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particulate matter in different Brazilian regions†

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Organophosphate esters (OPEs) in atmospheric

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Organophosphate esters (OPEs) have been widely used as plasticizers. Current studies have shown that these compounds may constitute atmospheric particulate matter (PM), but no research has focused on detecting

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Organophosphate esters (OPEs) have been widely used as plasticizers. Current studies have shown that these compounds may constitute atmospheric particulate matter (PM), but no research has focused on detecting these substances in Brazil. Hence, our research aims to determine OPEs concentrations in 26 PM₁₀ samples from three Brazilian cities in different regions. The substances found were tri-*n*-butyl phosphate (TNBP), tris(1,3-dichloro-2-propyl)phosphate (TDCIPP), triphenyl phosphate (TPHP), tris(2-butoxyethyl)phosphate (TBOEP), 2-ethylhexyl diphenyl phosphate (EHDPHP) and tris(2-ethylhexyl)phosphate (TEHP). In Catalão, the most abundant compounds were TNBP and TPHP, in Limeira TPHP and TEHP, and in Novo Hamburgo TEHP and TBOEP. Statistical analyses were performed to correlate the concentrations with meteorological variables. Results showed that temperature had a negative correlation with TBOEP, relative humidity had a negative correlation with TPHP and TEHP, atmospheric pressure had a positive correlation with TPHP, and wind speed had a significant influence on total and TPHP. Moreover, human exposure to these substances presents a negligible risk, for both children and adults.

Environmental significance

Atmospheric particulate matter (PM) is a pollutant that can cause damage to human health. Given this scenario, it is of extreme relevance to research PM concentration and composition, especially when it includes the identification of pollutants. Organophosphate flame retardants (OPFRs) are ubiquitous contaminants frequently detected in PM. However, no study evaluated their occurrence in PM samples from Brazil. Thus, it is necessary to determine the concentration of these compounds in the atmosphere to evaluate their principal emission sources and risks. In this study, PM samples were collected in three different regions in Brazil and several OPFRs were frequently detected, but at levels that do not pose a risk to humans. Also, the influence of meteorological variables and anthropogenic activities was assessed.

1 Introduction

Atmospheric particulate matter (PM) can cause major damage to human health, especially particles with smaller diameters, since the damage is inversely proportional to their size. One fraction that deserves attention is PM_{10} , a PM with an aerodynamic diameter less than or equal to 10 μm . Considered inhalable, PM_{10} is retained in the upper airways thus causing damage to human health, such as acute and chronic respiratory infections, aggravated asthma, difficulty breathing, as well as heart disease, which can lead to premature death.

PM composition is quite varied, especially regarding its organic fraction, due to the high number and diversity of

emitting sources.^{5–7} Organophosphate esters (OPEs), which are widely used in the industrial sector and in the manufacturing of household products to prevent the spread of fire, have been widely detected in PM.^{8,9}

OPEs emerged as an alternative to polybrominated diphenyl ethers (PBDEs), which were characterized by persistence and bioaccumulation. However, knowledge advancements on the environmental behavior and toxicity of these compounds indicated hazard to humans and biota, thus justifying the recent significant increase in studies focusing on these substances.

OPEs can present toxic¹² and carcinogenic¹³ characteristics and can cause adverse effects to the immune¹⁴ and neurological^{15,16} systems, altered sex hormone production,¹⁷ lung epithelial cell death,¹⁸ among other damages to human and animal health. Moreover, these compounds can undergo atmospheric chemical transformations that further increase the risks they can cause to the population's health.¹²

In addition to the atmosphere,^{19,20} these substances have also been detected in other environmental matrices, such as water,^{21,22} sewage,^{23,24} sediment^{25,26} and soil.^{17,27} Such

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widespread presence in different matrices increases human exposure to OPEs, thus justifying the various studies being conducted on the risks posed to individuals.

In Brazil, there is still a significant shortage of studies on OPES in particulate matter. Given this situation, this study determined the OPE concentration in PM_{10} samples from the cities of Catalão, Limeira and Novo Hamburgo using GC-MS, identifying their possible sources, and evaluating the human exposure to these compounds. To the best of our knowledge, this is the first study on the occurrence and risk of human exposure to OPEs in PM in these 3 Brazilian cities.

