Element / Ca ratios in Nodosariida (Foraminifera) and their potential application for paleoenvironmental reconstructions

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Abstract. The chemical composition of foraminiferal shells is a well-known tool in paleoceanography to reconstruct past environments and climate. Their application is based on the relation between environmental variables and the concentration of elements incorporated or stable isotope fractionation during calcification. The vast majority of these so-called proxy relationships are based on the foraminiferal order of the Rotaliida, which, for example, encompasses all living planktonic species. However, there are more orders of foraminifera with calcifying members, some of which have fundamentally different biomineralization pathways, such as the Nodosariida, the Polymorphinida and the Vaginulinida. All these belong to the class of the Nodosariata and produce calcite shells, which may serve as carriers of paleoenvironmental and climate signals. The microstructures of these shells and overall morphology of these foraminifera strongly deviate from the Rotaliida, suggesting that their elemental and stable isotopic composition do not necessarily respond similarly to environmental parameters. A potential advantage of the Nodosariata is that they appear considerably earlier in the fossil record (Carboniferous) than the Rotaliida (Jurassic), thereby possibly extending the range of foraminifer-based paleoceanographic reconstructions considerably. To test the potential application of Nodosariata foraminifera as paleoproxies, we investigated incorporation of 5 elements in 11 species as a function of environmental parameters from a transect sampled in the Gulf of Mexico. Their element composition (B / Ca, Na / Ca, Mg / Ca, Sr / Ca and Ba / Ca) shows a distinct geochemical signature for these foraminifera, different to that of members of other foraminiferal orders. Results also show an increase in Mg / Ca values with increasing temperature, similar to that known for the Rotaliida, which suggest that Nodosariata shells might be useful for paleotemperature reconstructions. The difference in Mg / Ca–temperature calibration in Nodosariata compared to Rotaliida, with the large differences in their morphology, shell microstructures and overall geochemical composition, suggests that the Mg / Ca–temperature relationship is partly independent of the exact calcification mechanism. We compare Mg / Ca–temperature sensitivities across foraminiferal orders and describe a relationship between the average Mg / Ca and the sensitivity of the Mg / Ca–temperature calibration. For other elements, the variability across orders is smaller compared to that in Mg / Ca, which results in more similar El / Ca–environmental calibrations.

1 Introduction

Reconstructing past climates is an integral part of predicting the impact of the ongoing rise in atmospheric CO₂ levels on the Earth’s future climate. The expected temperature increase for a doubling in pCO₂ (the so-called climate sensitivity) has, for instance, been estimated by comparing past seawater temperatures and carbon dioxide levels (Rosenthal et al., 2017; D’Arrigo et al., 2006; Mann et al., 1998). Reconstructions of such parameters rely on accurate and precise tools that can be applied to past episodes in Earth’s history with conditions similar to those expected in the future. In this context, foraminifera are popular tools as they are proxy signal carriers for constraining past seawater tem-

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temperature and pH. Field and culturing studies have shown the
dependence of the chemical composition of their shells on
the seawater chemistry and physics in which they calcified.
For example, the amount of incorporated Mg (expressed as
the shell’s Mg/Ca) increases exponentially with temperature (Nürnberg et al., 1996) and can hence be used to recons-
truct past changes in bottom water temperature using ben-
thic foraminifera (Lear et al., 2002) and sea surface temper-
ature using planktonic foraminifera (Hastings et al., 1998; Lea
et al., 2000).

The incorporation of Mg into the calcite of most foraminifera is, however, much lower compared to calcite
precipitated inorganically from seawater (Morse et al., 2007).
This offset and the observed differences in Mg/Ca val-
ues between species (Wit et al., 2012) are hypothesized to
be caused by the strong biological control that foraminifera exert on the chemistry of the calcifying fluid from which
they form their shells (Erez, 2003; de Nooijer et al., 2014).
This biological control and the resulting interspecies vari-
ability in calcite chemistry have highlighted the need for
species-specific calibrations (Wit et al., 2017; Allen et al.,
2016). With this in mind, Mg/Ca and other proxies based on
foraminiferal shell composition, including Na/Ca for salini-
ity (Dämmer et al., 2020; Bertlch et al., 2018; Wit et al.,
2013) and δ11B for seawater pH (Foster and Rae, 2016; Rae
et al., 2011; Spivack et al., 1993), have been developed and
successfully applied. Another complicating factor when ap-
plying foraminiferal proxy signals is the dependency of ele-
ment incorporation and isotope fractionation on more than
one environmental parameter. For example, shell Mg/Ca
values are also affected by the marine inorganic carbon sys-
tem (Evans et al., 2016), salinity (Raitzsch et al., 2010; Dis-
sard et al., 2010) and the [Mg2+] of the seawater (Evans and
Müller, 2012). Ideally, proxy application would therefore in-
clude multiple elements to reconstruct a single parameter or,
alternatively, use multiple proxy relationships to simulat-
aneously reconstruct multiple environmental parameters.

