High-precision measurements of seawater Pb isotope compositions by double spike thermal ionization mass spectrometry

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**HIGHLIGHTS**

- Precise and accurate determination of seawater Pb isotope compositions.
- Uncertainties (2sd) for $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{206}\text{Pb}$, $^{208}\text{Pb}/^{206}\text{Pb}$ are better than ±3‰.
- Analysis of a GEOTRACES depth profile from the South Atlantic Ocean.
- Pb isotopic compositions reflect the different water masses that were sampled.
- The $^{207}\text{Pb}/^{206}\text{Pb}$ and $^{206}\text{Pb}/^{204}\text{Pb}$ ratios display a correlation.

**ABSTRACT**

A new method for the determination of seawater Pb isotope compositions and concentrations was developed, which combines and optimizes previously published protocols for the separation and isotopic analysis of this element. For isotopic analysis, the procedure involves initial separation of Pb from 1 to 2 L of seawater by coprecipitation with Mg hydroxide and further purification by a two stage anion exchange procedure. The Pb isotope measurements are subsequently carried out by thermal ionization mass spectrometry using a $^{207}\text{Pb}$–$^{204}\text{Pb}$ double spike for correction of instrumental mass fractionation. These methods are associated with a total procedural Pb blank of 28 ± 21 pg (1sd) and typical Pb recoveries of 40–60%. The Pb concentrations are determined by isotope dilution (ID) on 50 mL of seawater, using a simplified version of above methods. Analyses of multiple aliquots of six seawater samples yield a reproducibility of about ±1 to ±10% (1sd) for Pb concentrations of between 7 and 50 pmol/kg, where precision was primarily limited by the uncertainty of the blank correction ($12 \pm 4$ pg; 1sd). For the Pb isotope analyses, typical reproducibilities (±2sd) of 700–1500 ppm and 1000–2000 ppm were achieved for $^{206}\text{Pb}/^{204}\text{Pb}$, $^{207}\text{Pb}/^{206}\text{Pb}$, $^{208}\text{Pb}/^{204}\text{Pb}$, $^{206}\text{Pb}/^{204}\text{Pb}$, respectively. These results are superior to literature data that were obtained using plasma source mass spectrometry and they are at least a factor of five more precise for ratios involving the minor $^{204}\text{Pb}$ isotope. Both Pb concentration and isotope data, furthermore, show good agreement with published results for two seawater intercomparison samples of the GEOTRACES program. Finally, the new methods were applied to a seawater depth profile from the eastern South Atlantic. Both Pb contents and isotope compositions display a smooth evolution with depth, and no obvious outliers. Compared to previous Pb isotope data for seawater, the $^{206}\text{Pb}/^{204}\text{Pb}$ ratios are well correlated with $^{207}\text{Pb}/^{206}\text{Pb}$, underlining the significant improvement achieved in the measurement of the minor $^{204}\text{Pb}$ isotope.

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1. Introduction

Investigations that map the distribution of trace metals and their isotopes in seawater and which study their relationship to and potential impacts on global biogeochemical cycles in the oceans are an overarching goal of modern marine geochemistry and a key objective of the international GEOTRACES program [1]. The global scale impact of human activities on natural Pb budgets was established more than 40 years ago by Chow and co-workers [2–5] and massive inputs of anthropogenic Pb into the oceans were soon thereafter identified by Patterson and co-workers [6–8], based on the first accurate and precise Pb concentration measurements for seawater. Studies of the anthropogenic impact on marine Pb budgets were further advanced with the first successful Pb isotope analyses of seawater [6,9–13]. Whilst these early measurements were all carried out by TIMS (thermal ionization mass spectrometry), the majority of seawater Pb analyses since the 1990s have been conducted using various types of ICP-MS instrumentation. At present, the most precise Pb isotope analyses of seawater apply either single collector high resolution or multiple collector ICP-MS (HR-ICP-MS and MC-ICP-MS, respectively), most commonly following low-blank pre-concentration of the element by co-precipitation with Mg(OH)₂ [14,15].

These methodological advances are a consequence of the particular importance of seawater Pb isotope data to studies of environmental pollution and marine geochemistry. For example, analyses of corals yielded a seawater Pb isotope record for the North Atlantic over the last 220 years [12,15,16], which revealed significant isotopic variability through time, in accord with known changes in the extent and type of anthropogenic emissions. Similarly, Pb isotope data for the modern ocean are used to infer the distribution of Pb from different anthropogenic and natural sources in surface seawater [9,13,17,18] and to determine the lateral mixing and advection of subsurface water masses [19–23]. Shen and Boyle [12], furthermore, highlighted that temporal changes in the Pb content and isotope composition of different water masses can be applied to trace ventilation times.

In this paper, we present a new method for the accurate and precise determination of seawater Pb concentrations and isotope compositions. The methodology involves pre-concentration of Pb by Mg(OH)₂ co-precipitation [24] followed by ion exchange chromatography [25,26], and subsequent isotopic analyses by TIMS using a 204Pb–207Pb double spike to achieve precise control of instrumental mass fractionation [27,28]. The technique is validated by analyses of both in-house and international seawater reference materials and its utility demonstrated by data obtained for a GEOTRACES seawater depth profile from the South Atlantic Ocean.

A number of previous investigations [28–34] have shown that the application of Pb double or triple spikes enables Pb isotope measurements that are significantly more precise and accurate than conventional TIMS or MC-ICP-MS techniques and less susceptible to the analytical artifacts, which can be generated by instrumental mass bias effects. Our study is the first to apply the double spike methodology to seawater Pb isotope analyses. Using these techniques, we are not only able to analyze the commonly measured ratios 207Pb/206Pb, 208Pb/206Pb, 207Pb/204Pb to high precision but also 206Pb/204Pb, 207Pb/204Pb and 208Pb/204Pb can also be determined to an uncertainty of better than about ±3%. Only very scarce seawater data are available for the latter ratios due to the analytical challenges but measurements of non-radiogenic 204Pb are deemed to be important to fully exploit the potential of the Pb isotope system.

2. Experimental

2.1. Samples

A gravimetric solution of the well-characterized NIST standard reference material SRM 981 Pb was employed for calibration of the Pb double spike and to evaluate the precision of the mass spectrometric methods. Four filtered seawater samples that were available in large quantities were utilized as in house reference materials to further assess reproducibility. These encompass a surface water sample (Atlantic – S) and three subsurface water samples, Atlantic – D1, Atlantic – D2 (both from 2000 m depth), and Weddell (400 m depth). These samples were not specifically collected for Pb isotope analyses, and potential contamination during sampling and initial handling hence renders them unsuitable for evaluation of accuracy or an assessment of water mass properties.

To enable a comparison of our results with literature data for assessment of accuracy, we analyzed the international seawater reference samples GSI and GDI. These were collected at water depths of 7 m and 2000 m, respectively, at the Bermuda Atlantic Ocean Time Series station (BATS) during the 2008 GEOTRACES intercalibration cruise on the R/V Knorr (KN193-6). The collection and handling of these samples employed materials and procedures that were specifically designed for the intercalibration of contamination-prone trace metals and isotopes [35].

