiburg, Germany; ^dEnvironmental Research Group, School of Public Health, Imperial College London, W12 0BZ London, United Kingdom; ^eSchool of Geography, Earth and Environmental Sciences, University of Birmingham, Edgbaston B15 2TT, United Kingdom; ^fDepartment of Environmental Sciences, King Abdulaziz University, 21589 Jeddah, Saudi Arabia; ^gCenter of Excellence in Environmental Studies, King Abdulaziz University, 21589 Jeddah, Saudi Arabia; ^hDepartment of European Projects, Foundation for the Promotion of Applied Scientific Research and Technology of Asturias, 33011 Oviedo, Spain; and ⁱDepartment of Civil and Environmental Engineering, Princeton University, Princeton, NJ 08544

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Although leaded gasoline was banned at the end of the last century, lead (Pb) remains significantly enriched in airborne particles in large cities. The remobilization of historical Pb deposited in soils from atmospheric removal has been suggested as an important source providing evidence for the hypothetical long-term persistency of lead, and possibly other pollutants, in the urban environment. Here, we present data on Pb isotopic composition in airborne particles collected in London (2014 to 2018), which provide strong support that lead deposited via gasoline combustion still contributes significantly to the lead burden in present-day London. Lead concentration and isotopic signature of airborne particles collected at a heavily trafficked site did not vary significantly over the last decade, suggesting that sources remained unchanged. Lead isotopic composition of airborne particles matches that of road dust and topsoils and can only be explained with a significant contribution (estimate of 32 ± 10 to $43 \pm 9\%$ based on a binary mixing model) of Pb from leaded gasoline. The lead isotopes furthermore suggest significant contributions from nonexhaust traffic emissions, even though isotopic signatures of anthropogenic sources are increasingly overlapping. Lead isotopic composition of airborne particles collected at building height shows a similar signature to that collected at street level, suggesting effective mixing of lead within the urban street canyon. Our results have important implications on the persistence of Pb in urban environments and suggest that atmospheric Pb reached a baseline in London that is difficult to decrease further with present policy measures.

lead isotopes | urban air pollution | aerosols | source tracing | contaminant persistence

Risk assessments of environmental lead (Pb) exposure have drawn the conclusion that it is not possible to identify a blood Pb level below which no adverse impact is detectable (1, 2). Even very low blood Pb levels in children are associated with a loss of full-scale intelligence quotient (IQ) score, and the curve steepens at lower blood Pb levels, with a greater loss of IQ points per unit of blood Pb in the lowest exposure groups (1, 2). Although currently available research has included subjects with very low blood Pb, it is not possible for such studies to go to zero Pb, but it is prudent to treat Pb as a nonthreshold toxin (3). Bellinger estimated full-scale IQ point losses in the early 2000s in the US child population associated with six medical conditions, four neurodevelopmental disorders, two socioeconomic, nutritional, and psychosocial factors, and three environmental chemical exposures (methylmercury, organophosphate pesticides, and Pb) (1). Pb exposure ranked second among all fifteen risk

factors, behind only preterm birth and first among the environmental chemical exposures. Much of the summed IQ loss occurred among the lowest-exposed groups due to their larger numbers, and even if population blood Pb levels have declined since the Bellinger study and more children have hence moved into the lower exposure groups (4), the total IQ loss across the population may remain substantial.

A major policy achievement in our efforts to reduce Pb in the environment has been the global phaseout of Pb from gasoline at the end of the last century, which resulted in a drastic decrease of atmospheric Pb concentration, especially in urban and remote areas of Europe and North America. While exposure to Pb from paints, Pb pipes, and Pb-containing toys are now recognized as the main cause of elevated blood Pb levels in children in non-industrial environments, Pb remains an environmental pollutant of great concern (5), as its ongoing existence in the environment affects environmental quality in cities and is raising particular

Significance

Lead is a highly toxic element, and low-level environmental lead exposure remains a prime concern due its effects on neurodevelopment in children. It provides a well-researched example of the impact of human activities on environmental quality and the effectiveness of mitigation policies. Using recent and historical data on lead isotopic composition in London airborne particles, we show strong evidence that sources have not changed substantially since the ban in 1999 of leaded gasoline, the main source of environmental lead during the 20th century, suggesting that historical gasoline-derived lead remains an important source of lead in the urban environment due to its persistence and effective remobilization. Novel abatement policies need to be considered if this cycle is to be interrupted.

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¹To whom correspondence may be addressed. Email: eleonore.resongles@ird.fr or d.weiss@imperial.ac.uk.

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concerns regarding long-term exposure (6). A significant association between Pb concentration in particulate matter smaller than 10 μm (PM_{10}) and the US population's blood was recently demonstrated (7), suggesting that inhalation and/or ingestion of coarse particles might be an important pathway for human exposure to Pb. Viewed in this context, Pb in the environment remains a significant threat to public health, and it is absolutely essential to accurately identify and monitor the sources and pathways of Pb in the urban environment.

While it has been suggested that Pb sources in urban environments today include local and/or distant emissions from industries, coal burning, and traffic (exhaust and nonexhaust emissions) (8–11), recent works studying Pb contamination of airborne particles in (peri-)urban environments (10, 12) have suggested that ongoing contributions of lead released into the environment by leaded gasoline combustion during the last century remain an important but underestimated source. If confirmed more widely, then abatement strategies aimed at eliminating Pb fully from the urban environment need to be adjusted. It is very important to note that these recent findings are in line with an early study of Pb in sediments and water of the San Francisco Bay, suggesting already in the early 2000s that recycling of legacy Pb from leaded gasoline may be a continuing threat to environmental health (13, 14). Furthermore, a shift in Pb concentrations within size distribution, from the fine range ($\text{PM}_{2.5}$) to the coarse range (PM_{10}), has been highlighted after the ban of leaded gasoline in the United States and Europe and ascribed to a change in dominant Pb sources consisting of industrial emissions and road dust resuspension instead of direct vehicle exhausts (15).

Alkyl Pb motor fuel additives were used in the United Kingdom from the 1930s until phaseout was completed at the end of 1999. Although added in an organometallic form, Pb emissions from vehicles were predominantly in the form of a fine aerosol of inorganic Pb salts (16). The maximum permitted level of Pb in UK motor fuel was 0.84 $\text{g} \cdot \text{L}^{-1}$. The limit fell successively to 0.40 $\text{g} \cdot \text{L}^{-1}$ in 1981 and 0.15 $\text{g} \cdot \text{L}^{-1}$ in 1986. At its peak in the early 1980s, of the order of 7,000 tons per year of Pb were emitted from road traffic exhaust in the United Kingdom, which had fallen to just 30 tons by the year 2000 (17). Until its final ban, leaded gasoline combustion remained the most important source of Pb emissions in the UK atmosphere (17). In the early 1980s, annual average airborne Pb concentrations at background sites in central London were around 500 to 600 $\text{ng} \cdot \text{m}^{-3}$, which fell to around 300 $\text{ng} \cdot \text{m}^{-3}$ in the second half of the 1980s and then dropped progressively to around 20 $\text{ng} \cdot \text{m}^{-3}$ in 2000. Airborne concentrations now are generally less than 10 $\text{ng} \cdot \text{m}^{-3}$ and have remained steady over the last decade (18).

