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Fast offline data reduction of laser ablation MC-ICP-MS Sr isotope measurements via

the interactive Excel-based spreadsheet 'SrDR'

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Abstract

Strontium isotopes are applied to a wide range of scientific fields and to different types of sample materials,

providing valuable information foremost about provenance and age, but also on diagenetic processes and

mixing relationships between different Sr reservoirs. The development of in-situ analytical techniques, such as

laser ablation ICP-MS, has improved our understanding of Sr isotope variability in several field of application,

because of the possibility to discriminate small-scale changes and their spatial distribution. However, large

outputs of Sr isotope data are produced by laser ablation MC-ICP-MS systems, which necessitate of multiple

offline steps to correct and assess the data. This requires the availability of simple and user-friendly tools,

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easily manageable also by non-specialists. With this in mind, we developed SrDR, an Excel-based interactive data reduction spreadsheet ('SrDR', Sr-Data-Reduction) for the processing of Sr isotopes measured by LA-MC-ICP-MS. The SrDR spreadsheet is easily customizable (a) to meet user-specific analytical protocols, (b) for different instruments (i.e. Nu Plasma vs. Neptune), and (c) for diverse target materials (e.g. Rare Earth Elements enriched or depleted samples). We include also several examples relevant to low and high temperature geochemistry fields - a fossil tooth, a modern seashell, a speleothem sample and plagioclase crystals - to show how different sample materials are corrected for different interfering masses.

Significance to JAAS: *In-situ* strontium isotope analyses are pivotal in several scientific fields; however, off-line data reduction can be complex and time-consuming. Here, we propose an interactive Excel-tool for the fast correction of Sr isotope measurements by LA-MC-ICP-MS.

1. Introduction

Strontium isotopes are one of the most versatile isotopic systematics due to the relatively high concentration of Sr in nature and its ubiquitous presence in geological and biological materials [1]. The Sr isotopic composition in a sample or reservoir changes continuously with time because of the radioactive decay of the parent ⁸⁷Rb to ⁸⁷Sr. Therefore, the ⁸⁷Sr/⁸⁶Sr ratio in a target material depends on the time elapsed since the closure of the system and the initial Rb/Sr ratio of the reservoir. The very first measurements of Sr isotopes have found application in geochronological and petrological studies of terrestrial and extraterrestrial rocks to determine their age and differentiation history [1 and references therein]. Nowadays Sr isotopes are applied with great success to a plethora of scientific fields, such as archaeology, anthropology, ecology, paleoclimatology and forensic sciences [e.g. 2-32]. In particular, ⁸⁷Sr/⁸⁶Sr ratios measured in tooth enamel have played a crucial role in understanding mobility patterns of ancient humans and animals [see 9 for a complete

review; 18,28), but also to trace several other unconventional evidences such as, e.g., water residence time and aeolian contribution in speleothem formation [33] or diagenetic processes affecting ancient fossil bones [34].

The presence of Sr in the low temperature realm is related to erosional and weathering processes of the crystalline bedrock where Sr ions enter the local ecosystem through soil and water [9]. Natural low temperature archives of Sr, suitable proxies of paleoclimate and/or landscape changes, include, among others, bioapatites (i.e. bones and teeth) and calcium carbonates (e.g. speleothems and seashells), where Sr²⁺ ions substitute Ca in their crystal lattice [9]. As a result, the ⁸⁷Sr/⁸⁶Sr ratios in biological tissues reflect the Sr isotope composition of the local available Sr [9]. The Sr of speleothems instead derives from cave drip water, which contains a mixture of the Sr from the soil above the cave, including potential external sources, and the carbonate host rock [33].

Sr isotopes are traditionally analysed by Thermal Ionization Mass Spectrometry after separation of Sr from the matrix by standard ion-exchange chromatographic techniques. However, more recently, laser ablation multi collector inductively coupled plasma mass spectrometry (MC-ICP-MS) systems have been widely employed to measure Sr isotopes *in-situ* in e.g. fossil teeth [13,29], speleothems [39], fish otolith [40] and mineral phases [41]. The intrinsic features of the LA-MC-ICP-MS methodology allow rapid analyses and generating large amounts of time-resolved data per analytical session in high spatial resolution bypassing laborious chemical work. Hence, to correct for matrix interferences and instrumental biases arising during LA analyses and to manage the large amount of data, a robust and reproducible method for offline data reduction and handling is highly desirable.

To date, several data reduction software to correct LA-MC-ICP-MS analyses is already accessible [42-45]. Nevertheless, adapting this software to meet the measurement criteria of each user requires advanced programming skills due to limited customizable settings in the current available versions.