Our knowledge of the controls on foraminiferal shell geo-
chemistry is almost exclusively based on results obtained
from Rotaliida. These foraminifera are characterized by mul-
tilocular shells composed of bilamellar calcite (Reiss, 1957,
1963) that can be optically radial or granular. The popularity of
using members of this order is partly due to the fact that
they encompass all extant planktonic foraminiferal species,
while the diversity and overall high abundance of benthic
Rotaliida add to their popularity for reconstructions of bot-
tom water conditions. Few studies have investigated element
incorporation in the Miliolida, which have a fundamentally
different calcification mechanism (ter Kuile et al., 1989; de
Nooijer et al., 2009; Debenay et al., 1998). The composition of
their calcite is markedly different from that of the Rotali-
ida, with for example markedly high Mg/Ca (Toyofuku et al.,
2000; van Dijk et al., 2017b) and more depleted δ25Mg
values (Dämmer et al., 2021).

Reconstructions based on Rotaliida could theoretically
span the last ∼190 Myr since they first occur in the fos-
sil record in the Pliensbachian (Haynes, 1981b). The order
of Nodosariata evolved calcification much earlier in the Per-
mian (Haynes, 1981a), and their application would therefore
roughly double the age for which paleoceanographic recon-
structions could be made using foraminiferal shell chemistry.
They separated from the Rotaliida and Miliolida likely be-
fore the Cambrian (Pawlowski et al., 2003) and are currently
found in many marine habitats and are easily recognizable by
their uniserial chamber arrangement (Haynes, 1981a). Their
walls are fibrous, composed of conical bundles of one to
tens of micrometers in length, that are only found in the
Nodosariata and therefore suggest that they have a unique biomineralization mechanism (Dubicka et al., 2018).
The isotopic composition of Nodosariata (Reolid, 2014; Dubicka
and Wierzbowski, 2021) and the isotopic difference between
Rotaliida and Nodosariida have been reported before (Du-
bicka and Wierzbowski, 2021); variability between species in
both δ13C and δ18O can be explained by differences in their
microhabitats. For instance, Lenticulina sp. is often
found deeper in the sediment, and such species are often
δ13C depleted compared to epifaunal foraminifera. Furth-
more, Lenticulina can live under dysoxic conditions, which
explains the overall lower δ18O in this genus.

So far, the Nodosariata elemental composition has not
been studied; therefore, we analyzed element incorporation
such as Na/Ca, Mg/Ca, Sr/Ca, B/Ca and Ba/Ca of dif-
f erent species collected along a depth transect in the Gulf
of Mexico. Accompanying environmental data (temperature,
salinity, etc.) allow us to detect any dependencies of the
incorporation of elements on these parameters and com-
pare them to existing calibrations for Rotaliida foraminiferal
species.

2 Material and methods

2.1 Sampling location

In February 2020, sediment samples were collected from
the continental margin in the northern Gulf of Mexico using
the research vessel Pelagia (expedition 64PE467). Samples
were collected along a transect close to the outflow of the
Atchafalaya River, at depths of 105, 272 and 619 m (Fig. 1).
From box cores, smaller subcores were collected on deck and
subsequently sliced (with a resolution of 0.5 cm for the upper
2 cm and in 1 cm slices down to a depth of 10 cm). The sedi-
ment was stored in ethanol, with rose bengal (rB) (2 g L−1)
added to stain the cytoplasm of living foraminifera.

From the overlying water of the box cores, vials were filled
for analysis of DIC (dissolved inorganic carbon) and TA (to-
tal alkalinity) after filtering over 0.4 µm filters. For both anal-
yses, 5 mL vials were filled with seawater and stored at 4 °C
after addition of 15 µL of HgCl2 to prevent biological al-
teration of the inorganic carbon system. The samples were analyzed after returning to the laboratory using a QuAA-
tro continuous flow analyzer. DIC samples were acidified, and the carbon dioxide that was formed was dialyzed over a
membrane that reduces the phenolphthalein indicator and was spectrophotometrically recorded at 550 nm (Stoll et al.,
2001). For TA, a slightly acid buffered solution of potassium hydrogen phthalate was added to the sample, after which
the intensity at 590 nm was recorded (Sarazin et al., 1999).
Values obtained for DIC and TA were consistent with the ones obtained from earlier expeditions (Sirois, 2017). Bot-
tom water temperature and salinity were taken from a device that measures conductivity, temperature and depth along the
seawater column (CTD) casts at the same station where the sediment samples were taken approximately 4 m above the
sea floor. Values for all inorganic carbon system parameters can be estimated using two measured parameters, since any
combination of two such parameters will allow calculating all others, including dissolved CO$_2$ (Zeebe et al., 1999), per-
formed with PyCO$_2$SYS (Lewis et al., 1998), using the recently published Python script (Humphreys et al., 2022) (Ta-
ble 1).

2.2 Sample preparation

Samples were washed using sieves with mesh sizes of 63 and 150 µm and dried in an oven at 60 °C. When selecting the specimens, rB-stained foraminifera were separated from non-stained specimens to allow detection of possible post-
mortem alteration of the primary geochemical signal. The foraminifera were cleaned after isolation from the sediment by
immersion in a solution of 1 % H$_2$O$_2$ and 0.1 M NH$_4$OH and three consecutive rinses with double-deionized water.
During the latter step, the Eppendorf tubes were placed in an ultrasonic bath to remove any particles adhering to the
shells. In this way, 188 individuals were prepared for single-
chamber geochemical analysis using laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS).