Using Pb isotope composition of atmospheric particles has been a crucial tool in tracing the origin of Pb in the environment and identifying leaded gasoline as a dominant source during the 20th century (19). Following the removal of Pb additives from gasoline in Europe, Pb isotope ratios of atmospheric particles generally changed toward more radiogenic values due to the increase in the relative contribution of other Pb sources (19). Such an evolution was recorded in London airborne particles between 1998 and 2001 during the final phasing out of leaded gasoline in the United Kingdom (20). However, this trend was no longer observed a decade later, leading to the suggestion that leaded gasoline remained an important source of atmospheric Pb (9).

The aim of this study was to test if remobilization of historical gasoline-derived Pb remains today an important and persistent source of Pb in London and the urban environment. To this end, we studied the Pb isotope composition of airborne particles collected in central London between 1995 and 2018 using new and historical data and quantified the possible contribution of historical gasoline Pb. London is representative of many large

cities in developed countries, where particle emissions from industries and coal combustion are now relatively low, and where traffic emissions and dust resuspension represent dominant sources of airborne particles. Therefore, it constitutes an ideal site to study the persistence of Pb in urban environments almost 20 y after the complete phaseout of leaded gasoline. The variability of Pb concentration and isotopic composition was determined in PM_{10} and total suspended particles ($\text{TSP}_{\text{passive}}$) during 1 mo at a heavily trafficked site in central London (Marylebone Road site [MR]), where particle emissions are dominated by traffic. The data were compared with the previously published isotopic composition of potential Pb sources and PM_{10} from this historically monitored site. In addition, the variability of Pb isotopic composition in $\text{TSP}_{\text{passive}}$ collected between 2014 and 2018 at building height in central London (Imperial College London site [IC]) is reported to determine mixing and source contributions within the urban canyon.

Results and Discussion

Pb Sources at a Heavily Trafficked Site in Central London. The PM_{10} concentration measured at the MR site ranged between 16 and 25 $\mu\text{g} \cdot \text{m}^{-3}$ (Table 1) and matched the daily means reported by the London Air Quality Network during the same period [$23 \pm 5 \mu\text{g} \cdot \text{m}^{-3}$ (21)]. Southerly to westerly winds prevailed over the sampling period, and only a few short rain showers occurred (*SI Appendix, Fig. S1*). Under such wind conditions, the airflow is coming from the nearby road junction or from adjacent traffic to the monitoring station, promoting high PM_{10} and air pollutant concentrations related to traffic emissions (22).

Table 1 shows Pb concentrations (PM_{10}) or deposition rates ($\text{TSP}_{\text{passive}}$), enrichment factors (EF_{Pb}), and isotope ratios in PM_{10} and $\text{TSP}_{\text{passive}}$ collected at the MR site during the summer of 2018. Pb deposition rates and EFs were determined using Eqs. 2 and 3, respectively (*Material and Methods*). Pb concentration in PM_{10} ranged between 3.9 and 19.4 $\text{ng} \cdot \text{m}^{-3}$ and was slightly higher than that measured at the MR site in July 2010 [1.1 to $8.8 \text{ ng} \cdot \text{m}^{-3}$, (9)]. Pb deposition rates for the $\text{TSP}_{\text{passive}}$ were between 11,700 and 45,800 $\text{ng} \cdot \text{m}^{-2} \cdot \text{d}^{-1}$, which represent 10 to 40 times and 2 to 9 times the Pb deposition rates recorded using the same sampling device at a busy boulevard in Strasbourg, France and a German highway, respectively (23). This is due to the high traffic-related particle load characteristic of the MR site. The EF for Pb, EF_{Pb} , ranged between 100 and 559 in PM_{10} and 66 and 91 in $\text{TSP}_{\text{passive}}$ (Table 1), indicating the high contribution of anthropogenic Pb in airborne particles at the MR site and showing an influence of the particle size on the enrichment of Pb. Elements commonly associated with traffic emissions from vehicle wear such as Ba, Cr, Cu, Fe, Ni, and Sb (24, 25) were highly enriched in PM_{10} ($\text{EF} > 10$) and to a lesser extent in $\text{TSP}_{\text{passive}}$ (Fig. 1 and *SI Appendix, Table S1*). This observation is in line with the known effect of the size contribution on the enrichment of these elements. In contrast, Zn presented similar high EF in both PM_{10} ($181 \leq \text{EF}_{\text{Zn}} \leq 318$) and $\text{TSP}_{\text{passive}}$ ($179 \leq \text{EF}_{\text{Zn}} \leq 749$), reflecting the important contribution of tire abrasion to the coarse fraction (PM_{10}) and the supercoarse fraction ($>10 \mu\text{m}$) as well (25, 26). Elements released by brake abrasion (Ba, Cu, Cr, and Fe) show strong correlations in PM_{10} ($r > 0.7$, *SI Appendix, Table S2*) and in $\text{TSP}_{\text{passive}}$ despite the lower enrichment values observed (Fig. 1). Surprisingly, EF_{Sb} in PM_{10} are five times lower in 2018 than in 2010 and no longer correlate well with other metals ($0.4 < r < 0.7$, *SI Appendix, Table S2*). This is likely the consequence of the progressive removal of Sb from brake pads by constructors because of its suspected carcinogenic effect (27), and our observation is consistent with the higher than usual Cu/Sb ratio reported for new brake pads in London in 2010 (9). EFs of Pb are not correlated with EFs of any other metal as already observed in a previous study at the site (9). This is suggesting that

Table 1. Pb concentrations, EF_{Pb}, and Pb isotope ratios in airborne particles collected at the MR site (PM₁₀ and TSP_{passive}) and at the IC site (TSP_{passive})

