

A traceable and continuous flow calibration method for gaseous elemental mercury at low ambient concentrations

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Abstract. The monitoring of low gaseous elemental mercury (GEM) concentrations in the atmosphere requires continuous high-resolution measurements and corresponding calibration capabilities. Currently, continuous calibration for GEM is still an issue at ambient concentrations $(1-2 \text{ ng m}^{-3})$. This paper presents a continuous flow calibration for GEM, traceable to NIST 3133 Standard Reference Material (SRM). This calibration approach was tested using a direct mercury analyser based on atomic absorption spectrometry with Zeeman background correction (Zeeman AAS). The produced continuous flow of GEM standard was obtained via the reduction of Hg²⁺ from liquid NIST 3133 SRM and used for the traceable calibration of the Zeeman AAS device. Measurements of atmospheric GEM using the calibrated Zeeman AAS were compared with two methods: (1) manual gold amalgamation atomic fluorescence spectrometry (AFS) calibrated with the chemical reduction of NIST 3133 and (2) automated gold amalgamation AFS calibrated using the mercury bell-jar syringe technique. The comparisons showed that a factory-calibrated Zeeman AAS device underestimates concentrations under 10 ng m^{-3} by up to 35 % relative to the two other methods of determination. However, when a calibration based on NIST 3133 SRM was used to perform a traceable calibration of the Zeeman AAS, the results were more comparable with other methods. The expanded relative combined uncertainty for the Zeeman AAS ranged from 8% for measurements at the 40 ng m^{-3} level to 91.6% for concentrations under 5 ng m^{-3} using the newly developed calibration system. High uncertainty for measurements performed under 5 ng m^{-3} was mainly due to instrument noise and concentration variation in the samples.

1 Introduction

Mercury (Hg) and its compounds are ubiquitous in the environment. Gaseous elemental mercury (GEM) is the main Hg species in the atmosphere (Enrico et al., 2016; Outridge et al., 2018). The relatively high chemical stability of GEM and its long lifetime of about 1 year (Ariya et al., 2015; Si and Ariya, 2018) facilitate its global transport (Koenig et al., 2022). Subsequent oxidation of GEM produces gaseous oxidised mercury (GOM) which can be readily deposited via dry or wet deposition (Phu Nguyen et al., 2019). Deposited GOM can be subsequently methylated to the most toxic Hg species, monomethyl mercury (Castellini et al., 2022). Assessment and monitoring of chemical transformations and physical processes require high measurement accuracy for all Hg species to identify transformation pathways and, consequently, possible effects on human health and biota (Burger and Gochfeld, 2021).

Although GEM is the most abundant Hg species in the atmosphere, its concentrations are still extremely low $(1-2 \text{ ng m}^{-3})$ (MacSween et al., 2022); thus, accurate and precise analytical methods are required for the determination of this species. Commonly used analytical methods are based on preconcentration techniques, where atmospheric Hg is sampled on sorbent traps (usually gold, gold-coated silica, or activated carbon traps) (Živković et al., 2020). Following preconcentration, mercury is thermally desorbed from the

traps and analysed using atomic fluorescence spectrometry (AFS) (Kamp et al., 2018; Skov et al., 2020) or atomic absorption spectrometry (AAS) (El-Feky et al., 2018). Alongside AFS and AAS, recent analytical methods are also based on laser techniques, including laser imaging, detection, and ranging (lidar) methods (Fantozzi et al., 2021; Ravindra Babu et al., 2022) and laser absorption spectroscopy (Srivastava and Hodges, 2022).

A more direct approach for Hg determination in atmospheric samples can be achieved using an atomic absorption spectrophotometer with Zeeman background correction (Zeeman AAS) with a high measurement resolution, such as a Lumex RA-915M. The high measurement resolution is achieved by direct, 1 s GEM measurements in the atmosphere. The principles of this method have been described elsewhere (Sholupov and Ganeyev, 1995). This technique is used worldwide for GEM measurements (Cabassi et al., 2020; Lian et al., 2018) due to its simplicity of operation and the absence of a preconcentration step. Zeeman AAS devices ought to be calibrated at the manufacturer, and the manufacturer suggests a yearly recalibration in the range from a few micrograms per cubic metre to milligrams per cubic metre. However, this might be inappropriate for ambient atmospheric measurements.