The seawater for a Pb isotope and concentration depth profile was collected in the South Atlantic Ocean off the African coast at 36.46°S, 13.39°E in October 2010 during the eastern part of the GEOTRACES GA10 transect along 40°S on the R.R.S. Discovery (D357; 18 October–22 November 2010) [1]. This depth profile from station 3 encompasses 12 samples that cover the entire 4500 m of the water column at the sampling location. Station 3 serves as a ‘cross over station’, which means that it was sampled by three separate GEOTRACES cruises (UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study; UK GEOTRACES cruise D357 – this study).

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2.2. Reagents

Water with a resistivity of >18.2 MΩ cm from a Milli-Q water system was used throughout for reagent preparation and cleaning of labware. Purified concentrated HCl (~28 M) and HBr (~8.5 M) were purchased from VWR as Optima (distilled) grade acids whilst concentrated (14.5 M) HNO₃ and 6 M HCl were prepared in-house by subboiling distillation of reagent grade acids in quartz stills. The Pb blanks of these acids were checked regularly and found to be consistently less than 2 pg/mL. The dilute mineral acids that were employed for the separation chemistry were freshly prepared for each batch of samples.

A purified aqueous solution of NH₃ was used to affect precipitation of Mg(OH)₂ from seawater and prepared by cold vapor phase equilibration of reagent grade 28% (w/w) aqueous NH₃ with Milli-Q water in a Savillex Teflon elbow. Approximately 2 volumes of 28% aqueous NH₃ were employed for each
volume of Milli-Q water, and these were equilibrated for about two weeks to obtain purified aqueous NH$_3$ (aq-NH$_3$) at a concentration of about 9 M. During this period, the Teflon elbow was stored double bagged in plastic, to limit contamination and loss of gaseous NH$_3$. The aq-NH$_3$ was collected from the still just prior to use and each batch of three samples, plus a full procedural blank, was treated with the same solution. The Pb blank of the aqueous ammonia was stable at less than 0.3 pg/mL during the course of the study.

2.3. Preparation and calibration of $^{207}$Pb–$^{204}$Pb double spike

Highly enriched (>99%) metallic $^{204}$Pb and $^{207}$Pb in wire form were purchased from Isoflex (USA). The wires were separately dissolved in 7.5 M HNO$_3$ and diluted with water to obtain solutions in 2 M HNO$_3$. A dilute double spike solution was then prepared with a total Pb concentration of ~10 ng/mL and an optimal molar ratio of $^{207}$Pb/$^{204}$Pb $\approx$ 1, based on the error propagation modeling of Galer [28] and Rudge [38].

Spike calibration was performed using a gravimetric solution of NIST SRM 981 Pb. In a first step, the concentration of the double spike was determined by two reverse isotope dilution runs that were carried out with a Nu Plasma HR MC-ICP-MS, using external normalization to added Tl for mass bias control [24,39,40]. Initially, a preliminary concentration was obtained, based on an approximate spike isotope compositions, which was calculated from the mixing proportions of the $^{204}$Pb and $^{207}$Pb solutions, and the respective isotope compositions, as supplied by Isoflex.

The isotope composition of the double spike (Table 1) was then calibrated by analyses of different gravimetrically prepared double spike – NIST SRM 981 mixtures using TIMS [38,41]. The instrumental mass fractionation correction [24,39,40], the spike-standard mixing line, and the double spike composition were calculated by over-determining the system of equations (number of double spike – NIST SRM 981 mixtures >2) and solving in the least-square sense, as described by Rudge et al. [38]. Finally, the calibrated double spike isotope composition was applied to the data from the initial reverse isotope dilution runs, to accurately define the double spike concentration.

2.4. Sample preparation for determination of Pb isotope compositions

2.4.1. Lead pre-concentration by co-precipitation with magnesium hydroxide

A number of previous studies have shown that co-precipitation of Pb with Mg(OH)$_2$ is an effective, low-blank method for pre-concentration of Pb from seawater [14,42] and this technique is retained in our procedure. Initial tests showed that Pb is efficiently and reproducibly extracted from seawater at yields >80% by precipitation with Mg(OH)$_2$ at pH $\approx$ 10. The yields were, furthermore, not correlated with the exact pH value that was adjusted or the amount of Mg(OH)$_2$ precipitated. The Mg(OH)$_2$ precipitate, however, was also found to scavenge a significant amount of sea salt, but the latter could be removed without noticeable reduction of the Pb yield by rinsing the precipitate with water.

In order to obtain sufficient Pb for precise analyses, even for Pb depleted seawater from the deep ocean, samples of up to about 2 L were processed. The seawater was transferred to 1 L pre-cleaned Teflon separating funnels, to aid the collection of the Mg hydroxide precipitate. Precipitation of Mg(OH)$_2$ was affected by introduction of the purified aq-NH$_3$ until a final pH value of about 10 was obtained. As each seawater sample had been acidified to a slightly different extent, and because the precipitate forms only gradually, the applied volume of aq-NH$_3$ was optimized for each sample by slowly adding successive portions of the reagent to the separating funnel. This was achieved by an initial addition of 2–4 mL aq-NH$_3$, whereupon the samples were stirred vigorously and then left to stand for at least 3 h to allow the Mg(OH)$_2$ precipitate to form. If precipitation had not occurred after this time, an additional volume of 0.5 mL aq-NH$_3$ was added every 2–3 h until formation of precipitate was observed.

Once about 5 mL of Mg(OH)$_2$ slurry had formed, the precipitate was drained through the bottom of the separating funnel, collected in a 15 mL Teflon centrifuge tube, centrifuged at 1700 rpm for about 10 min and the remaining solution poured off. This step was repeated several times until all precipitate was recovered from the funnel. The solids were then washed with 5 mL H$_2$O, centrifuged again and finally dissolved directly in the tubes by addition of 5 mL of a 2 M HBr–0.01 M HF acid mixture. To aid dissolution, the centrifuge tubes were shaken whereupon the formation of many small bubbles within the solutions was observed for seawater.

### Table 1

Lead isotope data of this study and reference values for NIST SRM 981 Pb. Also shown is the isotope composition of the Pb double spike used in this study.