Sample label	Date of collection	PM ₁₀ concentration (µg · m ⁻³)	Pb concentration (ng · m ⁻³)	EF _{Pb}	Pb isotope ratios						Leaded gasoline contribution (%)	
					²⁰⁶ Pb/ ²⁰⁷ Pb	2SD	²⁰⁸ Pb/ ²⁰⁶ Pb	2SD	²⁰⁶ Pb/ ²⁰⁴ Pb	2SD	Non-gasoline Pb: 100% Vehicle wear	Non-gasoline Pb: 100% UK coal and ore
MR site (PM ₁₀)												
MR1a	25/07/18 to 29/07/18	19	8.8	100	1.1407	0.0001	2.1195	0.0001	17.774	0.002	24	36
MR2a	29/07/18 to 02/08/18	21	3.9	118	1.1331	0.0001	2.1279	0.0001	17.636	0.003	32	43
MR3a	02/08/18 to 06/08/18	16	6.1	154	1.1333	0.0001	2.1255	0.0001	17.644	0.004	32	42
MR4a	06/08/18 to 10/08/18	25	7.6	155	1.1364	0.0001	2.1235	0.0002	17.694	0.003	29	40
MR5a	10/08/18 to 14/08/18	21	4.1	141	1.1322	0.0001	2.1282	0.0001	17.619	0.003	33	43
MR6a	14/08/18 to 18/08/18	21	19.4	559	1.1281	0.0001	2.1331	0.0002	17.541	0.003	37	47
MR7a	18/08/18 to 22/08/18	18	5.4	159	1.1370	0.0001	2.1220	0.0001	17.699	0.004	28	39
MR8a	22/08/18 to 26/08/18	16	5.1	197	1.1268	0.0001	2.1330	0.0001	17.523	0.003	39	48
Average ± SD		20 ± 3	7.6 ± 5.1	198 ± 149	1.134 ± 0.005		2.127 ± 0.005		17.64 ± 0.08		32 ± 5	42 ± 4
MR site (TSP _{passive})												
			Pb deposition rate (ng · m ⁻² · d ⁻¹)									
MR1b	25/07/18 to 29/07/18	n.a.	45,813	84	1.1383	0.0001	2.1232	0.0001	17.724	0.002	27	38
MR2b	29/07/18 to 02/08/18	n.a.	18,154	90	1.1355	0.0001	2.1257	0.0002	17.675	0.001	30	41
MR3b	02/08/18 to 06/08/18	n.a.	11,698	66	1.1319	0.0001	2.1289	0.0001	17.621	0.002	33	44
MR4b	06/08/18 to 10/08/18	n.a.	18,661	54	1.1399	0.0001	2.1212	0.0001	17.754	0.001	25	37
MR5b	10/08/18 to 26/08/18	n.a.	24,069	91	1.1375	0.0001	2.1241	0.0001	17.711	0.006	28	39
Average ± SD			23,679 ± 13,127	77 ± 16	1.137 ± 0.003		2.125 ± 0.003		17.70 ± 0.05		29 ± 3	40 ± 3
IC site (TSP _{passive})												
S1	14/02/14 to 14/03/14	n.a.	n.d.	n.d.	1.1482	0.0001	2.1153	0.0002	17.900	0.003	17	30
S3	11/07/14 to 20/08/14	n.a.	8,685	296	1.1354	0.0001	2.1281	0.0002	17.681	0.001	30	41
S4	20/08/14 to 26/09/14	n.a.	7,240	204	1.1392	0.0001	2.1244	0.0001	17.748	0.002	26	37
S5	26/09/14 to 27/10/14	n.a.	7,757	256	1.1403	0.0001	2.1217	0.0001	17.765	0.002	25	36
S6	27/10/14 to 27/11/14	n.a.	3,691	179	1.1419	0.0001	2.1203	0.0001	17.792	0.003	23	35
S7	27/11/14 to 27/12/14	n.a.	5,840	271	1.1358	0.0001	2.1263	0.0001	17.687	0.002	29	40
S8	27/12/14 to 22/01/15	n.a.	4,665	251	1.1331	0.0001	2.1290	0.0001	17.639	0.002	32	43
S9	22/01/15 to 25/02/15	n.a.	3,094	175	1.1240	0.0001	2.1380	0.0001	17.477	0.001	42	51
S10	25/02/15 to 22/04/15	n.a.	5,474	132	1.1357	0.0001	2.1267	0.0001	17.685	0.002	30	40
S11	22/04/15 to 03/07/15	n.a.	8,194	195	1.1403	0.0002	2.1223	0.0001	17.761	0.006	25	36
S12	03/07/15 to 26/08/15	n.a.	9,000	300	1.1334	0.0001	2.1293	0.0001	17.645	0.005	32	42
S13	27/03/17 to 02/05/17	n.a.	6,371	90	1.1343	0.0001	2.1281	0.0002	17.664	0.002	31	42
S14	02/05/17 to 02/06/17	n.a.	9,411	181	1.1303	0.0001	2.1317	0.0001	17.598	0.003	35	45
S15	02/06/17 to 06/07/17	n.a.	9,785	221	1.1304	0.0001	2.1327	0.0001	17.595	0.003	35	45
S16	06/07/17 to 03/08/17	n.a.	3,903	146	1.1351	0.0001	2.1269	0.0001	17.677	0.003	30	41
S17	03/08/17 to 04/09/17	n.a.	4,933	276	1.1219	0.0001	2.1412	0.0002	17.448	0.002	44	52
S18	04/09/17 to 06/10/17	n.a.	9,748	475	1.0977	0.0001	2.1636	0.0001	17.049	0.003	69	74
S19	06/10/17 to 10/11/17	n.a.	3,957	176	1.1355	0.0001	2.1276	0.0001	17.684	0.004	30	41
S20	31/07/18 to 07/08/18	n.a.	2,862	71	1.1373	0.0001	2.1253	0.0001	17.710	0.003	28	39
S21	07/08/18 to 30/08/18	n.a.	2,465	84	1.1327	0.0001	2.1285	0.0001	17.627	0.002	33	43
Average ± SD			6,162 ± 2,508	209 ± 95	1.13 ± 0.01		2.13 ± 0.01		17.64 ± 0.17		32 ± 10	43 ± 9

The percentage contribution of leaded gasoline estimated using the binary mixing model (Eq. 1) is indicated for the two extreme non-gasoline Pb end-members considered (100% vehicle wear and 100% UK coal and ore; sensitivity analysis shown in *SI Appendix, Fig. S7*). n.a., not applicable; n.d., not determined.

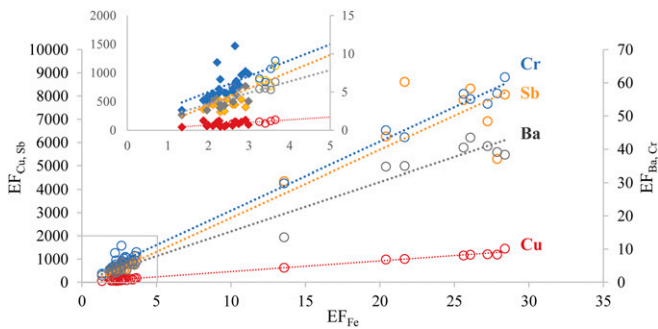


Fig. 1. Relationship between the EF of Fe (EF_{Fe}), Ba (EF_{Ba}), Cr (EF_{Cr}), Cu (EF_{Cu}), and Sb (EF_{Sb}) in PM_{10} and $TSP_{passive}$ at MR (open circles) and in $TSP_{passive}$ at building height at IC (filled diamonds).

its emission is not controlled by a dominant source such as brake (Ba, Cu, Fe, Cr) or tire wear (Zn) but rather results from another dominant source or, more likely, from a mixture of different sources. The lower EF observed in $TSP_{passive}$ can be explained by the collection of supercoarse particles less enriched with Pb and other metals (except Zn), which probably consist of road surface material besides tire particles. This is further supported by TLM (transmitted-light microscope) images of $TSP_{passive}$ samples showing opaque elongated and round/irregular-shaped supercoarse particles, which are typical for tire road wear (*SI Appendix, Fig. S2*, samples MR1b to MR3b). Recent high time-resolution measurements of Pb and Si in PM_{10} at the MR site showed that Pb and Si, a marker for soil-derived mineral dust, exhibited the same diurnal trend suggesting that resuspension with high traffic is associated with higher airborne Pb (*SI Appendix, Fig. S3*).

Average Pb isotope ratios in PM_{10} and $TSP_{passive}$ measured over the same time period at the MR site were not significantly different (t test, $P > 0.31$) ($^{206}Pb/^{207}Pb = 1.137 \pm 0.003$, $^{208}Pb/^{206}Pb = 2.125 \pm 0.003$, and $^{206}Pb/^{204}Pb = 17.70 \pm 0.05$ for PM_{10} [$n = 8$] and $^{206}Pb/^{207}Pb = 1.133 \pm 0.005$, $^{208}Pb/^{206}Pb = 2.127 \pm 0.005$, and $^{206}Pb/^{204}Pb = 17.64 \pm 0.08$ for $TSP_{passive}$ [$n = 5$]). Differences of Pb isotopic composition observed between PM_{10} and $TSP_{passive}$ at each sampling date were smaller than the variability observed within PM_{10} and $TSP_{passive}$ over the sampling period (Fig. 2). The results suggest that most of the Pb mass is associated with the PM_{10} fraction, which is supported by the lower EF_{Pb} observed in $TSP_{passive}$, indicating a “dilution” by less-contaminated supercoarse particles. Therefore, in this context, the continuous monitoring of Pb isotopic composition of $TSP_{passive}$ collected using the passive sampler Sigma-2 appears as a suitable low-cost technique to monitor long-term changes of Pb sources in coarse airborne particles. However, it should be noted that passive sampling does not allow investigation of Pb deposition changes with a high time resolution. For example, the sample MR6a showed the highest Pb concentration together with a relatively low $^{206}Pb/^{207}Pb$ ratio compared to the average value. This was likely due to a short-term event (e.g., construction works in the street, building demolition, etc.) and was not visible with passive sampling because the corresponding sample (MR5b) integrated a longer period of time.