2.3 Analytical procedure

Specimens were ablated for 50 s in an NWR193UC TV2 dual-volume chamber using circular spots of 80 µm set at a
repetition rate of 6 Hz, using an energy density of 1.00 ± 0.05 J cm$^{-2}$. The 193 nm wavelength of the LA-ICP-
MS used is deep ultraviolet (193 nm), which is excellently suited for ablating carbonates (Reichart et al., 2003) (Ta-
ble 2). The aerosol produced during the ablation was trans-
ported to a quadrupole ICP-MS (Thermo Fisher Scientific
iCAP-Q) on a helium flow with a flow rate of 0.6 L min$^{-1}$,
with 0.4 L min$^{-1}$ argon make-up gas being added before en-
tering the ICP torch. The background signal was measured for 60 s prior to ablation and 10 s after stopping the ab-
lation. Calibration was performed against USGS MACS-3 (synthetic calcium carbonate) pressed powder standard with
43Ca as an internal standard. Scanned masses include 11B,
23Na, 25Mg, 27Al, 43Ca, 88Sr and 138Ba. Standard reference
material for quality control was NFHS-2-NP (Boer et al.,
2022). Data reduction was performed using an adapted ver-
dion of the data reduction software SILLS (Signal Integration
for Laboratory Laser Systems, Boer et al., 2022; Guillong et
al., 2008) in MATLAB. Repeatability based on related stan-
dard deviations of measurements of NFHS-2-NP in this study
($n = 8$) is 4 % for Na/Ca, 1 % for Mg/Ca, 2 % for Sr/Ca,
5 % for B/Ca, 6 % for Al/Ca and 2 % for Ba/Ca. For all spec-
imens, two to three spot analyses were performed on the
final chamber (Fig. 2). The average element / calcium ratio
was calculated from the entire profile of the foraminifera
chamber wall with a delay of 3 s after firing the laser. The
end point of the profile, where the laser penetrates the cham-
ber wall, was calculated in the adapted MATLAB application
based on a drop of 30 % of the 43Ca intensity. This end point
of the profile was visually checked using laser ablation screen
shots continuously made and stored every 2 s.

Since they were not correlated and the occasional high
Mg/Ca was not accompanied by high Al/Ca, the elevated
Mg/Ca could not be attributed to contamination (e.g., by
clay particles or a recrystallized phase at the surface of the
shells; Figs. S1, S2, S3, S4), and we did not remove any of
the original data points. Instead, we statistically tested for
outliers to identify El/Ca ratios that are outside the expected
distribution given the data. These outliers ($n = 25$) are high-
lighted in the figures; including or excluding them was found
to have an insignificant effect on the reported regressions.

2.4 Statistical analyses

For the three species present at all depths sampled, we per-
formed an ordinary least sum of squares regression analy-
sis to test dependency of the elements incorporated on en-
vironmental parameters. For Mg/Ca and temperature, an
exponential response model was assumed, and for Sr/Ca and
temperature a linear model was assumed while assum-
ing a linear response model for Na/Ca and salinity and for
Ba/Ca and salinity. For B/Ca and the inorganic carbon pa-
rameters, a linear response model was assumed. Prior to re-
gression analysis, outliers were identified based on st uden-
tized residuals using the package “statsmodels” for Python
and applying the method “sidak” from the Holm–Šídák
method with a one-step correction (Seabold and Perktold,
2010). Identified outliers are highlighted in the figures: their
presence or absence had only a marginal effect on the regres-
sion analysis.

When plotting the results, the analytical error was plotted for individual analyses (which mostly falls within the size
of the sample marker), the standard deviation for the sample
to identify the variability within the sample as well as the
standard error (SE), to show the confidence interval for the
estimate of the average. Standard deviation (SD) and SE are
related according to SE = SD / $\sqrt{\text{no. measurements}}$.  

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Figure 1. (a) Sampling location in the Gulf of Mexico (GEBCO, 2022). (b) Salinity and (c) temperature data from CTD at the bottom sample.

Table 1. Chemical and physical seawater parameters at the stations where foraminifera were collected. Temperature, salinity, DIC and TA were measured; the other parameters ($pCO_{2}$ and all parameters to the right of $pCO_{2}$) were calculated using the software PyCO$_2$SYS (Lewis et al., 1998), using the recently published Python script (Humphreys et al., 2022).

