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Simultaneous measurement of nitrous acid, nitric acid, and nitrogen dioxide by means of a novel multipollutant diffusive sampler in libraries and archives

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Abstract

Introduction: A novel multipollutant diffusive sampler for HONO, HNO₃, and NO₂ was used and tested at four different libraries/archives in different seasons. Two were located in Switzerland in Bern (Swiss National Library) and Geneve (Bibliotheque de Geneve), both with HVAC system equipped with filters for pollutants removal, and the other two in the Czech Republic in Teplice (Regional Library) and in Prague (National Archives), where in this case the former is naturally ventilated and whereas the latter is equipped with HVAC system with filtration.

Results: The ratios between indoor and outdoor concentrations of NO_2 showed a greater penetration of pollutants indoors in the naturally ventilated library than in the filtrated archives. The indoor concentrations of HNO_3 were very low probably due to the high deposition velocity of nitric acid on available surfaces. HONO concentration values were usually lower outdoors, which indicated that HONO was produced by reactions on indoor surfaces.

Conclusion: The results revealed that the reproducibility of the new multipollutant sampler measurements was reasonable (according to EU directives) for NO_2 and HNO_3 and that the newly developed multipollutant sampler can be used in archives and libraries, allowing to map the pollutants distribution indoors. Due to high efficiency of insulating systems normally employed and of filtration systems, the values recorded for the pollutants indoors are often lower than the detection limits.

Keywords: Diffusive sampling, Nitrous acid, Nitric acid, Nitrogen dioxide, Archive, Library, Indoor/outdoor ratio

Background

Nitrogen oxides penetrate from the outdoor to the indoor environment and can be transformed, through complex reaction pathways, into gaseous nitric (HNO₃) and nitrous acids (HONO) [1]. HNO₃ is a very aggressive acid in contrast with HONO. The latter is not particularly active by itself, even if it is an important source of OH radicals, one of the most important air oxidants [2]. This air pollution cause oxidation and hydrolytic degradation of archive materials such as colour changes and reduction of degree of polymerisation of paper [3], decomposition

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of leather [4], corrosion of metals [5, 6], and damage of colorants [7–10].

Several works report measurements of nitrous and nitric acid in various indoor environments [11–13] and recent works highlight the importance of the former species as an emerging indoor pollutant [14], but studies in cultural heritage buildings are scarce [15, 16]. A well established and known technique to collect these trace gases is represented by denuder sequence based systems [17–19], but this technique is time consuming and labour intensive. Diffusive sampling can overcome these problems since it represents an easy to use technique which exploits the spontaneous diffusion of species collected by specific absorbing media. Diffusive samplers have been used for air quality monitoring of single gases in cultural

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heritage buildings [20-23]. Nevertheless, when conditions which promote the formation of HONO are present (high air moisture content and surface to volume ratio), especially indoors, monitoring of NO₂ by diffusive sampling can result in overestimation since the interference of the former pollutant is not negligible [24]. Therefore to overcome this problem and to account correctly for the different species a novel multipollutant diffusive sampler was designed for simultaneous sampling of three different pollutants (nitric acid, nitrous acid, and nitrogen dioxide) collected at separate sampling stages [25]. After the laboratory development, these devices were used indoors in different dwellings, as reported in [25]. The campaigns described in the following were the first applications to places of interest for cultural heritage conservation, as libraries and archives, where a certain kind of control on environmental parameters is commonly present. The aim of this study was, indeed, to test the newly developed multipollutant sampler for evaluating the air quality inside different libraries with or without HVAC (heating and ventilating and air conditioning systems). The exposure period, after these first sampling trials, could then be adjusted according to the typical range of values found for the different pollutants in the indoors of interest. These preliminary trials were necessary to avoid the saturation of the absorbing pads and the collection of insufficient analyte to be determined after the exposure period. Furthermore the sampler was also exposed outdoors to compare the pollutants concentration values and calculate indoor/outdoor ratios.

Methods

Sampling locations

Over the last 4 years the multipollutant sampler was used in different periods inside archives and libraries. Two seasonal campaigns were carried out at four different libraries: two libraries in Switzerland (the Swiss National Library in Bern and the Bibliotheque de Geneve in Geneva) and two libraries in the Czech Republic (the National Archives in Prague and the Regional Library in Teplice). All the indoor activities in the libraries are very limited.