In this study, we propose an Excel-based easily accessible and interactive data reduction spreadsheet ('SrDR', Sr-Data-Reduction), for fast correction of Sr isotope measurements by LA-MC-ICP-MS. Given that different instruments (i.e. Nu Plasma vs. Neptune) and different data acquisition protocols (e.g. measurement of *m/z* 83.5 and 86.5 or not) may require customization of the spreadsheet itself, we present several versions of the same protocol, based on analytical methods published in the literature, to meet the needs of a comprehensive group of users. The spreadsheet is easy to self-customize, and being an Excel-written code maintains its interactive features. With this sheet, the user is able to import all raw data files contained in a specific folder, to interactively select the background and the analyses cycles, to correct for the main Sr interferences (Kr, Rb, REE, Ca dimers and argides; Table 1) and instrumental biases and to automatically transfer the results in a final table. We also present and discuss several applications of the SrDR protocol for different materials such as teeth, shells, speleothems and mineral phases to illustrate how the spreadsheet works.

2. Data reduction philosophy

The extensive exploitation and the concurrent analytical advancement of isotope measurements with the output of extensive datasets have led to the development of software for fast data processing and correction of specific dataset and isotope systematics [e.g. 35-38]. Different procedures for data reduction of *in-situ* Sr isotope analyses are also currently present in the literature [46]. However, while there is a shared agreement on the fundamental corrections (as Kr, Rb and mass bias), several other adjustments applicable to the data are debated and applied depending on research group preferences. For this reason, we present several versions of the same spreadsheet with different data reduction procedures (as reported in Table 2) so that readers are able to choose the preferred version based on the mass spectrometer employed for data acquisition, masses collected during the analytical session and sample typology.

All spreadsheets include a Visual Basic macro for the import of raw data, a macro for the deletion of raw data (if needed) and a macro for fast copy & paste of the final results corrected for interferences and mass bias (Fig. 1). The workflow of the spreadsheet is summarized in Fig. 1 and discussed step by step in the following sections. Regardless of the data reduction protocol, we note here that each spreadsheet evidently yields different ⁸⁷Sr/⁸⁶Sr ratios, which result from the application of different processing steps (e.g. the ⁸⁷Sr/⁸⁶Sr ratio corrected for the sole Rb and mass bias vs. the ⁸⁷Sr/⁸⁶Sr ratio additionally corrected for the REEs and Ca dimers and argides).

3. Data reduction steps

3.1. Raw data import

Raw data can be manually copied into the spreadsheet (starting from cell 'O67') or automatically imported using the button 'Import raw data' within the Data sheet (Fig. 1). This macro imports all the raw data files included in the same folder of the SrDR file, generating a new sheet for each analysis and renaming it according to the file name. We suggest, prior to import the whole dataset, to check if the raw data template fits the SrDR length. If not, the user can add or remove lines to match the length. It is preferable that the first row of data coincides with the first data reduction row at row number 67. Although this operation is not strictly necessary, if the data start much lower, the spreadsheet can develop issues with graph visualization and data/background selection. The ⁸⁴Sr signal graph within the spreadsheet is conceived to help the user identify whether or not the copied data fit within the basic SrDR format, by visually checking if the analysis of the sample is plotted together with the preceding background. To overcome any issue, the user can also simply delete the excess calculation rows or add new ones, being careful to avoid cell miss-referencing. Any modification applied to the SrDR 'Data' spreadsheet is preserved when new data are imported 'Import raw data'. We also suggest exporting the raw data from the mass spectrometer without any alphabetic or numeric string at the end of the

data columns (i.e. the last row of the raw data sheet should be the last cycle of analysis). The presence of extra strings could clash with the proper visualization of signals and graphs. To avoid this issue altogether, the user can manually delete the last few rows in the raw data file prior to import.

3.2. Krypton correction as background subtraction

Given the presence of Kr on m/z 84 and 86, the interferences on these masses need to be properly assessed and corrected. The main sources of Kr in the LA-MC-ICP-MS system are the Ar and He used to generate the plasma and to carry the sample within the mass spectrometer [46]. The most common method to correct for the Kr presence is the so called 'on peak zero' method. This method requires the measurement of the gas background (generally acquired at the beginning of each analysis during laser warm-up) and the subsequent subtraction of the background signals from those recorded during analysis. This method also corrects for the potential minor presence of Sr and Rb impurities on the other masses.

Our spreadsheet exploits the 'on peak zero' method to correct for Kr, subtracting the average background signal from each single analysis cycle. The background selection is easy to perform thanks to the OFFSET Excel function. Within the 'Data selection' box the user writes the background start cycle of choice and the number of background cycles to use for the average calculation. This process is interactive: after this first operation, the user can try other background parameters by clicking the arrows up or down or by re-typing the parameters, while examining the graphs (e.g. 88Sr and 84Sr signals) and the final output (Fig. 1).