| Sample          | Pb used (ng) | $^{207}$Pb/$^{204}$Pb ±2sd | $^{208}$Pb/$^{204}$Pb ±2sd | $^{209}$Pb/$^{204}$Pb ±2sd | $^{206}$Pb/$^{204}$Pb ±2sd | $^{207}$Pb/$^{206}$Pb ±2sd |
|-----------------|-------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| This study      |             |                             |                             |                             |                             |                             |
| SRM 981 Pb      | 2           | 16.943 ± 16                 | 15.496 ± 15                 | 36.716 ± 38                 | 0.91460 ± 18                | 2.16710 ± 61                |
|                  | 42          | 920                         | 970                         | 1050                        | 200                         | 280                         |
|                  | ±2sd relative (ppm) | ±2sd of typical (ppm)$^a$ | ~500                        | ~500                        | ~60                         | ~80                         |
| SRM 981 Pb      | 10          | 16.9419 ± 31                | 15.4968 ± 38                | 36.7224 ± 91                | 0.91470 ± 11                | 2.16755 ± 20                |
|                  | 25          | 180                         | 240                         | 250                         | 120                         | 90                          |
|                  | ±2se of typical (ppm)$^a$ | ±2se relative (ppm)$^a$     | ~90                         | ~100                        | ~40                         | ~60                         |
| SRM 981 Reference values$^a$ | ~10          | 16.9405 ± 15                | 15.4963 ± 16                | 36.7219 ± 44                | 0.91475 ± 4                 | 2.16771 ± 10                |
| Galer and Abouchami (TIMS) [57] | ~20          | 16.9409 ± 22                | 15.4956 ± 26                | 36.7228 ± 80                | 0.91469 ± 7                 | 2.16770 ± 21                |
| Todt et al. (TIMS) [58] | 80           | 16.9356 ± 23                | 15.4891 ± 30                | 36.7006 ± 113               | 0.91459 ± 13                | 2.16701 ± 43                |
| Thirwell (MC-ICP-MS) [32] | ~50          | 16.9417 ± 29                | 15.4996 ± 31                | 36.724 ± 9                  | 0.91488 ± 8                 | 2.16770 ± 24                |
| Baker et al. (MC-ICP-MS) [29] | ~50          | 16.9416 ± 13                | 15.5000 ± 13                | 36.7262 ± 31                | 0.91491 ± 4                 | 2.16781 ± 12                |

$^a$ Mass of Pb loaded on filament and/or used for a single analysis.

$^b$ n = number of analyses.

$^c$ 2sd = 2 x standard deviation calculated from n separate runs; denotes long-term “external” reproducibility.

$^d$ 2se = 2 x standard error of the mean; denotes within-run (“internal”) precision of the data, based on individual results for measurement cycles.

$^e$ All reference values were obtained using either double or triple spike techniques for correction of instrumental mass fractionation.

$^f$ Isotope composition of the Pb double spike used in the current study.
samples that had not been filtered following collection. Bubble formation is hence presumably related to the presence of residual organic compounds. It is important to remove the bubbles at this stage because they can otherwise lead to the formation of larger bubbles within the resin bed during the subsequent column chemistry, effectively blocking the flow. For removal of the bubbles, the centrifuge tubes were left to stand overnight, sealed and then gently heated on a hotplate for 10 min at 50 °C. After cooling, bubbles should no longer be visible in the solutions, otherwise the heating was repeated.

2.4.2. Lead purification by anion exchange chromatography

The anion exchange purification procedure for Pb (Table 2) is based on previously published methods that apply strongly basic Mg(OH)₂ precipitate, which is used for the separation of Pb from seawater. Relatively high Si concentrations of about 10–150 μmol/kg are, furthermore, found in seawater samples from depths >1000 m [45,46]. During method development we found that such concentrations suffice to form precipitates of silica on top of the resin column, when HF is not present in sample solutions. Hence, the 1st stage column chemistry of our final method (Table 2) applies a 2 M HBr–0.01 M HF mixture for sample loading, whereby the small concentration of HF present enhances silica solubility to prevent precipitation of silicic acid. The same acid mixture is subsequently used to completely flush silica from the resin, prior to elution of other matrix elements (Table 2). In 2 M HBr, Pb has a distribution coefficient of about 200 toward the AG1X8 anion exchange resin that was used [25,47,48]. Elution experiments confirmed that, whilst the Pb Kᵢ value is reduced when HF is present, the use of trace HF in the elution scheme (Table 2) does not prevent retention of Pb on the resin bed during sample loading and matrix elution.

Following elution of Pb from the 1st stage columns, the samples were dried and further purified by a second column chemistry step, which is needed to eliminate remaining traces of Mg [49] and improve the ionization of Pb during the TIMS analyses. To this end, the dried samples were redissolved in 0.2 M HBr–0.5 M HNO₃ and processed using established HBr–HNO₃ procedures on small 20 μL resin columns (Table 2), which help to minimize the procedural blank.

2.5. TIMS analyses of Pb isotope compositions

The accurate determination of Pb isotope compositions with precise correction of instrumental mass discrimination requires two separate mass spectrometric runs – one of the pure sample and a second analysis of a double spike – sample mixture. To this end, the purified Pb was divided into two aliquots and the double spike was admixed to one, prior to the TIMS measurements. The addition of the double spike after, rather than before, the chemical separation of Pb has a number of advantages. Conveniently, each sample needs to be processed through the (time-consuming) pre-concentration and purification steps only once, rather than twice. Furthermore, the impact of contamination from reagents and handling is minimized and uncertainties arising from different blanks impacting on the spiked and unspiked analyses are reduced.

In detail, our procedure involves splitting the Pb solutions from the 2nd stage column chemistry into a larger and a smaller aliquot. The larger aliquot comprising 2/3 of the sample remained unspiked, whilst Pb double spike was added to the remaining aliquot with 1/3 of the solution. The double spike addition was thereby carried out to obtain a molar ratio of natural (N) to spike-derived Pb (S) that approximates the optimum S/N = 1.226 [38]. This is equivalent to a molar proportion of spike, S/(S + N), of 55.1%, corresponding to 206Pb/204Pb = 0.411 (assuming a natural seawater Pb/204Pb = 18.4). The amount of natural Pb present was thereby estimated from the Pb concentration measurements that were carried out first for all samples, and assuming a Pb recovery of 60% from the chemical separation (see below).

Following aliquoting and spiking, the Pb solutions were evaporated to dryness and redissolved in 1 μL 2 M HNO₃. The samples were loaded on a single, previously outgassed zone refined Re filament (99.999%; H. Cross, USA), dried at 0.4 A, 0.5 μL of silica–gel activator was then added and also dried at 0.4 A. The sample–activator mixture was finally heated at 2 A for ~1 s to evaporate phosphoric acid and fuse the silica gel.

Gerstenberger and Hasse [50] prepared a highly successful activator for Pb isotope analyses by TIMS but this applies silica particles which are no longer commercially available. Hence, a new SiO₂–H₃PO₄ activator mixture was made up for the current study. This mixture was prepared from SiO₂ nanoparticles (20–60 nm; Io-Li-Tec Inc.) and Merck Ultrapur phosphoric acid, whereby a SiO₂/H₃PO₄ weight ratio of 0.43% was applied with respect to the addition of the silica to dilute 0.16 M H₃PO₄. The amount of activator used for analysis was calibrated to optimize ionization yields, based on tests that were carried out with 2 ng NIST 981 Pb. These test revealed that use of 0.5 μL activator solution was preferable to loading with 1 μL of activator – whilst the latter yielded more stable Pb ion beams the former produced ion beam intensities that were typically about 2–3 times higher.

