

1 Monthly element/Ca trends and inter chamber variability in two planktic Foraminifera
2 species: *Globigerinoides ruber albus* and *Turborotalita clarkei* from a hypersaline
3 oligotrophic sea

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17
18
19 Abstract
20

21 Environmental and biological factors influence the trace element composition (element/Ca) of
22 planktic foraminifer shells. Consequently, the element/Ca measured in these shells (tests) are
23 utilized as proxies to reconstruct past oceanic and climatic conditions. As single shell analyses
24 are increasingly used in paleoceanographic research it is important to understand how proxy
25 systematics change between species, individuals of the same species in a given population, and
26 among chambers of a single individual during its life cycle. Here we present a time series of
27 the chemical composition of planktic foraminifers retrieved using sediment traps between June
28 2014 and June 2015 at the northern part of the Gulf of Aqaba (aka Gulf of Eilat). Laser ablation
29 ICP-MS element/Ca measurements were performed on single shells and chambers of
30 *Globigerinoides ruber albus* and *Turborotalita clarkei*, collected monthly from five water
31 depths (120 m, 220 m, 350 m, 450 m, and 570 m). Sediment trap samples were paired with
32 corresponding data on water column hydrography and chemistry. Pooled means of measured
33 element/Ca display species-specific and element-specific behavior, with generally higher

34 values for *T. clarkei* phenotypes ('big' and 'encrusted') in comparison to *G. ruber albus*. Some
35 element/Ca values measured in water column specimens, such as Al/Ca, vary significantly
36 from core-top specimens. A unique finding is a prominent increase in element/Ca around
37 March-April 2015, during maximum water column mixing, mostly apparent in *T. clarkei* and
38 to a lesser extent in *G. ruber albus*. This spring element/Ca increase is observed in most
39 measured elements and is further associated with an increase in inter-chamber variability
40 (ICV). Inter-chamber element/Ca patterns show element enrichment/depletion in the most
41 recently precipitated (final, F0) chamber in comparison to the older chambers (penultimate (F-
42 1), antepenultimate (F-2), etc.). Element/Ca in F0 may also be less sensitive to surrounding
43 environmental conditions. For example, the Mg/Ca of the F-1 and F-2 chambers of *G. ruber*
44 *albus* display a positive relationship with mixed layer temperatures while F0 does not. To
45 overcome this effect, we suggest using pooled means from non-F0 fractions as environmental
46 records and paleo proxies.

47 These results highlight the complexity of proxy systematics that rises from the variability in
48 element/Ca measured among different species and between chambers, caused by ecological
49 conditions and other processes in the water column including physical, chemical, and
50 biological effects.

51

52 1. Introduction

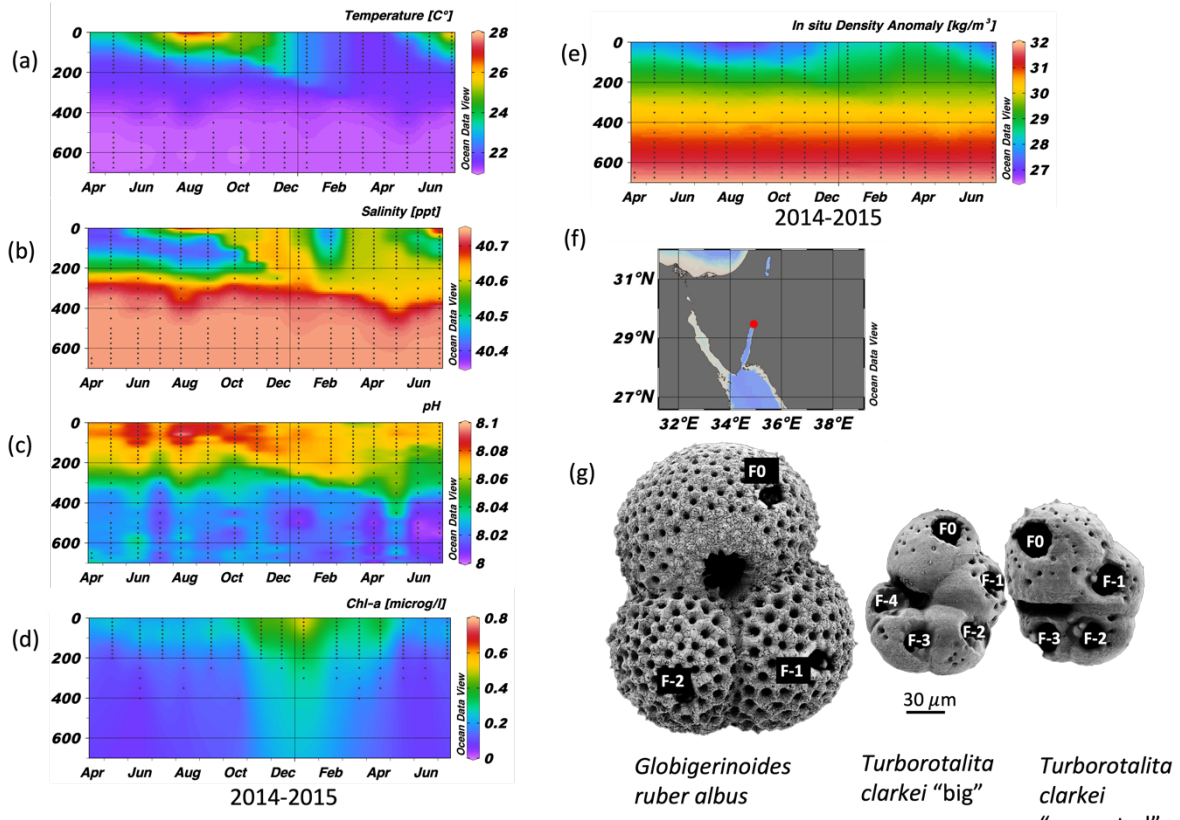
53 1.1 Planktic foraminifera as traces of the past environment