Pb isotope ratios of PM_{10} and $TSP_{passive}$ collected during the summer of 2018 were not statistically different to those of PM_{10} measured by Dong et al. (9) in 2010 at the same site (t test, $P > 0.30$), showing that the Pb isotope composition displayed a stable signature over the last decade and that Pb sources remained unchanged (Fig. 3). A source apportionment study of aerosols collected at the MR site based on a statistical factor analysis of the chemical composition suggested that coarse particles (2.5 to 10 μm) are composed of construction dust (6%), vehicle wear (13%), resuspended road dust (31%), and sea-salt (49%), while

fine particles ($<2.5 \mu m$) are composed mainly of construction dust (9%), sea-salt (10%), exhaust and nonexhaust traffic emissions (33%), and secondary inorganic aerosols (49%) (24). Pb was not included in the source apportionment of coarse particles due to concentrations being close to the relatively high detection limit of the analytical technique used (Particle-Induced X-Ray Emission). In the fine particles ($PM_{2.5}$), Pb was largely associated with a long-range transported “secondary inorganic aerosols” factor also containing crustal elements. In a separate sampling campaign during the same 2012 research program, Visser et al. (28) found using a different analytical technique (SR-XRF) an appreciable fraction of Pb (41% on average) associated with the coarse fraction showing a fairly balanced distribution of Pb between fine and coarse particles. During the sampling period, air masses generally traveled across the North Atlantic Ocean and crossed the southern United Kingdom before reaching the sampling site (*SI Appendix, Fig. S4*). Air trajectories coming from Northern France, Belgium, and the Netherlands were recorded only during the sampling of MR1. Under these conditions, the contribution of Pb transported from the European mainland must be minimal (29). The emissions inventory reported by the UK National Atmospheric Emission Inventory (NAEI) estimates that brake and tire abrasion could be the major current source of atmospheric Pb in London, representing about 60% of Pb primary emissions (17). However, this inventory is associated with high uncertainties and has been shown by Dore et al. to seriously underestimate Pb emissions as compared to Pb deposition measurements (30). Tire and brake material as well as road surface material have more radiogenic isotope ratios than $TSP_{passive}$ and PM_{10} (Fig. 3, ref. 9). However, the isotopic composition of road dust [$^{206}Pb/^{207}Pb = 1.133 \pm 0.011$, $^{208}Pb/^{206}Pb = 2.129 \pm 0.011$, and $^{206}Pb/^{204}Pb = 17.66 \pm 0.21$, $n = 6$ (9)] and urban topsoils of London ($^{206}Pb/^{207}Pb = 1.139 \pm 0.017$ and $^{208}Pb/^{206}Pb = 2.125 \pm 0.017$, $n = 50$) (31) overlaps with that of airborne particles (Fig. 3). As shown on Fig. 3, they fall on a mixing line between the signature of leaded gasoline formerly used in the United Kingdom (32) and of natural geogenic Pb in the United Kingdom (32), UK Pb ore (33), and nonexhaust emissions (9). A compilation of a $^{206}Pb/^{207}Pb$ ratio of potential Pb sources in London is presented in *SI Appendix, Table S3*. They all have a more radiogenic composition than airborne particles except for lead paint. Data for lead paint remain scarce; isotope composition published for paint in Scotland showed a $^{206}Pb/^{207}Pb$ ratio of 1.117 ± 0.003 (34), overlapping with the present-day signature of airborne particles. It is reasonable to assume that lead paint contributes to “Pb hotspots” in urban private and public gardens

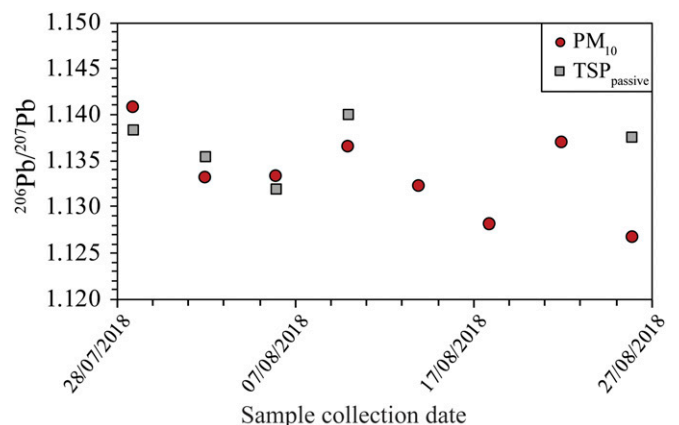


Fig. 2. $^{206}Pb/^{207}Pb$ ratio measured in PM_{10} and $TSP_{passive}$ collected over a 96-h period at the MR site (except the last sample, MR5b, which integrates $TSP_{passive}$ deposition over 16 d).

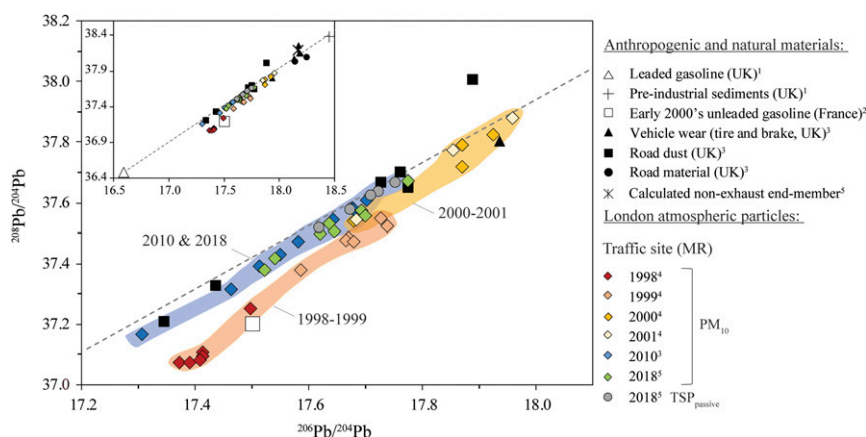


Fig. 3. $^{208}\text{Pb}/^{204}\text{Pb}$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ diagram of historic and current PM_{10} and $\text{TSP}_{\text{passive}}$ at the MR site and potential traffic-related and natural Pb sources. The gray dashed line represents mixing lines between the leaded gasoline end-member and a natural/nonexhaust end-member. References: ¹Monna et al. (32), ²Widory et al. (40), ³Dong et al. (9), ⁴Noble et al. (20), ⁵calculated in this study using data from (9).