2 Methods

2.1. PM sampling sites and methodology

Sampling was performed in three Brazilian cities, located in different regions: Catalão (CAT) – Goiás state, Limeira (LIM) – São Paulo state, and Novo Hamburgo (NH) – Rio Grande do Sul state; Midwest, Southeast, and South regions, respectively (Fig. 1). Catalão had an estimated population of 113 091 inhabitants in 2021 (ref. 28) and is characterized by mining, agriculture, and cultivated pastures.²⁹ Limeira had a population of 310 783 inhabitants in 2021 (ref. 28) and its economy is based primarily on the semi-jewelry industry and sugar cane and citrus plantations.³⁰ Novo Hamburgo had approximately 247 303 inhabitants in 2021 (ref. 28) and has a great diversity of industries, most notably the footwear sector.³¹ These three cities were selected because they are from different regions, have different meteorological characteristics, different possible emission sources and no previous studies on identifying OPEs in PM.

Twenty-six PM $_{10}$ samplings were conducted between June and July 2019 at the Federal University of Catalão (coordinates $18^{\circ}09'17.3''S$; $47^{\circ}55'38.7''W$), the School of Technology at the University of Campinas (coordinates $22^{\circ}33'43.81''S$; $47^{\circ}25'20.84''W$), and Feevale University (coordinates $29^{\circ}40'12.4''S$; $51^{\circ}07'15.7''W$), Novo Hamburgo.

All samples were collected using PM_{10} high volume samplers (Hi-Vol), with a constant flow of 1.13 $m^3~min^{-1}$ and the total sample volume of 1627.2 m^3 , always starting at midnight of one day and ending at midnight of the next day to complete a 24 h cycle. Fiberglass filters, porosity < 1 μm , pre-treated in a muffle furnace at 400 °C for 4 h to remove possible contaminants, were used for PM_{10} collection. Later, to remove moisture, they were kept in a desiccator for 24 h before sampling and only then weighed. After sampling, the filters remained for additional 24 h in the desiccator and were weighed again to obtain their mass by gravimetric analysis.

The blank filters were also kept in the muffle furnace and desiccator under the same conditions as the filters that were used for collection, but they remained 24 hours in a Hi-Vol with the device turned off.

2.2. Materials and reagents

We analyzed nine OPEs in total, including tri-*n*-butyl phosphate (TNBP), tris(chloroethyl)phosphate (TCEP), tris(1-chloro-2-propyl)phosphate (TCIPP), tris(1,3-dichloro-2-propyl)phosphate (TDCIPP), tris(2-butoxyethyl)phosphate (TBOEP), tris(2-ethylhexyl)phosphate (TEHP), triphenyl phosphate (TPHP), 2-ethylhexyl diphenyl phosphate (EHDPP) and trimethyl phenyl phosphate (TMPP). High purity analytical standards were acquired from Dr Ehrenstorfer (Augsburg, Germany) and the degree of purity of the standards is shown in Table S1.† Triphenyl phosphate-D15 (TPHP-D15) and tributyl phosphate-D27 (TNBP-D27) were purchased from Cambridge Isotope Laboratories (Andover, USA). HPLC grade hexane and acetone solvents were purchased from Honeywell Riedel-de-Haën (USA).

2.3. Sample preparation and extraction

For OPEs extraction, one eighth of the PM₁₀ collected area in each filter was chopped into small pieces and transferred to a glass centrifuge tube with a lid. Each sample was spiked with

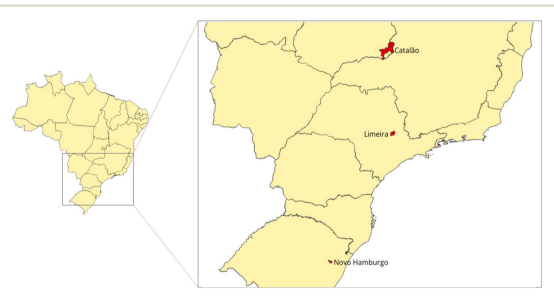


Fig. 1 Location of cities where PM_{10} samples were collected.