54 Planktic Foraminifera (PF) shells are useful archives for studying the history of Earth's
55 climate and oceans, as their calcareous shells reflect the environmental conditions during their
56 formation (Berggren et al., 1995; Rosenthal, 2007; Schiebel & Hemleben, 2017; Kucera, 2007;
57 Katz et al., 2010; Gupta, 1999; Davis et al., 2020,). Various element/Ca measured in PF tests
58 have been linked to ambient seawater temperature (e.g., Mg/Ca; Nurenberg et al., 1996;
59 Rosenthal et al., 2004), salinity (e.g., Na/Ca; Mezger et al., 2016; Gray et al., 2023), the
60 carbonate system (e.g., B/Ca; Babila et al., 2014; Henehan et al., 2015; Haynes et al., 2019),
61 productivity (e.g., Ba/Ca; Fritz-Enders et al., 2022), and chemical weathering (e.g., Ti/Ca;
62 Amaglio et al., 2025). In the past, the use of these proxies relied on bulk analysis of the entire
63 shell or multiple shells. However, although first attempts in single chamber LA-ICP-MS started
64 back in 2003 (Eggins et al., 2003, Reichart et al., 2003), only in recent years there has been an
65 increase in the use of high-resolution analytical techniques, such as Laser Ablation (LA) ICP-

66 MS and electron microprobe analyses in paleoceanographic studies (Davis et al., 2020). The
67 element/Ca measurements of single specimens (Individual Foraminifer Analysis, IFA)
68 revealed high variability between individuals of the same population as well as significant
69 intra-shell variability (i.e., inter chamber variability, ICV) (Sadekov et al., 2008; Fehrenbacher
70 et al., 2020; Hupp & Fehrenbacher, 2024; Fischer et al., 2024; Davis et al., 2020, and references
71 therein). Despite the analytical advancements, the variations in the geochemical signatures of
72 PF shells are poorly understood and while they are likely related to the life cycles and
73 reproductive modes of many species there are still knowledge gaps in our understanding of
74 proxy systematics in single shell and single chamber of PF species. There is also a lack of
75 detailed description and understanding of element/Ca systematics in description of small-sized
76 species such as *T. clarkei*, which have been largely overlooked in previous studies despite their
77 significant contribution to the settling PF tests (export flux), as observed in the northern Red
78 Sea (Chernihovsky et al., 2018). Furthermore, specific marine regions, such as in oligotrophic,
79 subtropical basins, particularly in deep-water environments, are not well-established in terms
80 of their spatial and temporal dynamics (Schiebel & Hemleben, 2017).

81 1.2 Planktic Foraminifer population dynamics in The Gulf of Aqaba

82 The Gulf of Aqaba (GOA) is considered an open ocean proxy environment (Chase et al.,
83 2011). It is an oligotrophic basin where the main lithogenic flux is derived from dust. During
84 summer (April-September), a ~200 m deep thermocline separates nutrient-depleted surface
85 waters (~25°C) from the nutrient-rich deep layer (~21°C). In winter-spring (October-April),
86 the thermocline gradually erodes due to surface cooling (Figs. 1a and 1e; Meeder et al., 2012),
87 which can lead to the development of a deep mixed layer. Although the depth of the mixed
88 layer varies annually with climatic conditions, the long-term mean mixing depth is
89 approximately 300-400 m, and deep mixing can extend to the sea floor while it typically
90 reaches maximum depth by late March. The regional terrestrial climate is hyper-arid (mean
91 annual rainfall <30 mm) and the main sources for terrigenous material to the GOA are dust
92 storms originating from the Sahara and Arabian Deserts, as well as rare localized floods (Katz
93 et al., 2015; Chase et al., 2011; Ganor et al., 2001; Torfstein et al., 2017).



94

95 Figure 1: Time series of temperature (a), salinity (b), pH (c), Chlorophyll-a concentration (d),
96 and in situ density anomaly (e), measured in the Gulf of Aqaba between April 2014 and June
97 2015 by the National Monitoring Program (NMP, Shaked & Genin, 2016). Y-axis is depth (m);
98 A map of the Gulf of Aqaba (f); and (g) scanning electron micrographs of the three morpho-
99 species (exhibiting ablation holes in each chamber (labelled), from Levy et al., 2023).

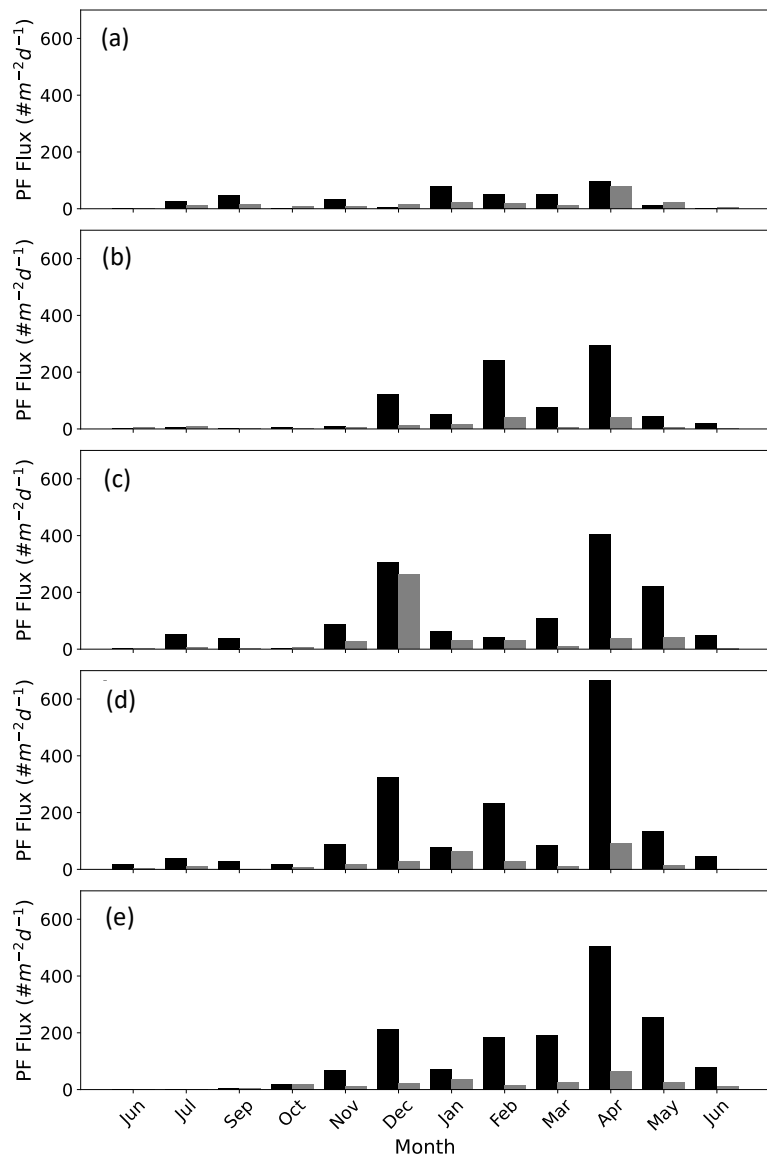
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101 Planktic foraminifera fluxes in the GOA demonstrate strong seasonality, with low fluxes
102 during the summer months, gradually increasing during the autumn-winter, coeval with
103 decreasing sea-surface temperatures and deepening of the mixed layer in the GOA that drives
104 advection of nutrient-replete subsurface waters into the mixed layer (Fig. 2). This in turn
105 triggers an increase in primary productivity, expressed by enhanced chlorophyll-a
106 concentrations and higher PF fluxes (Chernihovsky et al., 2018, 2020).