where old lead paint is deteriorating, but it is an unlikely major source in atmospheric particles in urban canyons. For example, a study in Tokyo, Japan showed that, while gasoline and paint chips were found to contribute to increased Pb in surface soils of playgrounds, the contribution of paint chips was significantly smaller than gasoline Pb (35). In New Orleans, United States, it was estimated that at least 10 times more Pb dust was released from vehicle exhaust than from lead paint (36). Therefore, the isotope signatures strongly suggest that only a contribution of historical Pb originating from leaded gasoline can explain the less radiogenic composition observed in soils, road dust, and airborne particles. This is also in line with previous results from a study that assessed that up to 43% of Pb in UK road dusts was derived from leaded gasoline in 2010 (37). Indeed, Dore et al. showed that national primary emissions of Pb estimated by the UK NAEI could only account for 7% of the measured Pb atmospheric concentration and suggested that wind-driven resuspension of surface dust could partly explain the gap between modeled and measured values (30). Amato et al. (38), in a study of dust resuspension on a Mediterranean freeway, calculated an emission factor of 16 to 22 $\mu\text{g Pb/veh} \cdot \text{km}^{-1}$. Using this value and road traffic statistics in London, we calculated that 450 to 650 kg Pb would be emitted in London every year by dust resuspension. Alternatively, Denier van der Gon et al. (39) provided emissions factors of PM_{10} for light and heavy-duty vehicles (48 and 432 $\text{mg PM}_{10}/\text{veh} \cdot \text{km}^{-1}$, respectively). Based on a Pb concentration in road dust in London of 22 to 410 $\mu\text{g} \cdot \text{g}^{-1}$ (9), we calculated that 45 to 800 kg Pb would be emitted yearly through resuspension. The latter upper-bounding estimate is in line with the former. Despite the uncertainty associated with this estimate, it shows that remobilization of Pb from road dust resuspension is likely of the same order as the current estimation of primary Pb emissions. This is consistent with our conclusions based on isotopic ratios.

Pb isotopic compositions in 2010 and 2018 were slightly different from those between 1998 and 1999, characterized by a less thorogenic composition (i.e., lower $^{208}\text{Pb}/^{204}\text{Pb}$), and those between 2000 and 2001, characterized by a more radiogenic composition (i.e., higher $^{208}\text{Pb}/^{204}\text{Pb}$ and $^{206}\text{Pb}/^{204}\text{Pb}$) (Fig. 3). During this period (1998 to 2001), a progressive shift of Pb isotope ratios is observed, indicating a declining contribution of leaded gasoline interpreted as the result of the final ban of leaded petrol in the United Kingdom in 2000 (20). The isotopic shift was accompanied by a rapid decline of Pb concentrations in airborne particles. The disparity observed between 1998 and 2001 and 2010 and 2018 data is likely explained by the contribution of a

different traffic-related Pb source at the end of the 20th century. We propose that this additional source is related to either unleaded gasoline combustion (20) and/or to increased contribution from vehicle wear (e.g., Pb wheel weights) before the introduction of a regulation in 2003 prohibiting the use of Pb in materials and components in vehicles sold in the European Union market (European Directive 2000/53/EC). Indeed, in the early 2000s, unleaded gasoline still contained low levels of Pb in Europe, and data available for an unleaded French gasoline could explain the observed trend (40). However, Pb isotope composition of unleaded gasoline and vehicle components may have varied in the past depending on the provenance of the original crude oil or Pb ore. The lack of isotope data for the materials used during this period undermines our ability to definitely identify the additional source(s) responsible for the observed offset. Recent European diesel ($n = 26$) and gasoline ($n = 64$) contained less than $1.5 \mu\text{g} \cdot \text{kg}^{-1}$ of Pb (41). This was confirmed by measuring lead concentration in gasoline ($n = 2$) and diesel ($n = 2$) collected in June 2018 from two different petrol stations in London. Pb content in gasoline and diesel was lower than 2 and $0.6 \mu\text{g} \cdot \text{kg}^{-1}$, respectively. Calculations accounting for fuel consumption in London (~ 1.5 Mt diesel and 0.9 Mt gasoline per year) and the Pb content in fuel measured in this study and others (41) demonstrate unequivocally that fuel combustion is no longer a relevant source of Pb (less than a few kilograms per year).

Pb Isotopic Composition Variation between 2014 and 2018 at Building Height in Central London. Pb deposition rates, EF_{Pb} , and Pb isotope ratios in $\text{TSP}_{\text{passive}}$ collected between 2014 and 2018 at the IC site are presented in Table 1. Pb deposition rates varied from $\sim 3,100$ to $9,800 \text{ ng} \cdot \text{m}^{-2} \cdot \text{d}^{-1}$, which represent, on average, 26% of the Pb deposition rates recorded at the MR site. EF_{Pb} varied from 71 to 475 with an average of 209 ± 95 ($n = 19$). Pb deposition rates were lower in autumn and winter than in spring and summer certainly due to suppression of resuspension in the wetter months of the year (*SI Appendix, Fig. S5*). Seasonality in coarse particles has been observed at both traffic (MR) and urban background sites in London and attributed to a more effective resuspension process during the drier seasons (42). However, EF_{Pb} does not exhibit such seasonal variations (Fig. 4 and *SI Appendix, Fig. S5*), showing that Pb level in $\text{TSP}_{\text{passive}}$ was not sensitive to seasonal emission sources like domestic fuel combustion, different to what has been found for Cu in 2014 in London (43). EF_{Pb} does not display a declining long-term trend, indicating that Pb reached a steady level in the atmosphere of central London. Similar EF_{Pb} values in $\text{TSP}_{\text{passive}}$ were recorded

at the IC site and at MR over the same period. EF_{Pb} does not correlate with enrichment factors of other metals as already observed at the MR site. Other elements studied (Ba, Cu, Cr, Fe, Ni, Sb, and Zn) show slightly lower EF values at the IC site compared to the MR site but still denote anthropogenic inputs (Fig. 1). Elements typical of brake-wear emissions fall on the same regression line as MR samples, demonstrating a common origin (Fig. 1).

The Pb isotopic composition of $TSP_{passive}$ at the IC site shows a larger range of isotopic ratios than at the MR site, with $^{206}Pb/^{207}Pb$, $^{208}Pb/^{206}Pb$, and $^{206}Pb/^{204}Pb$ ratios varying from 1.0977 to 1.1482, 2.1153 to 2.1636, and 17.049 to 17.900, respectively (Table 1). The sample S18 (09/2017) displays unusually low $^{206}Pb/^{207}Pb$ and $^{206}Pb/^{204}Pb$ ratios and has the highest EF_{Pb} and Pb deposition rate. A 3-d pollution episode (high $PM_{2.5}$ and PM_{10}) occurred during the S18 sampling interval. Polluted air arriving from the continent to the area of London might explain the unusual Pb signature (21). Alternatively, the anomaly could be related to local emissions and, possibly, to demolition works carried out next to the sampling site at that time. Apart from this event, as observed for EF_{Pb} , Pb isotope ratios do not exhibit a seasonal or long-term trend (Fig. 3), confirming that Pb sources remained the same throughout the year—this suggests that coal and fossil-fuel burning does not add significantly to the Pb burden. Pb isotope ratios at the IC site are not significantly different (t test, $P > 0.77$) to those measured in PM_{10} and $TSP_{passive}$ at the MR site (Fig. 5). Samples S20 and S21 collected during the summer of 2018 show similar EF_{Pb} and Pb isotope ratios as $TSP_{passive}$ collected during the same period at the MR site, indicating that the Pb sources are the same. These results suggest that Pb has a homogeneous origin in airborne particles of different sizes in urban canyons in central London and that there is no apparent additional source at the IC site compared to the MR site. This is consistent with TLM optical analysis of $TSP_{passive}$ samples of the IC site, showing that tire-wear particles emitted from traffic are, in small numbers, also present even at building height (SI Appendix, Fig. S2, sample S20). Changes in the $^{206}Pb/^{207}Pb$ ratio with possible varying contributions of coal have been modeled using a mixing equation between 1) the average composition of air particles at MR measured during the summer of 2018 (representative of Pb emissions from traffic and dust resuspension) and b) the average composition of coal used in the United Kingdom that has a $^{206}Pb/^{207}Pb$ ratio between 1.18 and 1.22 (SI Appendix, Table S3). An increase in Pb emissions due to coal combustion would be responsible for an increase of $^{206}Pb/^{207}Pb$ values in airborne particles toward a more radiogenic isotope composition (SI Appendix, Fig. S6). All samples (except S1) indicate a coal contribution to Pb signature lower than 10%, ruling out a significant contribution of coal combustion to Pb burden (SI Appendix, Fig. S6). Indeed, coal consumption has drastically declined in the United Kingdom during the last decade. Combustion of fuel (oil and coal) in relation to domestic, industrial, and commercial sectors has been estimated at ~ 300 kg Pb per year in the Greater London area, representing a maximum of 8% of total emissions (17) in accordance with our results.