3.3. Analysis selection

The number of analytical cycles to be included in the calculations are selected within the 'Data selection' box. Similarly to the background selection, the data can be interactively selected using the small arrows or by typing

the data interval. While looking at the ⁸⁷Sr/⁸⁶Sr profile graph, the user can choose different portions of the analysis (e.g. in the case of a laser ablation raster with variable isotopic domains) saving the average value for each sub-profile.

3.4. Rare Earth Element correction

Rare Earth Elements (REEs) can be abundant for example in fossil carbonate samples (in the order of tens or even hundreds of $\mu g/g$) and generally scarce in modern specimens ($\Sigma REE < ca.~10~\mu g/g$). Doubly charged ions of Yb and Er interfere with all the masses of interests, potentially biasing both the $^{87}Sr/^{86}Sr$ and the $^{84}Sr/^{86}Sr$ ratios. In particular, $^{176}Yb^{2+}$ interferes with mass 88, $^{174}Yb^{2+}$ with mass 87, $^{172}Yb^{2+}$ with mass 86, $^{170}Yb^{2+}$ and $^{170}Er^{2+}$ with mass 85 and $^{168}Yb^{2+}$ and $^{168}Er^{2+}$ with mass 84. Moreover, $^{166}Er^{2+}$ and $^{164}Er^{2+}$ are present on masses 83 and 82, respectively. Mass 82 is also interfered by $^{164}Dy^{2+}$. One of the most applied method to correct for these elements is the so called 'peak stripping' [46]. Doubly charged ions are measured on half-masses (as 86.5 for $^{174}Yb^{2+}$ and 83.5 for $^{167}Er^{2+}$), and subtracted from Kr, Sr and Rb masses using natural isotope ratios. For example: $^{174}Yb^{2+}$ is calculated by multiplying the signal on mass 86.5 ($^{173}Yb^{2+}$) by the ratio of ^{174}Yb ($^{32.025\%}$) and ^{173}Yb ($^{16.103\%}$) natural abundances. Other methods check and correct for the presence of Er using mass 83, after background subtraction and Ca dimers and argides correction [47] (see next paragraph).

3.5. Ca dimers and argides correction

Several authors [48] have observed that ArCa and CaCa molecules (see ESI Tables 1-3) interfere with Sr masses when high Ca/Sr material is analysed. In our experience, Ca dimers and argides may affect the accuracy of the 87 Sr/ 86 Sr ratio, but only within in-run analytical error, as also observed by others [43, 49]. These molecules are commonly monitored on m/z 82 (40 Ca 42 Ca and 40 Ar 42 Ca) after background subtraction. However,

if the sample contains high REE levels, most of the m/z 82 signal may derive from the presence of ¹⁶⁴Dy²⁺. Only few studies have considered this issue [11,50-52]: e.g. Horstwood et al. [11] corrected for this interference by employing a three-sequence peak jumping routine and acquiring, among others, mass 81.5 (¹⁶³Dy²⁺) to correct for mass 82 by peak stripping. However, this procedure is difficult/time-consuming given that it requires peak jumping mode or the exclusion of other masses (e.g. 83.5) from the analysis. Our correction protocol omits mass 81.5. Thus, in case of high-REE samples, we suggest to bypass the Ca dimers and argides correction, focusing on the sole REE correction. Later in this paper, we will show how mass 82 is strongly affected by high concentration of Dy in a Middle Pleistocene fossil tooth. Otherwise, if the REE concentration in the sample is low (e.g. in modern shell specimens), the ArCa and CaCa correction can be performed using the Dy-uncorrected mass 82 and ignoring the Dy interference.

Our spreadsheets (V1 and V2 spreadsheets of both Nu and Neptune, and V3 spreadsheet of Nu; see Table 2) correct for ArCa and CaCa molecules following the protocol presented in Weber et al. [39] for speleothem analyses and here briefly exemplified for mass 86:

$$86_{ArCaCORR} = 86_{bkg_REEcorr} - 82_{bkg_corr} * (\Sigma ArCa_{86}/\Sigma ArCa_{82})$$

where $86_{ArCaCORR}$ is the corrected signal on mass 86; = $86_{bkg_REEcorr}$ is the signal on mass 86 corrected for background and REE; 82_{bkg_corr} is the signal on mass 82 corrected for the background. Σ ArCa₈₆ is the sum of the relative portion of Ca argides on mass 86 and Σ ArCa₈₂ is the sum of the relative portion of Ca argides on mass 82, based on their natural occurrence [53]. The same procedure is also performed to correct for the Ca dimers interference.