 $10~\mu L$ of a solution containing the deuterated surrogates TPHP-D15 and TNBP-D27 at a concentration of $5~ng~\mu L^{-1}$. After adding 10~mL of hexane : acetone (1:1) to each tube, they were agitated in a vortex for 1 min, kept in an ultrasonic bath for 10 min, and centrifuged at 3500 rpm for 10 min. Each sample went through this complete cycle three times, but using 7 mL of hexane : acetone (1:1) instead of 10 mL in the second time and third time.

The extracts were then concentrated at a rotary evaporator until their almost complete evaporation. Clean-up used 1 g (6 mL) Florisil Cartridge (Supelco SupelcleanTM) conditioned with 12 mL of hexane: acetone (1:1). After removal from the rotary evaporator, the volumetric flask with the sample extracts was rinsed with two 1 mL portions of hexane: acetone (1:1), and the redissolved extract was transferred to Florisil Cartridges. Elution of the cartridges was performed using 6 mL of hexane: acetone (1:1). Subsequently, the samples were concentrated using ultrapure nitrogen gas flow (99 999%) in a water bath at 35 °C, until approximately 500 μL remained. The extracts were then transferred to 2 mL vials. For complete transfer of the analytes, the vial containing the extract was rinsed with two 400 μ L portions of hexane : acetone (1:1). Finally, the extracts were completely evaporated using nitrogen gas flow, and redissolved with 500 µL of ethyl acetate.

2.4. Quality assurance (QA) and quality control (QC)

To ensure quality control, powder-free nitrile gloves were used to handle the samples throughout the process. Some glassware used, such as Pasteur pipettes, vials, and glass centrifuge tubes, were muffled for 12 h at 400 °C, and the others were washed with 4% Extran® MA02 neutral detergent (Merck), and finally rinsed with acetone (99.5%). Scissors and rulers were cleaned with acetone between cutting one filter and another. For each batch, a procedural blank was performed using a glass filter and the same extraction process, to evaluate any possible contamination.

Recovery assays were performed by preparing 3 fortified samples (spiked before extraction with 50 μL of a solution containing 9 OPEs at concentration of 4 ng μL^{-1} and with 10 μL of surrogate solution at concentration of 5 ng μL^{-1}), 2 reference samples (spiked after extraction with the same amounts of standard solution and surrogate solution as described above) and 2 procedural blanks (spiked before extraction with 10 μL of surrogate solution). Since OPEs could be present in the PM samples used in the recovery assay, reference samples were used as a parameter for calculating recoveries, and the procedural blanks were used to evaluate selectivity and sample contamination during the extraction procedure. Extraction was performed as described in Section 2.2.

Analytical curve was obtained by preparing seven solutions at different concentrations, ranging from 0.005 to 0.5 mg L $^{-1}$, using procedural blank filter extracts. At each calibration solution, 10 μ L of a surrogate solution at a concentration of 5 ng μ L $^{-1}$ (TNBP-D27 and TPHP-D15) were added. Surrogate TNBP-D27 was used to quantify the compounds TNBP, TBOEP, EHDPHP and TEHP, whereas the surrogate TPHPD15 was used

to quantify the compounds TCEP, TCIPP, TDCIPP, TPHP and TMPP.

Table S1† shows the method detection limits (MDL), method quantitation limits (MQL), recovery (REC), and standard deviation (SD) for each compound. MDL and MQL were calculated as 3 and 10 times the signal-to-noise ratio obtained for the lowest calibration solution, except for the compounds EHDPHP, TCIPP and TPHP, which were detected in the procedural blanks. For these compounds, MDL was calculated based on the mean blank level plus 3 times the standard deviation and the MQL was calculated as the mean blank level plus 10 times the standard deviation.