<table>
<thead>
<tr>
<th>Position</th>
<th>Station</th>
<th>Depth (m)</th>
<th>Temperature (°C)</th>
<th>Salinity (µmol kg$^{-1}$)</th>
<th>DIC (µmol kg$^{-1}$)</th>
<th>TA (µmol kg$^{-1}$)</th>
<th>$pCO_{2}$ (ppm)</th>
<th>pH</th>
<th>$[CO_{2}^{−}]$ (µmol kg$^{-1}$)</th>
<th>$[HCO_{3}^{−}]$ (µmol kg$^{-1}$)</th>
<th>$fCO_{2}$ (ppm)</th>
<th>$\Omega$Ca</th>
</tr>
</thead>
<tbody>
<tr>
<td>91.812 W/ 28.052 N</td>
<td>A100</td>
<td>104.7</td>
<td>18.87</td>
<td>36.37</td>
<td>2151.7</td>
<td>2334.7</td>
<td>621.60</td>
<td>7.88</td>
<td>135.84</td>
<td>1995.29</td>
<td>619.46</td>
<td>3.16</td>
</tr>
<tr>
<td>91.862 W/ 27.812 N</td>
<td>A300</td>
<td>271.64</td>
<td>12.66</td>
<td>35.61</td>
<td>2144.9</td>
<td>2277</td>
<td>623.25</td>
<td>7.86</td>
<td>102.47</td>
<td>2017.51</td>
<td>620.93</td>
<td>2.32</td>
</tr>
<tr>
<td>91.92 W/ 27.665 N</td>
<td>A600</td>
<td>618.8</td>
<td>6.96</td>
<td>34.92</td>
<td>2202.1</td>
<td>2300.7</td>
<td>616.54</td>
<td>7.84</td>
<td>82.51</td>
<td>2089.71</td>
<td>614.07</td>
<td>1.75</td>
</tr>
</tbody>
</table>

Table 2. Characteristics and method for the LA-ICP-MS analyses.

<table>
<thead>
<tr>
<th>Laser ablation system sample cell</th>
<th>NIOZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wavelength</td>
<td>193 nm</td>
</tr>
<tr>
<td>Pulse duration</td>
<td>4 ns</td>
</tr>
<tr>
<td>Laser fluence</td>
<td>1 J cm$^{-2}$</td>
</tr>
<tr>
<td>Laser spot size</td>
<td>80 µm</td>
</tr>
<tr>
<td>Laser repetition rate</td>
<td>6 Hz</td>
</tr>
<tr>
<td>Carrier gas flow rate (He)</td>
<td>0.6 L min$^{-1}$</td>
</tr>
</tbody>
</table>

Since all data are derived from three locations, in addition to the regression analysis, a two-tailed $t$ test was performed to test whether the variances of the El/Ca between locations significantly differed.

3 Results

3.1 Average El/Ca in Nodosariata species

Combining the data from all stations shows that the average El/Ca in the individual Nodosariata species varies between 6.65 and 13.2 mmol mol$^{-1}$ for Na/Ca, between 5.94 and 20.1 mmol mol$^{-1}$ for Mg/Ca, and between 1.09 and 1.81 mmol mol$^{-1}$ for Sr/Ca (Table 3) (for more information about variability for two species within the same genus, see Table S2). For all measured Mg/Ca of a single species, the SD is on average 3.01 mmol mol$^{-1}$, where it is 1.07 mmol mol$^{-1}$ for Na/Ca, and 0.14 mmol mol$^{-1}$ for Sr/Ca. This translates to a relative variability in El/Ca...
within a species of 19.6 % for Sr / Ca, 17 % for Na / Ca and 56 % for Mg / Ca.

For the Nodosariata Ba / Ca varies considerably, between 2.5 and 4.6 µmol mol$^{-1}$, with an average SD of 1.2 µmol mol$^{-1}$, which corresponds to a 71 % variability. The B / Ca data vary between 51 and 83 µmol mol$^{-1}$, with a modest variability in the Nodosariata, with an average SD of 9.8 (or 31.2 %) per species (Table 3).

### 3.2 Impact of temperature on Mg / Ca and salinity on Na / Ca

Mg / Ca correlates positively with temperature in Nodosaria flintii, Dentalina spp. and Lenticulina calcar over the 12$^\circ$C range studied here (for more information about the variability between living and non-living species, see Table S1). The lowest and highest values for Mg / Ca were found in Dentalina spp., ranging from 3.09 mmol mol$^{-1}$ at 6.96 $^\circ$C to 29.2 mmol mol$^{-1}$ at 18.87 $^\circ$C (Fig. 3b and Table 3). For Nodosaria flintii, Mg / Ca ranges from 3.93 to 14.9 mmol mol$^{-1}$ (Fig. 3c and Table 3), and for Lenticulina calcar Mg / Ca increases from 6.6 to 18 mmol mol$^{-1}$ (Fig. 3 and Table 3).

Despite the differences in absolute values, the sensitivity of changes in Mg / Ca as a function of temperature is similar for the three species, with Dentalina spp. having a slightly higher Mg / Ca–temperature sensitivity than the other two species (Table 3).

Results also show a significant positive increase in Na / Ca with salinity for two species, Dentalina spp. and Nodosaria flintii, despite the relatively small range in salinities between sites (1.45 units). For N. flintii, average Na / Ca varied between 6.7 and 10 mmol mol$^{-1}$, which is similar to the increase of 5.6 to 13 mmol mol$^{-1}$ by Nodosaria flintii (Fig. 4).

