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Fast and high spatially resolved multielemental mapping of otoliths using IR&UV-femtosecond laser ablation-ICP-TOFMS

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A Femtosecond Laser Ablation-Inductively Coupled Plasma-Time of Flight Mass Spectrometer (fs-LA-ICP-TOFMS) was evaluated for multi-elemental mapping of complex otoliths; in particular, for the analysis of a sample of 8 year-old freshwater fish otolith with irregular growth ring distribution pattern. The fundamental wavelength ($\lambda \sim 1028$ nm) and the 4th harmonic ($\lambda \sim 257$ nm) of a diode pumped Yb:KGW fs-laser (<290 fs) were successfully employed for otolith elemental mapping (map area $\sim 6000 \mu\text{m}^2$), at high spatial resolution (spot sizes $\leq 5 \mu\text{m}$) and with negligible thermal effects. Fast, sensitive multi-elemental analysis was achieved combining the use of high laser repetition rates (≥ 200 Hz), high sample scanning speeds ($100 \mu\text{m s}^{-1}$), a fast response time ablation cell (<40 ms), and the use of a ICP-TOFMS equipped with a reaction/collision cell (operated in He and H₂) to remove plasma gas ion related species and polyatomic interferences. The multi-elemental (from ²³Na⁺ to ²³⁸U⁺) fs-LA-ICP-TOFMS mapping of the *Prochilodus lineatus lapillus* otolith allowed the identification of the appropriate growth axis based on the chronological chemical record associated with the endogenous and exogenous changes experienced by the fish throughout its life. Moreover, each of the detected and quantified elements provided useful biochemical information about environmental and endogenous factors. For instance, maximum P concentration values were observed during spring-summer, a favorable period for feeding and growth.

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Introduction

The chemical composition of biocarbonates has been widely used as an environmental indicator and to describe the life history of both terrestrial and aquatic organisms.^{1,2} Biocarbonates are calcified structures whose morphology and chemical composition is regulated by endogenous and exogenous factors.^{3,4} Among the most used are the otoliths of bony fish. These structures are housed in the inner ear of fish, are acellular and are composed of calcium carbonate crystals, precipitated mainly in the form of aragonite immersed in a protein matrix.⁵ The otoliths grow continuously, forming growth rings (similar to a tree), and they incorporate different elements from the surrounding environment, either through water or diet.⁵ Because they are metabolically inert, they keep

a chronological chemical record of the environments experienced by the fish throughout life and have been used as environmental proxies.^{6,7} The area of early growth (embryo-larva) is known as the core, while the edge of the structure represents the recent life span.

The use of Laser Ablation in combination with Inductively Coupled Plasma Mass Spectrometry (LA-ICP-MS) has allowed the determination of selected number of isotopes with high resolution, through core-to-edge transects.⁶ Typically, nanosecond pulsed laser ablation units (e.g., Nd:YAG and ArF* excimer) operated in the UV region have been employed for the ablation of otoliths at micrometer scale (e.g., laser spot size $\varnothing = 10-100 \mu\text{m}$).^{8,9} Shorter pulsed duration lasers, such as femtosecond lasers (fs-LA), are considered to generate laser-induced aerosols with a broad monomodal particle size distribution in the nanometer range (nano-sized agglomerates in the range of 50 up to 250 nm), resulting in higher ion signals and more stable elemental ratios when analyzed by ICP-MS.^{10,11} Nevertheless, it is considered that reduced elemental fractionation during the fs laser ablation might still be present.¹²

Multi-elemental ratios (e.g., Sr/Ca, Mg/Ca, Ba/Ca or Na/Ca) or isotopic (e.g., ⁸⁷Sr/⁸⁶Sr) data from otolith analyses can be provided by different LA-ICP-MS systems including quadrupole-based (QMS), sector-field-based (SFMS), or multicollector-based

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(MC-MS) mass spectrometers. It is also possible to couple a laser ablation system with multiple ICP-MS systems using an aerosol split to achieve simultaneous multi-elemental and isotopic data on a fish otolith surface.¹³ Quantitative results require the use of external certified reference materials (e.g., NIST 610/612 or MACS3), which are employed for drift corrections and to obtain calibration curves or response factors. Nevertheless, recent studies have also shown other quantitative approaches for the determination of trace elements in otoliths without the requirement of external reference material, making use of online isotope dilution (ID)-LA-ICP-MS. This method was based on two aerosols from an ablated sample and a nebulized isotopically enriched spike solution, which were online mixed using a dual-port spray chamber.¹⁴