2.5. Instrumental analysis

All analyses were performed on a Trace GC Ultra Gas Chromatograph (Thermo) coupled to an ITQ900 mass spectrometer (Thermo) with ion trap mass analyzer. The injector was a programmable temperature vaporization (PTV). The glass liner used was the PTV Straight Liner (120 \times 2.75 mm \times 1 mm, Thermo). Solvent split injection mode was used, with a 5 µL injection volume, as reported by Cristale et al. 2020.32 The sampler used was automatic (Triplus). The column was a Zebron ZB5-MS (30 m \times 0.25 mm \times 0.25 μ m) (Phenomenex, USA). The carrier gas was helium with a 1.5 mL min⁻¹ flow rate. The column oven was held at 60 °C (6 min), subsequently heated at a 10 °C min⁻¹ rate until it reached 230 °C (1 min), and finally heated at a rate of 20 °C min⁻¹ until 280 °C (14 min). Injections were performed in SIM and MS/MS mode under the detection conditions shown in Tables S2 and S3.† Compounds were quantified using the SIM mode, as it showed better accuracy than the MS/MS. The results obtained using MS/MS mode were used as criteria for confirming the presence of OPEs in the sample extracts.32 TCEP and TCIPP were quantified by MS/MS, whereas TNBP, TDCIPP, TPHP, TBOEP, TEHP, TMPP and EHDPHP were quantified by SIM.

2.6. Analysis of OPEPs concentration in relation to meteorological variables

Pearson's correlation analysis was performed comparing the OPEs data detected with the PM_{10} concentration and the meteorological data (temperature, relative humidity, wind pressure, and wind speed, all variables on the logarithmic scale)³³ obtained from the Meteorological and Monitoring Stations of each location. Correlation analysis was performed using the R programming language version 3.5.1.³⁴

2.7. Health risk assessment

Risk related to human exposure to OPEs was assessed by calculating the Estimated Daily Intake (EDI) *via* PM inhalation using eqn (1).

$$EDI = \frac{C \times IR \times t}{BW}$$
 (1)

in which C is the concentration of each OPEs in the PM₁₀ samples (ng m⁻³), IR is the inhalation rate (m³ d⁻¹),^{33,35} t is the outdoor exposure time (day), and BW is the average body weight

(kg). IR was set as 10 m³ d⁻¹ for children and 16 m³ d⁻¹ for adults,³6 the value of t assumed was 1 day, i.e., 100% of the time, assuming similar indoor and outdoor concentrations, and BW is considered as 70 kg for adults and 12 kg for children.³7 Hazard Quotient (HQ) was calculated according to eqn (2), in which RfD is the reference dose value of each OPEs. HQ \geq 1 indicates risk of adverse health effects.³4

$$HQ = \frac{EDI}{RfD}$$
 (2)

3 Results and discussion

3.1. OPEs concentration in the PM

We found mean PM_{10} concentrations of 28 $\mu g \ m^{-3}$ for Catalão, 40 $\mu g \ m^{-3}$ for Limeira, and 20 $\mu g \ m^{-3}$ for Novo Hamburgo, ranging from 13–51 $\mu g \ m^{-3}$, 6.6–70 $\mu g \ m^{-3}$, and 11–33 $\mu g \ m^{-3}$, respectively (Table S4†). Catalão city and the region where Novo Hamburgo is located have industrial activities and vehicular traffic as significant sources of PM emission. ^{38,39} In Limeira, intense vehicle traffic, industrial emissions and biomass burning are important contribution sources. ⁴⁰ The OPEs found in the samples analyzed were TNBP, TDCIPP, TPHP, TBOEP, EHDPHP, and TEHP. While TCEP and TCIPP were probably not found in the PM as, according to the gas-particle partitioning models, these compounds should be predominantly present in the gaseous phase. ⁴¹ Fig. 2 shows the concentrations; those at levels >MDL and <MQL were represented by half of the MQL value.

TNBP was detected only in samples from Catalão, compound with the highest concentrations in the locality, ranging from

0.19 to 1.3 ng m⁻³ (mean of 0.55 ng m⁻³). This compound has also been detected in PM in countries such as China, with mean concentration of 1.02 ng m⁻³, 18 and the United States, with 0.56 ng m⁻³. 42 TNBP can come from vehicle emissions, as it is used as a fluid in motor oil, 12,43 as well as in paints. 44

TPHP was also identified in all samples from the municipality, with a mean concentration of 0.17 ng m $^{-3}$. Other studies have found mean PM concentrations close to this value in different places worldwide, such as Poland 45 and China 46 with 0.11 ng m $^{-3}$, the United States with 0.2 ng m $^{-3}$ (ref. 42) and Pakistan with 0.17 ng m $^{-3}$. This compound can have, as some of its main emission sources, solid waste combustion, building materials, polymer additive and hydraulic fluids. 43,45

The high concentrations of TNBP and TPHP found in Catalão are associated with the presence of a factory of precast concrete-based items, widely used in civil construction, that is located less than 1 km northwest of the sampled site. To the southeast, about 2.5 km away, a mining-chemical industrial district⁴⁸ receives a daily high flow of heavy vehicles for loading and unloading inputs for manufacturing phosphate fertilizers. To the south/southeast, there is an industrial center with largescale production of concrete, light automobiles, agricultural machinery, and production of paints from resins.