The Pb isotope measurements were carried out on a TRITON TIMS instrument operating in static mode and using Faraday cups.

| Table 2 |
| --- |
| Column chemistry procedure for the separation of Pb from seawater for isotope compositions (IC) and isotope dilution (ID) concentration measurements by TIMS |

| Pb IC chemistry 1st stage | Pb IC chemistry 2nd stage | Pb ID chemistry |
| --- | --- | --- |
| Resin⁴ | 100 μL AG1X8, 100–200 mesh | 20 μL AG1X8, 100–200 mesh | 20 μL AG1X8, 100–200 mesh |
| Clean resin | 3 × 3 mL 0.1 M HNO₃ | 3 × 1 mL 0.1 M HNO₃ | 3 × 1 mL 0.1 M HNO₃ |
| Equilibrant resin | 2 × 0.1 mL | 2 × 0.1 mL | 2 × 0.1 mL |
| Load sample | 2 M HBr–0.01 M HF | 0.2 M HBr–0.5 M HNO₃ | 0.5 M HBr–0.2 M HNO₃ |
| Elute matrix | ~5 mL in | ~1 mL in | 2 mL |
| Elute Pb | 2 × 0.1 mL | 2 × 0.1 mL | 2 × 0.1 mL |
| 2 M HBr–0.01 M HF | 0.2 M HBr–0.5 M HNO₃ | 0.5 M HBr–0.2 M HNO₃ |
| 2 × 0.5 mL | 2 × 0.3 mL | 2 × 0.3 mL |
| 0.2 M HBr–0.01 M HNO₃ | 0.2 M HBr–0.5 M HNO₃ | 0.5 M HBr–0.2 M HNO₃ |
| 0.2 mL | 0.2 mL | 0.2 mL |
| 0.03 M HBr–0.5 M HNO₃ | 0.03 M HBr–0.5 M HNO₃ | 0.03 M HBr–0.5 M HNO₃ |
| Elute Pb | 3 mL | 1 mL | 1 mL |
| 0.03 M HBr–0.5 M HNO₃ | 0.03 M HBr–0.5 M HNO₃ | 0.03 M HBr–0.5 M HNO₃ |

⁴ The resin is used in columns prepared from shrink-fit Teflon tubing. The columns have a reservoir volume of ~3 mL and a resin column with an internal diameter of ~3 mm.
fitted with 1011 Ω resistors. Prior to each measurement session, a gain calibration was carried out for all amplifiers. For analysis, the filaments were heated to a target temperature of about 1300 °C within 30 min, and isotopic data were then collected in 15 blocks of 15 cycles with an integration time of 8.3 s. Half-mass baseline measurements (30 cycles × 1.05 s integrations following 5 s idle time) and automatic peak centering routine (followed by 3 s idle time) were carried out prior to each block. Loads of 2 ng NIST 981 Pb provided stable and very reproducible ion beam intensities of 200–300 mV 206Pb. For the seawater samples, the ion beam intensities were more variable and generally lower, whereby 2 ng Pb generally yielded 208Pb ion beams of about 100–200 mV.

The measured isotopic data for the spiked and unspiked sample aliquots were processed using an iterative solver that employs published methods [38] and which was implemented as a macro within a Microsoft Excel spreadsheet. The method uses the exponential law to obtain mass bias corrected Pb isotope data and is employed on a cycle-by-cycle basis to provide an accurate assessment of the final propagated data uncertainties.

Following Doucelance and Manhès [51], the TIMS source was baked for 3 h after the introduction of a new sample turret. This technique was observed to substantially reduce the formation of hydrocarbon species that can form inferring molecular ions during the Pb isotope runs [31]. The baking also significantly improved the vacuum within the ion source (to about 3.5 to 6 × 10⁻⁵ mbar, without use of a liquid N₂ cold trap) and enhanced Pb ion beam stability, thereby permitting analyses without interblock re-heating. The latter improvement is presumably due to the efficient drying of the silica gel, which is achieved during heating in vacuum condition.

Barium-Po₄ ions were mainly observed at temperatures, which are slightly higher compared to those employed for optimal Pb ionization [31]. The formation of such ions, and hence the presence of Ba in samples, is best avoided because they produce minor molecular interferences on 204Pb (from 138Ba³¹P₁⁷O₁⁸O⁻ 137-Ba²¹P₁⁸O⁻²), and, even more problematic, hinder the ionization of Pb whereby rapidly decaying Pb ion beams are observed. The presence of BaPo₄ ions was routinely monitored during the analyses at mass 201, which corresponds primarily to ¹³⁸Ba²¹P₁⁸O⁻². The measured beam intensities at mass 201 were observed to rise during Pb sample runs but only to maximum values of 1–3 mV. At this level, the BaPo₄ interference contributes less than 15 ppm to the 204Pb ion beam, such that a correction was deemed unnecessary.

2.6. Determination of Pb concentrations

Lead concentrations were determined by isotope dilution (ID), and using a chemical separation method similar to that employed for the isotope ratio measurements but optimized for the processing of smaller seawater samples. In detail, 50 mL of seawater were weighed into clean polyethylene centrifuge tubes. Following addition of the Pb double spike, the mixtures were left to stand for at least 10 days to ensure full spike–sample equilibration. The amount of Pb double spike was added based on an initial ‘guess’ of any information available for the Pb concentration of the samples, to achieve an optimal S/N ≈ 2.61 (equivalent to 206Pb/204Pb ≈ 0.188) for minimal error propagation during ID data reduction.

Co-precipitation of Pb with Mg(OH)₂ was affected by adding 150 μL of purified aq-NH₃. Following separation of the precipitate by centrifugation and dissolution in a dilute HNO₃–HBr mixture, Pb was further purified using a single stage anion exchange chemistry. This employed small resin columns of only 20 μL to minimize the procedural blank (Table 2). The anion exchange chemistry was slightly modified in comparison to the 2nd stage chemistry for Pb isotopes through use of a 0.5 M HBr–0.2 M HNO₃ acid mixture for sample loading and matrix elution, because this was observed to facilitate the dissolution of the Mg(OH)₂ precipitate whilst ensuring quantitative retention of Pb on the small resin columns.

The ID TIMS analyses for Pb used the same techniques that were applied for the isotopic measurements but with shorter runs comprising 5 blocks of 15 cycles. During filament loading of some samples for the ID TIMS analyses, the presence of impurities, in the form of salts and possibly also organic compounds, was detected. When present, such impurities were associated with poor Pb ion beam intensities during the TIMS runs and in some extreme cases the analyses were even abandoned. Such contamination may also produce interfering molecular ions that could generate inaccurate ID results. Subsequent test showed that the impurities can be reduced to negligible levels, by allowing the resin bed to drain and dry completely (for about 10 min) after each step of the column chemistry procedure for Pb separation (Table 2) and, in particular, prior to the elution of Pb.

3. Results and discussion

3.1. Blanks

The total blank for the Pb isotope measurements was determined by carrying out the complete analytical procedure without an initial seawater sample being present. Over a 24-month period, the mean procedural blank was 28 ± 21 pg (1sd, n = 25; ignoring two abnormally high blanks of ~100 pg encountered during a single two-week period). The individual contributions to this total from (i) filament loading and (ii) reagents were also determined and found to be 1.9 ± 0.5 pg for the former, 5.0 ± 1.4 pg for the mineral acids used, and 2.5 ± 0.2 pg for the aq-NH₃ solutions. The contribution of the anion exchange resin column chemistry (without reagents) to the total blank could hence be quantified as 17 ± 8 pg, by subtracting the reagent component from separately measured column chemistry blanks. The sum of the individual blank contributions is, therefore, 27 ± 10 pg, in accord with the full procedural blanks values determined during the study. Ignoring the individual contributions (and chemistry yields), a Pb blank of 10–50 pg contributes about 0.05–2.5% to the indigenous Pb budget of samples that were analyzed for isotopic compositions, assuming samples volumes and Pb concentrations of 1–2 L and 10–50 pmol/kg, respectively. The Pb isotope composition of the blank was also measured on several occasions with rather variable results, due to the low ion beam intensities and/or actual differences in blank compositions (e.g., 208Pb/206Pb = 18.34 ± 0.62, 207Pb/206Pb = 1.196 ± 0.028, 206Pb/208Pb = 2.42 ± 0.021; 2se, n = 10). Blank corrections were not applied to the isotopic data because such corrections did not improve the reproducibility of results from replicate sample analyses (see discussion below).