107 Spinoose species constitute the majority of the PF assemblage. The smaller size fraction,
108 63-125 μm , is 86% from the total flux and is dominated by *T. clarkei*. The 125-500 μm size-
109 fraction (~13 %) is dominated by the species *G. ruber albus*, while less than 1% of the shells
110 are in the range of 500-1000 μm , dominated by *O. universa* (Chernihovsky et al., 2018).

111 *Globigerinoides ruber albus* and *T. clarkei* inhabit different dwelling-depths and have
112 different life strategies. *Globigerinoides ruber albus* is a surface dweller and is photo-symbiont

113 bearing, while *T. clarkei* tends to dwell below the mixed layer depth and is barren of photo-
 114 symbionts (Rebotim et al., 2017; Schiebel & Hemleben, 2017; Levy et al., 2023). Furthermore,
 115 it has been suggested that *G. ruber albus* and *T. clarkei* do not share the same dietary
 116 preferences: *G. ruber albus* being more carnivorous than the detritivorous *T. clarkei* which may
 117 forage at the exported matter below the pycnocline (Schiebel & Hemleben, 2017). In the GOA,
 118 *T. clarkei* has two phenotypes: *T. clarkei* ‘big’, with all of its test chambers are fully
 119 recognizable and their surface is relatively smooth and *T. clarkei* ‘encrusted’ with a less smooth
 120 shell surface and is smaller than the ‘big’ type (Levy et al., 2023).



121
 122 Figure 2: PF fluxes in the Gulf of Aqaba between June 2014 and June 2015 presented according
 123 to the size fractions >63-125 μm (black bars) and >125-500 μm (grey bars) at the different
 124 sediment trap depths a) 120 m, b) 220 m, c) 340 m, d) 450 m, and e) 570 m. Data from
 125 Chernihovsky et al. (2018).

126 In this study, we investigate the range of element values, year-round trends and inter-
127 chamber element/Ca variability in *G. ruber albus* and *T. clarkei* tests collected in sediment
128 traps at various water column depths from the GOA. We assess whether the chambers record
129 temporal-seasonal patterns, and the implications for using single chamber data for geochemical
130 proxies (Mg/Ca, B/Ca, Na/Ca). Examining inter-chamber variability sheds light on how trace
131 elements are incorporated during calcification, distinguishing physiological controls (e.g.,
132 ontogenetic changes across successive chambers) from environmental influences such as
133 temperature, salinity and nutrient availability. Using *G. ruber* and *T. clarkei* from the GOA
134 provides a contrasting framework which together, these species enable us to refine element/Ca
135 proxy calibrations across vertical gradients in the water column in a warm and hyper-saline
136 oligotrophic environment.

137

138 2. Methodology

139 2.1. Sampling and oceanographic data

140 A bottom-tethered mooring has been deployed continuously since January 2014 near
141 Station A, northern GOA ($29^{\circ}28'00.0''N$, $34^{\circ}55'45.0''E$ ~605 m water depth) (Torfstein et al.,
142 2020). Five KC Denmark cylinder sediment traps were mounted vertically and located at
143 depths of 120 m, 220 m, 350 m, 450 m, and 570 m below the surface. The trap samples were
144 collected at a monthly resolution. Furthermore, PF samples from the sediment interface were
145 collected using a box core ('core top'). Further detailed description of the mooring, sampling,
146 sample processing, and trapping efficiencies can be found in Chernihovsky et al. (2018) and
147 Torfstein et al. (2020). Here, we report the findings derived from the PF tests collected between
148 June 2014 and June 2015. Water column physical and chemical parameters are routinely
149 collected at Station A by the Israel National Monitoring Program (NMP, Shaked & Genin,
150 2016). This includes sea surface and water column temperature ($^{\circ}C$), salinity, oxygen
151 concentration ($\mu\text{mol/l}$), alkalinity (meq/kg), *pH*, and chlorophyll-a concentration ($\mu\text{g/l}$). Mixed
152 layer depth (MLD) was defined as the shallowest depth where the TEOS-10 potential density
153 anomaly (σ_0), computed with gsw from Practical Salinity and in-situ temperature with pressure
154 from depth at Station A, exceeded a near-surface reference by $\Delta\sigma_0 = 0.03 \text{ kg m}^{-3}$; the reference
155 density was the median σ_0 within 0–10 m (or the shallowest 10 m available when the surface
156 was not sampled). Temperature, salinity, and *pH* within the mixed layer were then calculated
157 as thickness-weighted means by linearly interpolating each profile onto a 0.5 m grid and
158 averaging from the surface to the MLD.