3.6. Mass bias and Rb corrections

The two main sources of inaccuracy of the ⁸⁷Sr/⁸⁶Sr ratio are the mass bias and Rb content. While the former mostly derives from the instrumental discrimination of heavier vs. lighter masses (mass-dependent fractionation), the latter is related to the matrix composition of the sample. To obtain the interference-free ⁸⁷Sr, the isobar ⁸⁷Rb needs to be calculated starting from the signal of ⁸⁵Rb using the natural ⁸⁷Rb/⁸⁵Rb ratio equal to 0.38567. Mass bias discrimination on the ⁸⁷Rb/⁸⁵Rb ratio is calculated using an exponential law, assuming that instrument mass discrimination is the same for both Rb and Sr, and normalizing to an ⁸⁸Sr/⁸⁶Sr ratio of 8.3752. After ⁸⁷Rb signal is subtracted from mass 87, the ⁸⁷Sr/⁸⁶Sr ratio is corrected for mass bias as done for the Rb. The following equations summarize how the mass bias correction is done through the exponential law [as described in 44]:

$$\beta = \frac{\ln(\frac{nrR_{uv}^{true}}{nrR_{uv}^{meas}})}{\ln(\frac{M_u}{M_v})}$$

$$R_{ij}^{corr} = R_{ij}^{meas} \times (\frac{M_i}{M_j})^{\beta}$$

where β is the mass bias factor; $_{nr}R_{uv}^{meas}$ and $_{nr}R_{uv}^{true}$ are the measured and reference 'true' ratios of the chosen normalizing ratio (nr); M_u and M_v are the atomic masses of the normalizing ratio isotopes; R_{ij}^{meas} and R_{ij}^{corr} are the measured and corrected isotopic ratios (e.g. $^{87}Sr/^{86}Sr$); M_i and M_j are the atomic masses of the two isotopes considered for the ratio. At the end of all the mathematical corrections, the $^{87}Sr/^{86}Sr$ outliers are 2σ -filtered.

3.7.89Y monitoring

Some analytical protocols presented in literature [48] use m/z 89 (Y) as a proxy for the REE content of the sample. This mass, adjacent to mass 88, can be easily collected with both Neptune and Nu Plasma instruments.

Although we believe that most of the inferences on mass 89 can be easily deduced using other masses (e.g. 82, 83 or half-masses; see paragraph 5.1.), we present here some protocols where mass 89 is collected (V1 spreadsheet of both Nu and Neptune, see Table 2). In the Neptune V1 spreadsheet, the user can input or not mass 89 without interfering with the calculation (mass 89 is the last column of the dataset, so it can be easily left empty). In the Nu V1 spreadsheet, it is mandatory to have the first column of the dataset dedicated to ⁸⁹Y (or at least an empty column before the actual dataset). If the Nu user did not collect ⁸⁹Y, the V2 spreadsheet might be used (Table 2).

4. Spreadsheet outcome and specifications

All spreadsheets provide the ⁸⁷Sr/⁸⁶Sr ratio corrected for the sole background, Rb and mass bias (without REE and Ca dimers/argides and named 'Unc.' within the sheet); in addition, the different versions of the spreadsheet may yield additional ⁸⁷Sr/⁸⁶Sr values, based on the type of correction applied (e.g. REE only or both REE and Ca dimers/argides). Other parameters reported in the spreadsheet are the 2-standard error (2se) of the mean, the number of unfiltered-cycles considered for the analysis (*n*), the total Sr signal (V), the ⁸⁴Sr/⁸⁶Sr ratio, the ⁸⁵Rb/⁸⁸Sr ratio and the average signal collected for each mass. These values can be automatically copied in the 'Results' sheet using the corresponding export buttons ('Copy & Paste'), located below the result-column of interest.

The spreadsheets are compatible with Excel 2007 and newer versions, but also with LibreOffice and OpenOffice. Please note that to use the macros, the spreadsheet must be saved as an Excel Macro-Enabled Workbook.

5. Applications

5.1. Fossil tooth

The ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ ratio of calcium phosphates is relatively challenging to measure by LA-MC-ICP-MS, because of the formation of molecules with m/z 87, namely ${}^{40}\text{Ar}^{31}\text{P}^{16}\text{O}$ and ${}^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$, during analysis. The interference of these molecules has been observed by many authors [11,28,54,55], and yields higher ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ ratios than expected, in particular for samples with relatively low Sr concentration (<< 1000 µg/g). The two main methods to correct for the influence of these molecules are the following: 1) using N_2 to reduce oxide levels within the plasma [56]; 2) calibrate the daily formation of CaPO and ArPO through the analysis of a set of matrix-matched reference materials [11,28]. In this paper, we correct for the presence of these interferences (Ar-CaPO) using the protocol described in Horstwood et al. [11] and Lugli et al. [28], namely a calibration curve with in-house reference materials (i.e. teeth). This calibration is performed by analyzing a shark tooth ([Sr] ca. 1000 µg/g), a bovine tooth ([Sr] ca. 350 µg/g) and a swine tooth ([Sr] ca. 200 µg/g) [28]. A daily curve is built with the ${}^{88}\text{Sr}$ signals (as a proxy of the Sr content) and the accuracy of the Ar-CaPO uncorrected ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ ratios (i.e. LA data are normalized to the 'true' solution MC-ICP-MS results). Then, the resulting equation is employed to mathematically correct the unknown bioapatite data.