For Lenticulina sp., no regression was found between salinity and Na / Ca (panel c); “I” and “II” indicate significant differences in the averages between the three groups of data points (i.e., the average Na / Ca at lowest salinity is significantly different from the average Na / Ca in the other two sampled stations).
Table 3. Element incorporation average and SD for 11 species spanning 188 analyzed specimens of Nodosariata.

<table>
<thead>
<tr>
<th>Species</th>
<th>Na / Ca (mmol mol(^{-1}))</th>
<th>SD</th>
<th>Mg / Ca (mmol mol(^{-1}))</th>
<th>SD</th>
<th>Sr / Ca (mmol mol(^{-1}))</th>
<th>SD</th>
<th>Ba / Ca (µmol/mol)</th>
<th>SD</th>
<th>B / Ca (µmol/mol)</th>
<th>SD</th>
<th>Number of measurements/specimens</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amphycorina sp.</td>
<td>8.50 ± 0.74</td>
<td></td>
<td>5.94 ± 0.79</td>
<td></td>
<td>1.20 ± 0.07</td>
<td></td>
<td>3.18 ± 0.74</td>
<td></td>
<td>53.37 ± 5.67</td>
<td></td>
<td>52/19</td>
</tr>
<tr>
<td>Nodosaria flintii</td>
<td>8.22 ± 0.85</td>
<td></td>
<td>8.08 ± 2.91</td>
<td></td>
<td>1.30 ± 0.14</td>
<td></td>
<td>3.78 ± 1.85</td>
<td></td>
<td>53.09 ± 10.13</td>
<td></td>
<td>30/10</td>
</tr>
<tr>
<td>Dentalina spp.</td>
<td>9.28 ± 1.63</td>
<td></td>
<td>11.74 ± 6.49</td>
<td></td>
<td>1.33 ± 0.19</td>
<td></td>
<td>4.63 ± 2.88</td>
<td></td>
<td>66.73 ± 17.84</td>
<td></td>
<td>107/35</td>
</tr>
<tr>
<td>Fissurina radiata</td>
<td>6.65 ± 0.41</td>
<td></td>
<td>7.42 ± 2.92</td>
<td></td>
<td>1.09 ± 0.07</td>
<td></td>
<td>2.53 ± 0.25</td>
<td></td>
<td>53.98 ± 6.64</td>
<td></td>
<td>5/2</td>
</tr>
<tr>
<td>Grigelia semirugosa</td>
<td>10.85 ± 1.50</td>
<td></td>
<td>11.78 ± 4.04</td>
<td></td>
<td>1.57 ± 0.13</td>
<td></td>
<td>2.85 ± 0.79</td>
<td></td>
<td>62.21 ± 6.20</td>
<td></td>
<td>32/12</td>
</tr>
<tr>
<td>Lenticulina calcarea</td>
<td>10.71 ± 1.19</td>
<td></td>
<td>10.87 ± 2.43</td>
<td></td>
<td>1.58 ± 0.15</td>
<td></td>
<td>2.50 ± 1.10</td>
<td></td>
<td>82.72 ± 15.06</td>
<td></td>
<td>104/33</td>
</tr>
<tr>
<td>Lenticulina dentifera</td>
<td>10.03 ± 1.21</td>
<td></td>
<td>11.31 ± 2.82</td>
<td></td>
<td>1.54 ± 0.18</td>
<td></td>
<td>3.45 ± 1.40</td>
<td></td>
<td>67.29 ± 16.87</td>
<td></td>
<td>85/27</td>
</tr>
<tr>
<td>Vaginulinoidea baggi</td>
<td>9.50 ± 0.81</td>
<td></td>
<td>8.81 ± 1.88</td>
<td></td>
<td>1.54 ± 0.08</td>
<td></td>
<td>2.73 ± 0.64</td>
<td></td>
<td>51.12 ± 6.54</td>
<td></td>
<td>58/20</td>
</tr>
<tr>
<td>Procerolagenia gracilis</td>
<td>10.47 ± 1.35</td>
<td></td>
<td>8.64 ± 0.93</td>
<td></td>
<td>1.81 ± 0.25</td>
<td></td>
<td>2.85 ± 0.76</td>
<td></td>
<td>64.36 ± 3.84</td>
<td></td>
<td>6/2</td>
</tr>
<tr>
<td>Pseudoglandulina comatula</td>
<td>13.21 ± 0.93</td>
<td></td>
<td>20.06 ± 3.19</td>
<td></td>
<td>1.60 ± 0.07</td>
<td></td>
<td>4.28 ± 1.58</td>
<td></td>
<td>80.04 ± 9.38</td>
<td></td>
<td>38/13</td>
</tr>
<tr>
<td>Oolina spp.</td>
<td>10.01 ± 1.20</td>
<td></td>
<td>12.14 ± 4.75</td>
<td></td>
<td>1.39 ± 0.24</td>
<td></td>
<td>2.88 ± 0.66</td>
<td></td>
<td>58.12 ± 9.96</td>
<td></td>
<td>8/2</td>
</tr>
</tbody>
</table>

Figure 3. Correlations between Mg / Ca and temperature for three species of Nodosariata. Dots in blue show the outliers that were omitted before regression analysis based on studentized residuals. Red points are individuals living when sampled (i.e., stained with rose bengal), and green points are non-living individuals when sampled. Red line indicates the result of the exponential ordinary least squares (OLS) regression using both stained and non-stained specimens; cyan lines indicate the 95% confidence interval of the regression.