Otolith analyzes usually assume that the calcium carbonate matrix is deposited homogeneously, forming well-defined layers perpendicular to the axis of growth (approximately concentric). However, it has been documented that the heterogeneity in the elemental distribution within each growth ring is greater than assumed.¹⁵ In addition, both environmental and endogenous factors can affect the microstructure of the otolith causing structural discontinuities¹⁶ and even the form of crystallization of calcium carbonate,¹⁵ which could lead to interpretation errors if the readings are made in regions whose growth does not have a chronological order. This problem might be addressed by evaluating the distribution of elements in two dimensions (mapping the whole otolith). In this sense, in the last two decades different methods based on X-ray fluorescence (XRF) have been explored to perform 2D maps in calcified structures. For instance, nuclear microprobe combined with proton-induced energy emission spectroscopy (micro-PIXE)¹⁷ was employed to achieve high spatial resolution (5 μm) and moderate/high limits of detection (e.g., $>1-2 \mu\text{g g}^{-1}$ for Sr¹⁸ and $>160 \mu\text{g g}^{-1}$ for Ba¹⁹) for the detection of masses greater than ²⁷Al. More recently synchrotron X-ray fluorescence microscopy (SXFM)^{15,20} and wavelength-dispersive X-ray electron probe microanalyzer (EPMA)²¹ have been used. SXFM provides excellent limits of detection (near ng g⁻¹ for some analytes), but relatively long analytical times, ranging from a few hours to 24 hours in extreme cases (depending on the instrument configuration it can exceed 8 h for surfaces $>4 \text{ mm}^2$).²² Furthermore, this technique require large-scale installations and it has the limitation of mainly detecting masses greater than ⁴⁰Ca. On the other hand, EPMA achieves high spatial resolution ($<3 \mu\text{m}$),²³ but it provides relatively poor detection limits ($>10 \mu\text{g g}^{-1}$), not allowing the detection of some key analytes frequently present in otoliths (e.g., Li, Mn, Ba, Se).²⁴ LA-ICP-MS based on the use of sequential mass spectrometers might also allow multiple scans to cover the entire sample surface; however, depending on the size of the spot, the sample scanning speed, the required resolution, and the total integration times, it could take relatively long times. For instance, mapping an area of about 1500 \times 4000 μm (6 mm²) using a spot size of 5 μm and a scanning speed of 5 $\mu\text{m s}^{-1}$ (e.g., low speed for multi-elemental analysis (>20 isotopes)), might take more than 66 h (more than 2 days and a half), resulting in highly expensive and impractical. Nevertheless, the recent combination of advanced laser

ablation units (including fast laser repetition rate, high sample scanning speeds and ultra-fast wash-out ablation cells), with ICP-TOFMS systems (including collision/reaction cells and fast detectors that provide high mass spectra acquisition rates), has tremendously enhanced the analytical potential of LA-ICP-MS for fast multi-elemental mapping.^{25,26} Therefore, in this study, a IR&UV-fs-LA-ICP-TOFMS system is evaluated for fast and sensitive quantitative multi-elemental mapping of complex otoliths. In particular, a sample of 8 year-old *Prochilodus lineatus lapillus* otolith from the Bermejo River Basin (Argentina), which is subject to regular periods of floods and droughts (summer-winter), was considered as a model of a complex otolith with irregular growth ring distribution pattern.

Experimental

Instrumentation

A NWRFemto UltraCompact laser ablation system from Elemental Scientific Inc (ESL) was used in these experiments. This instrument consists of a diode pumped Yb:KGW fs-laser (<290 fs, Pharos, Light Conversion) that can be operated at the two different wavelengths ($\lambda \sim 1028 \text{ nm}$ (fundamental), and at $\lambda \sim 257 \text{ nm}$ (4th harmonic)). It provides repetition rates from 1 Hz up to 900 Hz, fluence at the sample surface up to 20 J cm⁻², and nominal spot sizes from 2 μm to 65 μm of diameter (\varnothing). A fast wash-out ablation cell (TwoVol 2) combined with a capillary inlet system provided washout times shorter than 40 ms. It should be remarked that ultra-fast wash-out ablation cells with response times even shorter than 1 ms are nowadays commercially available. Helium gas (99.999% minimum purity) from AirLiquide was used as cell carrier gas at a constant flow rate of 350 mL min⁻¹. The laser-induced aerosol was transported from the inner cup of the ablation cell to the ICP-TOFMS through a flexible peek tube ($<1 \text{ m}$ length and 1 mm i. d.).