In general, the calibration of instruments for GEM determination can be performed using traceable amounts of Hg⁰ obtained from (i) the reduction of Hg²⁺ Certified Reference Material (CRM), (ii) a mercury bell-jar calibrator unit, or (iii) GEM permeation tubes (Jampaiah et al., 2019). However, a stable and continuous flow of GEM calibration gas is required for a metrologically proper calibration of instruments for continuous GEM analyses in the atmosphere. Several calibration devices for elemental Hg already exist (albeit for higher concentrations), such as NIST PRIME (Long et al., 2020) and PSA 10.534/10.536 (Brown et al., 2008) from PS Analytical. They are dynamic continuous GEM calibrators which dilute mercury-saturated air originating from a temperature-controlled reservoir of liquid elemental mercury. Depending on the Hg reservoir temperature, reservoir flow, dilution flow, and pressure, the mercury concentration can be calculated using the Dumarey equation (Dumarey et al., 1985; Ebdon et al., 1989). Mercury saturation in air calculated using the Dumarey equation is an agreed standard used for calibration (ISO 6978-2:2003, ASTM D6850-03) and has been validated using Standard Reference Materials (SRMs) (Dumarey et al., 2010). In the literature, the lowest concentration of Hg⁰ validated using a dynamic calibrator is $0.501 \,\mu g \,m^{-3}$ (NIST Prime), with an expanded uncertainty of $0.018 \,\mu\text{g}\,\text{m}^{-3}$ (Long et al., 2020). The PSA 10.534/10.536 is normally used for concentrations in the microgram-percubic-metre range (Lopez-Anton et al., 2010; Wang et al., 2018), but the lowest reported concentration obtained from it is 26.4 $\mathrm{ng}\,\mathrm{m}^{-3}$ (Brown et al., 2010) by further diluting the output gas coming from the Hg reservoir. Recently, a primary mercury gas generator (PMGG) was developed for the calibration of GEM measurement systems (Ent et al., 2014). The working principle of the PMGG is based on a dilution of mercury in stripping gas from diffusion cells. The traceability of the PMGG to International System of Units (SI) units is established through the determination of diffusion rates, calculated by weighing the diffusion cells at regular time intervals (de Krom et al., 2021; Ent et al., 2014). Currently, the lowest concentration generated by the PMGG is $0.1 \,\mu g \,m^{-3}$ with a relative expanded uncertainty of 3 %.

The objective of this paper is to present a simple, cheap, and user-friendly calibration method for the continuous ambient GEM measurements using a Zeeman AAS device based on the reduction of Hg^{2+} from NIST 3133 SRM. This calibration approach can be used to generate traceable amounts of GEM in the low micrograms-per-cubic-metre concentration range. An uncertainty budget for the analytical procedure at low concentrations is presented. The performance of a Zeeman AAS device is compared with manual and automated gold amalgamation systems for GEM determinations.

2 Materials and methods

2.1 Chemicals and calibration standards

The acids used for the acidification of samples and standard solutions were 30 % HCl (Suprapur, Merck, Darmstadt, Germany) and 65 % HNO₃ (for analysis, Supelco, Darmstadt, Germany). Type-I purified water (electrical resistivity 18.2 M Ω cm; Milli-Q water, Merck, Darmstadt, Germany) was used in all experiments for the dilution of standards. Soda lime (Merck, Darmstadt, Germany) was used for drying the calibration Hg⁰ gas. The solution of SnCl₂ (made from SnCl₂ · 2H₂O max 10⁻⁶ % Hg, Merck, Darmstadt, Germany) in HCl was used for the quantitative reduction of Hg²⁺ from NIST 3133 SRM standard solutions.

NIST 3133 (lot no. 160921) SRM (National Institute of Standards and Technology, Gaithersburg, MD, USA) was used for the calibration of the Zeeman AAS and manual AFS devices. The Hg stock solution $(1.044 \,\mu g \, g^{-1})$ was prepared by two consecutive gravimetric dilutions of the initial NIST 3133 SRM $(10.004 \,m g \, g^{-1})$ in matching matrix $(10 \,\% \, \text{HNO}_3 \, v/v$ in Milli-Q water). From this stock solution, working standard solutions were prepared daily in $2 \,\% \, w/w$ HCl (exact concentrations are provided in the Supplement).

For automated AFS, a Hg-saturated gas standard was sampled using a gas-tight syringe (Hamilton gas-tight syringe 50 μ L, point style 2, 51 mm needle length) from a bell-jar calibrator unit (Tekran[®] Model 2505 mercury vapour primary calibration unit). The amount of injected mercury was calculated using the Dumarey equation (Dumarey et al., 1985).