159 2.2. Species classification and preparation for LA-ICP-MS

160

161 We examined the shell chemical properties of two flux dominating PF species *T. clarkei*
162 and *G. ruber albus* (i.e., sensu stricto, white). For *T. clarkei* we examined two morphotypes:
163 ‘big’ and ‘encrusted’. Identification and nomenclature of the PF taxa followed Schiebel &
164 Hemleben (2017), Morard et al. (2019), and Brummer & Kucera (2022). Three individuals
165 were picked from each sediment trap depth during each month between June 2014 and June
166 2015. Preliminary preparation and cleaning steps are detailed by Chernihovsky et al. (2018).
167 Reductive and oxidative cleaning had been avoided to retain original signals related to the
168 different encrustation processes and preserve all calcite layers added to the shell during
169 ontogeny (Schiebel & Hemleben, 2017; Jochum et al., 2019). Specifically, the shell of *T.*
170 *clarkei* is prone to loss of material during reductive and oxidative treatment as it has very thin
171 chamber walls (ranging between 1.9 and 3.6 μm ; Levy et al., 2023). Single chamber
172 measurements were performed to assess inter chamber variability (ICV), on individual shells
173 (individual foraminifer analysis; IFA) using Laser Ablation Inductively Coupled Plasma Mass
174 Spectrometry (LA-ICP-MS). We measured 156 specimens in total and 615 chambers: 57
175 individuals (168 chambers) of *G. ruber albus*, 52 individuals (242 chambers) of *T. clarkei* ‘big’
176 and 48 individuals (204 chambers) of *T. clarkei* ‘encrusted’. Samples were glued to glass slides
177 using a methyl-hydroxy-propyl-cellulose (MHPC 1:100), positioned with the umbilical side
178 up.

179

180 2.3. LA-ICP-MS and data processing

181 Analyses of the calcium-normalized elements for B, Na, Mg, Al, Ti, Mn, Fe, Co, Sr, Ba,
182 Nd, Pb, Th, and U were conducted using a 200 nm wavelength NWR femtosecond (fs) LASER
183 system from ESI, combined with a sector-field Thermo Element-2 ICP mass spectrometer
184 (Jochum et al., 2014). Measurements were performed using a 15 Hz pulse repetition rate (PRR),
185 at low fluence (0.1–0.6 J/cm^2), and 18 seconds dwelling time. A 30 μm diameter spot size was
186 selected, as it is the maximum diameter for analysis fitting in a single chamber of the small *T.*
187 *clarkei*. The microanalytical synthetic reference material MACS-3 for carbonate, NIST-612,
188 and NIST-610 were used for calibration. NIST-612 was used for the tuning of the ICP-MS
189 (Jochum et al., 2019). The average element-to-calcium ratio from the spot derived LA-ICP-MS
190 count data was calculated from count data immediately after the start of the ablation peak apex
191 until the point identified as the termination of calcite based on the Mg/Ca profile. This time

192 interval represents the stable internal material of the shell; excluding the noisy beginnings and
193 ends of the ablation event. For *G. ruber* the mean ablation time length used for calculation was
194 4.9 ± 2.3 secs, while for the smaller and thinner *T. clarkei* it was 2.6 ± 1.5 secs and 2.4 ± 1.4 secs,
195 for ‘big’ and ‘encrusted’ types, respectively.

196 The measurement precision (1 relative standard deviation in percent; 1 RSD) yield
197 uncertainties for references materials between ~ 5 -17 % for the calcium-normalized elements
198 (Supplementary table S1). Single spot measurements were made on each chamber of the
199 individual shells. Chambers are labelled F0 (final chamber), F-1 (final minus one), F-2, and so
200 on, for the penultimate, antepenultimate, and further chambers, respectively. We calculated
201 averages and standard deviations of element/Ca of single individuals (calculated from all single
202 chamber element/Ca in one shell) and relative standard errors of element/Ca of pooled
203 measurements for a specific morphotype.

204 2.4 Statistical Analysis

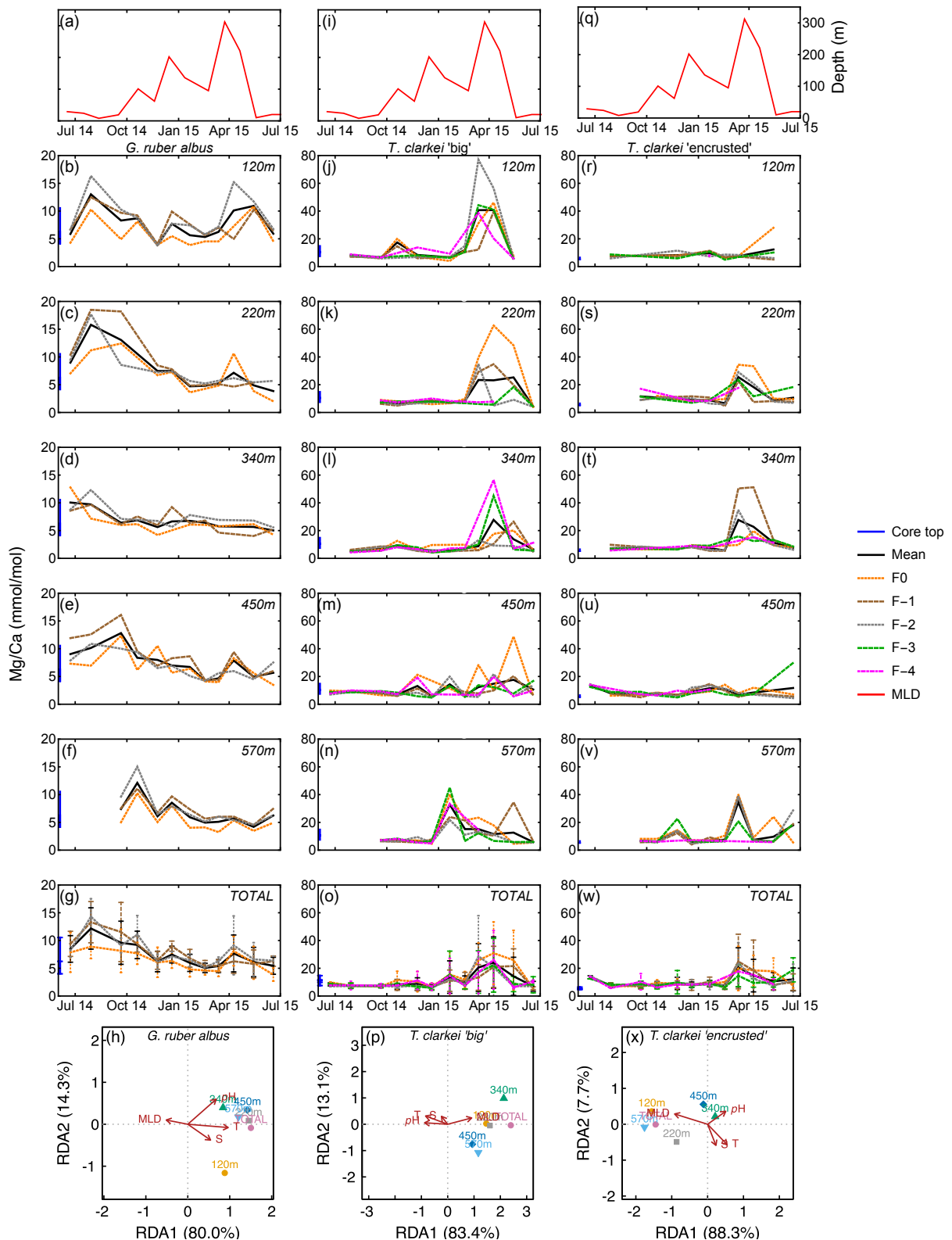
205 We used redundancy analysis (RDA) to quantify how variation in shell element/Ca
206 responses relates to environmental conditions (MLD, T, S, and pH). For each species, we
207 assembled two time-aligned matrices by date: (i) a multivariate response matrix containing the
208 element/Ca values at each sampled depth (120, 220, 340, 450, 570 m) plus the overall total,
209 and (ii) an environmental matrix containing environmental predictors. RDA was fit using the
210 vegan package in R (function rda), which is equivalent to multivariate multiple regression
211 followed by PCA of the fitted values. Biplots were produced with arrows show the direction
212 of increasing predictor values and their relative importance, and response points
213 (depths/TOTAL) project positively or negatively onto each arrow according to their alignment,
214 with the proportions of constrained variance carried by the first RDA axes (labels shown on
215 the biplot axes). As a measure of specimen inter-chamber variability (ICV), the standard
216 deviation (SD) was calculated for each individual. We calculated Spearman rank correlation
217 matrices in R for each species across element/Ca and SD, together with environmental
218 parameters, with the spearman correlation coefficients visualised as heatmaps (associated p-
219 values reported in supplementary table’s S2 and S3, respectively).