4 Discussion

4.1 Element incorporation in Nodosariata shells

The average values of Na / Ca, Sr / Ca and Ba / Ca of the Nodosariata are similar to those observed in planktonic (Barker et al., 2005) and many benthic Rotaliida (Lear et al., 2002; Elderfield et al., 2006), whereas Miliolida have considerably lower Na / Ca and higher Sr / Ca and Ba / Ca (van Dijk et al., 2017b). The coral’s aragonite Sr / Ca has been suggested as a proxy for temperature (Cohen et al., 2001; Reynaud et al., 2007), and this ratio has also been suggested to reflect temperature in the calcite of planktonic foraminifera (Cléroux et al., 2008). Here, we found no correlation between Sr / Ca and bottom water temperature (results not shown here). Nor did we find a correlation between Ba / Ca and salinity, which has been reported for benthic foraminifera from continental margins with noticeable freshwater input (Bahr et al., 2013). In addition, the Nodosariata’s Mg / Ca, is higher (5–30 mmol mol\(^{-1}\); Fig. 3) than those in plank-
Figure 4. Correlations between Na / Ca and salinity for three species of Nodosariata. Outliers are shown in blue that were omitted before regression analysis (panels a, b and c) based on studentized residuals. Red points are individuals living when sampled (stained with rose bengal), and data in green are non-stained individuals (panels a, b and c). Red lines in (a) and (b) indicate the result of the linear OLS regression using both the stained and non-stained specimens, and the two cyan lines indicate the 95% confidence interval of the regression (panels a and b).

This order- or class-specific signature of the shell’s composition support a fundamental difference in their calcification mechanisms (Dubicka et al., 2018; Dubicka and Gorzelak, 2017). Such a difference is also suggested with a fundamentally distinct morphology (i.e., chamber arrangement), as well as the micrometer-scale structures observed within the chamber walls. The chamber walls of the Nodosariata show a lamellar and fibrous texture, while Rotaliida show a granular texture (Dubicka et al., 2018) (Fig. S6). Such differences in the shell’s microstructure coincide with the contrasting Mg / Ca values observed here, similar to what was already reported for comparisons between other pairs of foraminiferal orders (van Dijk et al., 2016; Bentov and Erez, 2006). Deep evolutionary branching between Nodosariata and Rotaliida and the large difference in time of first fossil occurrence further support the hypothesis that they evolved their biomineralization mechanism independently. With different seawater chemical conditions (van Dijk et al., 2016; Tanner et al., 2020) at the time when Nodosariata and Rotaliida evolved, calcification biomineralization mechanisms may
well reflect contrasting selective pressures, which in turn is reflected by the shells’ Mg/Ca ratios.

Although they have a long geological and evolutionary history, El/Ca variability in the elemental-to-calcium ratio within the Nodosariata is remarkably small compared to that observed in, for example, the Rotaliida. The Mg/Ca values do not vary significantly between the different families: the Lagenidae, Nodosariidae, Ellipsolagenidae and the Vaginulidae (one-way ANOVA, p value > 0.05 and F = 1.082). Also, for the other elements no significant difference in the average elemental ratios is observed between species and, thus, between families. This does not exclude the existence of species or families within the Nodosariata that may have a different elemental signature than those reported here as we only investigated three species. Still, the analyzed species span three different orders within the Nodosariata. The relative uniformity in shell carbonate composition across the orders may indicate that the calcification mechanism invented by the Nodosariata is very well suited for a wide range for seawater chemical conditions. Alternatively, the relatively low species diversity of the present-day Nodosariata compared to that during the Jurassic (Haynes, 1981a) may reflect a selective loss of calcification mechanisms due to past changes in ocean chemistry and/or physics, possibly related to past climate variability. Such a hypothesis on the potential interplay between calcification and climate can be tested by comparing the El/Ca of extinct species from a suite of eras to that of species living today (Evans et al., 2013; Maeda et al., 2017; Toyofuku et al., 2011; Barrientos et al., 2018).

4.2 Effect of environment in the element incorporation

4.2.1 Na/Ca versus salinity

Na/Ca correlates with salinity in two of the Nodosariata species investigated: Dentalina spp. and Nodosaria flintii (Fig. 4). Sensitivities of Na/Ca to salinity relationships appear somewhat higher than those reported for Rotaliida species (Wit et al., 2013; Geerken et al., 2019; Allen et al., 2016; Mezger et al., 2016; Hauzer et al., 2021) (Table 4). Parallel to the increasing number of reports on the correlation between Na incorporation and salinity, there is discussion of what precisely controls foraminiferal Na/Ca, which could be [Ca$^{2+}$] (Hauzer et al., 2018) as well as the [Na$^+$]$_{sw}$ and/or salinity (Wit et al., 2013). In addition, it may be that Na incorporation is affected by precipitation rates as well, such as indicated by inorganic experiments showing that Na incorporation is affected by saturation state (Devriendt et al., 2021). Although poorly constrained in foraminifera (Geerken et al., 2022), environmental factors may affect the rate at which foraminifera precipitate their calcite, making the relationship between Na/Ca and an environmental parameter indirect. Still, the consistent increase in Na/Ca with salinity in many Rotaliida foraminifera and the correlations reported here for two Nodosariata species (Fig 4; Table 4) suggest a more direct coupling between seawater [Na$^+$] and [Ca$^{2+}$] and underscores the robustness of this proxy.