2.2 Analytical instrumentation

In this paper, we compare the analytical performance of a Zeeman AAS device with automated and manual gold amalgamation, followed by AFS quantification due to their widespread use in monitoring GEM concentrations in the atmosphere. For this purpose, we used a Lumex RA-915M (Lumex Analytics GmbH, 24558 Wakendorf II, Germany) (Mashyanov et al., 2022) for the Zeeman AAS, a Sir Galahad (PS Analytical, Arthur House, Main Rd, Orpington BR5 3HP, UK) (Rafeen et al., 2020) for automated AFS, and a Brooks Rand Model III (Souza et al., 2020) as the detector for manual AFS. An OLM30B sampler (Lumex Analytics GmbH, 24558 Wakendorf II, Germany), equipped with a high-accuracy flow meter, was used for GEM sampling on gold traps.

2.2.1 Manual AFS

The manual AFS (AFS-M) method is based on a double amalgamation procedure coupled with thermal desorption and AFS quantification. Atmospheric GEM samples were collected on a sampling gold trap connected to the OLM30B sampler. Air was pumped through the sampling gold trap for 30 min at a flowrate of 0.5 Lmin^{-1} , to match the sampling system of the automated AFS. In the measurement step, the sampling gold trap was immediately transferred to a double amalgamation AFS system. The sampling gold trap was heated for 30 s (ramp heating to a maximum of 600 °C), and mercury collected on this trap was released and purged with a flow of argon (Ar) onto a permanent analytical gold trap kept at room temperature. After heating the analytical gold trap, Hg was thermally desorbed into an argon stream that carried the released Hg vapour into the cell of an AFS detector (Brooks Rand Model III).

One-point calibration (100 or 250 pg, depending on the concentration in the samples) was used for the calibration of the AFS detector (Ma et al., 2012). Gaseous Hg⁰ standards were prepared by the reduction of the appropriate masses of NIST 3133 SRM working standards in 50 mL of Milli-Q water using 2 mL of 10% SnCl₂ (w/v). The produced Hg⁰ was purged with N₂ gas for 4 min at a flowrate of 50 mLmin⁻¹, dried through a soda lime trap, and quantitatively trapped on the sampling gold trap. The analytical signal for the calibration standard trapped on the sampling gold trap was obtained in the same manner as described for samples.

2.2.2 Automated AFS

The Sir Galahad is an automatic mercury analyser which works similarly to the manual double amalgamation AFS. However, GEM is preconcentrated on an incorporated single gold trap, thermally desorbed, and analysed in the AFS detector. Air sampling using an external pump was set to $0.5 \,\mathrm{L\,min^{-1}}$ for 30 min, resulting in a total volume of $15 \,\mathrm{L}$ of sampled air (the same as in the case of manual double amalgamation). The preconcentrated GEM is then thermally released in the Ar stream and detected using atomic fluores-

cence. The automated AFS (AFS-A) measures the volume of the sampled air and calculates the results automatically.

The calibration of the automated AFS was performed using eight calibration points of GEM-saturated gas standard. The gas standard was obtained from a bell-jar gas calibrator unit using a gas-tight syringe. Depending on the bell-jar temperature, the concentration of GEM was calculated using the Dumarey equation (Dumarey et al., 2010). The injected mercury was amalgamated on the gold trap, similarly to in the manual double amalgamation method. The signal for GEM standard was obtained at the AFS detector after thermal release from the gold trap.

2.2.3 Zeeman AAS

The Lumex RA-915M is a portable multifunctional AAS with Zeeman background correction. Atmospheric GEM is sampled using an in-built pump with a flowrate of 10 Lmin^{-1} (Lumex RA-915M Mercury Analyzer Operation Manual, Mission, Canada, 2015). The Zeeman AAS automatically performs baseline correction at the beginning and end of each analysis by sampling air through the incorporated zero filter. The baseline correction is used to alleviate possible detector drift. Throughout this work, the baseline correction time was set to 60 s. During air analysis, baseline correction was performed every 15 min.

The Zeeman AAS instrument is calibrated yearly at the manufacturer's facilities in the low microgram per cubic metre to microgram per cubic metre range. As this might not be appropriate for low atmospheric GEM measurements, we also performed external calibration as described in Sect. 2.3. The Zeeman AAS can report GEM values as often as every second. However, a reading resolution of 10 s was used in this work. Lower reading resolutions would greatly increase the uncertainty of integration, whereas higher resolutions would not significantly decrease the standard deviation. All 10 s signals were integrated using Rapid software for a period of 5 min. To eliminate any possible errors due to signal changes, the system blanks were analysed before each external calibration standard, while baseline correction was performed after each standard. Nevertheless, the Zeeman AAS proved to be a very stable detector.

