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Comparative high spatial resolution measurements of Sr isotopic composition in bio-apatite using different LA-MC-ICPMS configurations: application to faunal (sub)seasonal mobility studies†

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The effects of isobaric interferences on the precision and accuracy of *in situ* Sr isotopic composition measurements in bio-apatite were investigated using different plasma conditions, interface configurations and instrumentation. Analysis of modern shark enameloid via laser ablation MC-ICPMS (LA-MC-ICPMS) *Neptune* and *Neoma* using H-type cones show similar long-term $^{87}\text{Sr}/^{86}\text{Sr}$ reproducibilities of 0.709194 ± 0.000034 and 0.709189 ± 0.000027 (2SD), respectively. Neither instrument showed detectable interference from matrix-generated $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ or Ca dimer/argide isobars. The ratios agree with solution-based MC-ICPMS and TIMS analyses, yielding an average $^{87}\text{Sr}/^{86}\text{Sr}$ of 0.709176 ± 0.000008 (2SD). The use of a Jet sampler and a X-type skimmer cone with *Neoma* increases the sensitivity 2–3 times while maintaining the same level of accuracy. The formation of $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ polyatomic clusters can be avoided by tuning the plasma to a low oxide level. An inherent feature of high-sensitivity cones is the production of Ca dimers/argides, which affects $^{84}\text{Sr}/^{86}\text{Sr}$, additionally suffering from $^{68}\text{Zn}^{16}\text{O}$ isobars readily generated at higher oxide levels owing to the ubiquitously high Zn concentration in bio-apatite. The mass bias of $^{87}\text{Sr}/^{86}\text{Sr}$ analysis follows the exponential law across all instrument configurations, which is interpreted as being due to a small average mass difference between normalised and normalising ratios. We demonstrate the benefit of high-sensitivity cones by analysing the highly compositionally zoned enamel of a woolly mammoth. Higher ion beam intensity allowed improved spatial resolution down to 40 μm diameter, revealing compositional variability obscured by larger spots. Improved counting statistics allowed for shorter integration times, resulting in <10 days of temporal resolution for >10 years of enamel formation. Increased sensitivity is of particular value for analysing organisms with low Sr uptake, such as humans, and in slowly mineralizing bioapatite tissues. Another benefit is the enhanced ability to distinguish early *in vivo* enamel formation from subsequent maturation, which is crucial for reconstructing the palaeoecology of large mammals and human life histories.

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Introduction

Strontium isotopes have been widely used since the development of the modern mass spectrometer in the early 1950s.¹

Their immediate application was in earth and planetary sciences, where they have become a fundamental petrogenetic tracer and geochronometer using a ^{87}Rb – ^{87}Sr radioactive decay scheme.² However, owing to the relative ease of perturbation, Sr isotopes have played a subordinate role compared with more robust, immobile element-based systems such as ^{147}Sm – ^{143}Nd or later ^{176}Lu – ^{176}Hf .^{3–6} The high abundance of Sr and its compositional diversity in the natural environment, together with its common uptake by plants and animals, make Sr isotopes widely used in forensics, (palaeo)environmental studies, palaeoanthropology, (bio)archaeology and food or medical sciences. The development of laser ablation (LA) coupled with multi-collector inductively coupled plasma mass spectrometry (MC-ICPMS)⁷ has revitalized the application of Sr isotopes, enabling high spatial resolution analyses that reveal

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details obscured by bulk analyses. The quest to analyse minimum sample volume with maximum precision led to the development of various strategies in hardware, instrumental parameters and data reduction.^{8–14} Apart from technological improvements such as the reaction cell permitting chemical separation of the most problematic isobar ^{87}Rb from ^{87}Sr ^{14,15} or the application of special geometry cones enhancing instrument sensitivity, tuning conditions have been identified as an important factor in reducing or eliminating the effects impeding the accuracy of LA (MC) ICPMS measurements.¹⁶ The flexibility of the ICP source allows for a wide range of conditions influencing ionisation, isotopic and elemental fractionation, polyatomic cluster formation and doubly charged ions or oxide formation, which can be adjusted according to the properties of the element(s) of interest and the type of matrix. Bioapatite, the main compound of skeletal tissues, is one of the most commonly targeted matrices. Its composition can be approximated as a hydroxyapatite $\text{Ca}_5(\text{PO}_4)_3(\text{OH})$ with CO_3 substitutions. Although apatite is surprisingly flexible in incorporating various elements at the trace level, bio-mineralisation imposes some limitations, and relative abundances are linked to habitat, diet or trophic level. High Sr uptake is accompanied by modest Rb uptake and virtually no REE, whose doubly charged ions (particularly Er and Yb) interfere with Sr isotopes. The absence of doubly charged REE isobars makes bioapatite particularly suitable for the quantification of matrix effects such as Ca-argide and dimer or $^{40}\text{Ca}^{30}\text{P}^{16}\text{O}$ formation, whose influence on the accuracy of Sr isotopic composition measurements is debated.^{8,9,11,13,17–19} Some studies have shown a large magnitude of these effects, albeit correctable to a minor level,^{9,18} while others have not observed them at all.¹¹ An additional factor comes from the use of high-sensitivity cones, together with the increased ion beam results in different isotopic fractionation, mass bias or oxide production.^{14,20–27} In this study, we investigated the influence of plasma conditions, matrix effects and instrumental mass bias on the precision, accuracy and reproducibility of Sr isotopic composition measurements of bioapatite by LA-MC-ICPMS using dental enameloids of modern shark teeth. We quantify the influence of Ca dimer/argide, $^{40}\text{Ca}^{30}\text{P}^{16}\text{O}$ isobar, and Zn oxides on the accuracy of Sr isotopic ratio measurements. We present a decade-long record of analyses performed with the MC-ICPMS Neptune and compare it with data obtained using the newer generation MC-ICPMS Neoma. Additional experiments were performed using high-sensitivity cones. Except for $^{87}\text{Sr}/^{86}\text{Sr}$, we provide analyses of the rarely investigated $^{84}\text{Sr}/^{86}\text{Sr}$ ratio,^{9,13} whose accuracy commonly serves as the data quality check and is a sensitive monitor of robust Ca dimer/argide and Kr isobaric interference corrections. The LA-MC-ICPMS data were compared with a long-term record of solution-based analyses under wet and dry plasma conditions. In addition, we tested the current limitations of high-precision, high-spatial-resolution analyses using the dental enamel of a woolly mammoth molar. We demonstrate precise and accurate measurements at spatial resolution of 100 to 40 μm laser beam diameter and <10 days temporal resolution of Sr isotopic composition variations in about 11 year-long formation of a single dental plate. Enhanced

sensitivity and spatial resolution demonstrate a much improved ability to separate the *in vivo* record in the earliest formed enamel from the later maturation stage, which is critical for reconstructing the palaeoecology of large mammals and the life histories of the humans.

Material and methods

Solution analyses were performed on the SRM 987 Sr standard, which is most commonly used to monitor the performance of the analytical set-up and for interlaboratory comparisons. Periodically, modern shark enameloid was measured after the full chemical preparation procedure from digestion to ion exchange column separation as an additional quality check.²⁸ *In situ* laser ablation MC-ICPMS analyses were performed on modern shark teeth. An example of the application of the *in situ* LA-MC-ICPMS methodology is demonstrated on an approximately 30-ka-old woolly mammoth molar tooth from Torske, Chortkiv district, Ternopil region in Western Ukraine. The specimen (number D-14) was provided by the Lviv State Museum of Natural History, Ukraine. Sample preparation is described in ref. 29.

Solution mode MC ICPMS analyses

Analyses were performed using MC ICPMS Neptune with collectors connected to amplifiers equipped with $10^{11} \Omega$ resistors. The collector setup and the main analytical parameters are summarised in ESI Table S1 (S# – ESI†). Two introduction systems were used: (1) Stable Introduction System (SIS), which is a quartz glass dual-pass cyclonic spray chamber with ESI 100 $\mu\text{L min}^{-1}$ microconcentric, pneumatic PFA nebuliser; (2) Aridus II desolvating nebuliser fitted with the same type of PFA pneumatic nebulizer with either 50 or 100 $\mu\text{L min}^{-1}$ flow rate. Free aspiration was applied in both cases. The instrument was tuned in the hot plasma range (RF power = 1250–1300 W) to achieve a compromise between signal stability, sensitivity and low oxide production, monitored with the $^{140}\text{Ce}^{16}\text{O}^+/^{140}\text{Ce}^+$ ratio. For SIS, about 3–5% oxide was a typical value, but some measurements were made with an oxide level of about 10%. The Aridus II was tuned to an oxide level below 0.2%. The spray chamber and membrane temperatures were set to 160 °C and 120 °C, respectively. The flow rate of nitrogen used with the Aridus II was 4–8 mL min^{-1} .

The desolvating nebulisation increased the sensitivity of the mass spectrometers by about 4–8 times with respect to the SIS. For both introduction systems, SRM 987 was at a concentration providing 10–15 V of ^{88}Sr . The standard Ni sampler and H-type skimmer cones (hereafter jointly referred to as H-type cones) were used. The isotope ratios were measured in static mode. A single run consisted of 50 cycles of 4.048 s integration. The collector gains were measured prior to each analytical session using the factory setup. The peak centering was performed during tuning and verified every few hours. The baseline was measured on the masses of interest, “on peak zeroes” (OPZ), performed on 2% HNO_3 . Washouts between samples were conducted using 3% HNO_3 . The total integration time of the

baseline was similar to that of the sample. Krypton isobaric interferences associated with trace level contamination of the Ar plasma gas have been reported as an obstacle to accurate Sr isotopic composition measurements, and various correction methods have been proposed.^{7,10,13,30,31} In our setup, we observe less than 0.5 mV of ⁸⁴Kr, with a typical level of <0.2 mV, and ordinary gas blank subtraction efficiently removes Kr isobars from Sr isotopes. In solution-based analyses, the ⁸⁷Rb isobar is also a minor problem as samples are highly purified by ion exchange chromatography. The ⁸⁵Rb/⁸⁷Rb correction value was estimated from Rb–Sr mixed solutions as 0.38557, which is slightly lower than the natural reference value of 0.38571 due to the difference in mass bias between Rb and Sr. For the correction, the estimated ratio was de-normalised using the exponential mass bias law and within run ⁸⁸Sr/⁸⁶Sr ratio. After subtraction of the interfering isobars, the final ratios were normalized to ⁸⁶Sr/⁸⁸Sr = 0.1194, applying the exponential law.³² The data were filtered at a 2σ level.