The library in Bern is a modern building composed of a ground floor and four underground levels (1UG-4UG), the building is equipped with a heating, ventilation, and air conditioning (HVAC) system providing air filtered through a particulate filter and active carbon. An additional Purafil purification system is installed at the floor 3UG where newspaper collection is stored. Five indoor sites were selected at the different levels and an outdoor site (OUT) was chosen as well. One site was placed at the ground floor (A26) and the other four monitoring sites were placed both at the entrance, just outside of the double door insulation (WA2UG and WA3UG), and inside the second and third underground level (2UG and 3UG) in the area insulated by the double door system. The overall volume of the building in Bern is of approximately 4500 m³.

The library in Geneva is a historical building with a HVAC system equipped with particulate filters. Three indoor sampling sites were selected: the first in a compactor (number 51 in the following), the second at "La Reserve" and the third in a corridor. An outdoor site was chosen also in Geneva where the diffusive samplers were placed in a balcony (OUT). The library in Geneva has a volume of approximately 200 m³.

The archive in Prague is located in a modern building. The indoor measurements were carried out in a depository of "Bohemian tables (BT)" and in a depository of "Archives of the Czech Kingdom (ACK)". The BT and ACK depositories have volumes of approximately 550 and 220 m³, respectively. Each depository is equipped with its own HVAC system with air recirculation. The HVAC systems in both depositories contain HEPA filters for removing airborne particles. In addition the AKC depository is equipped with filters for removing NO₂ and SO₂. The outdoor measurements were performed on a balcony of the building orientated toward a busy street (OUT).

The library in Teplice is equipped with double glassed windows and an electrical heating system, i.e., the only ventilation is through cracks and small openings in the building, windows and doors. The measurements were performed inside the library (IN) and just outside of the window oriented to the chateau park (OUT). The depository has a volume of approximately 450 m³.

One monitoring location in each indoor and outdoor site was selected, because other measurements confirmed that the indoor air is well mixed [26].

Measurement campaigns

Seasonal campaigns were conducted in Switzerland during the spring/summer period of 2011 (S1) and during winter and spring/summer 2012 (W, S2). Later other measurements were performed in the Czech Republic during three campaigns in spring/summer 2013 (S1, S2) and winter 2014 (W) (Table 1).

Three multipollutant samplers were exposed indoors and three outdoors for during each campaign at every location.

In parallel the exposure of single stage Analyst diffusive samplers (Marbaglass, Italy) provided measurements of NO_2 and NO_x during the campaigns in Switzerland. Other species (HNO₃, SO₂, O₃ and NH₃) were also monitored by the Analyst samplers in the campaigns carried out in the Czech Republic, where in addition IVL

City	Location	Sites	Season	Campaign	Exposition period	Other pollutants measured
Bern	Swiss National Library (SNL)	1 level: site A26 2 level: sites 2UG; WA2UG 3 level: sites 3UG; WA3UG Outdoor: Balcony	Spring/Summer Winter	S1 W	10.06.2011–30.06.2011 16.01.2012–09.02.2012	NO ₂ ; NOx (Analyst)
			Spring/Summer	S2	31.05.2012-21.06.2012	
Geneva	Bibliotheque de Geneve (BdG)	Site 1: Compactor 51 Site 2: La Reserve Site 3: Corridor Outdoor: Balcony	Spring/Summer	S1	09.06.2011-23.06.2011	NO ₂ ; NOx (Analyst)
			Winter	W	23.01.2012-14.02.2012	
Prague	National archives	Site 1: BT Site 2: ACK Outdoor: Window	Spring/Summer	S1	8.4.2013-10.5.1013	HNO ₃ , SO ₂ , O ₃ and NH ₃
				S2	10.5.2013-11.6.2013	(Analyst)
			Winter	W	6.1.2014-6.2.2014	Acetic aciu, ionnic aciu (IVL)
Teplice	Regional library	Site 1 Outdoor: Window	Spring/	S1	9.4.2013-9.5.2013	HNO ₃ , SO ₂ , O ₃ and NH ₃
			Summer	S2	9.5.2013-10.6.2013	(Analyst)
			Winter	W	7.1.2014-12.2.2014	

Table 1 Monitoring scheme

diffusive samplers (Swedish Environmental Research Institute, Sweden) for monitoring formic and acetic acids were employed.

In both cases at least 10 % of the total number of samplers was used for each campaign as field blanks. The blanks were prepared and handled along with other samplers except for the exposition to pollutants. Additionally, basic meteorological parameters including ambient temperature and relative humidity were measured at both locations in Czech Republic, both indoors and outdoors, by Tiny Tag data loggers (Gemini, UK).