Additionally, in this paragraph, we show how the strong presence of REEs within a fossil sample may also considerably bias the final ⁸⁷Sr/⁸⁶Sr ratio.

We analysed the enamel ⁸⁷Sr/⁸⁶Sr ratio of a Middle Pleistocene (ca. 600ka) rhinoceros tooth (sample ID: RHIS14; stored at University of Modena and Reggio Emilia) using a 213 nm laser ablation system (New Wave UP) and a Neptune MC-ICP-MS (Thermo Scientific), both housed at the Centro Interdipartimentale Grandi Strumenti (CIGS) of the University of Modena and Reggio Emilia. We employed linear raster of 100 μm x 750 μm, a fluence of ~12 J/cm², a frequency of 10 Hz and a scan speed of 5 μm/s. A He flux of ca. 0.6 L/min was used to carry the sample into the mass spec. No additional gases, besides the Ar, were employed (e.g. N₂). During the session, the modern swine tooth was also measured several times as unknown. Its ⁸⁷Sr/⁸⁶Sr ratio

was previously determined by solution MC-ICP-MS and resulted equal to 0.70899 \pm 0.00001 (2se) [28]. The swine tooth, measured by LA-MC-ICP-MS, yielded an Ar-CaPO uncorrected $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.70992 \pm 0.00043 (2 σ , n = 16). After the Ar-CaPO calibration, the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio dropped to 0.70891 \pm 0.00031 (2 σ , n = 16). REE corrected and uncorrected values were identical within the error. This specimen shows in fact very low REE concentrations (SREE < 1 $\mu\text{g/g}$).

The fossil rhinoceros tooth ([Sr] ca. 400 μ g/g) shows very distinct isotopic values whether the REE correction is applied or not (Fig. 2). Step-by-step results are shown in Figure 3. The REE uncorrected ⁸⁷Sr/⁸⁶Sr ratio is 0.71036 \pm 0.00005 (2se), while the REE corrected ratio is 0.70966 \pm 0.00005. After the CaPO daily calibration, these two values become 0.71008 and 0.70937 respectively. Considering that the bulk Sr isotope ratios, measured by dissolution MC-ICP-MS, is equal to 0.70954 for enamel and to 0.70933 for dentine, the LA data seem more accurate after the REE correction is applied. The REE uncorrected ⁸⁴Sr/⁸⁶Sr ratio is 0.06031 \pm 0.00011, while the REE corrected ratio is 0.05556 \pm 0.0004. In this case, the ⁸⁴Sr/⁸⁶Sr is overly high before the correction, but seems overcorrected after the REE peak-stripping correction, similarly to what has been observed by other authors [9,47]. Moreover, compared to modern samples, this specimen shows quite high signals for all the masses, likely related to the presence of REEs (m/z 82 = 3.4E-03 V; m/z 83 = 1.4E-03 V; m/z 85.5 = 2.2E-04 V; m/z 86.5 = 3.0E-04 V; Fig. 2). The signal of mass 82 is correlated with the signal of mass 86.5 (¹⁷³Yb²⁺as proxy of REE content; $r^2 = 0.33$), but it is not with the signal of mass 88 (⁸⁸Sr as proxy of the ablation behavior of the sample; $r^2 = 0.004$). Thus, we suggest that the signal recorded on m/z 82 (Dy + ArCa and CaCa) reflects the presence of Dy, rather than the formation of ArCa and CaCa during the ablation of the specimen.

5.2. Modern seashell

We measured the Sr isotope composition of a modern seashell (sample ID: SS12; stored at the University of Modena and Reggio Emilia), collected from the Italian seashore, using the same instrumentation presented in

section 5.1. We thus employed the same parameters of the rhinoceros' tooth analysis, but using spots of 100 μ m in diameter (40 s of dwell time) instead of lines. Given that this sample is a modern calcium carbonate shell with low REE content, we did not employ any further correction (i.e. REE, Ca dimers/argides or Ca-ArPO calibration). Raw data files of 10 different ablation lines were automatically imported using the 'Import raw data' macro into the Neptune V3 Sr-DR worksheet. On average, the seashell yields an 87 Sr/ 86 Sr ratio of 0.70917 \pm 0.00003 (2 σ , n = 10), perfectly matching the modern seawater composition [57], and an 84 Sr/ 86 Sr ratio of 0.05650 \pm 0.00010 (2 σ , n = 10), in line with the expected natural ratio. Although few analyses present a slightly deviated 84 Sr/ 86 Sr ratio (see e.g. analysis #6 in Fig. 4), their 87 Sr/ 86 Sr ratio is still accurate. We note here that the entire processing of the data including data import, selection and export did not require more than 10 minutes.