The detection was carried out by a Vitesse ICP-TOFMS (Nu Instruments, UK). Operation conditions in the ICP-MS (e.g., ion optics, gas flows) are optimized on a daily basis. In particular, the ion optics are adjusted to obtain the maximum sensitivity monitoring ⁴⁴Ca⁺, ⁸⁹Y⁺, ¹³⁹La⁺, ²³²Th⁺, and ²³⁸U⁺ signals produced by the ablation of the standard reference material SRM NIST 610 in raster mode at 100 Hz, 10 μm spot size, 10 $\mu\text{m s}^{-1}$ and at a fluence of 3 J cm⁻². Additionally, the makeup gas flow is adjusted to keep the ratios ²³⁸U⁺/²³²Th⁺ below 120%. High purity Ar (99.999% minimum purity) from AirLiquide was employed as plasma and make up gas. Ar was coaxially mixed with the He carrier gas before entering the ICP. Moreover, the mass spectrometer is equipped with a collision/reaction cell operated with a flow rate of He (0.016 L min⁻¹) and H₂ (0.006 L min⁻¹) to remove polyatomic interferences, and to remove Ar⁺ related-ions, protecting the detector from this major plasma element. In this sense, it was possible to detect some polyatomic-interfered key-isotopes in otolith analysis, including ³¹P, or ⁸⁰Se. The TOFMS was operated at 30 kHz. Moreover, every two spectra, isotope ion signal are integrated, and background subtracted (considering the ion signals at both sides of the isotope-peak region). This process is repeated 40 times to achieve a full discrete spectrum from Na⁺ to U⁺ every 2.13 ms.



Table 1 ICP-TOFMS operating conditions

Applied power	1300 W
Coolant Ar flow rate	13 L min ⁻¹
Auxiliary Ar flow rate	2 L min ⁻¹
Nebulizer Ar flow rate	1080 mL min ⁻¹
Sample depth	17.2 mm
Entry focus lens	-120 V
Collision cell gas	He at 0.016 L min ⁻¹ and H ₂ at 0.006 L min ⁻¹
Pushout & pullout HV voltages	1630 V
Reflectron HV	2117 V
Detector HV	2492 V
Acquired mass to charge range	20–260 m/z

Additionally, blanking voltage (200 V) based on the use of Bradbury–Nielsen style timed ion gates were applied to remove selected isotopes from major elements before reaching the detector. ICP-TOFMS operating conditions are listed in Table 1.

Sample description

A sample of 8 year-old *Prochilodus lineatus lapillus* otolith from the Bermejo system (La Plata Basin, Argentina), which is deposited in the COLV (ID-3987) otolith collection (INPA-CICET-UBA, Argentina), was analyzed. This sample is a good analysis model because the Bermejo system is subject to regular periods of floods and droughts (summer–winter), which severely modifies the environments and physicochemical parameters of the water (e.g., salinity, temperature). These seasonal variations affect otolith growth, resulting in structures with an irregular growth ring distribution pattern. Furthermore, these seasonal variations produce alterations in the incorporation of elements to the otolith. Particularly for this species, including the bermejo river, a direct relationship in Sr (expressed as Sr/Ca ratio) between otolith, water and conductivity was reported, positioning this element in otolith as a good environmental proxy.²⁷

The otolith was embedded in epoxy resin (Cristal Tack, Argentina) and then cut a section at core level 800 µm thick using a low-speed saw (Buehler Isomet, China). Otolith slice was mounted on a glass slide using epoxy resin and manually polished with 800–2000 µm grid wet sandpapers. The otolith section was sonicated for 10 min submerged in ultrapure water (18.2 MΩ cm⁻¹) to remove particles generated during polishing.

Reference materials and data treatment software

NIST 610, NIST 612 (trace elements in glass) and MACS3 (trace elements in synthetic calcium carbonate) were employed as reference materials. These samples were analyzed at the beginning and at the end of the otolith mapping studies (bracketing method) to be used for drifts corrections and to calculate relative sensitive factors that allowed the conversion of qualitative data into quantitative multi-elemental maps (Ca was used as an internal standard).

Iolite software (v4.8.7)²⁸ was employed for data reduction to obtain multi-elemental maps and transects along different

directions in the otolith; and for calculations of LODs and concentration recoveries.

Results and discussion

IR-fs versus UV-fs for otolith analysis

The fs-LA-ICP-TOFMS experimental conditions for fast multi-elemental otolith mapping were optimized considering a compromise among spatial resolution, sensitivity and time of analysis. In particular, laser ablation conditions were selected to achieve the smallest detectable spot sizes in the otolith for each of the two available wavelengths ($\lambda \sim 1028$ nm and $\lambda \sim 257$ nm, respectively) and to apply the same laser spot overlapping percentage while scanning the sample at a constant speed (100 $\mu\text{m s}^{-1}$). When employing the fundamental wavelength of the Yb:KGW fs-laser (IR wavelength), the smallest circular spot size that provided a proper ablation of otoliths was $\varnothing 5$ µm, using a fluence of about 7.4 J cm⁻². For this spot size the fs laser was operated at 200 Hz to achieve a sample displacement of 0.5 µm per laser shot (90% overlapping requiring 10 laser shots per spot in a total time of 50 ms). However, using the 4th harmonic of the Yb:KGW fs-laser (UV wavelength), the smallest circular spot size that provided a proper ablation of otoliths was $\varnothing 4$ µm, using a fluence of about 6.6 J cm⁻². In that case, the fs laser was operated at 250 Hz to achieve a sample displacement of 0.4 µm per laser shot (90% overlapping requiring 10 laser shots per spot in a total time of 40 ms, which is approximately the response time of the TwoVol2 ablation cell combined with the capillary inlet system).