The total procedural blank for the Pb ID concentration measurements was determined using 100 μL seawater samples, as only negligible amounts of Pb (~0.2 to 1 pg) are present in such small volumes. These measurements revealed a relatively stable total Pb blank of 11.7 ± 4.3 pg (1sd, n = 36), which is equivalent to about 1–5% to the indigenous Pb in ID sample aliquots, assuming Pb abundances of ~10–50 pmol/kg (for simplicity, this ignores variable chemistry yields and individual blank contributions). Blank corrections were applied to all measured Pb concentrations based on the mean blank value. The uncertainties of the Pb concentration data propagate both the (within-run) precision of the isotopic analyses and the uncertainty of the blank correction.
3.2. Chemical separation of Pb

Stable isotope fractionation of Pb during anion exchange chromatography with HCl was recently documented and shown to vary systematically with Pb recovery [29,52]. Such fractionation can have a detrimental impact on the accuracy of subsequent Pb isotope measurements and may also occur during Pb purification with other anion exchange elution protocols. As part of this study, it was, therefore, investigated whether incomplete recovery of Pb from the anion exchange chemistry applied here (Table 2) also has a systematic impact on the measured Pb isotope compositions. To this end, multiple 10 ng aliquots of NIST SRM 981 Pb were processed through the complete 2-stage column separation procedure and the purified Pb fractions were then analyzed using standard methods. The data (Fig. 1) illustrate that the column chemistry routinely affords Pb yields of about 50–70%. Such recoveries are in accord with the total Pb yields of about 30–60% that were determined for multiple analyses of five different seawater samples (Tables 3 and 4). Assuming that the Mg(OH)2 co-precipitation achieves a recovery of >80% for seawater Pb (see above), the Pb yield of the column chemistry is constrained to be about 40–75%.

In a previous investigation, Reuer et al. [15] reported essentially complete recovery of seawater Pb pre-concentrated by Mg(OH)2 co-precipitation, during purification on AG1X8 resin (as in this study) but using HBr and HCl for elution [49]. Test conducted during the course of the present study showed, however, that the method of Reuer et al. [15] was associated with Pb recoveries similar to those reported here, when the procedure was modified to include additional use of HF to prevent precipitation of silicic acid during sample loading (Table 2). The use of dilute HF thus appears to be associated with either incomplete Pb retention on or elution from the anion exchange resin, regardless of the exact nature of the mineral acids, which are applied in the procedure. Ultimately, the current HBr-HNO3 method (Table 2) was preferred to the elution method of Reuer et al. [15], because the former applies only dilute mineral acids (which helps to minimize the blank) and routinely produced cleaner Pb separates for subsequent isotopic analyses.

![Fig. 1. The recovery of Pb and \(^{206}\text{Pb}/^{204}\text{Pb}\) ratios obtained for 10 ng aliquots of NIST SRM 981 Pb that were processed through the 2-stage column chemistry (Table 2) are shown as full red circles with error bars that denote a ±2sd uncertainty of about 240 ppm (Table 3). Shown for comparison are literature data for cumulative recovery and isotope composition of SRM 981 Pb that was collected in successive fractions from anion-exchange resin which was eluted with HCl: full blue squares and solid line–elution with 6 M HCl; open blue squares with hashed line–elution with 2 M and 8 M HCl [Baker 29]. The dashed black lines indicate the \(^{206}\text{Pb}/^{204}\text{Pb}\) reference value determined for SRM 981 Pb (15.496 ± 16; Table 1). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)](image)

**Table 3.** Lead concentration and isotope composition data for the four in-house seawater reference materials and other key information relating to samples.

| Sample | Lead concentration (pmol/kg) | Lead isotope composition \(\delta^{206}\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb}/\delta^{204}\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb} \text{Pb}^\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb}/\delta^{204}\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb} | Pb yield (%) | Measured Pb (mean) ng | Sample size Le (g) | Pb yield (40–60) % | Pb yield (30–50) % | Pb yield (10–70) % | Pb yield (8–20) % | Pb yield (5–10) |
|--------|-----------------------------|-------------------------------------------------|---------------|----------------------|-------------------|-------------------|-------------------|------------------|-----------------|-----------------|
| Atlantic D1 | 6.0 ± 3.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Atlantic D2 | 2.0 ± 1.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Wedde | 8.0 ± 4.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Meas. Pb (mean) ng | Pb yield (%) | Measured Pb (mean) ng | Sample size Le (g) | Pb yield (40–60) % | Pb yield (30–50) % | Pb yield (10–70) % | Pb yield (8–20) % | Pb yield (5–10) |
| Atlantic D1 | 6.0 ± 3.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Atlantic D2 | 2.0 ± 1.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Wedde | 8.0 ± 4.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |

- **Sample**: Sample ID for individual seawater reference materials.
- **Lead concentration (pmol/kg)**: Lead concentration in the samples.
- **Lead isotope composition \(\delta^{206}\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb}/\delta^{204}\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb} \text{Pb}^\text{Pb}^\text{Pb}^\text{Pb}/\delta^{204}\text{Pb}^\text{Pb}^\text{Pb}^\text{Pb} | Pb yield (%) | Measured Pb (mean) ng | Sample size Le (g) | Pb yield (40–60) % | Pb yield (30–50) % | Pb yield (10–70) % | Pb yield (8–20) % | Pb yield (5–10) |
| Atlantic D1 | 6.0 ± 3.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Atlantic D2 | 2.0 ± 1.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |
| Wedde | 8.0 ± 4.0 | 207 ± 14 | 15.637 ± 0.015 | 2.097 ± 0.025 | 25 | 0.84862 ± 0.004 | 40 | 60 | 10–70 | 8–20 | 5–10 |

- **Pb yield (%)**: Percentage of Pb recovery in the chemical separation process.
- **Measured Pb (mean) ng**: Mean Pb concentration in individual aliquots used for isotopic analyses.
- **Sample size Le (g)**: Size of the sample leachate used for Pb isotope analyses.
- **Pb yield (40–60) %**: Estimated Pb yields ranging from 40% to 60%.
- **Pb yield (30–50) %**: Estimated Pb yields ranging from 30% to 50%.
- **Pb yield (10–70) %**: Estimated Pb yields ranging from 10% to 70%.
- **Pb yield (8–20) %**: Estimated Pb yields ranging from 8% to 20%.
- **Pb yield (5–10) %**: Estimated Pb yields ranging from 5% to 10%.