2.2.4 External GEM calibration for the Zeeman AAS

The principle of external continuous flow GEM calibration was based on the continuous reduction of NIST-traceable Hg^{2+} standard solution with SnCl₂. The general scheme of the external continuous flow GEM calibration system is shown in Fig. 1. NIST 3133 calibration standard (ranging from 2.5 to 1000 pg g⁻¹) and 3 % w/v SnCl₂ in 3 % HCl are transferred using peristaltic pumps through a first T-piece to a 45 cm Teflon mixing coil to ensure the quantitative reduction of Hg²⁺ to Hg⁰. The mixed solution is then pre-purged in the second T-piece with a known flow of Hg-free nitrogen



Figure 1. Scheme of external GEM calibration for the Zeeman AAS device.

to facilitate gas-phase Hg⁰ transfer from the liquid solution. The N₂ gas carries the solution to an impinger, where the liquid is vigorously purged with the same carrier gas through a quartz frit. The impinger is made of quartz and has a height of 46 cm and a diameter of 13.5 cm. All the T-pieces used in the set-up are made of Teflon, along with all of the connections until the impinger. Between the impinger and the analyser, the connections are made with Tygon tubing, and the last T-piece is made of quartz. The quartz frit is placed 2 cm above the bottom of the impinger. The flows of NIST 3133 liquid standards used in this work were between 2.2 and $3.0 \,\mathrm{g\,min^{-1}}$ and were determined by weighing the waste from both the peristaltic pump and the impinger. No noticeable difference was caused by the N₂ flow from the impinger. The amount of Hg in the waste exiting the impinger was always below the limit of detection of the manual double amalgamation system, thus ensuring quantitative transfer to the gaseous phase. In order to obtain the required sampling flow for the external calibration of the Zeeman AAS, the output from the impinger was diluted with a known flow of Hg-free air through a third T-piece (Fig. 1). As the N_2 flow is lower than the flow of the pump inside the Zeeman AAS, a T-piece was placed between the impinger and the device. This allows for the air from outside to enter the system in order to compensate for differences in flows and pressure. The air passes through a Hg filter to ensure that it does not interfere with the analysis. The Hg filter is a carbon trap similar to the one that the Zeeman AAS device uses for baseline correction. A soda lime trap placed between the last T-piece and the detector removed any moisture coming from the impinger or the air outside the system.

The flow of N₂ used was between 3.0 and 8.3 Lmin^{-1} , depending on the concentration range of the standards used. The flow of N₂ was set with the aid of a mass flow controller. For the highest NIST 3133 liquid standard used (1 ng g⁻¹), a N₂ flow of 8.27 Lmin⁻¹ was necessary in order to ensure the quantitative release of Hg⁰ from the liquid phase. As long as the N₂ flow was high enough to quantitatively purge the Hg⁰ from the standard, changes in its flow did not bring noticeable changes to the signal.

2.3 Uncertainty of the measurement results for the Lumex RA-915M

The uncertainty of the Zeeman AAS measurement results, calibrated with the external GEM calibration system, was calculated according to the ISO-GUM (International Organization for Standardisation – Guide to the Expression of Uncertainty in Measurement) approach (ISO/IEC 17025, Switzerland, 2017). The uncertainty of each component was calculated following the ISO-GUM rules for the calculation of uncertainty sources. The combined standard uncertainty (at a coverage factor of 1) was calculated following the general rules for the propagation of uncertainty for a mathematical model expressed in the form of additions and/or multiplications. The identification of uncertainty components was based on the mathematical model (Eqs. 1 and 2) used for the calculation of GEM concentrations from the detector's analytical signal:

$$C_{\rm S} = \frac{A_{\rm S}' - i}{m},\tag{1}$$

$$C_{\rm std\ gas} = C_{\rm NIST\ 3133} \times \frac{Q_{\rm aq}}{Q_{\rm inflow}}.$$
 (2)

Here, C_S is the concentration of the air sample $[\text{ng m}^{-3}]$, *m* is the best-fit gradient of the calibration curve (slope), *i* is the intercept of the calibration curve, A'_S is the signal of the sample after blank subtraction $[\text{ng m}^{-3}]$, $C_{\text{std gas}}$ is the concentration of the produced gas standard and was used to calculate *m* and *i* $[\text{ng m}^{-3}]$, $C_{\text{NIST 3133}}$ is the concentration of the liquid standard $[\text{ng g}^{-1}]$, Q_{aq} is the flow of liquid SRM which is introduced by the peristaltic pump in the T-piece before the impinger $[\text{g min}^{-1}]$, and Q_{inflow} is the flow of air and nitrogen that enters the detector $[\text{m}^3 \text{min}^{-1}]$.