The TOFMS acquisition method was adapted for the different matrices employed in this study to avoid the saturation of the detector. When analyzing NIST 610 and NIST 612 reference materials, blanking was employed to remove ²³Na⁺, ²⁸Si⁺ and ⁴⁰Ca⁺ ion signals considering the high concentration of these elements in these glass samples. In this case, mass spectra were collected from 23.2 to 260 m/z, and a blanking region was applied between 27.8 and 40.2 m/z (also removing ³¹P⁺ ion signal). During the analysis of MACS3 reference material and the otolith (both CaCO₃ matrices), mass spectra were collected from 20 to 260 m/z, and a blanking region was applied between 33.8 and 42.8 m/z to remove ⁴⁰Ca⁺ ion signals. It should be highlighted that the minimum mass blanking window increase, from 5 m/z width to 15 m/z width, at increasing masses because blanking is based on the application of fast time ion gates (≥ 50 ns). Moreover, blanking process is calibrated to completely remove the signals of the blanked masses while not affecting the sensitivity of the nearby masses.

The otolith multi-elemental mapping analysis covered a region of 1540 µm (height) \times 4000 µm (length). This area was divided into two different mapping regions: the upper region (height \times length = 768 \times 4000 µm²) was ablated using the UV-fs laser (192 parallel raster lines) and it took about 2 h 10 min; while the lower region (height \times length = 770 \times 4000 µm²) was ablated using the IR-fs laser (154 parallel raster lines) and it took about 1 h 30 min. Therefore, the total multielemental map of the otolith took about 3 h 40 min, which was up to 20 times faster than using typical operating conditions employed in



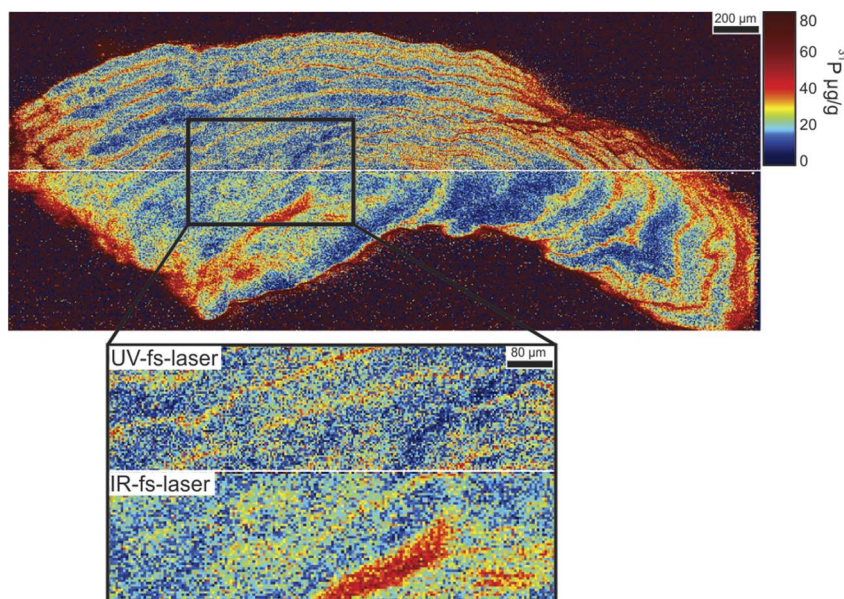


Fig. 1 Quantitative spatial distribution of P within the otolith. Upper map was obtained using UV-fs-LA; while the lower map was obtained using IR-fs-LA. Figure inset shows in higher detail the spatial distribution of P in the otolith (zoom of the region highlighted with a dashed rectangle). Scale based on non-linear ECDF (Empirical Cumulative Distribution Function).

multielemental (≥ 20 isotopes) LA-ICP-QMS analysis and even faster than the SXFM.²⁰ As an example, Fig. 1 shows the quantitative spatial distribution of P within the otolith (calculated using MACS3 as external reference material), which demonstrates the analytical potential of this method. It is noticed that

using both wavelengths at the selected operating conditions it is possible to resolve the growing rings from the otolith core and to determine the heterogeneous distribution of this minor element (concentrations lower than $80 \mu\text{g g}^{-1}$). Scale based on non-linear ECDF (Empirical Cumulative Distribution Function)