**Note:**
- \(\delta^{206}\text{Pb}/\delta^{204}\text{Pb}\) values are corrected for mean total blank of 11.7 ± 0.3 ppm.
- Uncertainties in isotope data are reported as 2 standard deviations (2sd).
- The uncertainty in Pb concentration data is reported as 2 standard deviations (2sd).
| Sample | n-ID | [Pb] ± 1sd pmol/kg | n-IC | 206Pb/204Pb ± 2sd | 207Pb/204Pb ± 2sd | 208Pb/204Pb ± 2sd | 208Pb/206Pb ± 2sd | Sample size L² | Total Pb ng | Pb yield (%) |
|--------|------|-------------------|------|---------------------|---------------------|---------------------|---------------------|----------------|-------------|-------------|
| GSI    | 6    | 27.9 ± 1.2        | 3    | 18.359 ± 26         | 15.635 ± 7          | 38.246 ± 30         | 0.85162 ± 93      | ~2             | ~10         | 45–60       |
|        | ±2sd relative (ppm) | 1430 | ~160 | 1430 ± 40          | 980 ± 70             | 3820 ± 200         | 0.849 ± 10        | ~8             | ~20         | 40–60       |
| Ref. – MIT | 5   | 45.7 ± 2.6        | 2    | 18.475 ± 41         | 15.624 ± 10         | 38.276 ± 60         | 0.84568 ± 134     | ~2             | ~17         | 50–55       |
|        | ±2sd relative (ppm) | 9320 | ~190 | 1430 ± 60          | 980 ± 1570           | 3820 ± 1570        | 0.849 ± 10        | ~8             | ~20         | 40–60       |
| Ref. – UCSC | 4  | 29.5 ± 2.1        |      | 18.46 ± 0           | 15.68 ± 1           | 38.36 ± 7           | 0.84929 ± 50      | ~2             | ~17         | 50–55       |
| Ref. – Consensus | 4  | 42.2 ± 1.3        |      | 18.54 ± 3           | 15.69 ± 2           | 38.43 ± 10          | 0.84640 ± 42      | ~8             | ~20         | 40–60       |

The Pb isotope reference data (Ref.) are from the labs of Ed Boyle at the Massachusetts Institute of Technology (MIT) and Russ Flegal at the University of California Santa Cruz (UCSC) [35]. Results in italics were calculated from data given in the literature. The ‘Consensus’ Pb concentrations are from [53].

- Number of individual sample aliquots analyzed for Pb concentrations.
- Pb concentrations are corrected for a mean total blank of 11.7 ± 4.3 pg. The uncertainties of the concentration data propagate both the (within-run) precision of the isotopic analyses and the uncertainty of the blank correction (see text).
- Number of individual sample aliquots analyzed for Pb isotope compositions.
- 2sd = 2 × standard deviation calculated from results of n separate runs; denotes long-term “external” reproducibility.
- Volume of individual aliquots used for Pb isotope analyses.
- Original mass of Pb in sample aliquots used for Pb isotope analyses.
- Pb recovery for the chemical separation process.
- 2se = 2 × standard error of the mean; denotes within-run (“internal”) precision of a run, based on individual results for measurement cycles.
- Duplicate analyses of Pb separated from a single seawater sample.
Importantly, the isotope data of Fig. 1 demonstrate that incomplete recovery of Pb from the column separation is not associated with significant shifts in isotope composition. These results do not, however, rule out small, but barely resolvable, enrichments of heavier Pb isotopes following chemical purification. This follows from the observations of previous studies, which detected only minor Pb isotope fractionations, comparable to or smaller than the reproducibility of our results, as a result of incomplete elution of the element from anion exchange columns (Fig. 1).

3.3. Analyses of NIST SRM 981 Pb

Multiple measurements of 2 ng (n = 42) and 10 ng (n = 25) loads of NIST SRM 981 Pb were carried out for a basic validation for our double spike techniques of Pb isotope analysis. Notably, the mean isotopic data acquired for both the 2 ng and 10 ng runs are in excellent agreement with recent reference values that were acquired using various Pb double or triple spikes in conjunction with analyses by either TIMS or MC-ICP-MS (Table 1).

The 2σ between-run (‘external’) precision of the 10 ng data is ~100 ppm for 207Pb/206Pb, 208Pb/206Pb and ~250 ppm for 204Pb/206Pb, where x = 206Pb, 207Pb, 208Pb hereafter. This is similar to or only slightly worse than the reproducibility achieved by other workers with TIMS or MC-ICP-MS for comparable or somewhat larger Pb samples (10–80 ng; Table 1). Our data for the 2 ng loads, a sample size that is more relevant for seawater analyses (Table 3), is less precise than the 10 ng results, with a reproducibility (~2σ external) of ~250 ppm for 207Pb/206Pb, 208Pb/206Pb and ~1000 ppm for 204Pb/206Pb (Table 3). The significantly larger uncertainty of the 2σ between-run precision (quantified by 2σ; standard error of the mean) of both the 2 and 10 ng Pb measurements are about a factor of 2–3 better compared to the between-run uncertainties for all Pb isotope ratios (Table 1).

3.4. Analyses of in-house seawater reference samples

Multiple Pb concentration and isotope ratio analyses were carried out on four different in-house seawater reference samples (Table 3). These were available in sufficient quantity such that 5–7 repeat isotopic analyses could be carried out to rigorously investigate the reproducibility of the methods.

For the Pb contents, the precision (expressed as ±1σ, as is common for concentration data) varies between less than ±1% to about ±12% (Table 3). Notably, the reproducibility is worse for samples with low Pb contents. As similar samples sizes (of only ~50 ml) were employed for these measurements, the higher uncertainties for the Pb-depleted samples are most likely related to inaccuracies of the Pb blank correction, possibly due to spurious high blanks.

The isotopic results for the in-house seawater reference samples (Table 3) were acquired on individual aliquots of 1–2 L, from which about 2–10 ng of Pb were separated. The data display a ±2σ reproducibility of ~500–1500 ppm for 207Pb/206Pb, 208Pb/206Pb and ~500–2000 ppm for 204Pb/206Pb (Table 3). Clearly, this is inferior to the 2σ precision that was obtained for the comparable 2 ng and 10 ng loads of pure NIST SRM 981 Pb (Table 1). At the same time, the internal (within-run) ±2σ precision of the seawater measurements is similar to the NIST SRM 981 results, with uncertainties of ~40–100 ppm for 207Pb/206Pb, 208Pb/206Pb and ~300–700 ppm for 204Pb/206Pb (Table 3). Given this discrepancy, it is most likely that the external 2σ reproducibility of the seawater Pb isotope data is limited by contamination of the samples by blanks of both variable magnitude and isotope composition. This interpretation is supported by two observations. First, the Weddell seawater, which has the lowest Pb concentration and is also the sample with the smallest Pb sample sizes, generally features the largest 2σ uncertainties (Table 3). Second, the application of a blank correction did not improve the 2σ reproducibilities for any of the samples, regardless of whether the corrections used mean blank values or only blank data that were linked to a specific batch of samples. In addition, it is also conceivable that the between-run 2σ uncertainties of the samples (Table 3) are, at least in part, limited by minor isotope fractionations that occur during the column chemistry (Fig. 1).