Multipollutant diffusive sampler

The body of the multipollutant diffusive sampler is similar to the Analyst [27]. The internal design of the sampler, on the other hand, was changed to collect HNO_3 , on the first filter and NO_2 and HONO in the successive absorbing pads.

A correct speciation of these compounds, intertwined through the hydrolysis mechanism according to the known pathway [28–30]:

$$2NO_2 + H_2O \rightarrow HONO + HNO_3$$
 (1)

could, indeed, only be achieved by the subsequent collection of the two species on successive reactive substrates.

Most frequently NO_2 , which is the predominant species in the atmosphere also due to the photolysis of HONO in the outdoor ambient air, can affect the measurements of the less abundant HONO. On the other hand, particularly indoors, when moisture and an elevated surface/ volume ratio is present, NO_2 measurements can be overestimated as reported in [24]. Therefore there is a mutual interference on the measurement of these two species. In this case, since nitrite is the species collected and analytically determined to get to the atmospheric concentration of these pollutants, the selectivity in the sampling phase was achieved through the implementation of successive sampling stages. After the first filter used to collect HNO₃, other two successive pads were used to collect these species, assuming that HONO and part of the NO₂ would both react on the second, whereas only NO₂, normally the most abundant, is collected on the successive filter. During the development of this diffusive sampler trials, aimed at quantifying the relative amounts of nitrite collected on the second and third filter at different HONO/NO₂ concentration ratios, were performed. The ratio of the amounts of analyte collected on the two filters at RH = 0 was determined and used in the calibration of the device.

Hence three polyethylene discs housing three active glass microfiber filters were placed along the same diffusive path of the former sampler. The first two filters perform the sequential collection of the two acidic species and the third filter is used to estimate the interference of NO₂ on the HONO measurement and vice versa.

Before impregnation with the reactive solution, a washing procedure of the microfiber filters with sodium carbonate (5 %) and successive rinse (for three times) with deionized water was implemented to improve the quality of the blanks. The filters (Whatman GF/A) inside the first polyethylene disc were then coated by using a 0.1 % (w/v) aqueous solution of sodium chloride then dried in an oven at 80 °C.

The following other two glass microfiber filters, used to collect HONO and to account for the interference of NO₂, were impregnated by using a 1.8 % (w/v) aqueous solution of sodium carbonate containing 1.8 % (w/v) of glycerine, they were then dried in an oven at 80 °C. Filters are placed along the diffusive path in front of the alkaline carbon filter prepared according to the procedure already described [27] and positioned at the bottom of the device for the sole NO_2 sampling. An enlarged scheme of the sampler is reported in Fig. 1.

Chemical analysis procedure

As for the other diffusive samplers, the multipollutant samplers at the end of collection time were sealed and successively analysed according to a procedure similar to the one reported elsewhere for NO_2 [27]. In this case the filters were removed from the sampler, then they were extracted by adding a solution of sodium bicarbonate and carbonate, 0.3 mM $\rm NaHCO_3$ and 2.7 mM $\rm Na_2CO_2$ in a plastic vial. The solution obtained was stirred with VIBROMIX 203 EVT (from Tehtnica, Železniki, Poland) and analysed through ion chromatography (IC) (Dionex ICS 1000 equipped with AS12A column. The concentrations of analytes such as nitrate and nitrite were determined referring to calibration curves constructed with water solutions prepared by opportune dilution of stock standards (Certipur from Merck, Milan, Italy) containing 1000 mg/L of each analyte.

Results and discussion

Multipollutant diffusive sampler calibration

Some information will be given concerning the calibration of the new device performed during the laboratory trials in the development phase. The prototypes of the new sampler were exposed to standard atmospheres containing the pollutants of interest in a smog chamber. The