220

221 3. Results:

222 3.1. Shell-bound element/Ca time series trends in *G. ruber albus* and *T. clarkei* shells
223 Single chamber Mg/Ca over water column depths in *G. ruber albus* range between 2.01
224 mmol/mol (340 m; June 2015) and 18.49 mmol/mol (340 m; July-August 2014), with
225 lower/higher values during winter/summer months, respectively (Figs. 3b-3f). A unique
226 observation is an increase in Mg/Ca seen during spring (March-April), i.e., months with
227 maximum surface water column mixing, at some water depths (220 m, 340 m, 450 m; Figs. 3c-
228 3e). Accompanied with the Mg/Ca increase is a clear increase in ICV as evident by the
229 divergence of chamber values. The SD of *G. ruber albus* ranges between 0.14 in February 2015
230 at 450 m water depth and 7.27 during April 2015 at 120 m water depth. Generally, it appears
231 that Mg/Ca is lower in F0 chambers (orange dotted line) compared to preceding chambers,
232 especially during months with very shallow MLD (3a-3g). Mg/Ca in *T. clarkei* ‘big’ range
233 between 4.00 mmol/mol (340 m; June 2015) and 77.02 mmol/mol (220 m; March 2015) and
234 between 4.06 mmol/mol (570 m; December 2014) and 51.22 mmol/mol (120 m; April 2015)
235 in *T. clarkei* ‘encrusted’, respectively. For both *T. clarkei* ‘big’ and *T. clarkei* ‘encrusted’ there
236 are high excursions in all chambers during months of water column mixing while MLD is
237 deepest (circa April; 3j-3w). The SD ranges between 0.43 and 25.38 (120 m; September 2014
238 and, 220 m; April 2015 respectively) for *T. clarkei* ‘big’ and for *T. clarkei* ‘encrusted’ is up to
239 18.52 (340 m; March 2015).

240



241

242 Figure 3: Time series of Mg/Ca values measured from the shells of *G. ruber albus* (b-f), *T.*
 243 *clarkei 'big'* (j-n) and *T. clarkei 'encrusted'* (r-v), derived from sediment traps located at
 244 different water depths (120-570 m). Mg/Ca of core top are marked by a blue bar along the left
 245 y-axes. Top panels a, i, and q show the depth (meters below surface layer) of the mixed layer