Laser ablation MC ICPMS analyses

Spatially resolved (*in situ*) Sr isotopic composition measurements were conducted using an excimer laser (ArF, 193 nm) RESOLUTION by Resonetics (now Applied Spectra) equipped with the M-50 double-volume sample cell, later replaced by the larger S-155 two-volume cell, both by Laurin Technic. Modern shark teeth serve as a standard for bioapatite treatment. Nearly all analyses were conducted with the preferred energy density of about 7–8 J cm⁻², but the data include experiments with energy down to 5 J cm⁻². The laser pulse frequency was varied from 5 to 30 Hz. Ablation was conducted as a line scan with spot size diameter varying from 40 to 130 μm and stage speed of either 0.5 or 1.0 mm min⁻¹. Ablation occurred when He was mixed in the internal funnel with the Ar sample gas. The laser-generated aerosol was delivered to the MC-ICPMS source after passing through a signal smoothing manifold. To enhance MS sensitivity, 4 to 8 mL per min nitrogen was added to the sample carrier gas (Table S1†). The sensitivity determined on the NIST 614 glass standard varied from 0.8 to 1.5 mV ppm⁻¹ of ⁸⁸Sr for the 60 μm diameter spot and 5 Hz repetition rate, and depended largely on the set of cones used (same type but different sets). Solution analyses were performed using a standard sampler and H-type skimmer cones. The analyses were conducted using *Neptune*, followed¹¹ by the modifications described below. Additional analyses were performed using MC ICPMS *Neoma* coupled with the same laser ablation system. The experiments were conducted in two modes: (1) with the H-type cones and (2) with the X-type skimmer cone and Jet sampler cone (hereafter referred to as Jet-X cones). The collector setup and the main instrument parameters for both modes are presented in Table S1.† Data reduction was analogous to that performed for the solution-based analyses, with the difference confined to ⁸⁷Rb correction. Due to the differing properties of Rb and Sr ablation, this was corrected for elemental fractionation using the fractionation factor ⁸⁵Rb/⁸⁶Sr of 1.155 for *Neptune*¹¹ and 1.138 estimated for *Neoma* with the Jet-X cones by ablation of BHVO-2G glass standard. The polyatomic isobaric interferences related to

the ablation of the apatite matrix, such as Ca-dimers, Ca-argides, or ⁴⁰Ca³¹P¹⁶O, vary depending on the instrumental setup, and their treatment is discussed below.

Results and discussion

Solution-based MC ICPMS analyses under wet and dry plasma conditions

The isotope ratios determined by thermal ionisation mass spectrometry (TIMS) are commonly used as target values for MC-ICPMS measurements. This is chiefly due to much reduced problems related to baseline corrections, the considerably smaller mass bias, and the lack of matrix effects. In the classic paper by ref. 33, the long-term reproducibility of SRM 987 was determined as ⁸⁷Sr/⁸⁶Sr = 0.710248 ± 0.000011 and ⁸⁴Sr/⁸⁶Sr = 0.056492 ± 0.000016 (uncertainties are 2SD). Our more limited dataset obtained with TIMS *Triton* gave ⁸⁷Sr/⁸⁶Sr = 0.710244 ± 0.000007 and ⁸⁴Sr/⁸⁶Sr = 0.056493 ± 0.000005 (*n* = 32, 2SD), which agrees well with the above values (ratios normalised to ⁸⁶Sr/⁸⁸Sr = 0.1194 using exponential law³²). Although the published range of ⁸⁷Sr/⁸⁶Sr is much broader, the aforementioned ratios have long been used as reference values and are preferred in this study.

Data obtained by MC ICPMS *Neptune* for SRM 987 shows long-term reproducibility expressed as 2SD (standard deviation) for >10 years period of ⁸⁷Sr/⁸⁶Sr = 0.710260 ± 0.000011 (*n* = 213) and 0.710251 ± 0.000013 (*n* = 107) for the wet and dry plasma, respectively (Fig. 1a). The external precision is similar to the internal precision of an individual run in both cases, which is expressed as 2SE (standard error). Notably, the dry plasma conditions provide slightly more accurate data (relative to TIMS values), although, in a strict sense, within uncertainties, the wet and dry plasma results are indistinguishable. The corresponding ⁸⁴Sr/⁸⁶Sr ratios are 0.056492 ± 0.000023 and 0.056499 ± 0.000030, which are also accurate. The internal precision of ⁸⁴Sr/⁸⁶Sr is nearly three times better (absolute error) than that of long-term reproducibility. Although the latter values are practically the same and agree well with the TIMS ratio, the long-term reproducibility of data obtained using *Aridus II* is worse than that of *SIS*. This is interpreted as being due to a low ⁸⁴Sr signal (17 to 170 mV) and the higher importance of accurate baseline correction. In the case of the desolvating nebuliser, it requires a lot of patience due to the excessively long washout time, and thus, incomplete washout likely affected some measurements. Some contribution to lowering the precision could be associated with ⁸⁴Kr and ⁸⁶Kr interferences. However, as mentioned above, the Kr content in our analyses is minor, and the baseline subtraction practically corrects it to zero (mean measured ⁸³Kr/⁸⁶Sr = 0.000001 ± 0.000007). In the absence of any major isobaric interferences, the main factor potentially affecting the accuracy of isotope ratio determination is the mass bias correction, if it departs from the exponential law of ref. 32, as demonstrated for Nd isotopes *e.g.* ref. 34–37. Ionisation in the ICP source generates approximately 10 times larger isotope fractionation than the thermal source of TIMS, but unlike TIMS, the bias is relatively stable throughout a single measurement. The prediction of an exponential mass bias law is that log

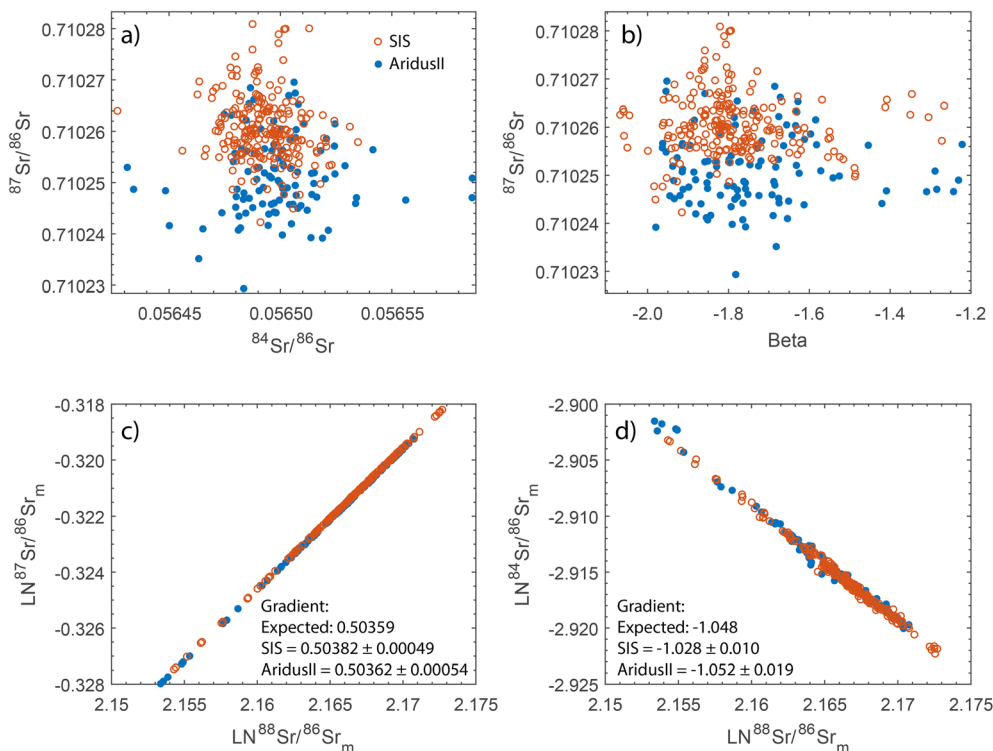


Fig. 1 Results of SRM 987 Sr standard measurements under wet (SIS) and dry (Aridus II) plasma conditions: (a) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. $^{84}\text{Sr}/^{86}\text{Sr}$; (b) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. beta fractionation factor; (c) natural logarithm of the measured (amplifier gain-, baseline- and interference-corrected) $^{87}\text{Sr}/^{86}\text{Sr}_m$ vs. natural log of the raw $^{88}\text{Sr}/^{86}\text{Sr}_m$ ratio; (d) natural logarithm of the measured $^{84}\text{Sr}/^{86}\text{Sr}_m$ vs. natural log of raw $^{88}\text{Sr}/^{86}\text{Sr}_m$ ratio. Plots (c) and (d) compare the estimated gradients for wet and dry plasma conditions, with the slope predicted using the exponential mass bias law. Uncertainties are expressed as 2SD. See main text for details.

natural $^{88}\text{Sr}/^{86}\text{Sr}$ versus $^{87}\text{Sr}/^{86}\text{Sr}$ ratios corrected for baseline and interferences form a linear trend with a gradient equal to 0.50359 (Fig. 1b and c). The linear regression parameters estimated for ratios in $\text{LN}(^{87}\text{Sr}/^{86}\text{Sr})_m$ vs. $\text{LN}(^{88}\text{Sr}/^{86}\text{Sr})_m$ coordinates yield 0.50382 ± 0.00049 (MSWD = 0.8) and 0.50362 ± 0.00054 (MSWD = 1.0) for wet and dry plasma conditions, respectively (Fig. 1c and d). Both values are in excellent agreement with the theoretical prediction, demonstrating that the exponential law perfectly describes the mass discrimination of Sr isotopes during MC-ICPMS measurements. This remarkable agreement makes the Sr isotope system unique. However, in our view, this relationship arises from the small average mass difference between the normalised $^{87}\text{Sr}/^{86}\text{Sr}$ and the normalising $^{88}\text{Sr}/^{86}\text{Sr}$ ratios. As shown for Nd isotopes, the smaller the mass difference, the closer the agreement with the exponential law; conversely, larger mass differences result in greater deviation from the exponential law.^{35–37} Our data suggest that this model is also valid across lower mass ranges. Analogous assessment for $^{84}\text{Sr}/^{86}\text{Sr}$ gave -1.028 ± 0.010 (MSWD = 3.6) and -1.052 ± 0.019 (MSWD = 9.3) gradients for the wet and dry plasma, respectively (Fig. 1d). Both values approximate the expected -1.023 gradient, however, the lower precision of the $^{84}\text{Sr}/^{86}\text{Sr}$ measurements limits a more detailed assessment of mass bias effects. Nonetheless, under wet and dry plasma conditions, the short- and long-term reproducibility of Sr isotope ratios is accurate at high precision.