Fig. 1 Exploded scheme of the multipollutant sampler. *1* Plastic cap (before and after exposition) or air barrier (during exposition), *2* Ring, *3* Polyethylene holder of absorbing pad impregnated with NaCl (HNO₃ sampling), *4* Polyethylene holder of absorbing pad impregnated with Na₂CO₃ (HONO sampling), *5* Polyethylene holder of absorbing pad impregnated with Na₂CO₃ (HONO sampling), *5* Polyethylene holder of absorbing pad impregnated with Na₂CO₃ (interference of HONO and NO₂), *6* Carbon Filter (NO₂ sampling), *7*, *8*, *9* Plastic spacer, *10* Body of the sampler, *11* Absorbing pad (containing Na₂CO₃ or NaCl). *Geometric features A* cross sectional area (330 mm²), *R* maximum length of the plastic support (10 mm), *r* plastic support radius (6.5 mm), *a* diffusive path to NaCl impregnated filter (11.3 mm), *c* diffusive path to second Na₂CO₃ impregnated filter (17.1 mm), *d* diffusive path to impregnated carbon filter (24.0 mm)

experimental setup was equipped with reference technique for the measurement of each species, the data were used to calibrate the new device. The calibration was achieved through comparison with denuder data: NaCl coated denuders were the reference for HNO₃, whereas two Na₂CO₃ coated denuders [31] accounted for nitrous acid.

Successively the comparison between the two different techniques gave good results also in field trials (for HNO₃ $R^2 = 0.97$; for HONO $R^2 = 0.96$). The chemiluminescent analyser, used as a reference for NO₂, was also in agreement with the newly developed diffusive sampler ($R^2 = 0.97$).

A paired *t* test was performed to compare the data obtained by denuder technique and diffusive sampling [HONO: $t(10) = 0.31 \ p = 0.75$; HNO₃: $t(10) = 2.18 \ p = 0.053$], and by chemiluminescent analyzer and diffusive sampling [NO₂: $t(10) = 1.59 \ p = 0.14$]. At a *p* value of 0.05, in all cases the results are consistent with the null hypothesis that there is no difference between techniques would be accepted, and certainly for HONO there is very good agreement; for HNO₃ more measurements might in retrospect have been appropriate to form a more considered view.

A comparison with the Analyst sampler for NO_2 was also performed. The Analyst differs from the multipollutant sampler for the geometry, since it is basically a cylinder open on one end, whereas the multipollutant has a rather complex structure (Fig. 1). The uptake rate (*Ur*), which is defined as:

$$Ur = \frac{D \cdot A}{L} \tag{2}$$

where *D* is the diffusion coefficient (cm²/s) of the gaseous species, *A* is the cross-sectional area (cm²) and *L* the diffusive path (cm), is quite different for the two samplers. The ratio between the uptake rate for NO₂ for the Analyst and the multipollutant is 3, being the $Ur_{Analyst}$ about 12 cm³/min and $Ur_{Multipollutant}$ about 4 cm³/min (measured at RH = 0 %, in a dry atmosphere containing NO₂ without HONO). This feature of the newly developed diffusive sampler can explain the lower sensitivity to NO₂ if compared to the Analyst, since in the same lapse of time the latter can collect a higher amount of analyte (nitrite), on the other hand the selectivity of multipollutant sampling is increased as already explained.

In field use of Analyst and multipollutant sampler

As a first approach, the agreement of the measurements of NO_2 obtained by the Analyst sampler and by the multipollutant sampler was checked (Fig. 2). The correlation coefficient was quite good ($R^2 = 0.87$), taking into account the whole set of measurements (both Swiss and Czech libraries and archives).



The new multipollutant samplers and standard Analyst samplers were also compared by estimation of sensitivity given as limits of detection (LODs) and uncertainty given as relative standard deviations. The LODs were calculated as three times the standard deviation of blank samples [32]. The sensitivity of multipollutant and Analyst samplers was comparable for HNO₃, but the Analyst samplers were more sensitive to low concentrations of NO₂. The lowest detectable concentrations for the multipollutant samplers after one month exposure were found to be 1.4 μ g/m³ for NO₂, 0.2 μ g/m³ for HNO₃, and 0.5 μ g/m³ for HONO, and for the Analyst samplers 1.0 μ g/m³ for NO₂ and 1.0 μ g/m³ for HNO₃.

To assess the reproducibility [33] relative standard deviations were considered. Standard deviations were calculated for each set of replicates during the different campaigns. The standard deviations for multipollutant samples were found to be 4 % for NO₂ and 20 % for HNO₃ and for Analyst samplers 5 % for NO₂ and 26 % for HNO₃. These results were in agreement with the variations found by other studies [34, 35] and also fulfil the ± 25 % uncertainty requirement of the European Directive for indicative monitoring with diffusion samplers [36]. Higher values of the standard deviations were found for HONO measured by the multipollutant samplers (approximately 68 %). HONO measurements reproducibility was likely affected by the really low values found indoors, which in many cases were below the detection limit, therefore the dataset was eventually composed by a narrow number of data.