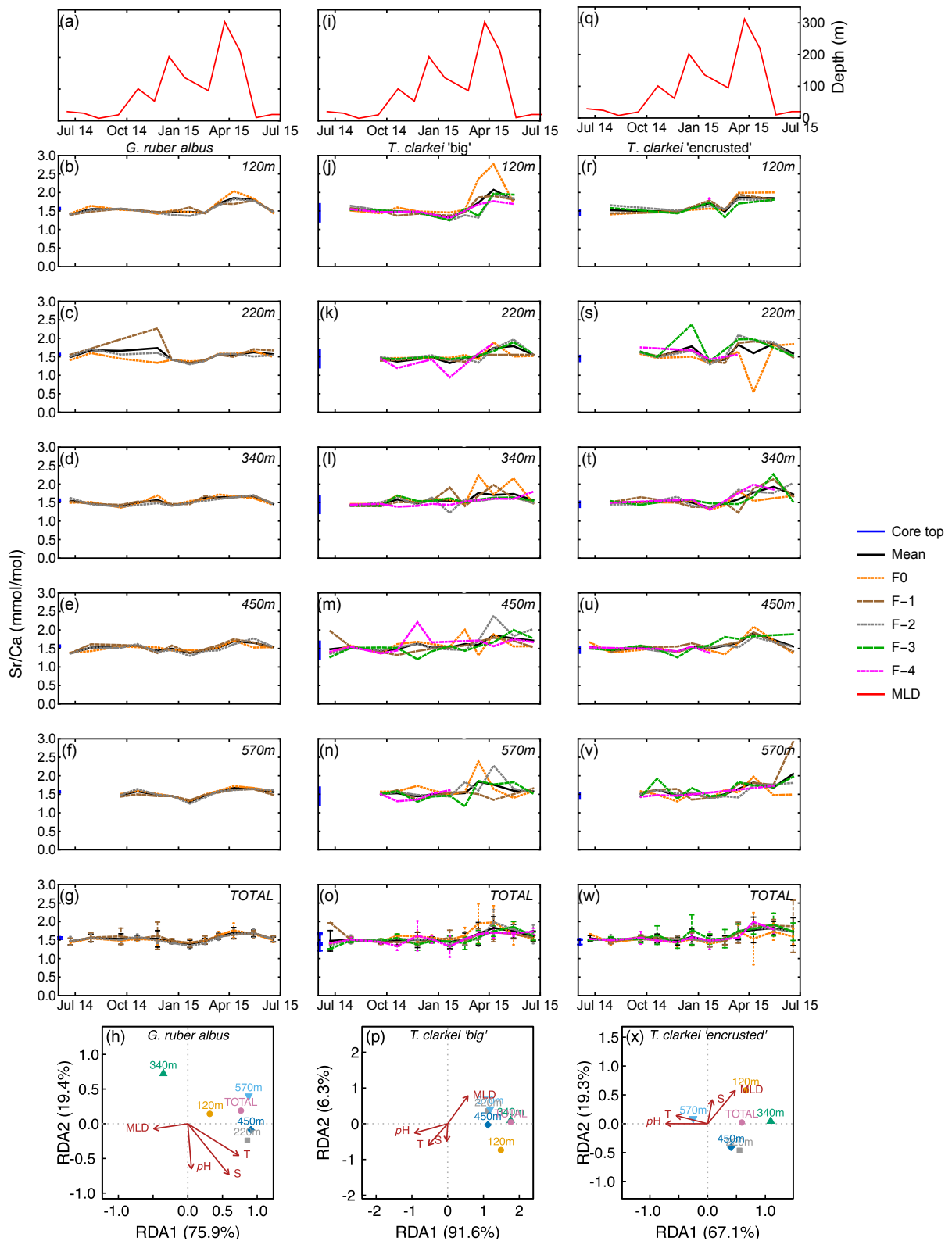
246 depth (MLD). Panels g, o, and w represent the average values of all depths for each chamber
247 for each species. Panels h, p and x show the redundancy analysis (RDA) per species for each
248 water depth with MLD depth, temperature, salinity and pH as explanatory variables.

249

250 The *G. ruber albus* RDA analysis of Mg/Ca show that most water depth samples, except for
251 the 120 m are clustered together and align with temperature and MLD, positive and negative
252 respectively. *Turborotalita clarkei* 'big' exhibits a contrasting image compared to *G. ruber*
253 *albus* and positively correlates with MLD while negatively correlating with temperature,
254 salinity, pH. All three morpho-species show a distinct dominance of RDA1 explaining >80%
255 of the variance (Figs. 3h, 3p and 3x).

256

257 Sr/Ca in *G. ruber albus* range between 1.25 mmol/mol (570 m; January 2015) and 2.27
258 mmol/mol (340 m; November 2014) (Figs. 4b-4f). The SD in *G. ruber albus* ranges reaches up
259 to 0.48 (220 m; November 2014). Single chamber Sr/Ca in *T. clarkei* 'big' range between 0.94
260 mmol/mol (340 m; January 2015) and 2.76 mmol/mol (220 m; April 2015) and for *T. clarkei*
261 'encrusted' between 0.54 mmol/mol (340 m; April 2015) and 2.92 mmol/mol (570 m; June
262 2015), respectively (Figs. 4j-4n, and 4r-4v). *Turborotalita clarkei* 'big' and *T. clarkei*
263 'encrusted' display more ICV than *G. ruber albus*, with peaking Sr/Ca in numerous chambers
264 around April 2015 (Figs. 4j-4w). During the spring months of 2015, Sr/Ca values range
265 between 1.45-2.04 mmol/mol in *G. ruber albus*, 1.32-2.76 mmol/mol in *T. clarkei* 'big' and
266 0.54-2.27 mmol/mol in *T. clarkei* 'encrusted', respectively (Fig. 4; Fig. S1). Additionally, the
267 SD of *T. clarkei* 'big' is the highest at 0.43 in March 2015 at 120 m water depth, and lowest at
268 0.02 in June-July 2014, at 340 m water depth. For *T. clarkei* 'encrusted' SD reaches 0.7 (220
269 m; April 2015). While the RDA analysis of *T. clarkei* 'big' show a high RDA1 dominance
270 (91.6%), the RDA1 of *G. ruber albus* and *T. clarkei* 'encrusted' are lower (75.9% and 67.1%
271 respectively). In both *G. ruber albus* and *T. clarkei* 'big' the environmental factors (pH, salinity
272 and temperature) point together and opposite to MLD, while in *T. clarkei* 'encrusted' salinity
273 and MLD are more closely related.