Quantification of Leaded Gasoline's Contribution to Pb Contamination of Airborne Particles. We made an attempt to estimate the historical contribution of Pb originating from leaded gasoline (X in %) using a binary mixing model (Eq. 1) considering two end-members, that is, 1) “gasoline Pb,” using the signature of leaded gasoline used in the past in the United Kingdom (32) and 2) “non-gasoline Pb,” representative of other possible sources (present-day nonexhaust emissions and historical UK coal and ore). A natural crustal source can reasonably be neglected considering the high EF values ($54 < EF_{Pb} < 559$) observed in airborne particles both at the traffic MR site and the IC site (Table 1).

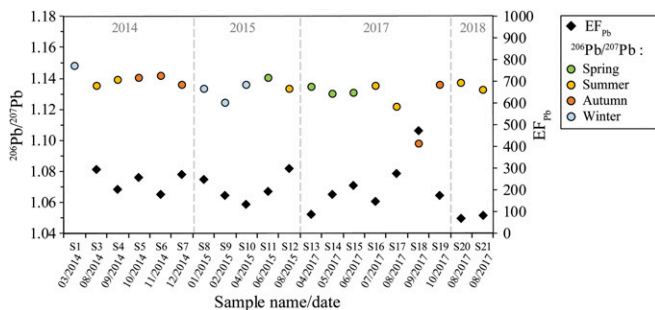


Fig. 4. Temporal variations of the $^{206}Pb/^{207}Pb$ ratio of $TSP_{passive}$ collected at the IC site. EF_{Pb} are represented by black diamonds and the $^{206}Pb/^{207}Pb$ ratio by colored circles indicating the season (green: spring, yellow: summer, orange: autumn, and blue: winter).

$$X = \frac{(^{206}Pb/^{207}Pb)_{sample} - (^{206}Pb/^{207}Pb)_{non-gasoline Pb}}{(^{206}Pb/^{207}Pb)_{gasoline Pb} - (^{206}Pb/^{207}Pb)_{non-gasoline Pb}} \times 100. \quad [1]$$

To test the validity of this approach, a sensitivity analysis was performed to assess the contribution of leaded gasoline, using different isotopic compositions for the “non-gasoline Pb” end-member calculated using different proportions of Pb from non-exhaust emissions, UK ore, and UK coal. To this end, the Pb isotopic composition of the “vehicle wear” end-member was calculated using recent Pb emission factors associated with tire and brake abrasion (SI Appendix, Table S4), indicating a contribution of 17% from tire wear and 83% from brake wear and the isotopic composition of tires and brakes used in London as reported by Dong et al. (9). Results do not significantly change if the average isotope composition of all nonexhaust emissions is used or that of vehicle wear alone (SI Appendix, Fig. S7). The results obtained using the two extreme “non-gasoline Pb” sources (i.e., 100% vehicle wear and 100% UK coal and ore) are presented in Table 1 and SI Appendix, Fig. S7. Other ratios ($^{206}Pb/^{204}Pb$ and $^{208}Pb/^{206}Pb$) gave similar results with a maximum bias of 8%.

At the heavily trafficked MR site, leaded gasoline could contribute to $32 \pm 5\%$ of Pb in PM_{10} , which is similar to the contribution of $37 \pm 8\%$ determined using 2010 data from Dong et al. (9) when using vehicle wear as the “non-gasoline Pb” source end-member. An equivalent average gasoline contribution was found in $TSP_{passive}$ at the MR site ($29 \pm 3\%$) and at the IC site ($32 \pm 10\%$). The estimated contribution of leaded gasoline to airborne Pb reaches $42 \pm 4\%$ (PM_{10} , MR site), $40 \pm 3\%$ ($TSP_{passive}$, MR site), and $43 \pm 9\%$ ($TSP_{passive}$, IC site) when using UK coal and ore as the “non-gasoline Pb” source end-member. These results highlight a continuous and significant contribution of historical Pb from gasoline in central-London airborne particles over the last decade.

It should be noted that this approach does not allow estimation of the relative contribution of other historical sources (UK coal and ore) or present-day nonexhaust emissions due to the similarity of their isotopic signatures, nor the contribution of lead paint because its signature is not well constrained and is probably overlapping that of airborne particles. However, vast amounts of Pb have been emitted from leaded gasoline combustion in the London atmosphere during the 20th century. It is estimated that $\sim 140,000$ tons of gasoline-derived Pb were released in the UK atmosphere between 1970 and 2000 (17). Consequently, Pb has been deposited in urban soils in which it can have a long residence time (44–46) as reflected by the high median Pb concentration ($175 \mu g \cdot g^{-1}$) reported for topsoils in the Greater London area (31, 47–49). Soil isotope composition has been explained by the long-term accumulation of

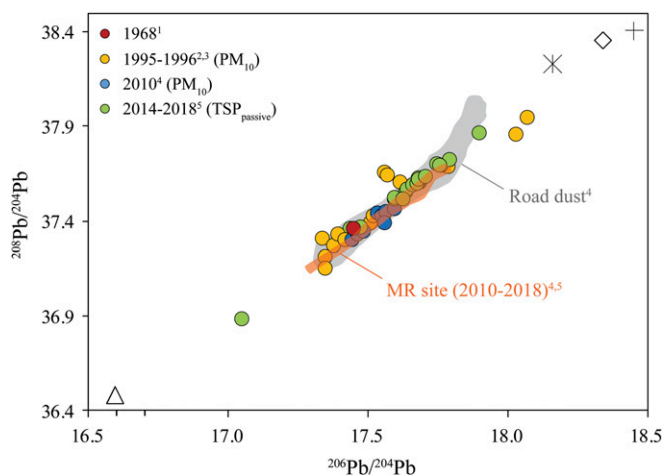


Fig. 5. $^{208}\text{Pb}/^{204}\text{Pb}$ versus $^{206}\text{Pb}/^{204}\text{Pb}$ diagram of historic and current PM_{10} and $\text{TSP}_{\text{passive}}$ collected at various urban sites in London (Teddington², North Kensington⁴, the IC site⁵, and the MR site^{4,5}). The variation range observed in road dust³ (dashed line) and in airborne particles at the MR site (2010 to 2018) (blue area) are represented. The isotopic composition of natural lead² (+), leaded gasoline² (Δ), vehicle wear⁴ (X), UK ore⁶ (\diamond) end-members are indicated. References: ¹Hamilton and Clifton (61), ²Monna et al. (32), ³Véron et al. (62), ⁴Dong et al. (9), ⁵this study, ⁶Rhol (33).