 $\rm NO_2$ and $\rm HNO_3$ concentrations values at the different locations in Swiss libraries, in particular at the SNL where abatement of the pollutants was achieved through the combination of different filters, were often lower than the detection limit.

Nitric acid concentrations were very low probably due to the very high deposition velocity on available surfaces [1, 37].

The extension of the exposure period was implemented to increase the amounts of analyte collected, optimizing the sampling length according to the sensitivity of the technique. Therefore the duration of the campaigns in Switzerland was of about 20 days, whereas the later campaigns in Czech Republic were extended to about a month.

Indoor and outdoor concentrations

Indoor pollutant concentrations are influenced by indoor sources and sinks and penetration from the outdoor environment. As it can be seen from a comparison of the different environments I/O (indoor/outdoor) ratios (Tables 2, 3, 4, 5), the efficiency of the filtration systems (turned off at Bern during the second and third campaign) in removing the pollutants does not seem to produce large differences in the environmental parameters measured. For HONO, when it was possible to perform the calculation, these ratios were always higher than 1 indicating that the formation of the pollutant happens indoors. In this case the filtration systems removing NO₂ should avoid the formation of this pollutant, rather than removing it from infiltrating air.

The reaction (1) is negligible in the gas phase but occurs in the presence of surfaces, where produced HNO_3 remains in the surface water film, whereas HONO forms an equilibrium shifted towards the gaseous phase [38].

According to (1) the equilibrium concentration of HONO should depend on NO_2 concentration and relative humidity RH.

The surface/volume ratio is also known [30] to affect both the formation of HONO and NO at different concentrations of NO_2 and for different values of relative humidity.

It is in agreement with higher NO_2 and HONO outdoor concentration in Prague than in Teplice. Indoor concentrations of HONO were found in the BT depository and the Teplice library, with similar indoor conditions: indoor NO_2 concentrations (Tables 4, 5), RH (Table 6), and a surface/volume ratio 2.6/m and 2.9/m in Teplice and BT, respectively.

Moreover, HONO was not detected in the ACK depository, where NO_2 concentrations were lower than the detection limit of both types of dosimeters. These results are in agreement with the production of HONO indoors from NO_2 by reactions on exposed surfaces which include books and manuscripts, giving rise to acid deposition of HNO₃.

Pollutant	Campaign	Campaign Indoor					Outdoor	IN Average	I/O
		3UG	WA3UG	2UG	WA2UG	A26-Coll	OUT		
NO ₂	S1	bdl	bdl	bdl	bdl	1.5 ± 0.1	6.4 ± 0.3	1.5	0.2
	W	bdl	1.6 ± 0.1	bdl	1.9 ± 0.1	1.7 ± 0.1	31.1 ± 1.2	1.7	0.1
	S2	4.3 ± 0.2	2.8 ± 0.1	2.5 ± 0.1	3.0 ± 0.1	9.2 ± 0.4	12.0 ± 0.5	4.3	0.4
HONO	S1	bdl	2.5 ± 1.7	0.7 ± 0.5	2.7 ± 1.8	1.8 ± 1.2	bdl	1.9	
	W	1.2 ± 0.8	1.7 ± 1.2	bdl	2.2 ± 1.5	bdl	bdl	1.7	
	S2	5.7 ± 3.9	3.3 ± 2.3	2.5 ± 1.7	bdl	11.0 ± 7.5	1.4 ± 0.9	5.6	4.2
HNO3	S1	0.3 ± 0.1	0.2 ± 0.0	bdl	bdl	bdl	0.5 ± 0.1	0.2	0.4
	W	0.4 ± 0.1	0.3 ± 0.1	0.3 ± 0.1	bdl	bdl	bdl	0.3	
	S2	0.2 ± 0.0	0.2 ± 0.0	0.2 ± 0.0	0.2 ± 0.0	0.3 ± 0.1	2.1 ± 0.4	0.2	0.1

Table 2 The Swiss National Library in Bern campaigns results (µg/m³), bdl—below detection limit

Table 3 The Bibliotheque de Geneve campaigns results (µg/m³), bdl—below detection limit