Laser ablation MC-ICPMS of bioapatite

LA-MC-ICPMS Neptune. The teeth of contemporary sharks are an ideal material for the quality control of laser ablation MC-ICPMS Sr isotopic composition analyses of dental enamel. They have a high Sr concentration (about 1750 ppm, as determined by LA ICPMS) and a homogenous isotopic composition that reflects that of sea water. Some variations may arise depending on the shark's origin. From the batch of 50 purchased teeth, we randomly selected 9 whose dental enameloid was micro-sampled (using a microdrill) and subjected to the solution mode analyses. Measurements were performed using MC-ICPMS *Neptune* and *Neoma*. To compensate for the slight variations in reproducibility of SRM 987 at different instruments, the obtained $^{87}\text{Sr}/^{86}\text{Sr}$ shark values were normalised to the preferred SRM 987 value of $^{87}\text{Sr}/^{86}\text{Sr} = 0.710248$.³⁸ The measurements ranged from 0.709164 ± 0.000005 to 0.709186 ± 0.000008 with an average of 0.709173 ± 0.000014 (2SD) (weighted average 0.709172 ± 0.000006). This value was additionally verified by 5 replicate analyses using TIMS *Triton*, which gave a mean value of $^{87}\text{Sr}/^{86}\text{Sr} = 0.709178 \pm 0.000010$ (2SD). The results are in perfect agreement with the average Sr isotopic composition of the present-day sea water of 0.709175 ± 0.000019 (ref. 39) and with the solution-based MC-ICPMS analyses of bull shark (0.709205 ± 0.000035) and spot-tail shark (0.709170 ± 0.000025) determined by ref. 40.

In Fig. 2, we compiled the LA MC ICPMS analyses of shark dental enameloid collected over 10 years. An individual analysis represents a mean ratio for the integration time between 60 and 600 seconds, with the vast majority lasting from 150 to 250 s (Fig. 2a and b). The intensity of ^{88}Sr varied from *ca.* 1 to 20 V, indicating variable spot size, laser pulse repetition rate, and, to a lesser degree, MC-ICPMS settings. The $^{87}\text{Sr}/^{86}\text{Sr}$ ranges from 0.70915 to 0.70922, with the internal precision expressed as 2RSE (relative standard error) ranging from 120 ppm for signals ≤ 1 V of ^{88}Sr to approximately 20 ppm for signals >10 V. It is noteworthy that low-intensity measurements, although less precise, remain accurate. The internal precision well approaches or occasionally even matches that achieved for the SRM 987 solution-based measurements (*ca.* 15 ppm 2RSE) for

comparable integration time and intensities $> ca.$ 10 V of ^{88}Sr . However, the long-term reproducibility (0.709195 ± 0.000025 ; 2SD) is approximately two times lower than that of the solution analyses. This is attributed to the instability of the transient signal and imperfections of natural solid materials (defects, zonation, *etc.*). Overall, the long-term reproducibility, expressed as arithmetic mean with 2SD (standard deviation) uncertainty for all data, yielded $^{87}\text{Sr}/^{86}\text{Sr} = 0.709194 \pm 0.000034$ (Fig. 2a and b), which is satisfactory for resolving compositional differences in a broad range of applications requiring high spatial resolution.

Sr isotope analyses of shark teeth are unaffected by isobars of doubly charged REE ions simply because there is no detectable uptake of these elements during the animals' lives. The Kr level

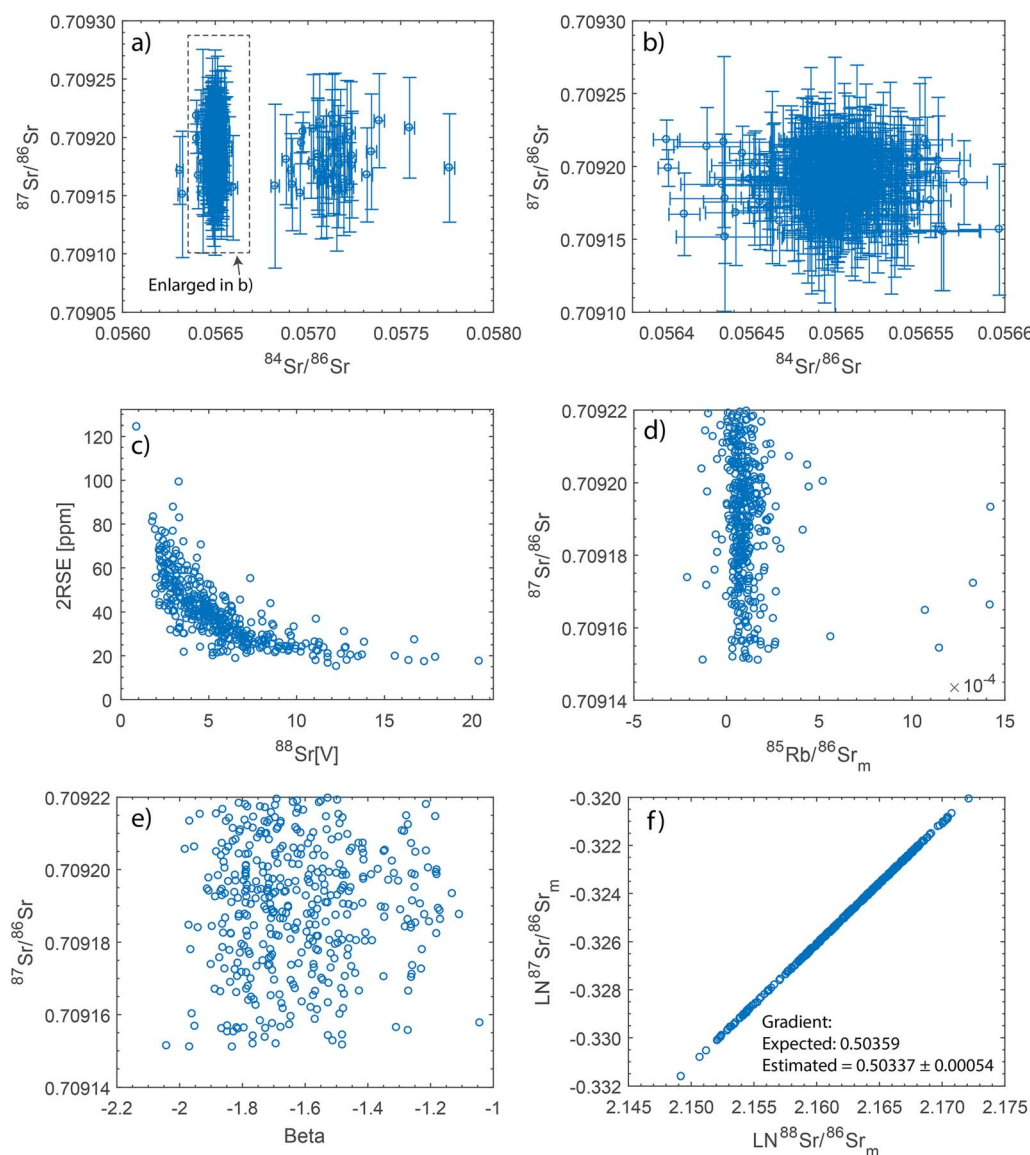


Fig. 2 Summary of LA-MC-ICPMS *Neptune* measurements of the Sr isotopic composition of a modern shark enameloid: (a) $^{84}\text{Sr}/^{86}\text{Sr}$ vs. $^{87}\text{Sr}/^{86}\text{Sr}$. (b) Close-up of the dataset highlighted by the black rectangle in (a); (c) 2RSE (relative standard error) as a function of ^{88}Sr ion beam intensity; (d) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. measured $^{85}\text{Rb}/^{86}\text{Sr}_m$ (*m* denotes ratios corrected for amplifier gain, baseline, and interferences); (e) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. beta fractionation factor; (f) natural logarithm of measured $^{87}\text{Sr}/^{86}\text{Sr}_m$ vs. natural logarithm of raw $^{88}\text{Sr}/^{86}\text{Sr}_m$, comparing the observed slope to the gradient predicted using the exponential mass bias law. The error bars in (a) and (b) represent 2SE (standard error).

in our setup is the same as that in the solution mode analyses; thus, ordinary gas blank subtraction efficiently strips the interference from the peaks. Thus, the primary concerns are the ^{87}Rb and matrix-related isobars. The Rb concentration in shark tooth is negligible. The measured $^{85}\text{Rb}/^{86}\text{Sr}$ (Fig. 2d) normalized using Sr mass bias spanned from below the detection limit of Rb to 0.0004 with the average 0.000088 ± 0.000103 (2σ outliers rejected). Such small interference can be effectively corrected using the standard peak stripping method. Some of the data are slightly overcorrected (negative $^{85}\text{Rb}/^{86}\text{Sr}$ values, Fig. 2d) due to a “more intense washout” of ^{85}Rb and contrasting cone memory effects affecting Rb vs. Sr such that ^{85}Rb can drop below the corresponding baseline level observed in some runs. This is illustrated in Fig. 3, which shows a decrease in the ^{85}Rb signal below the blank level during ablation. Such inadequate baseline correction may change the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio by as much as 15 ppm. This effect is undetectable in materials with higher Rb abundance because the baseline correction does not lead to negative values. Only long pre-conditioning of the cones with apatite matrix seems to prevent more intense Rb washout during the ablation.