4.2.2 Mg/Ca versus temperature

The Mg/Ca–temperature relationships found for the Nodosariata species reported here (Fig. 3) are likely also affected by different bottom water [CO$_3^{2-}$] at the sampled stations (Sadakov et al., 2014). The effect of saturation state on Mg incorporation was found to be approximately 40 mmol mol$^{-1}$ for a difference between the lowest and the highest of ∼ 1000 µmol [CO$_3^{2-}$] kg$^{-1}$ seawater in culturing experiments (Dissard et al., 2010; Yu et al., 2019; van Dijk et al., 2016). This would amount to a change of approximately 2 mmol mol$^{-1}$ Mg/Ca over the total change at the three locations studied here, assuming that the sensitivity of Mg incorporation as a function of [CO$_3^{2-}$] in the Nodosariata is similar to that in Rotaliida. This would reduce the change in Mg/Ca as a function of temperature by less than 10%, and hence this would only have a very modest impact on the Mg/Ca–temperature sensitivities reported here (Fig. 3).

For each of the three species that were found at all three stations, Mg/Ca increased exponentially with temperature (Fig. 3) (Rosenthal et al., 1997). The application of these calibrations for Nodosariata for reconstructing past temperature can be challenging since using the chemical composition of fossil shells this far back in time requires careful assessment of the calcite diagenetic overprints (Stainbank et al., 2020), although the case of Lenticulina is reported to have a thick calcite and high fossilization potential (Dubicka and Wierzbowski, 2021). Furthermore, proxies for the chemical composition of the seawater such as fluid inclusion in evaporite minerals indicate that the concentration of major ions such as [Mg$^{2+}$]$_{sw}$ and [Ca$^{2+}$]$_{sw}$ changed significantly during the Phanerozoic (Fante and DePaolo, 2005), and these changes can affect the incorporation of Mg into the calcite. Nevertheless, few studies have developed new proxies to better understand these changes for both [Mg$^{2+}$]$_{sw}$ (Pogge Von Strandmann et al., 2011; van Dijk et al., 2016) and [Ca$^{2+}$]$_{sw}$ (Nambari et al., 2023).

The Mg/Ca–temperature sensitivities are slightly lower than those reported for most Rotaliida species. On average, Mg/Ca increases by 6% per degrees Celsius in the Nodosariata species analyzed here (Fig. 3), while in many planktonic species Mg/Ca increases by 10% per degrees Celsius (Barker et al., 2005). Low-Mg/Ca benthic rotaliids display an increase of approximately 8% in Mg/Ca per degrees Celsius (Lear et al., 2002; Raitzsch et al., 2010; Russell et al., 2004). High-Mg/Ca benthic Rotaliida species, on the other hand, increase by only 2% (Maeda et al., 2017), which is similar to the slopes of those reported for Milolida (de Nooijer et al., 2017; Toyofuku et al., 2000).

Combining sensitivities for the different groups of foraminifera and their average Mg/Ca and comparing them to those of inorganically precipitated calcites (Morse et al.,
Table 4. Comparison of sensitivities of Na/Ca versus salinity of benthic foraminifera from this study and Rotaliida with different Mg incorporation ratios.

<table>
<thead>
<tr>
<th>Order</th>
<th>Species</th>
<th>Sensitivity (mmol mol$^{-1}$)</th>
<th>Paper</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nodosariida</td>
<td>Nodosaria flintii</td>
<td>1.27</td>
<td>This study</td>
</tr>
<tr>
<td>Nodosariida</td>
<td>Dentalina sp.</td>
<td>1.19</td>
<td>This study</td>
</tr>
<tr>
<td>Rotaliida low Mg</td>
<td>Ammonia tepida</td>
<td>0.22</td>
<td>Wit et al. (2013)</td>
</tr>
<tr>
<td>Rotaliida low Mg</td>
<td>Ammonia tepida</td>
<td>0.064</td>
<td>Geerken et al. (2019)</td>
</tr>
<tr>
<td>Planktonic Rotaliida</td>
<td>G. ruber</td>
<td>0.074</td>
<td>Allen et al. (2016)</td>
</tr>
<tr>
<td>Rotaliida mid Mg</td>
<td>Amphistegina lessonii</td>
<td>0.077</td>
<td>Geerken et al. (2019)</td>
</tr>
<tr>
<td>Planktonic Rotaliida</td>
<td>G. ruber</td>
<td>0.66</td>
<td>Mezguer et al. (2016)</td>
</tr>
<tr>
<td>Planktonic Rotaliida</td>
<td>G. sacculifer</td>
<td>0.6</td>
<td>Mezguer et al. (2016)</td>
</tr>
<tr>
<td>Rotaliida high Mg</td>
<td>Operculina ammonoides</td>
<td>0.33</td>
<td>Hauzer et al. (2021)</td>
</tr>
</tbody>
</table>