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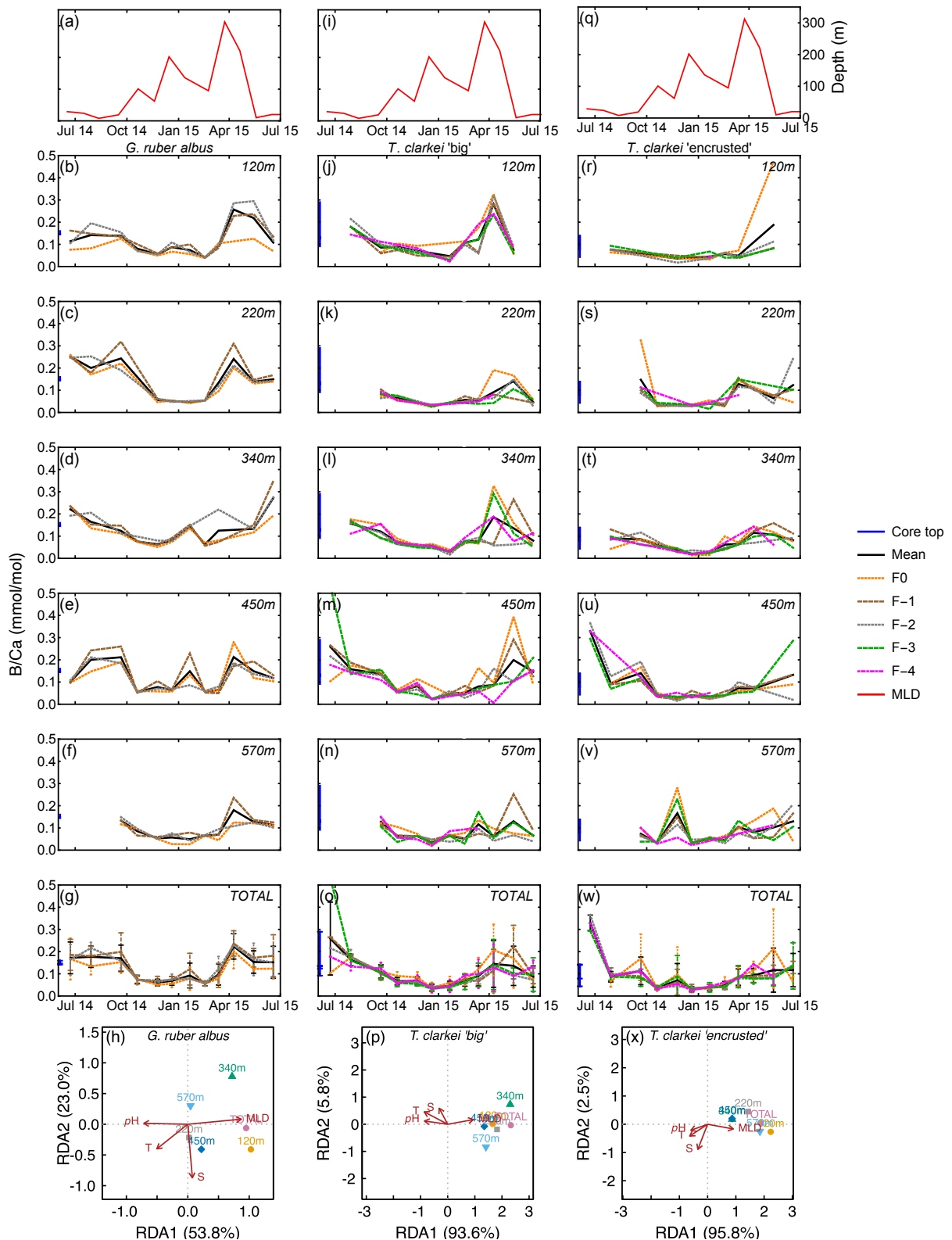
275 Figure 4: Time series of Sr/Ca values measured from the shells of *G. ruber albus* (b-f), *T.*
276 *clarkei* 'big' (j-n) and *T. clarkei* 'encrusted' (r-v), derived from sediment traps located at
277 different water depths (120-570 m). Sr/Ca of core top are marked by a blue bar along the left
278 y-axes. Top panels a, i, and q show the depth (meters below surface layer) of the mixed layer

279 depth (MLD). Panels g, o, and w represent the average values of all depths for each chamber
280 for each species. Panels h, p and x show the redundancy analysis (RDA) per species for each
281 water depth with MLD depth, temperature, salinity and pH as explanatory variables.

282

283 B/Ca values range between 0.03 mmol/mol (570 m; January 2015) to 0.35 mmol/mol (120 m;
284 June 2015) in *G. ruber albus*, with higher values during summer and spring and lower values
285 during the winter (Figs. 5b to 5f). B/Ca measured in the final chamber, F0, are systematically
286 lower compared to F-1 and F-2 values. Unlike most other element ratios, B/Ca values in both
287 phenotypes of *T. clarkei* are similar to the range measured in *G. ruber albus*. In both *T. clarkei*
288 phenotypes, lower B/Ca values were measured during the winter months, most prominently in
289 January. The B/Ca values of *T. clarkei* ‘big’ range between 0.01 mmol/mol to 0.53 mmol/mol
290 with some higher values during spring (Figs. 5k to 5n). B/Ca values in *T. clarkei* ‘encrusted’
291 range between 0.01 mmol/mol to 0.47 mmol/mol (Figs. 5r to 5v). Generally, B/Ca ICV is
292 higher in *T. clarkei* than *G. ruber albus*, especially during spring (Figs. 5g, 5o, and 5w). The
293 SD of *G. ruber albus* is highest at 0.086 during May 2015, at 120 m, and for *T. clarkei* ‘big’
294 and *T. clarkei* ‘encrusted’ the SD is 0.164 (450 m; June-July 2014) and 0.19 (120 m; May 2015)
295 respectively. RDA analysis on B/Ca reveal a distinct RDA1 dominance in both *T. clarkei* ‘big’
296 and *T. clarkei* ‘encrusted’ (93.6% and 95.8%, respectively) compared to only 53.8% in *G. ruber*
297 *albus*. In both *T. clarkei* phenotypes, the sediment trap data cluster together, showing a positive
298 correlation with MLD and negative correlations with temperature, salinity, and pH. In contrast,
299 *G. ruber albus* exhibits a more scattered distribution across water depths, with the 120 m and
300 total samples showing a stronger alignment with MLD.