In Limeira, TPHP is the only compound to appear on all sampling days and present the highest concentrations, with a mean of 0.55 ng m⁻³ (range from 0.10 to 0.90 ng m⁻³). TEHP appears in all samples, with one exception, having a mean concentration of 0.10 ng m⁻³. Other research has found similar concentrations for this compound, such as in China with 0.106 ng m⁻³,⁴⁹ Pakistan with 0.156 ng m⁻³ (ref. 47) and Spain with 0.12 ng m⁻³.⁵⁰ This substance is commonly used as a plasticizer, in PVC, cellulose, lubricant, rubber and foam, and solvent.^{12,51}

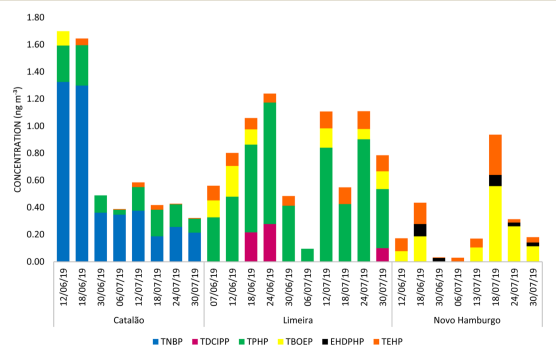


Fig. 2 OPEs concentration in PM₁₀ from Catalão, Limeira and Novo Hamburgo.

Table 1 Correlation coefficient of OPEs with meteorological factors. including temperature, relative humidity, wind pressure and wind speed, and PM₁₀ concentration

	TNBP	TPHP	TBOEP	TEHP	ΣOPEs
PM_{10}	0.071	0.765^{a}	-0.161	0.387	0.608^{a}
Temperature	0.047	-0.063	-0.312	-0.025	0.240
Relative humidity	-0.048	-0.739^{a}	-0.429	-0.833^{a}	-0.281
Pressure	-0.357	0.483^{b}	0.478	0.136	-0.388^{c}
Wind speed	-0.241	-0.709^{a}	-0.046	-0.457^{b}	-0.624^{a}

^a Means correlation was significant at 0.01 level. ^b Means correlation was significant at 0.05 level. ^c Means correlation was significant at 0.10 level.

Specifically in Limeira, TEHP concentration may have tire wear as its main source, considering that the sample site is located near a roundabout with high vehicular traffic.

The two predominant compounds in Novo Hamburgo are TBOEP and TEHP, with means of 0.22 and 0.10 ng m⁻³, respectively. TBOEP has also been detected in PM in Spain with a 0.21 ng m⁻³ concentration⁵⁰ and in China with 0.37 ng m⁻³.⁵² This substance is widely used in plastic and rubber products, wax, grease, paint, coating, and can have road traffic as an emission source. 43,51 The sampling site is located near a federal highway and a state highway, as well as urban roads with heavy vehicular traffic.

3.2. Correlation between OPEs concentration and meteorological variables

PM₁₀ concentration is directly correlated with the total number of OPEs and TPHP (Table 1).

Temperature was not significantly correlated with any OPEs. The temperature variation in Catalão was 4.6 °C, in Limeira 12.1 °C and Novo Hamburgo 15 °C during the collection period.

Conversely, relative humidity showed a significant negative correlation (p < 0.001) with TPHP and TEHP. Higher TEHP values occurred on days when relative humidity was below 35%, whereas higher TPHP values occurred on days when humidity was below 40%.

TPHP showed a positive correlation with atmospheric pressure. However, it was not identified in Novo Hamburgo (mean pressure of 762 mmHg). Between the cities of Limeira (mean of 708 mmHg) and Catalão (mean of 689 mmHg), the former had the highest TPHP values.