Based on the model, a fishbone diagram (Ishikawa diagram) was constructed and is presented in Fig. 2. Proper construction of the fishbone diagram was an important step in the uncertainty evaluation, as it showed the influences of different components on the combined uncertainty. Fishbone diagrams for the manual and automated AFS systems are presented in the Supplement (Figs. S1 and S2).

The uncertainty of the calibration curve was estimated following Eurachem guidelines (Ellison and Williams, 2012) as follows:

$$u_x = \frac{S_y}{m} \times \sqrt{\frac{1}{k} + \frac{1}{n} + \frac{(y - \overline{y})^2}{m^2 \sum (x_i - \overline{x})^2}}.$$
 (3)

Here, u_x is the standard uncertainty brought by calibration for the value x, S_y is the standard deviation on the y axis, m is the calculated best-fit gradient of the calibration curve, k is the number of replicated measurements on the sample, n is the number of paired calibration points, y is the mean



Figure 2. Fishbone diagram of the uncertainty of measurement results in the case of sample analysis with the Zeeman AAS device.

value of the replicates of the sample, \overline{y} is the mean of the y values for the calibration standards, x_i is the position of the standards on the x axis, and \overline{x} is the mean of the x_i values.

All uncertainty components were calculated as type-A uncertainty, except for the weight and temperature, which were calculated as type-B uncertainties. The expanded combined relative standard uncertainty was calculated by multiplying the relative combined standard uncertainty by a coverage factor of 2:

$$U_{\rm ex,r} = u_{\rm r,c} \times k. \tag{4}$$

The contribution index $(\% u_i)$ of individual uncertainty components was calculated as follows:

$$\% u_i = \frac{u_i^2}{\sum_{i=1}^{n} u_i^2},$$
(5)

where n is the number of uncertainty components used to calculate the relative combined standard uncertainty.

2.4 Data analysis and statistical evaluation

Uncertainty evaluation was performed following ISO-GUM guidelines (ISO/IEC Guide 98-3:2008) and was calculated using Microsoft Excel 2019. Appropriate statistical tests (analysis of variance – ANOVA, repeated-measures ANOVA, and regression analysis) were performed using SigmaPlot 14.0. These tests are mentioned throughout the paper along with the data on which the tests were applied. The statistical significance was set to an α value of 0.05.

3 Results

3.1 External calibration for the continuous mercury analyser

The stability of the external calibration of the Zeeman AAS signal is shown in Fig. S3 in the Supplement. Although the values on the y axis are expressed as Hg concentrations and not as raw analytical signals in all of the following figures, we use the expression "calibration plot" in these figures to emphasise the necessity of additional external calibration of already-calculated Hg concentrations. Therefore, the values on the y axis can be considered equivalent to the analytical signal. An external calibration, presented in Fig. S3, was performed using corresponding liquid standards with concentrations between 5 and 55 pgg^{-1} as described in Sect. 2.2.4. The system was calibrated from the highest to the lowest concentration to test whether leftover Hg remained in the system after each calibration standard. This was confirmed by the absence of systematic differences between the blanks after each standard. Parallel-line analysis (PLA) was performed between two calibration curves which were in the same concentration range (one with randomised calibration points and one with calibration points analysed from the lowest to the highest concentration), and there is no statistical difference between them, with P = 0.1995 for the slopes and P = 0.5493 for the intercepts. The equation for the randomised calibration curve is y = 0.6616x - 0.0088, whereas the equation for the other curve is y = 0.6412x + 0.22.

The repeatability of both blanks and low-concentration standards are comparable. The standard deviation varied between 0.32 and 0.43 ng m⁻³ for a reading resolution of 5 s, which indicates that there was a stable outflow from the calibration set-up (Fig. 1). The background correction was done

after each standard in order to avoid any uncertainty arising from signal drift. For each blank and standard, an integration time of 5–6 min was used. As all of the results reported throughout this paper were calculated based on multiplepoint calibration plots, the blank measured before each standard was used for subtraction. The concentration output was also highly dependent on the autozero done by the Zeeman AAS system. In order to eliminate any drifts that may occur during calibration, autozero was applied after each pair of blank–standard detections. Recalibration was also performed when the standards were introduced in a random order of concentrations to demonstrate the absence of bias due to hysteresis. The slope and regression coefficients of the calibration plots did not differ. An example of this randomised calibration curve was used to calculate the results in Sect. 3.3.3.