5.3. Speleothem

A stalagmite sample from the Hüttenbläserschachthöhle (Germany) [58,59] has been analysed at the Max Planck Institute for Chemistry, Mainz, following methods described in Weber et al. [39]. The sample mainly consists of aragonite and is therefore highly suited for *in-situ* LA-MC-ICP-MS analyses of Sr-isotopes due to the high Sr concentration in aragonite compared to calcite. Three sections of the stalagmite (samples ID: HBSH-1-9, HBSH-1-11 and HBSH-1-20) have each been analysed with three line scans of 750 μ m positioned directly next to each other within a single growth layer (see Fig. 5). The ⁸⁷Sr/⁸⁶Sr ratio was determined using a New Wave UP 213 nm laser ablation system coupled to a Nu Plasma MC-ICP-MS. Each line scan was performed using a circular spot size of 100 μ m, a scan speed of 5 μ m/s, a frequency of 10 Hz and a fluence of 12 – 14 J/cm². The stalagmite sections are store at the Institute of Geosciences, Mainz.

Correction of Ca argides and dimers does not significantly affect the resulting 87 Sr/ 86 Sr ratios , unlike the REE correction. Stalagmite slab HBSH-1-20 yielded a REE uncorrected average 87 Sr/ 86 Sr ratio of 0.70961 \pm 0.00003 (2 σ , n = 3). Nevertheless, after REE and Ca argide and dimer correction, the 87 Sr/ 86 Sr ratio decreased to an

average ratio of 0.70953 \pm 0.00003 (2 σ , n = 3) and a ⁸⁴Sr/⁸⁶Sr ratio of 0.05621 \pm 0.00003 (2 σ , n = 3). The same is true for the two other stalagmite slabs HBSH-1-11 (uncorrected ⁸⁷Sr/⁸⁶Sr = 0.70932 \pm 0.00004, REE and Ca argide and dimer corrected ⁸⁷Sr/⁸⁶Sr = 0.70916 \pm 0.00004, each 2 σ , n = 3) and HBSH-1-9 (uncorrected ⁸⁷Sr/⁸⁶Sr = 0.70944 \pm 0.00003, REE and Ca argide dimer corrected ⁸⁷Sr/⁸⁶Sr = 0.70932 \pm 0.00003, each 2 σ , n = 3). The significant effect of the REE correction might be related to the age of the sample, which dates back to MIS 7 [58]. Thus, an old sample with high REE content needs to be properly corrected and cannot be simply considered as modern calcium carbonate specimens. Please note that all speleothem ⁸⁷Sr/⁸⁶Sr ratios have been corrected by standard bracketing after data evaluation with the SrDR spreadsheet, using JCt-1 as reference material to a ⁸⁷Sr/⁸⁶Sr ratio of 0.70917 \pm 0.00001 [52].

5.4. Magmatic plagioclase

Fifteen plagioclases from a gabbroic dyke sampled in the Finero Phlogopite-Peridotite mantle unit (western Alps, Italy) were analyzed *in-situ* through LA-MC-ICP-MS at the CIGS laboratories (see section 5.1). Analytical parameters are identical to the fossil tooth analysis in section 5.1, with the exception that different plagioclase grains were analyzed each with one line.

Plagioclase contains ca. 3100 ug/g of Sr, only 0.6 ug/g of Rb and 19 ug/g of Σ REE. The sample was previously analyzed at the Marine Environmental Sciences Laboratory (LEMAR) of the Institut Universitaire Européen de la Mer (IUEM), Université de Bretagne Occidentale (Brest, France) after conventional dissolution and chromatographic separation using a TRITON Thermal-Ionization Mass Spectrometer (TIMS) as described in [60]. TIMS analysis yielded an 87 Sr/ 86 Sr ratio of 0.704519 \pm 0.000008 (2SE).

REE correction did not affect the 87 Sr/ 86 Sr ratio of *in-situ* LA-MC-ICP-MS Sr isotope analyses, being the uncorrected and corrected values equal within the error. The average uncorrected 87 Sr/ 86 Sr ratio is 0.704514 \pm 0.000057 (n=15; 2 σ), identical to the 'true' value obtained by TIMS.