Figure 5. Comparison between the sensitivity and the average Mg/Ca for different groups of foraminifera. Results for the Nodosariata are from this study; all other Mg/Ca–temperature calibrations are from previous studies (Toyofuku et al., 2011; Douglas and Staines-Urias, 2007; Barrientos et al., 2018; Rosenthal et al., 1997; Quillmann et al., 2012; Raitzsch et al., 2008; Lea et al., 1999; Anand et al., 2003; Barker et al., 2005; Rosenthal et al., 2011; Kristjánssóttir et al., 2007; Lear et al., 2002; Evans et al., 2013; Maeda et al., 2017; Wit et al., 2012; Morse et al., 2007; Toyofuku et al., 2000; de Nooijer et al., 2017; Knorr et al., 2015). Species towards the lower right corner are increasingly affected by biomineralization. These calibrations were used to calculate and plot the Mg/Ca at 20°C.

2007; Wit et al., 2012) suggest a negative relation between Mg incorporation and sensitivity to temperature (Fig. 5). The relative increase in Mg/Ca with temperature is smaller for species incorporating a relatively large amount of Mg in their calcite and vice versa. Highest Mg/Ca ratios are found in inorganically precipitated calcites (Morse et al., 2007) in which the increase of Mg/Ca is approximately 2% for a 1°C temperature increase. For species incorporating equally as much Mg (e.g., Operculina ammonoides; Evans et al., 2013), the slope of the Mg/Ca–temperature calibration is similar, while for species like Ammonia tepida (incorporating 50–100 times less Mg in their shell), the increase is approximately 7% per degrees Celsius temperature increase (Fig. 5).

This suggests that the observed high sensitivity of Mg/Ca to temperature in the low Mg/Ca species actually consists of two factors: an inorganic temperature-dependent fractionation and a biomineralization-related partitioning, which is also temperature dependent. The large difference in Mg/Ca between foraminifera (Wit et al., 2012) has been suggested to reflect the efficiency to lower the Mg/Ca in the calcifying fluid, either achieved by active Mg$^{2+}$ removal (Elderfield et al., 1996; Spero et al., 2015) or by selective inward Ca$^{2+}$ transport (Toyofuku et al., 2017). Foraminiferal
species with calcite Mg/Ca ratios close to those found in inorganic precipitation experiments may well lack such a mechanism, and the increase in Mg/Ca with temperature hence matches that found in inorganic precipitation experiments. The species that are capable of lowering the Mg/Ca in the fluid from which they calcify incorporate consistently more Mg at increased temperatures (Fig. 5). This suggests that foraminiferal Mg/Ca–temperature relationships are determined by two components (Dämmer et al., 2021). The first component is the biological control on Mg partitioning, and the second component is the thermodynamic effect of temperature on Mg/Ca. In foraminiferal species where the first component is absent (i.e., when they precipitate from a seawater like fluid), the second component determines the Mg/Ca–temperature sensitivity. For species that lower the Mg$^{2+}$ in the fluid from which they precipitate their calcite, the biological component dominates the Mg/Ca–temperature calibration. The relatively large variability in the low-Mg/Ca species may be explained by small environmental factors (e.g., salinity or water depth) or by processes that are part of the calcification mechanism (e.g., Rayleigh fractionation, organic templates) that may vary slightly between species. The Nodosariidae studied here have similar Mg/Ca but differ in their sensitivities (Fig. 5), which may well reflect the environmental and/or calcification-related differences between species. (Elderfield et al., 1996; Branson et al., 2018).

5 Conclusions

The chemical composition of the shells of various Nodosariida species collected in the Gulf of Mexico was found to be clearly different from those of other foraminiferal orders. Their Mg/Ca was between 6 and 20 mmol mol$^{-1}$, and their Na/Ca was relatively high compared to ratios for most planktonic Rotaliida species. Sr/Ca and B/Ca were comparable to those found in other foraminiferal species. In two of the species studied, the Na/Ca increased linearly with salinity. Between families of the Nodosariidae analyzed, the El/Ca was relatively similar. The Nodosariida’s Mg/Ca is correlated to temperature and could thus serve as a seawater temperature proxy. Compared to Rotaliida and Miliolida orders, our analysis shows a relation between the Mg/Ca of the species and its sensitivity to changes in temperature. When more Mg is incorporated, it is less sensitive to changes in temperature and vice versa. This suggests the interaction between two components that together determine the Mg/Ca: the capacity of a species to control the Mg/Ca of the calcite and the thermodynamic effect of temperature on Mg incorporation.

Data availability. All data used in this paper are available at https://doi.org/10.25850/nioz7/b.b.lf (Pacho, 2023).
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