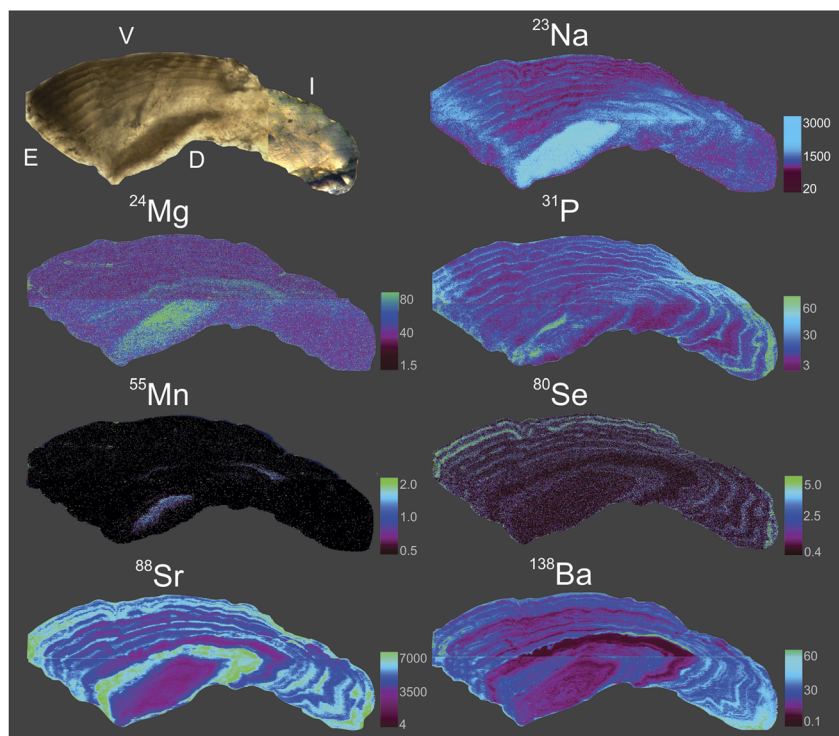


Fig. 2 Quantitative spatial distribution of elements in a cross section of a *Pochilodus lineatus lapillus* otolith determined by UV&IR-fs-LA-ICP-TOFMS (upper and lower maps were merged). V: ventral; D: dorsal; I: internal; E: external face. Scale based on non-linear ECDF (Empirical Cumulative Distribution Function); except for Mn and Se trace analytes, whose scale is linear. Scale concentration units: $\mu\text{g g}^{-1}$.



that maximizes contrast of the image.²⁹ Fig. 1 inlet shows in higher detail the spatial distribution of P in the otolith (zoom of the region highlighted with a dashed rectangle). Mass spectra could be exported every 2.13 ms; however, maps were built up considering an average mass spectra per spot in the sample in order to minimize the amount of data providing comprehensive information. Therefore, considering 10 laser shots per spot, approx. 1520 spectra (# spectra measured in 40 ms) were average per 4 μm spot when using UV-fs-LA, and approx. 1920 (# spectra measured in 50 ms) spectra were average per 5 μm spot when using the IR-fs-laser. Furthermore, the use of both femtosecond laser wavelengths (IR and UV) provided analogous P concentrations, highlighting the potential use of both of them for the analysis of otoliths.

Otolith elemental distribution

The usefulness of otolith features as indicators depends, at least in part, on knowledge about the dynamics of incorporation into the otolith. Various elements present in otoliths can be used as

environmental (*e.g.*, temperature and salinity) or physiological (*e.g.*, hatching, reproduction, growth) indicators. Trace element incorporation into the otolith is governed by exogenous and endogenous factors. Among all the different isotopes that were measured from Na to U, ²³Na, ²⁴Mg, ⁵⁵Mn, ⁸⁰Se, ⁸⁸Sr, and ¹³⁸Ba are shown in detail as they show spatial heterogeneity, and their concentrations are consistently above the LOD. Fig. 2 shows an image of the otolith morphology, and the full quantitative spatial distributions of the other analytes within the otolith, merging the analysis carried out using both laser wavelengths (scale based on non-linear ECDF, except for the trace analytes ⁵⁵Mn and ⁸⁰Se where a linear scale was employed for clarity purposes). Qualitative data (analyte ion signals measured in cps) were converted into concentration values ($\mu\text{g g}^{-1}$) using ⁴⁴Ca as internal standard (38.8 wt%),³⁰ and using MACS3 as external reference material, except for ⁸⁰Se where NIST 610 was used as external reference material (Se concentration is not certified in MACS3). Moreover, Table 2 lists the LODs estimated for the selected isotopes using Iolite software. Calculations were performed on MACS3 reference material and using Longerich *et al.* criteria based on background noise.³¹ In particular, minimum and maximum values of calculated LODs from the $n = 8$ analysis of MACS3 reference material (4 before and 4 after otolith analysis) are listed. It is observed that UV-fs-LA provided slightly improved LODs *versus* IR-fs-LA. Furthermore, it is remarked that the applied methodology based on fs-LA-ICP-TOFMS showed lower detection limits than those reported for other methods such as micro-PIXE ($>160 \mu\text{g g}^{-1}$ for Ba¹⁹) and EPMA ($>10 \mu\text{g g}^{-1}$ for Mn, Ba and Se²⁴), which is optimal for the detection of elements present at relatively low concentrations.