6. Conclusion

Given the rapid spread of *in-situ* LA-MC-ICP-MS Sr isotope analyses to several fields of the applied sciences (from paleoclimate to anthropology to forensic science), we present here a fast and accurate data reduction protocol exploiting the flexibility offered by Microsoft Excel. As exemplified by our case studies, several parameters need to be considered during LA-MC-ICP-MS Sr isotope analysis and data reduction, with some differences depending on the diverse matrices that can be used to measure Sr isotopes. The SrDR spreadsheet provides a user friendly and unique way to interactively evaluate the LA-MC-ICP-MS Sr isotope data for any kind of specimen, offering different solutions based on the preferred analytical protocol and considering various possible corrections (as REE and Ca dimers/argides).

Conflicts of interest

There are no conflicts to declare.

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Data availability

All data generated or analysed during this study are included in this paper (and its Electronic Supplementary Information files).

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Figure 1. (a) Overview of the SrDR spreadsheet workflow; (b) data reduction philosophy, including the proposed corrections.

Figure 2. Analyses of a Middle Pleistocene rhinoceros tooth. (a) Signal (V) of m/z 86.5 (173Yb++) is correlated with signal of m/z 82. (b)

Signals of m/z 82 and 83 are strongly correlated. (c) The increase of the m/z 82 signal during the ablation of the fossil tooth (and not

during the ablation of the modern swine tooth) suggests its correlation with the presence of REEs within the sample, in particular with

¹⁶⁴Dy⁺⁺. (d) The REE-uncorrected ⁸⁷Sr/⁸⁶Sr ratio and the REE-corrected ⁸⁷Sr/⁸⁶Sr ratio are significantly different (see text for details). Error

bars are 2se. The small red square over the tooth photograph represents the area where the analyses were performed (both solution

and LA).

Figure 3. 87Sr/86Sr and 84Sr/86Sr data of RHIS14 (Middle Pleistocene rhinoceros tooth) after each single correction step using Sr-DR sheet.

The blue line is the reference 84Sr/86Sr value (0.0565); while the red line is 87Sr/86Sr ratio of the tooth enamel (0.70954) obtained by

solution MC-ICP-MS.

Figure 4. 87Sr/86Sr and 84Sr/86Sr of a modern seashell. As example, analysis #6 reveals a slightly deviated 84Sr/86Sr ratio (0.05637), even if

the ⁸⁷Sr/⁸⁶Sr perfectly match the modern seawater (0.70918). Error bars are 2se.

Figure 5. Stalagmite sample from the Hüttenbläserschachthöhle (Germany). In-situ Sr isotope analyses (red squares) were performed

using a Nu plasma mass spectrometer and a 213 nm laser ablation system. Numbers (20, 11, 9) represent the slab ID (HBSH-1-20, HBSH-

1-9 and HBSH-1-11). See text for details.

Table 1: Sr isotopes and possible sources of interference.

m/z	88	87	86.5	86	85.5	85	84	83.5	83	82	81.5
Strontium	⁸⁸ Sr	⁸⁷ Sr	-	⁸⁶ Sr	-	-	⁸⁴ Sr	-	-	-	-
				⁸⁶ Kr			⁸⁴ Kr		⁸³ Kr	⁸² Kr	
		⁸⁷ Rb				⁸⁵ Rb					
	¹⁷⁶ Yb	¹⁷⁴ Yb	¹⁷³ Yb	¹⁷² Yb	¹⁷¹ Yb	¹⁷⁰ Yb	¹⁶⁸ Yb				
	¹⁷⁶ Lu					¹⁷⁰ Er	¹⁶⁸ Er	¹⁶⁷ Er	¹⁶⁶ Er	¹⁶⁴ Er	
Interferences	¹⁷⁶ Hf									¹⁶⁴ Dy	¹⁶³ Dy
	⁴⁰ Ca ⁴⁸ Ca	⁴³ Ca ⁴⁴ Ca		⁴⁰ Ca ⁴⁶ Ca		⁴² Ca ⁴³ Ca	⁴⁰ Ca ⁴⁴ Ca		⁴⁰ Ca ⁴³ Ca	⁴⁰ Ca ⁴² Ca	
	⁴⁰ Ar ⁴⁸ Ca	⁴⁰ Ca ³¹ P ¹⁶ O		⁴² Ca ⁴⁴ Ca			⁴⁰ Ar ⁴⁴ Ca		⁴⁰ Ar ⁴³ Ca	⁴⁰ Ar ⁴² Ca	
		⁴⁰ Ar ³¹ P ¹⁶ O		⁴⁰ Ar ⁴⁶ Ca							