3.2 Comparison of factory calibration against a known concentration generated by NIST 3133 at different concentration levels

The linearity of the detector's response was checked using a wide range of concentrations, ranging from 1.76 to 176.8 ng m^{-3} , as shown in Fig. 3a. The corresponding concentrations of Hg in the NIST 3133 liquid standards were between 10.4 and 1044 pg g^{-1} . All standard solutions were analysed daily under the same experimental conditions for external GEM calibration of the Zeeman AAS (Sect. 2.2.4). The overall slope of the calibration plot was close to 1 (Fig. 3). However, in the lower calibration range, the slope was considerably lower (0.681; Fig. 4b), thus confirming the requirement for additional external calibration at low concentration levels ($< 15 \text{ ng m}^{-3}$). Five of the lowest calibration standards (Fig. 3b) fell within the 95 % confidence interval of the overall calibration plot calculated based on all 10 calibration standards. If only one calibration standard was measured in this low-concentration range, samples measured in this region would be underestimated.

This difference in response at lower concentrations was reproducible and constant throughout this work. Even though it is not known why the difference occurred at low concentrations, the lower response was observed during sample analysis of GEM. This indicates that there was not an issue with the external calibration but rather a bias from the Zeeman AAS. Factors such as spectral emission profiles, concentrations of analyte, and magnetic field strength can influence the performance of Zeeman background correction (Ganeyev and Sholupov, 1992).

Furthermore, the external calibration curve (Fig. 3) was plotted along with the internal calibration curve determined by the manufacturer (Fig. 4) during yearly service. The calibration standards used by the external calibration system were well within the confidence interval of the manufacturer calibration, and the corresponding slopes were parallel. Therefore, the application of the internal calibration curve for the determination of high GEM concentrations was valid and gave accurate results. For the lower-concentration region in Fig. 3b, proper external calibration must be performed.

3.3 Comparison of AAS, automated double amalgamation AFS, and manual double amalgamation AFS

In order to validate the external calibration method, the Zeeman AAS device was compared with automated AFS and manual AFS at different calibration ranges. The measurements were performed at high, medium, and low concentrations, defined by the following criteria. High-concentration (around 40 ng m^{-3}) measurements were those measurements where no difference was observed among the three methods, with comparable results and combined extended uncertainties. For the medium-concentration ranges $(5-10 \text{ ng m}^{-3})$, there were differences between the results obtained with the manufacturer-calibrated Zeeman AAS and the other two methods, with the combined extended uncertainties being low enough to back up this statement. Low-concentration ranges ($< 5 \text{ ng m}^{-3}$) were concentrations at which differences between the methods were observed, but the combined extended uncertainties were too high to actually state that there was a difference.

3.3.1 High-concentration measurements

Measurements at approximately 40 ng m^{-3} were performed using the above-mentioned methods. No noticeable differences among the three methods were observed, with the combined expanded uncertainties varying between 2.2% and 9%. Even after recalculating the results based on the external calibration, the results from all three methods were comparable. The average difference between the raw results from the Zeeman AAS system and those recalculated using the external calibration method for this set of data was 3.85%. The main contributor to the combined uncertainty in the case of the Zeeman AAS was the repeatability of the sample, bringing a contribution of 76.4% on average.

Through evaluating the uncertainty ranges at a coverage factor of 2, it cannot be stated that there was a difference among the three methods. The same statistical evaluation performed in the previous section for low concentrations as well as the one-way repeated-measures ANOVA with a post hoc Bonferroni test for pairwise multiple comparison also showed no difference among the three methods. The same test was performed with the data calculated with the external GEM calibrator system, and there was no statistical difference.

3.3.2 Medium-concentration measurements

Measurements under 10 ng m^{-3} showed that the Zeeman AAS was underestimating the results, even though the manual AFS and automated AFS systems were comparable, with relatively low uncertainties (Fig. 6). The long-term measure-



Figure 3. Panel (a) shows the analysis of NIST 3133 standards in the $1.76-176.8 \text{ ng m}^{-3}$ range. Panel (b) presents the analysis of NIST 3133 standards in the $1.76-12.5 \text{ ng m}^{-3}$ range. The *x* axis is the theoretical concentration of the NIST 3133 SRM; the *y* axis is the calculated concentration by the Zeeman AAS device, proportional to the signal.