As seen in Table 1, wind speed had a significant influence on total (p < 0.001) and TPHP (p < 0.001), as observed for PM₁₀. However, while PM₁₀ correlation with total and TPHP was positive, with wind speed, it was negative. This is because the highest PM₁₀ concentrations, and hence TPHP, are associated with low wind speeds, less than 0.5 m s⁻¹. Low wind speed, low relative humidity, and low height of the mixing boundary layer hinder pollutant dispersion.40 TEHP also showed a significant negative correlation (p < 0.05) with wind speed, differently from the correlation between PM₁₀ and TEHP.

3.3. Human exposure assessment

Table 2 presents the minimum, maximum, and mean values of the calculated EDI and HQ. Mean EDI for adults ranged from 0.009 to 0.13 ng kg per bw per day considering all municipalities. For children, mean EDI ranged from 0.033 to 0.46 ng kg per bw per day. Note that EDIs are higher for children than for adults due to lower body mass. The maximum HQs found illustrate the worst case scenario among the three cities and are well below 1, ranging from 1.5 \times 10⁻⁵ to 10.7 \times 10⁻⁵ for children and 0.42×10^{-5} to 3.0×10^{-5} for adults, indicating negligible health risks due to exposure to PM₁₀-associated OPEs in Catalão, Limeira, and Novo Hamburgo. Other studies have also concluded that exposure to these compounds through

Table 2 Reference doses (RfD, ng kg⁻¹ d⁻¹), estimated daily intake (EDI, ng kg per bw per day) and hazard quotient (HQ) of OPEs for children and adults by inhalation exposure^a

Compound	RfD	EDI Catalão		EDI Limeira		EDI Novo Hamburgo		HQ (×10 ⁻⁵)	
		Child Mean (range)	Adult Mean (range)	Child Mean (range)	Adult Mean (range)	Mean (range)	Adult Mean (range)	Child Maximum	Adult Maximum
TDCIPP	15 000	_	<u> </u>	0.17 (0.084-0.23)	0.045 (0.023-0.063)	_	_	1.5	0.42
TPHP	70 000	0.14 (0.031–0.25)	0.039 (0.008-0.068)	0.46 (0.080-0.75)	0.13 (0.022-0.21)	_	_	11	3.0
TBOEP	15 000	0.088 (—)	0.024 (—)	0.11 (0.064-0.19)	0.031 (0.017-0.052)	0.18 (0.066-0.47)	0.050 (0.018-0.13)	3.1	0.85
EHDPHP	_	_	_			0.072 (0.068-0.076)	0.020 (0.019–0.021)	_	_
TEHP	_	0.033 (0.029–0.041)	0.009 (0.008-0.011)	0.085 (0.055-0.11)	0.023 (0.015-0.030)	0.084 (0.020-0.25)	0.023 (0.006-0.068)	_	_

^a Ref. 18.

atmospheric particulate matter does not represent a health risk, considering the current reference values. 18,33

4 Conclusions

Our study showed the occurrence of OPEs in atmospheric PM_{10} from three Brazilian cities: Catalão, Limeira, and Novo Hamburgo. OPEs profile and concentration differed between the cities, indicating that the anthropic activities characteristic of each city and the meteorological variables influenced the presence of OPEs in the local atmosphere. Despite the high detection rate of various OPEs observed here, based on currently practiced reference values for non-carcinogenic risk, the levels found do not pose a threat to human health.

Author contributions

Priscila B. Gonçalves: data curation; formal analysis; investigation; methodology; validation; visualization; writing – original draft, writing – review & editing. Joyce Cristale: formal analysis; methodology; validation; writing – original draft. Amanda A. da Silva: formal analysis; methodology. Danilo C. Nogarotto: data curation; software. Daniela M. M. Osório: methodology; data curation. Lincoln L. Romualdo: methodology; data curation. Simone A. Pozza: conception; funding acquisition; investigation; methodology; project administration; resources; supervision; validation, visualization, writing – review & editing.

Conflicts of interest

The authors declare that they have no knowledge of competing financial interests or personal relationships that could have influenced the work reported in this paper.

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