A debated obstacle in obtaining accurate $^{87}\text{Sr}/^{86}\text{Sr}$ ratio is the $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ isobar generated during the ablation of apatite matrix.^{9,18,41,42} This, however, seems specific to the instrument or setup, as some studies essentially do not observe the formation of this compound.¹¹ The production of this cluster is strongly linked to the oxide level in the plasma.^{9,18} We de-tuned the torch position and gas flows to achieve $^{232}\text{Th}^{16}\text{O}/^{232}\text{Th}$ of approximately 10%, but we observed no detectable effect on the accuracy of $^{87}\text{Sr}/^{86}\text{Sr}$ ratio determination, indicating that even relatively high oxide production is insufficient to generate enough $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ ions to affect $^{87}\text{Sr}/^{86}\text{Sr}$ ratio within the obtained precision, which agrees with the observation of ref. 11. Other apatite matrix-related isobars include Ca-dimers and argides.^{9,13,31,43,44} The Ca dimers and argides affect all Sr isotopes, but due to the low abundance of ^{84}Sr (*ca.* 0.59%), the

$^{84}\text{Sr}/^{86}\text{Sr}$ ratio is a particularly sensitive indicator of these interferences. This is well illustrated with the anomalous group of $^{84}\text{Sr}/^{86}\text{Sr}$ ratios showing values from 0.0568 to 0.0578 (Fig. 2a). They show a clear positive linear correlation with mass ^{82}X used as a monitor of Ca argide/dimer ($^{42}\text{Ca}^{40}\text{Ca}$, $^{42}\text{Ca}^{40}\text{Ar}$) (Fig. S1a and b†). The measured intensity of ^{82}X was 20–120 μV , with the majority of measurements ranging from 20 to 60 μV , which translates into max. 0.2 mV of ^{44}Ca interference on ^{84}Sr ref. 13 report about 100 mV of total Ca-argide/dimer signal and shift in $^{84}\text{Sr}/^{86}\text{Sr}$ ratios similar to those in our study. The anomalous $^{84}\text{Sr}/^{86}\text{Sr}$ ratios show a positive correlation with $^{82}\text{X}/^{86}\text{Sr}_m$ (Fig. S1a and b†). Linear regression defines a zero intercept at 0.05658 ± 0.00018 , which agrees with the reference value of 0.05650 (both ratios normalized using Sr mass bias). The slope of the regression defines $^{84}(\text{CaCa}, \text{CaAr})/^{82}(\text{CaCa}, \text{CaAr})$ as 3.2 ± 1.1 (MSWD = 6.3), which broadly agrees with the expected value of *ca.* 3.2241 for $^{44}\text{Ca}/^{42}\text{Ca}$. The data scatter may be due to the small instability of the plasma conditions that change the dimer/argide production or the contribution from other isobar(s) affecting $^{82}\text{X}/^{86}\text{Sr}$ ratio (Fig. S1a†). Although it is impossible to differentiate between the Ca dimer and argide clusters, as pointed out by ref. 13, the relative abundances of key isotopes are similar, and the same correction will accommodate both interfering compounds. The group of inaccurate $^{84}\text{Sr}/^{86}\text{Sr}$ data was generated by a single user during several consecutive sessions. We investigated all the operator-controlled parameters of laser ablation and MC-ICPMS, but no correlation was found with the anomalous data. Moreover, we did not intentionally tune the MC-ICPMS to stimulate the Ca-dimer/argide production to a visible level. The accidental use of the X-type skimmer cone is the most probable cause of the observed systematics. Nonetheless, even such significant dimer/argide production does not affect the corresponding $^{87}\text{Sr}/^{86}\text{Sr}$ ratio, which, uncorrected for dimer/argide isobars, is 0.709181 ± 0.000032 (2SD, $n = 28$) and perfectly agrees with the reference value. All the remaining $^{84}\text{Sr}/^{86}\text{Sr}$ measurements, uncorrected for dimer/argide, yielded an accurate mean value of 0.056501 ± 0.000049 (2SD, $n = 328$). The small positive correlation with $^{82}\text{X}/^{86}\text{Sr}$ (Fig. S1b†) is certainly not because of the dimers/argides, as the relationship in the $^{82}\text{X}/^{86}\text{Sr}$ and $^{84}\text{Sr}/^{86}\text{Sr}$ coordinates does not follow the argide/dimer trend (Fig. S1a†). A potential reason could be a “residual”, uncorrected by blank subtraction, Kr interference. However, assuming that all mass 82 is ^{82}Kr , the performed correction reduces the slope of the observed trend from 0.96 ± 0.14 (2SD, MSWD = 6.2) to 0.52 ± 0.17 (2SD, MSWD = 6.7). This changes the mean $^{84}\text{Sr}/^{86}\text{Sr}$ ratio to 0.056496 ± 0.000053 , resulting in a negligible difference in accuracy and precision.

The determination of the mass bias trajectory, as for the SRM 987 (Fig. 2f and S1c, d†), in the case of $^{87}\text{Sr}/^{86}\text{Sr}$ ratio, shows a gradient of 0.050337 ± 0.000054 (MSWD = 0.96), which closely follows the exponential mass bias law (Fig. 2f). In the case of the $^{84}\text{Sr}/^{86}\text{Sr}$ ratio, the fractionation trajectory with a slope of -0.999 ± 0.010 (data affected by dimer/argide interferences excluded) significantly deviates from the expected value of -1.048 (Fig. S1d†) and the data obtained in solution mode (Fig. 2d). This discrepancy is likely due to the low precision of

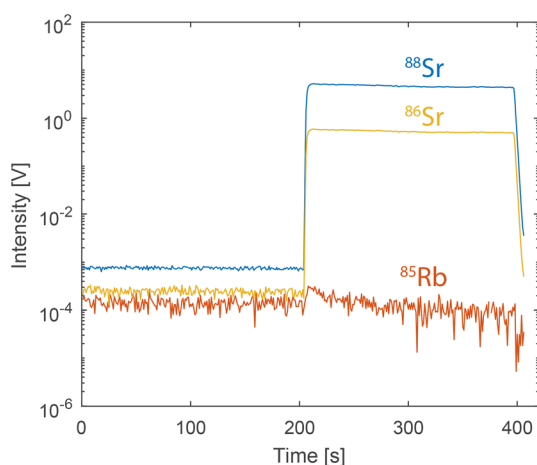


Fig. 3 Isotopic composition measurement of shark enameloid demonstrating intense washout of ^{85}Rb during ablation, dropping below the corresponding gas blank and leading to inaccurate ^{87}Rb isobaric interference correction.

the measurements (0.1% 2RSD), which limits the detection of mass bias effects. Nevertheless, from a practical perspective, the measured $^{84}\text{Sr}/^{86}\text{Sr}$ ratio agrees with the TIMS value within uncertainty, indicating that any deviation from the exponential law did not significantly distort the results.

MC-ICPMS *Neoma*, H-type cones

The experiments conducted with H-type cones are analogous to the *Neptune* measurements. At the same RF plasma power set to 1250 or 1300 W, we typically achieve about 2% of oxide production ($^{232}\text{Th}^{16}\text{O}/^{232}\text{Th}$), and exceptionally, about 0.5%, which was accompanied by $^{232}\text{Th}/^{238}\text{U}$ fractionation of *ca.* 0.8 (determined ablating NIST 612 glass standard). On *Neptune*, achieving low oxide levels (*ca.* 0.5%) was rather effortless. However, in the case of *Neoma*, oxide levels are typically higher, and their reduction is often at the expense of a higher loss of sensitivity. As demonstrated above, a very low oxide level is not a prerequisite for precise and accurate Sr isotope measurements. Similar to *Neptune*, the Kr level was essentially negligible, with ^{84}Kr gas blank below 0.2 mV and the mean gas blank corrected $^{83}\text{Kr}/^{86}\text{Sr}_m = 0.000009 \pm 0.000020$. We verified the $^{85}\text{Rb}/^{87}\text{Rb}$ ratio applied to the ^{87}Rb isobar correction using solution Rb–Sr mixtures. Dry plasma was achieved using a desolvating nebuliser, the Aridus II introduction system. The resultant $^{85}\text{Rb}/^{87}\text{Rb}$ ratio normalized to $^{88}\text{Sr}/^{86}\text{Sr}$ was 0.38573. The elemental fractionation was determined by ablation of BHVO-2G $^{85}\text{Rb}/^{86}\text{Sr} = 1.148$, which is similar to that observed with *Neptune*.¹¹ The quality of data obtained for shark teeth over 4 months is comparable to that of *Neptune* data. The internal precision was 20 to 60 ppm (2RSE) for the ^{88}Sr ion beam, ranging from 2 to 9 V and 150–200 s integration time (Fig. S2a†). The accuracy of $^{87}\text{Sr}/^{86}\text{Sr} = 0.709189 \pm 0.000027$ (2SD, $n = 57$) and $^{84}\text{Sr}/^{86}\text{Sr} = 0.056513 \pm 0.000027$ (2SD, one gross outlier rejected) was almost identical to that of the *Neptune* data. The *Neoma* data has long-term reproducibility, which may be due to the shorter data acquisition period. The normalized ratios showed no correlation with the isobaric interference monitors (^{82}X and ^{85}Rb) or with the changes in the instrumental mass bias (Fig. S2c and d†). The extent of the instrumental mass bias expressed as a beta factor is similar to that observed for *Neptune* for the H-type cones and follows the exponential law as indicated by the estimated gradient 0.5019 ± 0.0037 (MSWD = 0.7), which within uncertainty follows the predicted trajectory (Fig. 2e, f and 5e, f). Matrix-generated ion clusters, such as Ca dimers/argides or $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$, were not detected within the obtained precision.

The internal precision achieved for LA-MC-ICPMS shark analyses approaches that of SRM 987 solution-based analyses when comparing data acquired with signals of similar intensity and integration time. For $^{88}\text{Sr} > 6$ V, $^{87}\text{Sr}/^{86}\text{Sr}$ ratios show an internal precision of 35 to 20 ppm 2RSE and yield a mean ratio of 0.709182 ± 26 (36 ppm, 2RSD). The solution mode analyses have similar internal and external precision at the level of approximately 15 ppm (for measurements at about 15 V of ^{88}Sr). Lower external precision is expected in the case of transient signal analyses of a natural standard with small-scale

heterogeneities, local mechanical defects, or zonation in element concentration, matrix effect, *etc.* Nonetheless, the long-term reproducibility of LA-MC-ICPMS at the level presented in this study is satisfactory for most applications and can certainly be improved.