Figure 4. Panel (a) shows the manufacturer calibration in the 2532–41 476 ng m⁻³ range. Panel (b) presents the NIST 3133 standards in the 1.76–176.8 ng m⁻³ range.

ments showed that the difference between the Zeeman AAS system and two AFS methods was, on average, 35.1 %.

In the low-concentration range for the Zeeman AAS, the relative standard deviation varied from 5.6% at the highest concentration (8.5 ngm^{-3}) to 9% at the lowest one (4.3 ngm^{-3}) . In all cases, the contribution from sample re-

peatability was the highest contributor to the combined uncertainty, which can be seen in Fig. 5. The uncertainty contribution coming from the gaseous standard was very low (< 4% for this series of measurements).

The uncertainty intervals overlapped, which indicates that there was no difference between the manual and automated



Figure 5. Comparison between the Zeeman AAS, AFS-A, and AFS-M. The error bars are the combined uncertainties for k = 2.



Figure 6. Comparison between the Zeeman AAS, AFS-A, and AFS-M. The error bars are the combined uncertainties for k = 2. The "*" represents the externally calibrated Zeeman AAS. For the direct mercury analysers calibrated by the manufacturer, the error bars represent twice the standard deviation, whereas they are represented by the combined expanded uncertainty for a coverage factor of 2 in the other cases.

AFS methods. It was the same for the AAS system calibrated with our set-up, although not for the raw data obtained with the manufacturer calibration. To further confirm whether differences existed, a one-way repeated-measures ANOVA was performed, with a post hoc Bonferroni test for pairwise multiple comparison. The results indicated that there was no difference between automated and manual AFS (p = 1), but there was a difference between the results from the first two methods and the results from the Zeeman AAS calibrated by the manufacturer (p < 0.001). Between the first two methods and the results from the Zeeman AAS system calibrated using the dynamic set-up, there was no difference (p = 1).



Figure 7. Comparisons between two direct mercury analysers and the atmospheric mercury analyser. The "*" represents the externally calibrated Zeeman AAS device. For the direct mercury analysers calibrated by the manufacturer, the error bars represent twice the standard deviation, whereas they are represented by the combined expanded uncertainty for a coverage factor of 2 in the other cases.



Figure 8. Relative contribution indexes with respect to the uncertainties for the externally calibrated Zeeman AAS systems.

3.3.3 Low-concentration measurements

Another comparison was made at a lower concentration using two Zeeman AAS systems of the same series and the automated AFS system (Fig. 7). The AAS devices showed similar responses. The uncertainty due to sample repeatability was much higher due to the lower concentrations, and it increased the combined uncertainty by up to 98.6% for a coverage factor of 2. As seen in Fig. 8, the repeatability of the sample was still the main contributor to the extended combined uncertainty (contribution up to 95%), like in previous cases. It is important to note that the high uncertainty contribution brought by the repeatability of the sample in the case of the Zeeman AAS was not entirely due to its performance but also due to the variability in the concentration of the sample, which (in this case) was atmospheric air. Even slight changes at these low concentrations result in very high standard deviations. The standard deviations of calibration standards and blanks in this concentration range were always under 0.5 ng m^{-3} , whereas the standard deviation was as high as 1.1 ng m^{-3} for samples. This high contribution from the repeatability of the sample is to be expected, as the limit of quantification is 3.2 ng m^{-3} .

Due to these high uncertainties with respect to the measurement results, it cannot be stated that there was a difference between the results obtained from the manufacturercalibrated Zeeman AAS systems and the automated AFS system or between the results obtained from the manufacturercalibrated Zeeman AAS systems and the externally calibrated Zeeman AAS. This is because the uncertainty ranges overlapped. Even so, the previously used one-way repeatedmeasures ANOVA shows that there was a difference between the results obtained with the manufacturer-calibrated Zeeman AAS and the automated AFS (P < 0.001) and that there was a difference between the results obtained with the manufacturer-calibrated Zeeman AAS and the externally calibrated Zeeman AAS systems (p < 0.001). There was no difference between the automated AFS and the externally calibrated Zeeman AAS systems (p = 1).