3.5. GEOTRACES intercalibration samples GSI and GDI

Multiple Pb isotope and concentration measurements were also carried out for the two GEOTraces intercomparison samples GSI and GDI (Table 4). For the Pb concentrations, GEOTRACES ‘consensus values’ were recently published, which are based on analyses of more than 10 independent laboratories [53]. The mean Pb concentrations determined for the samples in this study from analyses of five or separate aliquots have ±1σ external reproducibilities that are similar to those obtained for the in-house seawater reference samples (Table 3), at about ±4 to ±6σ (Table 4). The results are, furthermore, identical to the reference values within the quoted uncertainties (Table 4).

The Pb isotope data for GSI and GDI were obtained on 2–3 sample aliquots of 2 L each, with originally about 10–15 ng Pb (Table 4). Following chemical separation, about 5–10 ng Pb were available for isotopic analysis. As for the seawater results of Table 3, the GSI and GDI measurements achieved (within-run) 2σ repeatabilities that are similar to those obtained for similar loads of pure SRM 981 Pb, at 30–60 ppm for 207Pb/206Pb, 208Pb/206Pb and 150–200 ppm for 204Pb/206Pb (Table 4). In contrast, the external 2σ uncertainties, of ~650–1500 ppm for 207Pb/206Pb, 208Pb/206Pb and ~500–2200 ppm for 204Pb/206Pb, are typically significantly larger (Table 4). Again, this is most likely due to small Pb blank contributions of variable magnitude and isotope compositions and, potentially, minor isotope fractionation during Pb purification. Despite of this, our results for the two samples compare favorably and are generally in agreement with the published reference results.

The reference Pb isotope data for the GEOTRACES samples were obtained at MIT (Massachusetts Institute of Technology) using MC-ICP-MS and at UCSC (University of California at Santa Cruz) by HR-ICP-MS [33]. These analyses did not apply a Pb double spike and were carried out using significantly smaller samples sizes (of about 0.25–0.5 L) compared to those utilized here.

For GSI, all Pb isotope ratios determined are identical to the MIT results within the 2σ uncertainties. There is also good agreement with the UCSC results for 207Pb/206Pb, 208Pb/206Pb but the UCSC data are higher for 204Pb/206Pb (Table 4). The uncertainties of the UCSC data are only poorly characterized, however, as they are based on only two analyses of a single seawater aliquot. Our new isotopic data for GDI are in excellent agreement with the MIT and UCSC results for 207Pb/206Pb, 208Pb/206Pb and in reasonable agreement, just within uncertainty, with the UCSC 208Pb/206Pb and 204Pb/206Pb ratios (Table 4). Hence, there are small but significant discrepancies for the remaining 204Pb/206Pb data with the reference values, whereby our ratios are slightly lower. There is, however, also only marginal agreement between the MIT and UCSC results for these Pb isotope ratios, which are difficult to determine due to the low natural abundance of 204Pb. This indicates that further analyses are needed to establish more reliable reference data for the 204Pb/206Pb ratios of the GEOTRACES intercalibration samples.
The particular difficulty of $^{204}$Pb measurements is also apparent in a comparison of the reproducibilities for the Pb isotope data obtained in this study and previously [35]. For the $^{206}$Pb/$^{204}$Pb ratios, the multiple MIT analyses reveal a ±2sd uncertainty of ~500–10,000 ppm, whilst our data for GSI, GDI (Table 4) and the in-house test samples (Table 3) have uncertainties which are at least a factor of 5 better, at ~500–2000 ppm. For $^{207}$Pb/$^{206}$Pb and $^{208}$Pb/$^{206}$Pb the difference is less apparent, however, as the reproducibility of the MIT results varies between 500 and 2500 ppm, whilst the data of this study (Tables 3 and 4) typically have an uncertainty of about 500–1500 ppm.

3.6. Depth profile for Pb isotopes and concentrations in the eastern South Atlantic Ocean

Twelve seawater samples from a depth profile in the eastern part of the GEOTRACES GA10 section in the South Atlantic Ocean were thus analyzed (Fig. 2a and Table 5), to (i) demonstrate that our new methodology is suitable for routine acquisition of high precision seawater Pb isotope data and (ii) highlight the important insights, which can be gained from such results. Lead concentrations are highest in the uppermost part of the water column with a surface (~3 m water depth) concentration of 17.7 pmol/kg, increasing to a relative maximum of 25.2 pmol/kg at 49 m and decreasing again to 22.4 pmol/kg at 397 m depth. From 400 m downwards, the Pb concentrations decrease, until they reach relatively constant and low values of 5–10 pmol/kg below 2000 m (Fig. 2a). A comparison with the dissolved Pb concentrations for a depth profile collected at the exact same location during the second expedition along the South Atlantic GA10 transect (cruise JC068; same station; Schlosser et al., pers. comm.) demonstrates excellent agreement for the entire water column. In contrast, data reported from IPY cruise MD166 (Boye et al. [36]) for the same location show significant deviations from our results, as the dissolved Pb concentrations are elevated by up to a factor of 2 and they define a pattern which is less smooth with depth.

Overall, our Pb concentrations for seawater from the southeast Atlantic Ocean, sampled in 2010, are significantly lower than the values of 29.5 pmol/kg and 42.2 pmol/kg that were reported for 2008 samples from the surface and 2000 m depth at the BATS station in the northwest Atlantic Ocean (Table 4). Lead concentrations in seawater have been strongly affected by anthropogenic Pb emissions over the past century. Such emissions were generally higher in the northern Hemisphere compared to southern regions but peaked in the 1970s. Since then, they have decreased by almost a factor of 10 due to the phasing out of leaded gasoline, first in the US and then in Europe (e.g., Kelly et al. [16]). Consequently, northwest Atlantic surface waters now contain less anthropogenic Pb (and exhibit a different isotopic composition) than deeper North Atlantic waters, which were contaminated in the past [13,19].

Table 5

| Sample | Depth (m) | [Pb] pmol/kg | ±1sd | $^{206}$Pb/$^{204}$Pb | $^{207}$Pb/$^{204}$Pb | $^{208}$Pb/$^{204}$Pb | $^{207}$Pb/$^{206}$Pb | $^{208}$Pb/$^{206}$Pb |
|--------|-----------|--------------|------|---------------------|---------------------|---------------------|---------------------|---------------------|
| Fish 113 | 2–3 | 17.7 | 0.9 | 18.073 | 15.620 | 38.030 | 0.86428 | 2.1040 |
| 183 | 25 | 20.5 | 0.4 | 18.111 | 15.624 | 38.017 | 0.86266 | 2.0991 |
| 180 | 49 | 25.2 | 0.5 | 18.013 | 15.618 | 37.922 | 0.86707 | 2.1053 |
| 177 | 99 | 22.4 | 0.8 | 18.044 | 15.614 | 37.941 | 0.86533 | 2.1027 |
| 174 | 198 | 17.3 | 0.6 | 18.125 | 15.627 | 38.015 | 0.86221 | 2.0975 |
| 171 | 397 | 22.3 | 0.6 | 18.087 | 15.618 | 37.970 | 0.86350 | 2.0993 |
| 170 | 594 | 15.7 | 0.5 | 18.031 | 15.600 | 37.885 | 0.86518 | 2.1011 |
| 169 | 989 | 11.5 | 0.5 | 18.007 | 15.604 | 37.911 | 0.86555 | 2.1052 |
| 167 | 1975 | 9.6 | 0.4 | 18.234 | 15.623 | 38.159 | 0.86577 | 2.0927 |
| 166 | 2955 | 5.3 | 0.4 | 18.418 | 15.717 | 38.452 | 0.85331 | 2.0879 |
| 164 | 3931 | 6.1 | 0.4 | 18.370 | 15.625 | 38.266 | 0.85050 | 2.0830 |
| 163 | 4724 | 7.1 | 0.4 | 18.540 | 15.664 | 38.537 | 0.84492 | 2.0785 |

a Pb concentrations are corrected for a mean total blank of 11.7 ± 4.3 pg.