	le 2. Diff		•			,		n the	1						
alla	m/z	82	83	83.5	oyed for the analysis. 83.5 84 85 85.5 86 86.		86.5	87	88	(89)*					
	Isotop	82	83	¹⁶⁷ Er	84	⁸⁵ R	¹⁷¹ Yb	86	¹⁷³ Yb	⁸⁷ Sr	⁸⁸ Sr	89γ	Correcti	Notes	Literature
	е	Kr	Kr	++	Sr	b	++	Sr	++				ons		examples
	V1	Х	х		х	х		х		х	х	х	ArCa+Ca Ca	Best option for modern samples containing no REE. ArCa and Caca correction assumes no REE content.	[13] + ⁸⁹ Y
	V2	х	х	x	x	х		x	x	x	x		REE and ArCa+Ca Ca	Best option for fossil samples containing high REE.	[8] + ⁸² Kr
	V3	Х	х		х	х	х	х	х	x	x		REE	Alternative option for both fossil and modern samples. Not possible to properly check for ArCa and CaCa. Er corrected using m/z 83.	[28]
e e	V4	х	х		х	х		х	х	х	х		REE	Alternative option for both fossil and modern samples. Not possible to properly check for ArCa and CaCa.	[47] with Nu
Neptune	V5		х	х	х	х	х	х	х	х	х		REE	Alternative option for both fossil and modern samples.	[49]
	V1	х	х		х	х		x		х	х	х	ArCa+Ca Ca	Best option for modern samples containing no REE. ArCa and CaCa correction assumes no REE content.	[45] + ⁸⁹ Y
E.	V2	х	х		х	х		х		x	x		ArCa+Ca Ca	Best option for modern samples containing no REE. ArCa and CaCa correction assumes no REE content.	[45]
Nu Plasma	V3	х	х	×	х	х	х	х	x	x	x		REE and ArCa+Ca Ca	Best option for fossil samples containing high REE.	[39]
	ass 89 is reported in brackets because is not involved in the data reduction calculations; it is only employed to check for the diagenetic status of specimen.											enetic status of			

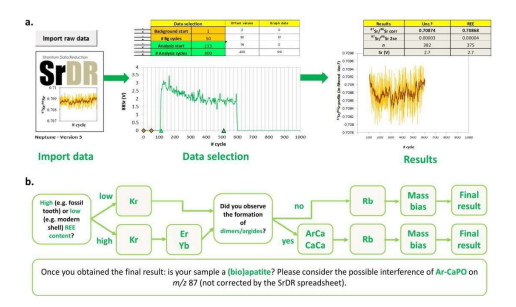


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349x209mm (223 x 223 DPI)

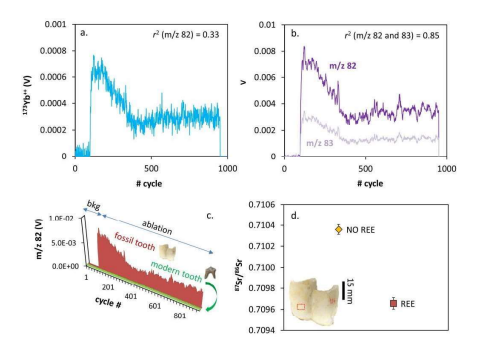


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254x190mm (300 x 300 DPI)

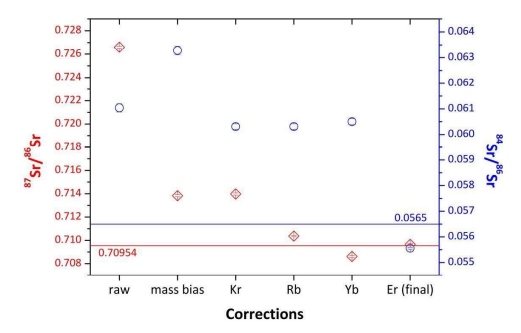


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269x169mm (600 x 600 DPI)

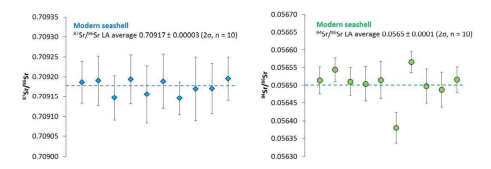


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254x90mm (300 x 300 DPI)

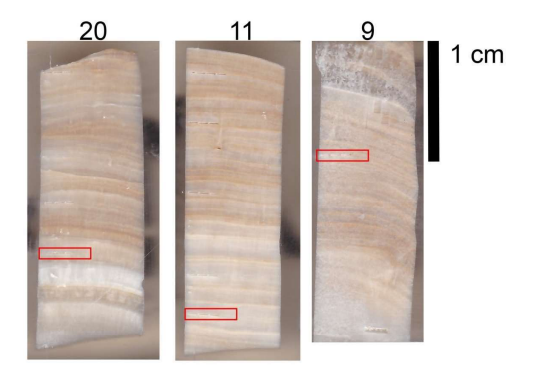


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254x190mm (300 x 300 DPI)