Additionally, a one-way ANOVA with a Holm-Šidák pairwise comparison was performed, and, in most of the cases, there was no statistical difference between the externally calibrated Zeeman AAS systems and the automated AFS, as can be seen in the Supplement (Table S1). We tested statistical differences between corresponding individual values that were averaged to calculate the results shown in Fig. 8. The results of the one-way ANOVA showed a significant difference between the manufacturer-calibrated and externally calibrated Zeeman AAS (p < 0.001 in all cases, except between Zeeman AAS no. 2 and Zeeman AAS no. 2* from the fourth series of measurements, p = 0.004). There was no significant difference between the two externally calibrated Zeeman AAS systems (p > 0.05). The AFS-A was not considered for this statistical test, as each result is based on a single data point.

4 Discussion

It is difficult to assess why the Zeeman AAS system underestimated the results for low concentrations, as shown in Sect. 3.3.1. It is most likely associated with the extrapolation of the regression function obtained from the calibration plot performed by the manufacturer. In the calibration certificate received for the device used in this work, it is stated that the calibration was performed in the $2.63-42.4 \,\mu g \,m^{-3}$ range. An

extrapolation from these high concentrations to those presented in Sect. 3 can lead to substantial systematic errors. Even so, the Zeeman AAS system performed very well for concentrations of around 40 ng m^{-3} .

For the comparison experiments, the repeatability of the samples could not be considered for the two preconcentration methods. This is due to the fact that sample homogeneity was not assured from one sampling period to the next. For this reason, additional statistical tests were performed (as stated in Sects. 3.3.1 and 3.3.2).

The preconcentration methods used for comparisons are sampling and analysing all mercury species in the atmosphere. Along with GEM, GOM and particle-bound mercury (PBM) are also present in the atmosphere (Schroeder and Munthe, 1998). During the thermal desorption of mercury from gold traps, GOM and PBM species are reduced and end up being analysed as GEM as well. Reactive mercury (RM) is the sum of GOM and PBM, and its concentration is usually under 100 pg m^{-3} in continental areas (Pierce et al., 2018; Zhou et al., 2018). Although speciation analysis has not been done, RM concentrations higher than 100 pg m^{-3} were not observed at the sampling site when using a 2537B/1130/1135 Tekran speciation system (Gustin et al., 2013). These concentrations would not affect the results of the comparisons done in this work, as they are almost 3 orders of magnitude lower than was analysed.

Future work should be focused on improving the calibration set-up in order to obtain higher concentrations of GEM. The limiting factor in this work was the high flow needed for the quantitative removal of the dissolved gaseous mercury from the solution introduced in the impinger. The highest purging flowrate used was $8.27 \text{ L} \text{ min}^{-1}$, which managed to quantitatively purge the mercury from a $1 \text{ ng g}^{-1} \text{ Hg}^{2+}$ solution introduced at a rate of $2.26 \text{ g} \text{ min}^{-1}$ into the impinger. Higher flows were not tested due to safety reasons, but, at the moment, quartz is the best material for handling mercury, as it does not retain it, compared with other materials which are more durable, such as stainless steel.

5 Conclusions

Continuous flow calibration for GEM was established through a simple, cheap, and easily assembled set-up. All measurements are traceable to NIST 3133 SRM, and other liquid CRMs of Hg^{II} can be used. The set-up ensured stable and repeatable signals, even at very low concentrations suitable for the atmospheric continuous measurements of GEM. The main drawback of the calibration set-up is that it cannot be used for very polluted measurements, as higher flows of purging gas would be required, which implies considerable pressure inside the system. The calibration method was successfully tested using a commercially available Zeeman AAS system, and the results were compared with an automated gold amalgamation AFS system calibrated with a Dumarey bell-jar gas standard and with a manual AFS calibrated with NIST 3133 SRM. Although the Zeeman AAS system calibrated by the manufacturer underestimated concentrations under 10 ng m^{-3} , the newly developed calibration system corrected this. The two AFS methods were in good agreement, with relatively low uncertainties (< 9 % for a coverage factor of 2). Improvements to the continuous flow calibration set-up may result in obtaining a broader range of concentrations for the gas standard obtained from NIST 3133 SRM.

Data availability. The dataset used in this paper is available upon reasonable request.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/amt-17-1217-2024-supplement.

Author contributions. TDA was responsible for conceptualisation, methodology, formal analysis, investigation, data curation, writing – original draft, and visualisation. WTC was responsible for conceptualisation, validation, resources, writing – review and editing, and supervision. IŽ was responsible for data curation, writing – review and editing, and visualisation. SWA was responsible for formal analysis and writing – review and editing. SVN was responsible for formal analysis, writing – review and editing, and visualisation. MH was responsible for conceptualisation, writing – review and editing, supervision, project administration, and funding acquisition. All listed authors were responsible for the review and editing process.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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