Analyte concentrations and the recoveries were also calculated for NIST 610 and for NIST 612, using MACS3 as calibrating material (non-matrix matched calibration). Tables 3 and 4 list the certified analyte concentrations of MACS3, and the calculated analyte concentrations and recoveries in NIST 610 and

Table 2 Estimated LODs (minimum and maximum values) in a CaCO_3 matrix, at the selected fs-LA-ICP-TOFMS operating conditions, using MACS3 as external reference material (** except for ⁸⁰Se were NIST 610 was employed instead)

Analyte	Limit of detection ($\mu\text{g g}^{-1}$) using UV-fs-LA-ICP-TOFMS	Limit of detection ($\mu\text{g g}^{-1}$) using IR-fs-LA-ICP-TOFMS
²³ Na	15–20	15–45
²⁴ Mg	1.5–3.5	5–9
³¹ P	3–7	3–4
⁵⁵ Mn	0.55–0.65	0.55–0.7
⁸⁰ Se**	0.4–1.4	1.5–3
⁸⁸ Sr	0.5–4	1.5–14
¹³⁸ Ba	0.1–0.4	0.3–0.5

Table 3 Calculated concentrations and recoveries for different analytes from the analysis of NIST 610 and NIST 612, at the selected UV-fs-LA-ICP-TOFMS operating conditions, using MACS3 as external reference material

Analyte	Certified concentration MACS3 ($\mu\text{g g}^{-1}$)	Calculated – nominal concentrations ($\mu\text{g g}^{-1}$) and recoveries from NIST 610	Calculated – nominal concentrations ($\mu\text{g g}^{-1}$) and recoveries (%) from NIST 612
⁵⁵ Mn	534 ± 24	387–444 87%	28–39 72%
⁵⁶ Fe	9987 ± 1728	205–458 45%	14–51 28%
⁵⁹ Co*	54 ± 4	406–410 99%	26–35 73%
⁶³ Cu*	113 ± 4	393–441 89%	37–38 98%
⁸⁸ Sr	6675 ± 245	593–515 115%	82–78 105%
¹³⁸ Ba	58 ± 3	480–452 106%	37–39 94%
²⁰⁸ Pb*	60 ± 3	247–426 58%	30–39 79%
²³² Th*	54 ± 2	568–457 124%	38–38 101%



Table 4 Calculated concentrations and recoveries for different analytes from the analysis of NIST 610 and NIST 612, at the selected IR-fs-LA-ICP-TOFMS operating conditions, using MACS3 as external reference material

Analyte	Certified concentration MACS3 ($\mu\text{g g}^{-1}$)	Calculated – nominal concentrations ($\mu\text{g g}^{-1}$) and recoveries (%) from NIST 610	Calculated – nominal concentrations ($\mu\text{g g}^{-1}$) and recoveries (%) from NIST 612
^{55}Mn	534 ± 24	346–444 78%	22–39 57%
^{56}Fe	9987 ± 1728	153–458 34%	1–51 2%
$^{59}\text{Co}^*$	54 ± 4	385–410 94%	20–35 56%
$^{63}\text{Cu}^*$	113 ± 4	362–441 82%	27–38 71%
^{88}Sr	6675 ± 245	513–515 99%	58–78 74%
^{138}Ba	58 ± 3	441–452 98%	28–39 73%
$^{208}\text{Pb}^*$	60 ± 3	292–426 69%	20–39 52%
$^{232}\text{Th}^*$	54 ± 2	688–457 150%	41–38 108%

NIST 612 obtained using UV- and IR-fs-LA, respectively. Data related to analytes that were not detected during the analysis of the otoliths (highlighted with “*”) were included in the Tables to expand the mass spectra range of the study. It is noticed that most recoveries showed improved values when using ultraviolet radiation. Additionally, it is observed that worse values were obtained for the quantification of NIST 612 (more transparent glass), except for analytes such as Pb and Th, whose concentration was very similar in NIST 612 and in MACS3. The worst recovery case was found for the quantification of Fe in NIST 612, probably due to the fact that Fe is highly concentrated in MACS3 and also heterogeneously distributed.