MC-ICPMS *Neoma* and Jet-X cones

The jet sampler cone and X-type skimmer cone enhance the sensitivity of MC-ICPMS but also oxide formation, polyatomic interferences, and changes in isotopic fractionation.^{20–22,24–26,45} While keeping the plasma conditions similar to those described above for the H-type cones, oxides increase to several tens of percent, resulting in elevated $^{87}\text{Sr}/^{86}\text{Sr}$ ratios due to $^{40}\text{Ca}^{30}\text{P}^{16}\text{O}$ formation (Fig. 4a). Additional contribution to deteriorating Sr isotope ratio accuracy is from Ca dimers/argides. Applying near-cold plasma conditions (RF power *ca.* 1020 W) and reducing oxide formation by adjusting Ar gas flows and sampling depth appears sufficient, if not to eliminate, then at least to suppress the $^{40}\text{Ca}^{30}\text{P}^{16}\text{O}$ isobar to a negligible level, while maintaining approximately 2–3 times signal enhancement relative to H-type cones. Although considerably reduced, the oxide level was still high, up to approximately 14%. Bringing the $^{232}\text{Th}^{16}\text{O}/^{232}\text{Th}$ ratio down to approximately 0.5% had no detectable benefit regarding $^{87}\text{Sr}/^{86}\text{Sr}$ ratio accuracy, and further reduction was at the cost of a large loss of sensitivity, making the use of Jet-X cones worthless. Despite lower, in comparison to the H-type cones, short-term signal stability (*ca.* 3% 2RSE for ^{88}Sr), the benefit of higher signal allows to obtain internal precision improved by at least a factor of 2 (20–30 ppm 2RSE) for the same laser ablation parameters as used with the H-type cones (Table S1†). Because of the appearance of invisible effects with H-type cones, the correction parameters used for ^{87}Rb interference for the Jet-X cones and low RF power were verified on Rb/Sr standard solution mixtures using an Aridus II desolvating nebuliser. Tune parameters were selected to closely resemble those of LA data acquisition as closely as possible. The estimated $^{85}\text{Rb}/^{87}\text{Rb} = 0.38572$ is practically the same as for the H-type cones, while the elemental fractionation determined on the BHVO-2G glass standard was $^{85}\text{Rb}/^{86}\text{Sr} = 1.138$, which is slightly lower than that observed for the H-type cones. The reproducibility for data collected over 4 months, $^{87}\text{Sr}/^{86}\text{Sr} = 0.709193 \pm 0.000028$ (2SD, $n = 51$), is the same as that obtained with the H-type cones for much larger sample volumes (Fig. 4b and c). Although accurate within uncertainty, the ratio is slightly higher than the TIMS or MC-ICPMS solution-based value. Although we cannot exclude some minor production of $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$, its contribution to mass 87 seems unlikely at low oxide levels. We did not find any interference that could be attributed to the slight “excess” of the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio. Calcium dimers/argides do not have a detectable effect on this ratio, as supported by the lack of correlation between the $^{82}\text{X}/^{86}\text{Sr}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ (Fig. 4d). A similar conclusion was reached in the studies of ref. 42, 44 and 46. Naturally, at this minor level of “inaccuracy,” small details play an important role. Incomplete or extra washouts for ^{85}Rb will affect the baseline correction and hence the accuracy of the ratio estimate, especially when the measurements are at low signal

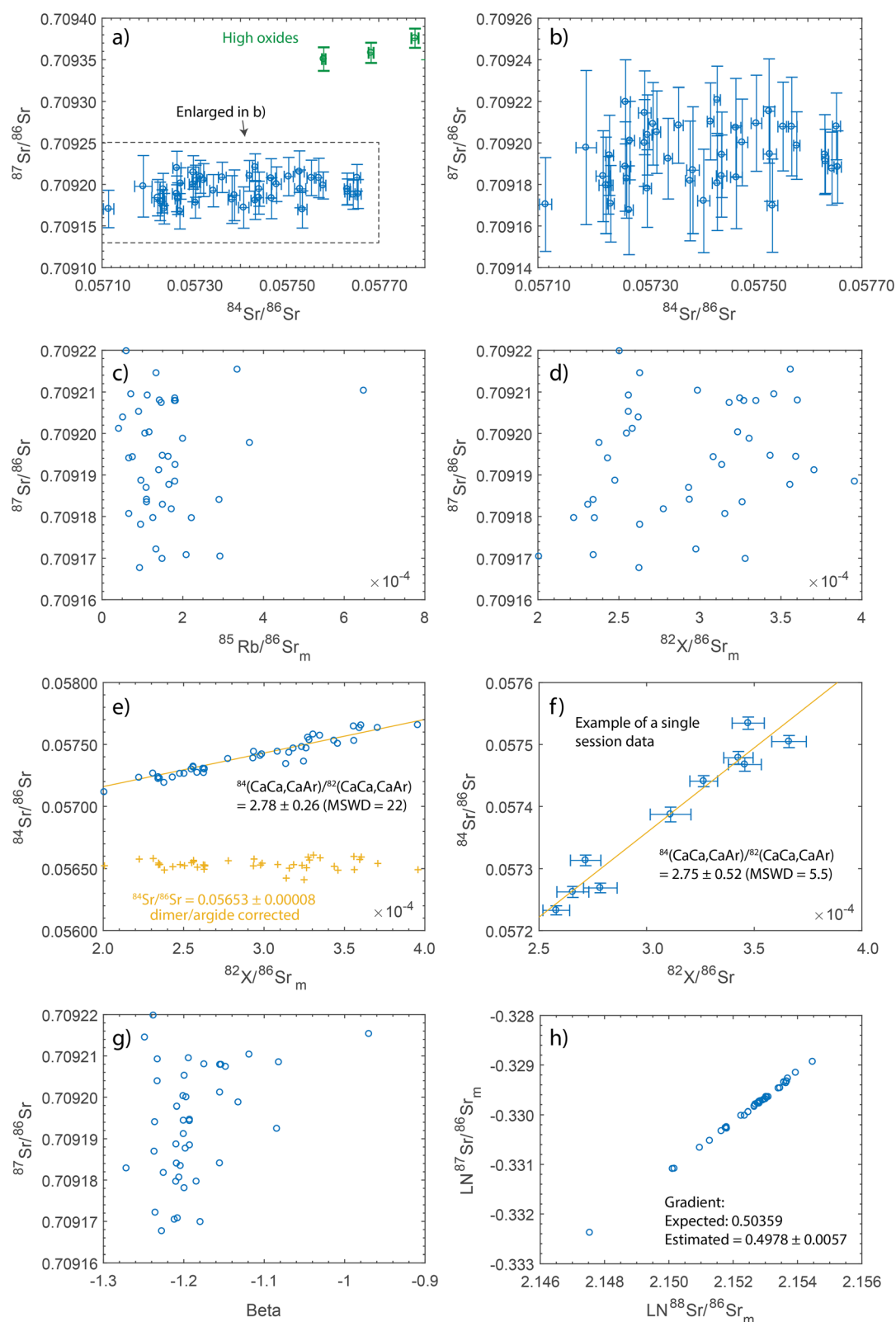


Fig. 4 Summary of laser ablation MC-ICPMS Neoma measurements of Sr isotopic composition of a modern shark enameloid with Jet-X cones: (a) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. $^{84}\text{Sr}/^{86}\text{Sr}$; (b) enlarged view of the region highlighted in (a); (c) measured $^{85}\text{Rb}/^{86}\text{Sr}_m$ vs. $^{87}\text{Sr}/^{86}\text{Sr}$; (d) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. $^{82}\text{X}/^{86}\text{Sr}$ (Ca dimer/argide monitor); (e) $^{84}\text{Sr}/^{86}\text{Sr}$ vs. $^{82}\text{X}/^{86}\text{Sr}_m$; (f) example of $^{87}\text{Sr}/^{86}\text{Sr}$ vs. $^{82}\text{X}/^{86}\text{Sr}$ behaviour over a course of a single session; (g) $^{87}\text{Sr}/^{86}\text{Sr}$ vs. beta fractionation factor; (h) natural logarithm of measured $^{84}\text{Sr}/^{86}\text{Sr}_m$ vs. natural logarithm of measured $^{88}\text{Sr}/^{86}\text{Sr}_m$, comparing the observed gradient with that predicted by the exponential mass bias law. Uncertainty on the gradient is given as 2SD. Subscript *m* denotes ratios corrected for amplifier gain, baseline, and interferences.

intensities. We observed a small deterioration in the data quality when the cones were not well coated with the apatite matrix, which is probably a prerequisite for accurate high-precision analyses. Although the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio remains accurate within uncertainty, it tends to be slightly higher. Larger magnitude changes are occasionally observed during long runs for the $^{84}\text{Sr}/^{86}\text{Sr}$ ratio, which also tends to be higher at the beginning of the run and reaches the reference ratio at a later stage. We observed such behaviour only with the Jet-X cones. This suggests that some minor discharge occurs between the plasma and the sampler cone, which enhances the dimer/argide production. Adjusting the plasma parameters does not eliminate dimers/argides, which appear to be an inherent feature of the cone geometry. The uncorrected mean for matrix-generated isobars $^{84}\text{Sr}/^{86}\text{Sr} = 0.05728 \pm 0.00028$ is inaccurate but notably, the data scatter is smaller than that in the “accidental dataset” produced with *Neptune* (Fig. 4e). The relationship is linear with a gradient in normalized $^{82}\text{X}/^{86}\text{Sr}$ vs. $^{84}\text{Sr}/^{86}\text{Sr}$ coordinates 2.78 ± 0.26 (MSWD = 22). Over a single session (Fig. 4f), the relationship remains the same with the slope defining $^{84}(\text{CaCa}, \text{CaAr})/^{82}(\text{CaCa}, \text{CaAr})$ ratio as 2.75 ± 0.52 (MSWD = 5.5). This is lower than the expected $^{44}\text{Ca}/^{42}\text{Ca}$ ratio of 3.22,⁴⁷ which is likely due to a large difference in the mass bias between Sr and Ca, but possibly also due to the contribution of other, yet unrecognized sources of interferences (see below). The inference of the latter observation is that the correction ratio will vary with varying plasma conditions and possibly also compositional variations in trace elements in sharks' enameloid. Correction with the use of the estimated ratio (Fig. 4e) yields $^{84}\text{Sr}/^{86}\text{Sr} = 0.05653 \pm 0.00008$ (2SD).