b The uncertainties of the concentration data propagate both the (within-run) precision of the isotopic analyses and the uncertainty of the blank correction (see text).

c For the Pb isotopes, assumed external 2sd reproducibilities of ±1.250 ppm for $^{206}$Pb/$^{204}$Pb and of ±800 ppm for $^{207}$Pb/$^{206}$Pb, $^{208}$Pb/$^{206}$Pb are larger than the ±2se internal uncertainties in all cases. Based on this, appropriate 2sd uncertainties are ±0.0023 for $^{206}$Pb/$^{204}$Pb, ±0.020 for $^{207}$Pb/$^{204}$Pb, ±0.047 to 0.048 for $^{208}$Pb/$^{206}$Pb, ±0.00068 to 0.00069 for $^{207}$Pb/$^{206}$Pb and ±0.0017 for $^{206}$Pb/$^{204}$Pb.
comparison of our (sub-) surface Pb concentrations (18–25 pmol/L) with data for eastern South Atlantic surface waters sampled in 1990 (with [Pb] = 11–29 pmol/kg at 21–32° S and 7–13° E; [47]) reveals similar results. This implies that there was no major change in the Pb supply to the southeast Atlantic Ocean between 1990 and 2010. The 207Pb/206Pb isotope ratios for our seawater profile from the South Atlantic exhibit the highest values in the subsurface and at mid depth (0.867 at 49 m and 989 m water depth) and slightly lower values at the surface and at 198 m (207Pb/206Pb = 0.862–0.863; Fig. 2b). Below ~1000 m water depth, 207Pb/206Pb decreases all the way to the bottom of the profile (207Pb/206Pb = 0.845 at 4723 m water depth). These trends in 207Pb/206Pb are mirrored by the 206Pb/204Pb ratios, which are lowest at the subsurface (206Pb/204Pb = 18.01; 49 m) and at intermediate depth (206Pb/204Pb = 18.01; 989 m), and increase to 206Pb/204Pb = 18.54 at the bottom of the profile (Fig. 2b). The correlation between the seawater 207Pb/206Pb and 206Pb/204Pb ratios (r² = 0.97) observed here is significantly improved compared to the results of previous studies [13,19,20,23,54]. We assign this improvement to the accurate and precise determination of the minor isotopes 204Pb and (hence 206Pb/204Pb) ratios with our new methodology, and surmise that previous claims of poor correlations may be due to analytical artifacts and should hence be assessed with care.

The major deep and intermediate water masses encountered at station 3 in the eastern South Atlantic are indicated by the salinity contouring in Fig. 2b. In brief, most of the water column is dominated by water masses sourced from high southern latitudes – cold and dense Antarctic bottom water (AABW) at ~4500 m depth, and fresh Antarctic intermediate water (AAIW) at ~500–1100 m – whilst saline, nutrient-poor North Atlantic Deep Water (NADW) is present at ~1700–3500 m depth (see Wyatt et al. [37] for further details on the transect hydrography). Since we are presenting the first Pb isotope depth profile for the eastern South Atlantic, there are no data available for direct comparison. However, our results confirm previously suggested differences [23] in the Pb isotope fingerprints of two main Southern Ocean water masses, AAIW (207Pb/206Pb = 0.865–0.867, 206Pb/204Pb = 18.01–18.03; n = 2) and AABW (207Pb/206Pb = 0.845, 206Pb/204Pb = 18.54; n = 1). These isotopic differences are a consequence of different water mass formation mechanisms and regions (e.g., Orsi et al. [55]), and hence different proportions of anthropogenic versus natural Pb, prior to ventilation into the deeper ocean. The Pb isotopic composition of NADW in the southeast Atlantic Ocean (207Pb/206Pb = 0.853–0.857, 206Pb/204Pb = 18.42–18.23 at ~2000–3000 m water depth; n = 2) appears distinct from NADW in the northwestern North Atlantic (BATS Station at 2000 m; Table 4), and values previously reported for the equatorial western Atlantic Ocean [23]. This points to older ventilation ages in the eastern basins of the Atlantic Ocean compared to the western basins (e.g., England [56]). Further interpretation of the results will, however, have to await the acquisition of Pb isotope and concentration data for additional profiles from the GEOTRACES GA10 transect.

4. Conclusions

A new analytical procedure has been developed for Pb isotope analysis of seawater and associated Pb concentration measurements by isotope dilution. This involves Mg hydroxide co-precipitation and anion exchange chromatography for Pb pre-concentration and isotopic analysis by TIMS in conjunction with a Pb–Cd–Pb double spike. Analyses of multiple aliquots of six seawater samples demonstrate typical reproducibilities (±2σd) of about 700–1500 ppm and 1000–2000 ppm for 207Pb/206Pb, 208Pb/206Pb and 206Pb/204Pb, 207Pb/204Pb, 208Pb/204Pb, respectively. With this precision, the new data are superior to literature results that were obtained by plasma source mass spectrometry. In particular, they are at least a factor of five more precise for ratios involving the minor 204Pb isotope. Analyses of two GEOTRACES intercalibration samples from the Atlantic Ocean (GDI and GSI) display good agreement with recent results of two other laboratories.

Our methods were applied to samples from a seawater depth profile from the eastern South Atlantic Ocean at ~40° S. For these samples, both Pb concentrations and isotope compositions show a smooth pattern with depth and no obvious outliers. Furthermore, 208Pb/206Pb shows a clear correlation with 207Pb/206Pb, underlining the improvement that was achieved for the challenging Pb isotope analyses, particularly of 207Pb/206Pb. The depth profile results permit the identification and characterization of several key water masses, including AAIW, AABW and NADW. Considering that our profile is not located far from the formation regions of AAIW and AABW, the data provide the most precise Pb isotope characterization of these water masses published to date, but substantiation of the results by analyses of additional seawater samples from this region is needed.

Acknowledgements

We thank all of the MAGIC team for helping to keep spirits high, the mass spec running and the clean labs clean, and are particularly grateful for the vital support provided by Barry Coles and Katharina Kreissig. Ed Boyle volunteered helpful tips during method development, first suggesting the use of trace HF during column chemistry. The scientific party (principal scientist Gideon Henderson), technicians and crew of the R.R.S. Discovery on GEOTRACES cruise D357 are thanked for collection of the South Atlantic seawater profile. The supportive and helpful comments of two anonymous referees and swift editorial handling were also highly appreciated. Financial support for this study was provided by NERC consortium grant NE/H006095/1 to the UK GEOTRACES program and D. Weiss, T. van de Flierdt and M. Reh(2,4),(996,993)