Fig. 2 provided elemental-dependent information obtained from the high-spatially resolved multi-elemental maps. It is observed that these analytes are heterogeneously distributed, both in the transversal axis to the direction of growth (temporal variation) and within the same period or growth ring. Specifically, ^{23}Na showed a heterogeneous distribution within the growth rings, with the quantity being greater in the external edge and internal area of the otolith. ^{24}Mg showed the highest levels around the core ($<36.5 \mu\text{g g}^{-1}$). Moreover, it showed discontinuous rings (approximately at the first annual mark) and discrete areas (e.g., external-ventral area) with relatively high values ($<75.9 \mu\text{g g}^{-1}$). ^{55}Mn distribution pattern similar to Mg. It showed a temporal distribution with relatively low heterogeneity, except for the core where high values ($<1.15 \mu\text{g g}^{-1}$) were recorded with a distribution parallel to the dorsal edge. The rest of the structure showed discontinuous bands with relatively high concentration ($<0.73 \mu\text{g g}^{-1}$). ^{80}Se showed a weak annual periodicity. Two discontinuous rings (age 7 and 8 years) of relatively high concentration ($5.1\text{--}5.6 \mu\text{g g}^{-1}$) were observed. A cyclic temporal variation was observed for ^{88}Sr , with higher concentrations ($<6850 \mu\text{g g}^{-1}$) at the inner and external area of the otolith. Annual periodicity was also found for ^{138}Ba , with heterogeneous distribution within the growth marks.

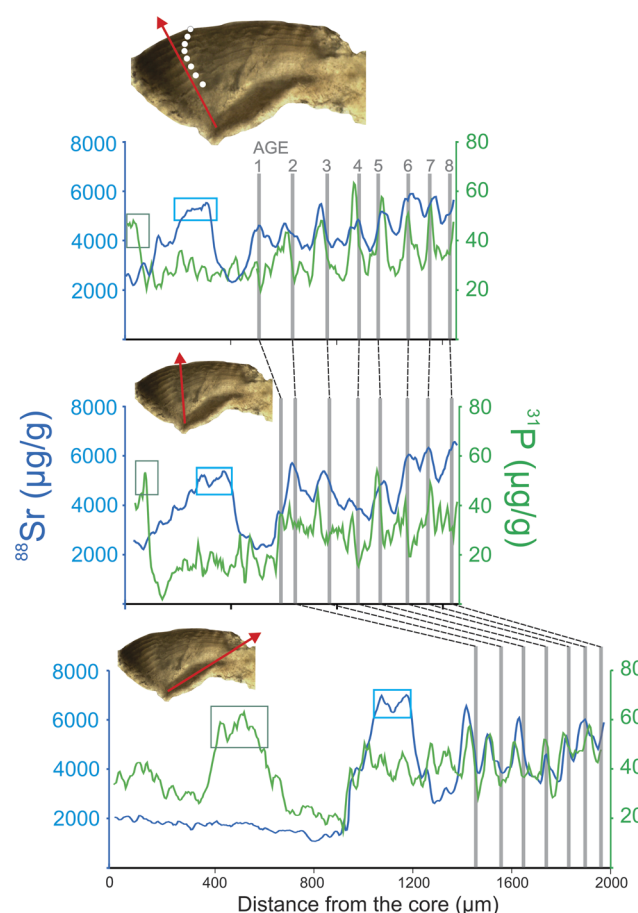


Fig. 3 Core-to-edge Sr and P profiles for transects performed in three different sections of the otolith. The red arrow on the otolith indicates the trajectory of the core-to-edge analysis and the white dots the position of the annual growth rings. The position of the rings (age) is also represented with gray vertical lines, which can be followed between the different analytical trajectories. The colored boxes within the first year of life indicate the same period represented in the different trajectories.



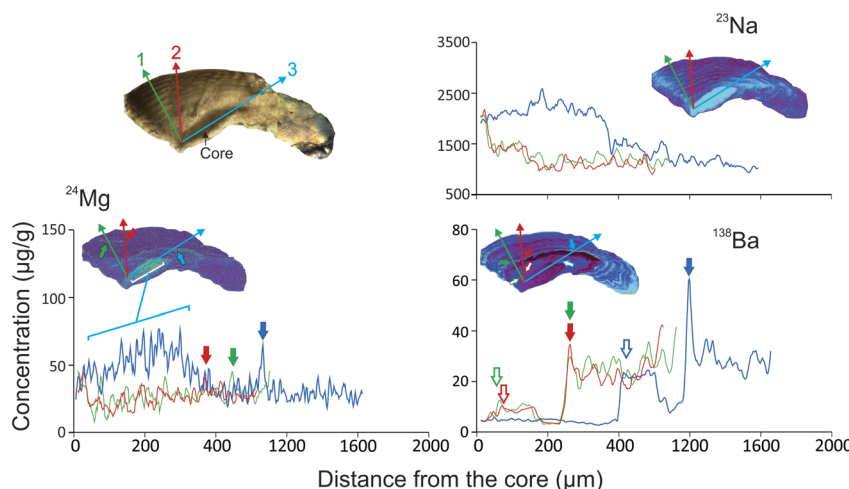


Fig. 4 Core-to-edge elemental profiles for transects performed in three different sections of the otolith. The arrows indicate the same period represented in the different trajectories.