The effect of Zn oxide on Sr isotopes has rarely been explored and was found to be insignificant in solution-based work with Zn/Sr ratios up to approximately 1.⁴⁴ Dental tissues are rich in Zn, especially in the outer enamel,^{48,49} and their abundance and Zn/Sr ratio are typically strongly zoned. In sharks, the Zn concentration is on the order of several hundred ppm, reaching even *ca.* 0.2 wt%, approaching the Sr concentration. Oxygen and zinc form a range of interfering isobars on masses 83, 84, 85, 86, 87 and 88. The most abundant ^{66}Zn and the most abundant oxygen isotope ^{16}O form the highest interference, making mass 82 a sensitive monitor of ZnO isobars. The most significant consequence is on ^{84}Sr due to the formation of $^{68}\text{Zn}^{16}\text{O}$, which is the second most abundant Zn isotope (18.5%). The accurate $^{84}\text{Sr}/^{86}\text{Sr}$ ratio measured with H-type cones, both with *Neptune* and *Neoma*, indicates that ZnO oxide, if present, is negligible in such a configuration. In the case of high-sensitivity cones, where minor ^{84}Sr isotope and mass 82 are additionally affected by Ca-dimer/argide, assessing the potential effect of superposed ZnO isobars is more difficult. Hence, we performed experiments in solution mode, aspirating only the NIST 683 Zn standard using Aridus II connected with *Neoma*. Experiments were conducted for different oxide levels controlled by lowering the sampling depth while keeping all other parameters unchanged (Fig. 5). At $^{232}\text{Th}^{16}\text{O}/^{232}\text{Th}$ of *ca.* 2–3% or lower, the ion beams at 82 and 84 masses are at the acid blank level (or the effect is too small to be detected with a Faraday collector). Increasing oxides to about 10% causes a significant rise at mass 82 ($^{66}\text{Zn}^{16}\text{O}$) and

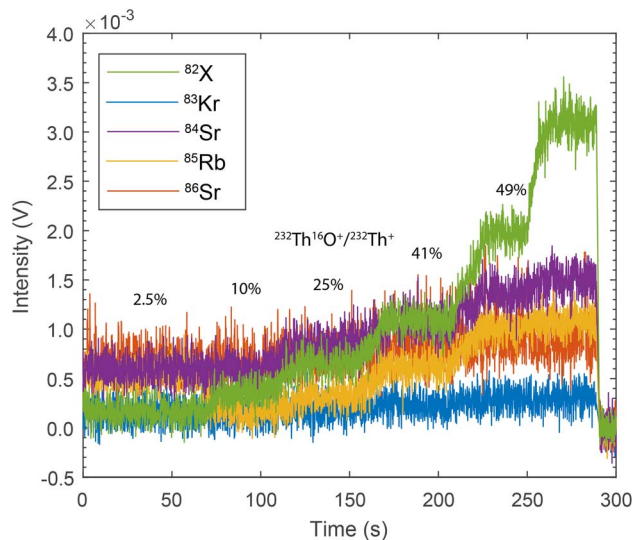


Fig. 5 Contribution of ZnO polyatomic isobars on Sr, Rb and Kr masses in dependence on the oxide level using MC-ICPMS *Neoma* with a Jet sampler cone and an X-type skimmer cone. A NIST 683 Zn standard solution was aspirated using Aridus II, and the oxide level was increased by lowering the sampling depth, while all other parameters were kept constant.

a smaller rise at mass 84 ($^{68}\text{Zn}^{16}\text{O}$). This also changed the signal on mass 85 to the level below the corresponding blank, yet had no effect on masses 86 and 83, the latter being the most reliable Kr interference monitor. Higher oxides caused the rise of interferences in all monitored isotopes, with the biggest change for masses 82 ($^{66}\text{Zn}^{16}\text{O}$) and 84 ($^{68}\text{Zn}^{16}\text{O}$). A further increase in the number of oxide magnifies the effect, which is best manifested by the changes in mass 82 (Fig. 5). Noteworthy, the effect of $^{70}\text{Zn}^{16}\text{O}$ isobars on mass 86, although present, is negligible as ^{70}Zn constitutes only 0.6% of total Zn. The effect of $^{70}\text{Zn}^{17}\text{O}$, involving a minor oxygen isotope, on mass 87 is even smaller. Hence, the effect of ZnO is expected to be minor or even negligible on the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio. However, it is important for the $^{82}\text{X}/^{86}\text{Sr}$ ratio used for argide/dimer correction, if Zn concentration or oxide level is high. A complementary test for potential ZnO interferences was directly performed on the dental enamel. High Zn enrichment in the outer enamel layer is an expression of amelogenesis.^{48,50,51} We measured the invariant in nature, the $^{84}\text{Sr}/^{86}\text{Sr}$ ratio, as a line scan from the inner to outer enamel of a woolly mammoth (see below). Subsequently, we measured the intensity of ^{66}Zn as a measure of element concentration along the same line. The estimated Zn/Sr ratio changes from approximately 0.2 in the inner enamel to nearly 4 in the outer enamel. The first scan was conducted with an oxide level of *ca.* 2%. We observe a strong increase in $^{82}\text{X}/^{86}\text{Sr}$ and $^{84}\text{Sr}/^{86}\text{Sr}$ in the outer enamel layer, where the zinc concentration abruptly rises. A small effect is also visible on $^{83}\text{Kr}/^{86}\text{Sr}$ ratios (Fig. 6a and b). Applying only the correction for dimers, the $^{84}\text{Sr}/^{86}\text{Sr}$ ratio is accurate, but notably, it gently rises in the Zn-rich zone, and the data is more scattered. Setting up oxides to a very high level of approximately 50% did not reveal any extra features and resulted only in the magnification of the changes observed for the

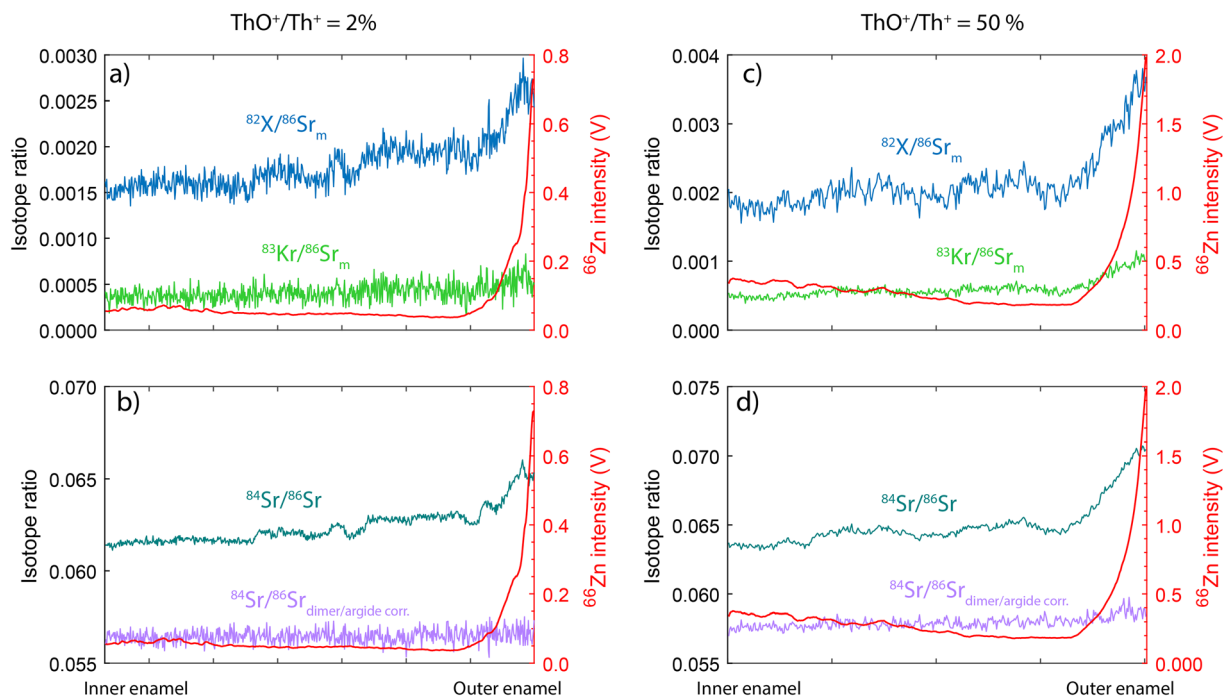


Fig. 6 Laser ablation MC-ICPMS line scans from inner to outer enamel in a mammoth molar showing the $^{84}\text{Sr}/^{86}\text{Sr}$ ratio along with $^{82}\text{X}/^{86}\text{Sr}_m$, $^{83}\text{Kr}/^{86}\text{Sr}_m$ isobaric monitors, compared with zinc concentration measured along the same line and expressed as ^{66}Zn ion beam intensity. Experiments were conducted with $^{232}\text{Th}^{16}\text{O}/^{232}\text{Th} = 2\%$ (a and b) and 50% (c and d).

low oxide production (Fig. 6c and d). Under such conditions, the final $^{84}\text{Sr}/^{86}\text{Sr}$ ratio becomes highly inaccurate. A “by-product” of our experiments is the observation of a rise in the $^{82}\text{X}/^{86}\text{Sr}$ ratio across the enamel, even when the Zn content gently decreases. We did not observe any chemical change in the studied profile that could explain this observation, which certainly deserves further investigation. Overall, our experiments show that even at fairly low oxide levels, ZnO is a significant obstacle in obtaining accurate $^{84}\text{Sr}/^{86}\text{Sr}$ ratios in materials with elevated Zn/Sr ratios.

Better plasma sampling and ion extraction with the Jet-X cones also results in smaller isotopic fractionation, which was also observed for the wet plasma, solution mode Sr isotopic composition analyses.²⁰ The latter authors also reported a good agreement between the fractionation trend and the exponential law. Our data show a narrow range of the beta factors from -1.25 to -1.08 for nearly all our measurements (Fig. 4g). The fractionation trend has a gradient of 0.4978 ± 0.0057 (MSWD = 0.6) and deviates from the expected exponential law prediction by 1.15% but overlaps with it within uncertainty and does not have any bearing on the accuracy of $^{87}\text{Sr}/^{86}\text{Sr}$ measurements (Fig. 4h). The higher error of the estimated gradient is due to the smaller amount of data and the smaller range of isotope fractionation. Smaller isotopic fractionation is also reflected in the slightly smaller elemental fraction between Rb and Sr, with $^{85}\text{Rb}/^{86}\text{Sr} = 1.138$.

Spatial and temporal resolution of seasonal Sr isotopic composition record in woolly mammoth enamel

Unlike sharks, a marine open-sea water animal living in an environment with virtually no variation in Sr isotopes, the

variability of bioavailable Sr in terrestrial animals is considerable. This is the foundation of the application of Sr isotopes in studies of animal or human mobility.^{52–55} A woolly mammoth molar, with its prolonged mineralisation time (>10 years), large size and structure that helps to better withstand diagenetic changes, provides an insight into a significant period of an animal's life recorded in a single dental plate.^{29,56–58} A single molar comprises several dental plates, whose formation in the mesio-distal direction is slightly displaced in time, but adjacent plates reflect the uptake of virtually the same Sr, and their measured isotopic composition should show differences only related to different enamel formation rates. The details of the tooth enamel histology are beyond the scope of this article. For a practical summary in the context of Sr isotopic measurements, the reader is referred to *e.g.* ref. 11, 29, 48, 56 and 58–60. In this study, we used a woolly mammoth molar from the Torske region of Western Ukraine and compared its remarkable compositional variation in adjacent plates to test the limits of spatial and temporal resolution, along with the precision and accuracy of LA-MC-ICPMS analyses.