Pollutant	Campaign	Indoor	Indoor			IN Average	I/O
		Depot 51	La reserve	Corridor	OUT		
NO ₂	S	3.4 ± 0.1	bdl	4.5 ± 0.2	16.1 ± 0.6	3.9	0.2
	W	10.8 ± 0.4	5.7 ± 0.2	11.4 ± 0.5	38.9 ± 1.6	9.3	0.2
HONO	S	4.0 ± 2.7	bdl	10.5 ± 7.1	0.0	7.2	
	W	1.9 ± 1.3	1.4 ± 0.9	3.9 ± 2.6	2.0 ± 1.4	2.4	1.2
HNO ₃	S	0.3 ± 0.1	bdl	0.5 ± 0.1	0.7 ± 0.1	0.4	0.6
	W	0.3 ± 0.1	0.5 ± 0.1	0.5 ± 0.1	0.7 ± 0.1	0.4	0.6

Table 4 The National Archives in Prague campaigns results $(\mu g/m^3)$, bdl—below detection limit

Pollutant	Campaign	Indoor	Outdoor	I/O OUT	
		ВТ	ACK		
NO ₂	S1	5.1 ± 0.2	bdl	44.5 ± 1.8	0.1
	S2	4.7 ± 0.2	bdl	31.0 ± 1.2	0.2
	W	5.6 ± 0.2	bdl	42.0 ± 1.7	0.1
HONO	S1	bdl	bdl	bdl	
	S2	0.9 ± 0.6	bdl	bdl	
	W	2.2 ± 1.5	bdl	6.6 ± 4.5	0.3

Table 5 The library of Regional Museum in Teplice campaigns results (μ g/m³), bdl—below detection limit

Pollutant	Campaign	Indoor	Outdoor	I/O
		IN	OUT	
NO ₂	S1	3.1 ± 0.1	19.9 ± 0.8	0.2
	S2	bdl	12.2 ± 0.5	0.2
	W	4.0 ± 0.2	29.1 ± 1.2	0.1
HONO	S1	1.0 ± 0.6	0.9 ± 0.6	1.1
	S2	bdl	bdl	
	W	5.2 ± 3.5	1.9 ± 1.3	2.7

The outdoor concentrations of the pollutants measured in the four cities reflect the seasonal trend, being higher everywhere during wintertime due to low atmospheric layers mixing. The cities of Geneva and Prague showed the highest values (about 40 μ g/m³).

The concentration values of NO_2 were always lower indoors than outdoors, which indicates that there were no indoor sources. Higher indoor values of NO_2 were measured at WA3UG and WA2UG, the two rooms at the library in Bern where the insulation from double doors is not provided. In these two rooms the HONO levels were also higher than in the others. The A26 Collection, where a window is sometimes opened, the concentrations of the pollutants monitored were the highest at the library in Bern. The same can be observed for the corridor at the library in Geneva.

Relatively constant values of NO_2 were observed at the first indoor site in Prague (BT) during all sampling periods. Concentrations measured inside the second indoor site (ACK) were always below detection limit. The difference in NO_2 concentrations was probably caused by the efficiency of filters for NO_2 in the HVAC system of the AKC depository compared to the BT depository equipped only with filters for particles. Indoor NO_2 concentrations measured at Teplice were comparable to concentrations

Table 6 Temperature and relative humidity

	Campaign	Prague			Teplice	
		BT	ACK	OUT	IN	OUT
Relative humidity (%)	I	56	55	69	46	66
	II	56	55	73	47	70
		57	55	84	55	92
Temperature (°C)	I	14.3	14.9	14.5	20.2	15.0
	Ш	15.0	15.0	14.8	22.5	16.0
	III	12.9	14.8	3.8	12.0	2.2

in BT though outdoor concentrations were lower. This was probably due to the higher natural ventilation in this library as compared to the isolated archives.

Conclusions

The results revealed that the reproducibility of the new multipollutant sampler measurements was reasonable (according to EU directives) for NO_2 and HNO_3 . The measured data were comparable with the standard Analyst sampler results, but differ for lower NO_2 concentrations, probably due to differences in sampling and analytical procedure.

The results showed that the newly developed multipollutant sampler can be used in archives and libraries, allowing to map the pollutants distribution indoors. Due to high efficiency of insulating systems normally employed (such as double doors etc.) and filtration systems, the values recorded for the pollutants indoors are often lower than the detection limits. Hence HONO and HNO_3 measurements were carried out in very "clean" environments, and the dataset eventually available was likely not wide enough to truly assess the reproducibility of the measurements, especially for the former pollutant.

Authors' contributions

FV design of the monitoring project and discussion of the results. LM samplings organization and revision of the manuscript. MF technical support in the laboratory. Al analytical activity. JS revision of the manuscript. All authors read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

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