Concentration variations from three core-to-edge profiles (transects) were extracted as a representation of three classic scenarios of elemental analysis in otoliths. For instance, Fig. 3 shows the concentration variations for ^{88}Sr and ^{31}P along these three core-to-edge transects. The three core-to-edge transects showed very different profiles for Sr: transect 1, which was carried out in the portion where the growth rings showed a homogeneous structuring, revealed a dominant peak ($5408 \mu\text{g g}^{-1}$) in the core area (age < 1 year) and eight oscillations with annual periodicity starting from the first year; transect 2 also showed an annual periodicity, but with a different pattern, especially in the core where a double top is evident. These peaks are also of lesser magnitude in relation to the first profile ($5489 \mu\text{g g}^{-1}$); in transect 3 the first annual mark is lost, and 7 oscillations are observed. A more pronounced double top is observed in the core, with highest values ($< 6854 \mu\text{g g}^{-1}$) than the other transects. In relation to P, only transect 3 registered the maximum concentration values due to the high heterogeneity in the core.

Moreover, Fig. 4 shows the concentration variations for ^{23}Na , ^{24}Mg , and ^{138}Ba , along these three core-to-edge transects. Sodium was the element with the highest concentration ($808\text{--}2790 \mu\text{g g}^{-1}$) and showed a temporally heterogeneous distribution with periodic oscillations mainly in the ventral-external region. Like Mg, the core was the area with the highest concentration. In relation to Ba distribution, within the first year of life, a plateau was observed in the three transects, with the distribution being greater in the third (transect 1: $< 11.7 \mu\text{g g}^{-1}$; transect 2: $< 11.0 \mu\text{g g}^{-1}$, transect 3: $< 23.8 \mu\text{g g}^{-1}$). Transect 3 also shows an area of high concentration of ^{138}Ba ($\sim 63.0 \mu\text{g g}^{-1}$) in the first annual rings and restricted to the ventral-internal area of the otolith.

Otolith chemistry as a bioindicator

The elements that make up metalloproteins in the hemolymph of the inner ear (such as ^{31}P) are physiologically regulated and reflect endogenous processes (e.g., reproduction, growth,

metamorphosis) rather than environmental (e.g., physicochemistry of water).³² Only one study reported a pilot analysis of the seasonal distribution of P in the otolith, suggesting that the incorporation of this element would be linked to the deposition of proteins in the bicarbonate matrix and would be favored by periods of rapid growth.³³ Our results support this hypothesis because maximum P values were observed during spring-summer, a favorable period for feeding and growth.

Other elements such as Mg, Mn, Sr, Ba are mainly associated with the salt-fraction of the endolymph and would be the best option to resolve unknowns associated with habitat use.³² Reports on Se in otoliths are scarce, with one study suggesting that this could be a natural marker in environments with heterogeneous distribution of this element.³⁴ In the present study, two marks with relatively high Se concentration were reported, so it is worth exploring the use of this element as an environmental proxy.

Conclusions

IR- and UV-fs-LA-ICPTOFMS has demonstrated a great potential for fast, high-spatially resolved, multi-elemental analysis of otoliths, allowing the visualization of the growing rings from the otolith core and the determination of heterogeneous distribution of major and minor elements. IR- and UV-fs-LA-ICPTOFMS can be up to 20 times faster than using typical operating conditions employed in multi-elemental LA-ICP-QMS analysis and even faster than SXFM. Both methods also provided lower detection limits than those reported for other micro-PIXE (e.g., Ba) and EPMA (e.g., Mn, Ba and Se) resulting in a good analytical option for elements that have a relatively low concentration. Despite the advantages of reduced analysis times, high lateral resolution and low LODs, access to fs-LA-ICPTOFMS system is relatively limited. Furthermore, it was found that the use of UV-fs-LA, at the selected operating conditions, provided slightly improved results in terms of limits of detection and recoveries calculated using non-matrix matched calibrations.

Elements such as Sr, Ba and P revealed a heterogeneous growth pattern with overlapping and even discontinuous growth rings. In addition to the heterogeneity in the microstructure, a heterogeneous distribution of elements within the same growth ring was observed. For example, Sr, Ba, and P showed higher values in the inner portion of the otolith, while discrete or discontinuous areas with relatively high values of Se, Mg, and Mn were observed. This fact highlights the usefulness of mapping to decide which area should be analyzed and avoid errors in interpretation.

The heterogeneous distribution of growth rings and elements (even within continuous growth rings) suggests differential crystallization mechanisms. Many species exhibit vaterite incrustations in addition to aragonite. These areas usually present rather disorganized growth that can be difficult (or impossible) to identify using optical equipment and divergent element concentrations with respect to aragonite.³⁵ In addition to the form of crystallization, the structural heterogeneity may be due to cessation of calcium carbonate deposition that affects the “rhythmic growth”,¹⁶ which could be due to different factors such as environmental stress (very cold winters, temperature changes, flow) or reproduction, among others.³⁶

Conflicts of interest

There are no conflicts to declare.

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