The analyses were performed with the setups described above, namely, MC-ICPMS *Neptune* with H-type cones and MC-ICPMS *Neoma* with (1) the H-type and (2) the high sensitivity (Jet-X) cones for different laser beam diameters. As the S-155 laser sample cell was large enough to fit an entire dental plate of a mammoth molar, the measurements were taken as a single line scan (Fig. 7a). The ablation lines were placed near the enamel–dentine junction (EDJ) and proceeded from the cusp (early forming enamel) to the cervix (late forming enamel). The results obtained with the *Neptune* for the longest centrally

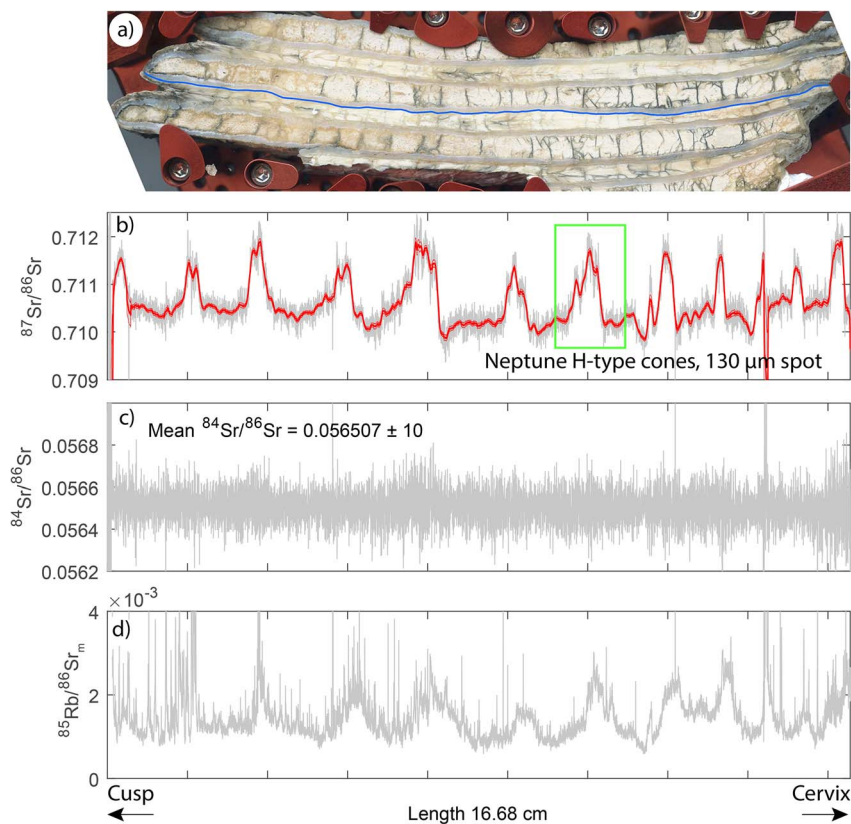


Fig. 7 Cusp to cervix Sr isotopic composition of the dental enamel of a woolly mammoth from the Torske region of Western Ukraine. Measurements performed with the LA-MC-ICPMS *Neptune* using H-type cones; (a) scan of a molar tooth fragment with a marked ablation line (blue colour) within the internal enamel layer; (b) $^{87}\text{Sr}/^{86}\text{Sr}$, (c) $^{84}\text{Sr}/^{86}\text{Sr}$, and (d) measured $^{85}\text{Rb}/^{86}\text{Sr}_m$. Individual 1 s integration is shown in grey. The red lines represent smoothed data using a moving average with a span of 50 s. The line thickness corresponds to $\pm 2\text{SE}$ uncertainty.

located tooth plate (Fig. 7a), summarised in Fig. 7b–d, show twelve cyclic changes in the $^{87}\text{Sr}/^{86}\text{Sr}$ ratio ranging from 0.7099 to 0.7119 and invariant $^{84}\text{Sr}/^{86}\text{Sr}$ with a mean value of 0.056507 ± 0.000010 (2SD). The latter ratio is accurate and indicates the absence of significant dimer production, with the mean of the interference monitor ratio $^{82}(\text{Kr} + \text{CaCa} + \text{CaAr})/^{86}\text{Sr}_m = 0.000025 \pm 0.000014$ (2SD). The accurate $^{87}\text{Sr}/^{86}\text{Sr}$ ratio obtained for the accompanying matrix-matched shark tooth analyses proves $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ to be negligible, as expected for low-oxide plasma. This leaves ^{87}Rb as the only significant isobaric interference. The $^{85}\text{Rb}/^{86}\text{Sr}$ ratio in the shark (≤ 0.0001) is more than an order of magnitude lower than that observed in terrestrial animals (Fig. 7c), and the shark tooth cannot serve as an optimal verification of the accuracy of the ^{87}Rb interference correction. Instead, we relied on the reference ratio determined using the Rb–Sr mixed solutions, which demonstrates the reliability of the ^{85}Rb isobar correction well above $^{85}\text{Rb}/^{86}\text{Sr}_m$ of 0.025. Even a slightly higher interference can be accurately corrected.^{9,11}

The cusp to cervix variations along the EDJ in Sr concentration show an ^{88}Sr ion beam from 3.5 to 8.2 V. Smoothing the $^{87}\text{Sr}/^{86}\text{Sr}$ data using a moving average with a span of 50 seconds, corresponding to 0.8 mm of ablated enamel, gives an internal precision of 40 to 280 ppm (2RSE), with the vast majority of data (about 90%) falling within the range of 40 to 120 ppm (excluding locally mechanically damaged zones). The precision

varies greatly depending on the compositional gradient of the Sr isotopic composition (Fig. 7b–d). Our histomorphometric record, with a close-to-a-week resolution chronology of enamel secretion, suggests that the observed compositional cyclicality reflects 12 annual cycles interpreted as diet changes during the cold and warm seasons. The decreasing ‘wavelength’ of the cycles towards the cervix represents the slowing down of the recruitment rate of new ameloblasts in the secretion front along the EDJ with time. At each stage of enamel formation, the precision obtained is more than satisfactory for resolving temporal variations in $^{87}\text{Sr}/^{86}\text{Sr}$ at the sub-seasonal level.

The results obtained using *Neoma* with the H cone for different dental plates from the same tooth are in excellent agreement with the data obtained using *Neptune*. The compositional details can be easily followed and correlated with the analyses of both instruments (Fig. 7 and 8). The non-radiogenic $^{84}\text{Sr}/^{86}\text{Sr}$ ratio is also accurate and shows no indication of the presence of polyatomic Ca dimer/argide interferences. The internal precision obtained for the time equivalent fragments of dental enamel is comparable for both instruments, ranging from 70 to 110 ppm (2RSE). Measurements of contemporaneously mineralised enamel on the opposite side of the same tooth plate, with the same laser parameters but using Jet-X high-sensitivity cones, are in perfect agreement with the H-type cones (Fig. 8c). Higher sensitivity (4.5 to 8.0 V of ^{88}Sr) led to improved

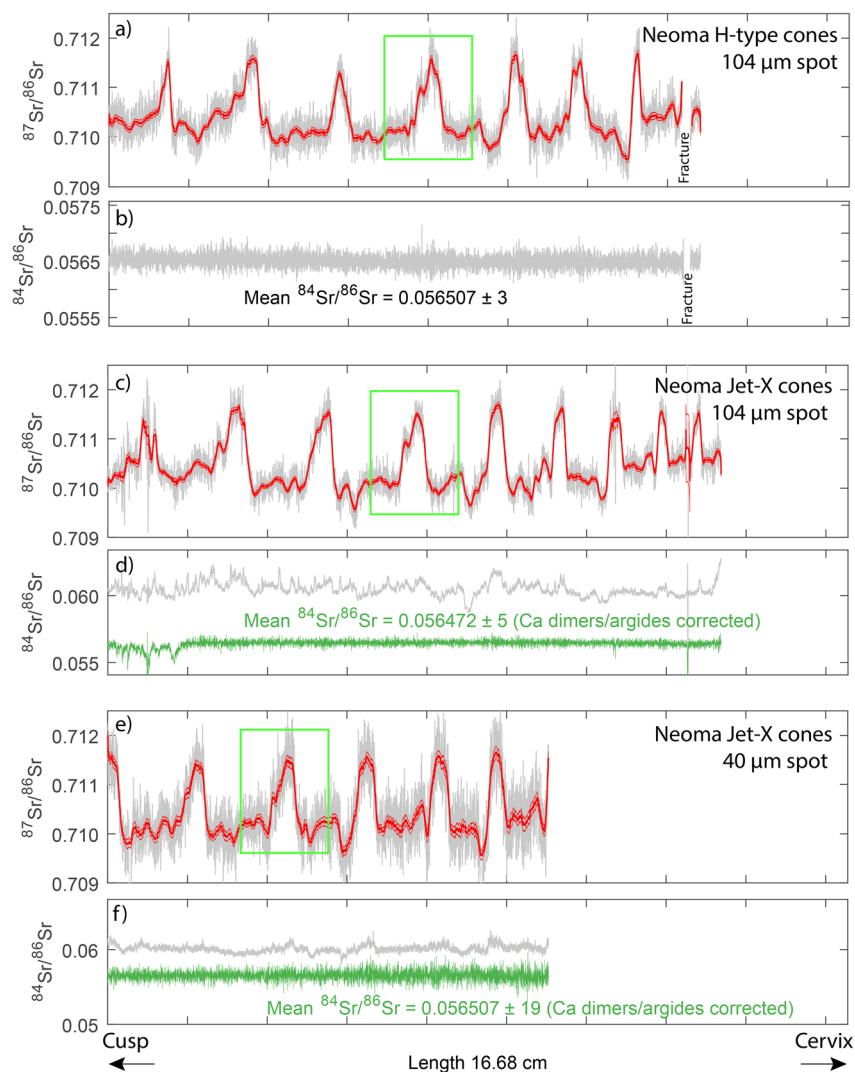


Fig. 8 Cusp to cervix Sr isotopic composition measurements of woolly mammoth enamel performed using LA-MC-ICPMS *Neoma*. Measurements were conducted with the H-type cones (a and b) and high-sensitivity Jet-X cones on the opposite side of the same dental plate (c and d). A shorter dental plate was analysed using Jet-X cones with a 40 μm laser beam diameter (e and f). Green rectangles show equivalent regions mineralized at approximately the same time (see also Fig. 9).