anthropogenic emissions from several sources including leaded gasoline combustion, coal burning and demolition of buildings during the Second World War (paint, pipes) (31, 48). However, higher median Pb concentrations are measured in topsoils located near roads (<100 m) probably due to accumulation of a greater amount of Pb from leaded gasoline (48). A recent study of a sediment core sampled in the Thames in London showed that sediments were already enriched with Pb in the 1940s ($^{206}\text{Pb}/^{207}\text{Pb}$ ratio ≈ 1.16), but the isotope composition shifted later with the lowest $^{206}\text{Pb}/^{207}\text{Pb}$ values recorded between the 1950s and 1980s, when leaded gasoline consumption was at its highest level (50). Pb isotope composition in the most recent sediments did not return to the <1940s value ($^{206}\text{Pb}/^{207}\text{Pb}$ ratio ≈ 1.14), providing further evidence of the long-term contribution of leaded gasoline. This is consistent with the results of Ayrault et al. (51), who found that only 1.4% of Pb emitted annually in the Seine river basin during the use of leaded gasoline was exported in the river, implying that most of the gasoline Pb accumulated in soils of the basin (51). This conclusion is supported by several studies showing that urban and roadside soils preserved the gasoline signature (37, 52). Old buildings in cities kept the signature of past Pb atmospheric emissions as well (53, 54), demonstrating the general “impregnation” of the urban environment with historical gasoline-derived Pb.

Environmental Significance. Pb concentrations measured in PM_{10} in London over a decade were stable and low compared to the concentration recorded during the last century [annual average of $8.8 \pm 0.9 \text{ ng} \cdot \text{m}^{-3}$ for the period between 2010 and 2017 at London Westminster (18)]. Nevertheless, this study shows that historical Pb stored in the urban environment represents a persisting source of Pb in airborne particles in London via the remobilization of contaminated dust. Noncombustion traffic-related emissions represent a major source of atmospheric Pb through direct emissions from road and vehicle wear and indirect emissions consisting of the resuspension of road dust, which is a mixture of abrasion particles (vehicle and road wear), and of mineral dust deriving from historically contaminated soils and urban surfaces. Pb isotopic composition of airborne particles collected at building height showed a similar signature to those

collected at street level, suggesting that Pb is well mixed within the urban environment in central London and has reached a steady baseline. Even though health policy needs to solve the problem of high-level exposure of children as a matter of priority, our study clearly shows the persistence of Pb historically emitted from leaded gasoline combustion in urban environments and may be important from an environmental health perspective. A positive correlation has been found between soil resuspension, airborne Pb, and children’s blood Pb level in several large cities in the United States (10, 36, 55). Our data strongly suggest that soils act as a persistent source of airborne Pb in London, implying that Pb remobilization might be an overlooked exposure pathway in London. Future research is now needed to investigate the seasonal variations of airborne Pb in relation to soil resuspension and to assess its possible relationship with children’s blood Pb levels. If such a link exists, historical Pb remobilization could only be interrupted by new abatement strategies. Possible remediation strategies to reduce mobilization of soil Pb include covering contaminated urban soils with low-Pb soil, driven by a high-resolution Pb-soil mapping, which has been effective in reducing children’s blood Pb levels in New Orleans (6).

Material and Methods

Sample Collection. Eight samples of PM_{10} and five samples of $\text{TSP}_{\text{passive}}$ were collected during the summer of 2018 at street level (height of 3.5 m) at a heavily trafficked site in central London (MR). This site belongs to the London Air Quality Network and has been used extensively for air quality studies, including for Pb isotopic composition monitoring during the period from 1998 to 2001 and in 2010 (9, 20). The station is located on the curbside of a street canyon characterized by high traffic (>80,000 vehicles per day) and frequent congestion. PM_{10} (MR1a to MR8a) were collected over 96 h on acid-cleaned PVDF (polyvinylidene fluoride) filters (Ø 47 mm, 1 μm) over 32 d in July and August of 2018 using a Partisol-Plus Model 2025 Sequential Air Sampler. In parallel, two passive samplers (Sigma-2 device, Deutscher Wetterdienst [DWD]) were installed next to the active sampler inlet (distance of ~ 1 m and height of ± 25 cm) to collect $\text{TSP}_{\text{passive}}$ [defined as particulate matter with a diameter up to 50 μm deposited via sedimentation in the passive sampler device (26, 56)]. One passive sampler collected TSP in an acid-cleaned receptor dish (Ø 65 mm) and was dedicated to geochemical analyses (Pb and metal concentrations and Pb isotope ratios). Standard DWD transparent sticky foil was placed in the second passive sampler and used for TLM. The receptors of both passive samplers were exposed for 96 h and exchanged at the same time as PM_{10} filters (MR1b to MR4b) except the last sample (MR5b), which integrated a longer sampling period (4×96 h). The passive sampling method was used in addition to the conventional active sampling of PM_{10} to determine the influence of the size of particles (coarse 2.5 to 10 μm and “supercoarse” 10 to 50 μm) on Pb isotopic composition due to the possible contribution of different Pb sources to the different size fractions.

In addition, the variability of Pb isotopic composition in $\text{TSP}_{\text{passive}}$ at building height in central London was assessed by analyzing a series of samples collected using a passive sampler Sigma-2 installed on the rooftop (24 m above street level) of the Royal School of Mines at the IC site. A total of 20 samples (S1 to S21) were taken at intervals of 4 to 8 wk between February 2014 and August 2018. $\text{TSP}_{\text{passive}}$ were collected in an acid-cleaned receptor dish (Ø 65 mm). Metal deposition rates, EFs (Al, Zn, Cu, Fe, Sb and Cd), and Cu and Zn isotopic signatures of samples S1 to S12 have been presented in Ochoa Gonzalez et al. (43).

Sample Digestion. All the experimental work was carried out in a Class 1000 clean laboratory and under Class 10 laminar flow hoods in the MAGIC (MASS Spectrometry and Isotope Geochemistry at Imperial College London) laboratories at Imperial College London. All material was acid-cleaned before use. Acids were of Suprapur quality or purified by distillation. For each set of digestion, a blank and two certified reference materials (20 mg granite USGS G-2 from the United States Geological Survey and road dust BCR 723 from the Community Bureau of Reference) were processed in the same way as samples.

PM_{10} filters were weighed before and after the collection to determine the PM_{10} concentration. The filters were placed in PFA (perfluoroalkoxy) vials, and particles were recovered by refluxing the filter with 4 mL 15 M HNO_3 (nitric acid) and 2 mL 70% vol/vol HClO_4 (perchloric acid) at 100 $^\circ\text{C}$ over

24 h, using an ultrasonic bath for 1 h every 12 h. Then filters were removed using clamps and rinsed with 5 mL 0.5 M HNO₃. Solutions were evaporated to dryness at 230 °C, refluxed with 1.5 mL 15 M HNO₃ and 1 mL 40% vol/vol HF (hydrofluoric acid), evaporated again, and redissolved in 2 mL 0.5 M HNO₃.