internal precision (50–90 ppm 2RSE). However, the normalised $^{84}\text{Sr}/^{86}\text{Sr}$ ratio fluctuates strongly throughout a run, ranging from approximately 0.060 to 0.062 (Fig. 8c and d). An additional correction for Ca dimer/argide isobars gives a stable and accurate $^{84}\text{Sr}/^{86}\text{Sr} = 0.056472 \pm 0.000005$ (2SE). A small region at the beginning of the scan deviates from this value because of intense fracturing, which also affects the corresponding $^{87}\text{Sr}/^{86}\text{Sr}$ ratio (Fig. 8d). Reducing the spot size to 40 μm reduces the ablated sample volume almost 7 times. The intensity of ^{88}Sr ranged from approximately 1.0 to 2.5 V. Despite the lower internal precision, all compositional details are visible (Fig. 8e and f). The lower precision did not obscure the high spatial resolution advantage. As expected, it revealed more compositional details. This is well illustrated by magnifying the same peak measured with the 3 different settings (Fig. 9a–c). The region with the highest $^{87}\text{Sr}/^{86}\text{Sr}$ ratio, visible in the 40 μm spot measurement, is only detected as a sharp peak in the 104 μm

spot measurements. Similarly, smaller peaks are much better defined in the regions of less radiogenic Sr when scanned with a smaller ablation spot diameter (Fig. 9c). Despite some loss of precision, the increased spatial resolution is certainly beneficial in terms of obtaining a more detailed reconstruction of the temporal changes in the mineralised enamel. The increased sensitivity provided by the Jet-X cones not only allows the use of a smaller laser beam diameter but also reduces the integration time owing to the improved counting statistics, resulting in improved spatial and temporal resolution of measurements even with a larger laser beam. Reducing the integration region from 50 to 25 seconds, which represents 0.4 mm of ablated enamel, corresponding to approximately 10 days of enamel growth, reveals much more detail, similar to analyses using the smaller laser spot (Fig. 9e). The same reduction of the integration region in the case of the 40 μm beam gives no improvement, which is simply a consequence of the poorer counting

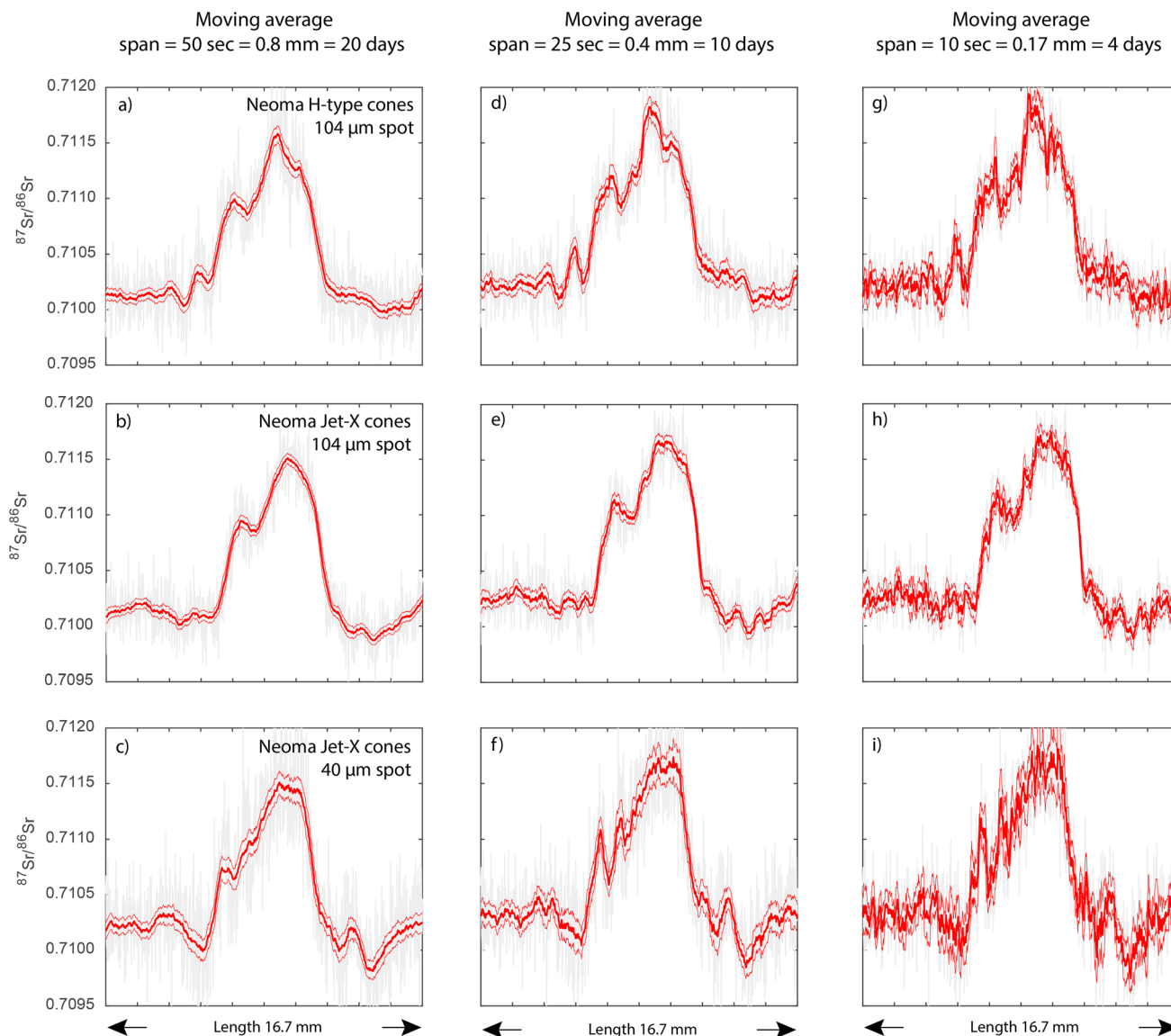


Fig. 9 Comparison of measurement precision using LA-MC-ICPMS *Neoma* with H-type (a) and Jet-X cones (b and c) in an enamel region corresponding to approximately 1 year of mineralization. Laser beam diameters of 104 μm (a and b) and 40 μm (c) were applied. Precision is shown for 50, 25, and 10 s integrations corresponding to enamel mineralization times of approximately 20 (a–c), 10 (d–f) and 4 (g–i) days, respectively. The optimal compromise between precision and highest temporal resolution is represented by the analyses with Jet-X cones, 104 μm diameter and 25 seconds integration time (e). The highest spatial resolution was achieved with a 40 μm diameter and 50 second integration time (c). The selected region is marked with a green rectangle in Fig. 8a, c and d. The heavy red line represents the moving average, and the light red lines represent $\pm 2\text{SE}$ uncertainty.

statistics. Reducing the integration region even further, to 10 seconds (0.17 mm of mineralised enamel = 4 days), does not bring any improvement, although the major compositional changes can still be depicted and might still be useful for mobility studies. Although the high spatial and temporal resolution obtained cannot generally be interpreted with the same level of detail when reconstructing migratory paths (the regional coverage of Sr isotopic composition is not known in sufficient detail), such high-resolution data provide additional insight into the habits of the animals, especially when combined with other techniques such as dental histology, stable isotope, trace element data and isoscapes. The use of

high-sensitivity Jet-X cones will be even more appreciated in the analysis of enamel with low extension (formation) rates and in the analysis of higher trophic level animals, particularly humans, whose dental enamel contains considerably less Sr. The reduced laser beam diameter also allows the separation of different stages of enamel formation and the recovery of the least disturbed *in vivo* record of bio-apatite mineralisation.

Conclusions

The long-term measurements of SRM 987 using MC-ICPMS Neptune demonstrate the high accuracy and reproducibility of

$^{87}\text{Sr}/^{86}\text{Sr}$ and $^{84}\text{Sr}/^{86}\text{Sr}$ under both wet and dry plasma conditions, with the dry plasma values more closely matching the TIMS reference data. Solution-based analyses of contemporary shark dental enameloid, used as a bioapatite reference material for LA MC-ICPMS, yield an $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.709176 ± 0.000008 (2SD), consistent with present-day seawater. Laser ablation MC-ICPMS analyses of shark enameloids using the *Neptune* and *Neoma* instruments with H-type cones yielded consistent $^{87}\text{Sr}/^{86}\text{Sr}$ ratios, with internal precision ranging from 50 to 25 ppm (2RSE) and long-term reproducibility of ~ 50 ppm for *Neptune* and ~ 40 ppm (2RSD) for *Neoma* cones. Mean $^{87}\text{Sr}/^{86}\text{Sr}$ values were 0.709194 ± 0.000034 (*Neptune*) and 0.709189 ± 0.000027 (*Neoma*, 2SD), which agreed well with the solution-based reference data. No significant interferences from $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ or Ca-dimer/argides were observed. The use of a Jet skimmer and X-type sampler cones on *Neoma* increases oxide production, introducing $^{40}\text{Ca}^{31}\text{P}^{16}\text{O}$ interference. This can be effectively suppressed by plasma tuning while preserving a 2–3 times signal enhancement, improving $^{87}\text{Sr}/^{86}\text{Sr}$ precision by at least a factor of two. Elevated Ca dimer/argide species bias the $^{84}\text{Sr}/^{86}\text{Sr}$ ratio but are correctable to an accurate value. Minor $^{68}\text{Zn}^{16}\text{O}$ interference on ^{84}Sr is minimized at low oxide levels ($<2\%$ $^{232}\text{Th}^{16}\text{O}/^{232}\text{Th}$). A notable feature of $^{87}\text{Sr}/^{86}\text{Sr}$ analysis by MC-ICPMS is its mass bias behaviour, which follows the exponential law across all instrument configurations. This is attributed to the small average mass difference between the normalised and normalising ratios, as also seen for isotopic measurements of Nd.^{35,36}

We tested different LA-MC-ICPMS configurations to examine seasonal variations in Sr isotopic composition in woolly mammoth enamel, obtaining high-precision measurements with sub-monthly resolution. The use of Jet-X cones enhanced the sensitivity, enabling improved spatial resolution. Reducing the laser spot diameter to 40 μm results in lower precision, yet still sufficiently high, allowing for the detection of finer compositional details that would otherwise be averaged out with a larger beam size. Similarly, shortening the integration time at higher signal intensities resulted in a resolution of <10 days for moderate enamel extension rates. Enhanced instrument sensitivity, leading to higher spatial and temporal resolution, is vital for analysing organisms at higher trophic levels, incorporating much lower amounts of Sr, such as humans, and bio-apatite tissues mineralizing at a slower pace. Another advantage is the improved ability to separate the *in vivo* record in the earliest formed enamel from the later maturation overprint, which is critical for reconstructing the palaeoecology of large mammals and the life histories of humans.

Data availability

Data for this article are available at <https://database.ing.pan.pl/> and <https://dataportal.ing.pan.pl/>.

Conflicts of interest

There are no conflicts to declare.

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