Samples of TSP_{passive} collected with the passive sampler were prepared following the protocol described in Ochoa Gonzalez et al. (43). The samples were transferred from the receptor dish into PFA vials with 15 mL 0.1 M HNO₃ and then dried down. The samples and reference standard materials were digested using a mixture of 3 mL 15 M HNO₃, 1 mL 40% vol/vol HF, and 0.5 mL 70% vol/vol HClO₄ over 4 d on a hot plate at 150 °C. Solutions were evaporated to dryness at 230 °C, refluxed in a 1:3 mixture of 15 M HNO₃ and 6 M HCl (hydrochloric acid), evaporated again, and redissolved in 2 mL 0.5 M HNO₃.

Aliquots of 0.5 mL were used for concentration measurements using ICP-MS (inductively coupled plasma mass spectrometer), and the remaining volume of each solution was prepared for Pb isotope ratios measurement by MC-ICP-MS (multiple collector inductively coupled plasma mass spectrometer).

In addition, samples of unleaded gasoline and diesel were collected in two different petrol stations in London to determine the present-day concentration of Pb in fuel. The sample digestion procedure is described in *SI Appendix*.

Determination of Pb Concentrations, Deposition Rates, and EFs. Small amounts of digest solutions were available for samples S1 to S12, already analyzed for metal concentrations and Cu and Zn isotopes during a previous study (43). The Pb concentrations of samples S3 to S8 were not available, and they were analyzed during this study using an isotope dilution technique and MC-ICP-MS analysis (protocol in *SI Appendix*). All other PM₁₀ and TSP_{passive} samples and fuel samples were analyzed for Pb and metal concentrations after an adequate dilution in 1% HNO₃ by ICP-MS, either at Imperial College London or at the University of Montpellier. Concentrations were determined with external calibration using In and Bi as internal standards to correct potential sensitivity drifts. Digestion blanks represented less than 1% of Pb analyzed in the samples and less than 5% for other elements (Al, Ba, Cd, Cu, Cr, Fe, Ni, V, Zn, and Sb). The concentrations determined for the reference material BCR 723 were within 10% of the indicative values for all studied elements.

The weight of TSP_{passive} collected in the receptor dish using the Sigma-2 passive sampler device was too small (<5 mg) for precise weighing. Therefore, Pb concentration is expressed as a deposition rate (DR) corresponding to the elemental mass deposited on a specific surface per day (ng · m⁻² · d⁻¹) and calculated using Eq. 2 (23),

$$DR = W_{Pb} / (S \times t), \quad [2]$$

with W_{Pb} as the total mass of Pb recovered in the receptor dish (ng), S as the surface of the receptor dish (m²), and t as the collection time (in days).

Metal EFs in airborne particles (PM₁₀ and TSP_{passive}) were calculated (Eq. 3) using Al-normalized metal concentration and the upper continental crust (UCC) value as reference (57).

$$EF = (Me/Al)_{\text{sample}} / (Me/Al)_{\text{UCC}}, \quad [3]$$

with Me and Al representing the metal and aluminum concentrations (in μg/g).

Al and Si concentrations are highly correlated in airborne particles at the MR site, and the Si/Al ratio is very close to the ratio in average crustal material (22, 24). In addition, Si and Al concentrations at the MR site are similar to those observed at a background site in London (22, 24). This shows that Al has no significant traffic-related sources and is thus a good proxy of resuspended particles made of soil and other crustal material. Al is preferred as a geological proxy due to its more uniform distribution in geological material (aluminum silicates) than Ti or Sc. The UCC value was used as a reference to facilitate comparison with results from previous studies (9, 43), although it is important to note that it is not representative of the local geochemical background. An EF_{Pb} value of 3 was calculated using median concentrations

of topsoils ($n = 1,599$) sampled in rural areas in the London region showing a slightly higher regional background than UCC composition (49), but it is widely recognized that EFs <5 are not reliable to be interpreted as an anthropogenic enrichment.

Determination of Pb Isotope Ratios. Pb was separated from the sample matrix by ion exchange chromatography (Sr-Spec resin), following a procedure adapted from Weiss et al. (58) and described in the *SI Appendix, Table S5*. The isotope ratios were determined using a Nu Plasma multicollector ICP-MS (Nu Instruments) equipped with an Aridus II Desolvating Nebulizer System (Teledyne CETAC Technologies). The instrumental mass bias was corrected using an exponential law with Tl doping (58).

The total procedural blank was <75 pg Pb. To assess the precision and accuracy of Pb isotope-ratio measurements, analyses of the NIST SRM 981 (Standard Reference Material of the National Institute of Standards and Technology) and USGS G-2 reference materials were performed. Average values of ²⁰⁸Pb/²⁰⁶Pb = 2.16761 ± 52 (2σ), ²⁰⁶Pb/²⁰⁷Pb = 1.09301 ± 14 (2σ), ²⁰⁶Pb/²⁰⁴Pb = 16.9335 ± 53 (2σ), ²⁰⁷Pb/²⁰⁴Pb = 15.4925 ± 58 (2σ), and ²⁰⁸Pb/²⁰⁴Pb = 36.7052 ± 181 (2σ) were obtained for NIST SRM 981 standards treated as samples ($n = 51$) corresponding to an accuracy <45 ppm for ²⁰⁸Pb/²⁰⁶Pb and ²⁰⁶Pb/²⁰⁷Pb ratios and <450 ppm for ²⁰⁸Pb/²⁰⁴Pb ratios relative to reference values (59). Ratios measured for USGS G-2 reference material were ²⁰⁸Pb/²⁰⁶Pb = 2.11422 ± 49 (2σ, $n = 7$), ²⁰⁶Pb/²⁰⁷Pb = 1.17639 ± 26 (2σ), ²⁰⁶Pb/²⁰⁴Pb = 18.3949 ± 34 (2σ), ²⁰⁷Pb/²⁰⁴Pb = 15.6369 ± 44 (2σ), and ²⁰⁸Pb/²⁰⁴Pb = 38.8911 ± 144 (2σ), which are in good agreement with published data [²⁰⁶Pb/²⁰⁴Pb = 18.3992 ± 219 (2σ), ²⁰⁷Pb/²⁰⁴Pb = 15.6357 ± 38 (2σ), and ²⁰⁸Pb/²⁰⁴Pb = 38.9006 ± 154 (2σ) (60)].

Long-Term Dataset of Pb Isotopic Composition in London Airborne Particles.

New data determined in this study were interpreted in the context of the long-term evolution of Pb isotopic composition in London airborne particles using data published in the literature. Hamilton and Clifton (61) provided the first measurement of Pb isotopes in London airborne particles in 1968. Later, Monna et al. (32) and Véron et al. (62) measured Pb isotopes in PM₁₀ collected in London in 1995 and 1996, when leaded gasoline was still on the market. Noble et al. (20) and Dong et al. (9) reported Pb isotopic composition for PM₁₀ sampled at the MR site between 1998 and 2001 and in 2010, respectively. Literature also provided reference values for Pb isotopic composition of natural geogenic Pb in the United Kingdom (61), UK leaded gasoline (32, 63), UK ore (33), UK coal and imported coal (64, 65), UK lead paint (34), topsoils in London (31), and modern nonexhaust emissions from vehicle and road wear in London (9). All data are summarized in *SI Appendix, Table S3*.

Data Availability. All study data are included in the article and/or supporting information. Previously published data were used for this work (all references to previously published data by other authors used in the paper can be found in the section *Long-Term Dataset of Pb Isotopic Composition in London Airborne Particles* and in the legends of figures and tables wherever applicable).

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