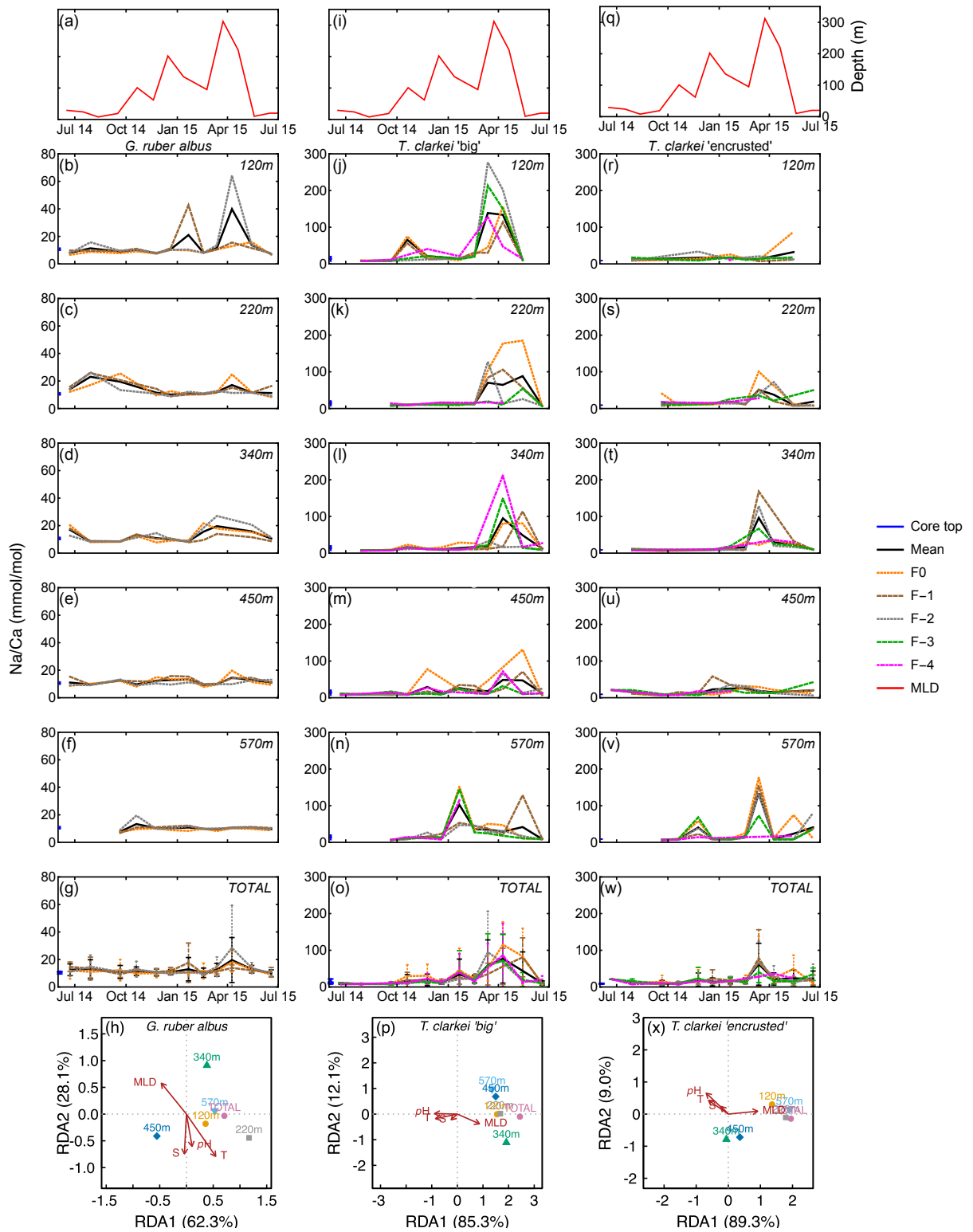
301



307 Panels g, o, and w represent the average values of all depths for each chamber for each species.
308 Panels h, p and x show the redundancy analysis (RDA) per species for each water depth with
309 MLD depth, temperature, salinity and pH as explanatory variables.

310

311 Na/Ca in *G. ruber albus* ranges between 6.60 mmol/mol (220 m; June-July 2014) to 64.14
312 mmol/mol (220 m; April 2015) with a median value of 10.43 mmol/mol (Fig. 6; Fig. S1). The
313 SD of Na/Ca ranges between 0.06 (570 m; February 2015) to 34.37 (120 m; April 2015). Na/Ca
314 in *T. clarkei* ‘big’ ranges from 6.23 mmol/mol (570 m; September 2014) to 426.54 mmol/mol
315 (220 m; March 2015) with a median value of 12.33 mmol/mol and SD range between 0.44 (570
316 m; September 2014) to 106.33 (120 m; March 2015). Na/Ca in *T. clarkei* ‘encrusted’ ranges
317 between 5.43 mmol/mol (570 m; September 2014) to 176.91 mmol/mol (570 m; March 2015)
318 with a median value of 12.41 mmol/mol and SD reaches up to 64.64 (340 m; March 2015).
319 *Globigerinoides ruber albus* has a low ICV during spring, while *T. clarkei* ‘big’ and ‘encrusted’
320 phenotypes display higher ICV during the same time interval. All morphotypes include
321 significant excursions in Na/Ca with high values in *G. ruber albus* during January and April at
322 220m (Fig. 6c), and high Na/Ca in both *T. clarkei* phenotypes at multiple depths and seasons
323 (Figs. 6i-6m and 6p-6t). In particular, *T. clarkei* phenotypes show significant Na/Ca excursions
324 during March-April and ICV (Figs. 6i-6t). RDA analysis on *T. clarkei* ‘big’ show, like B/Ca, a
325 clustering of sediment trap data aligned with MLD and negatively correlated to pH, salinity
326 and temperature positioned on the RDA1 axis (85.3%). The sediment trap data of *T. clarkei*
327 ‘encrusted’ show two distinct groups: 120 m, 220 m, 570 m and ‘Total’ together with MLD
328 and the 350 m and the 450 m groups in the middle between MLD and pH, salinity and
329 temperature, albeit slightly negative on the RDA2 axis (9.0%), while the explanatory
330 parameters are positive on RDA2. The *G. ruber albus* Na/Ca displays a similar distribution to
331 B/Ca however, the environmental parameters are aligned differently: MLD negative on RDA1
332 axis (62.3%) and positive on RDA2 axis (28.1%); and, temperature, salinity and pH negative
333 on RDA2 axis and positioned more to the center of RDA1.



334

335 Figure 6: Time series of Na/Ca values measured from the shells of *G. ruber albus* (b-f), *T.*
336 *clarkei* 'big' (j-n) and *T. clarkei* 'encrusted' (r-v), derived from sediment traps located at
337 different water depths (120-570 m). Na/Ca of core top are marked by a blue bar along the left
338 y-axes. Top panels a, i, and q show the depth (meters below surface layer) of the mixed layer
339 depth (MLD). Panels g, o, and w represent the average values of all depths for each chamber

340 for each species. Panels h, p and x show the redundancy analysis (RDA) per species for each
341 water depth with MLD depth, temperature, salinity and pH as explanatory variables.

342

343 Ba/Ca in *G. ruber albus* ranges from 0.73 $\mu\text{mol/mol}$ (120 m; November 2014) to 36.81
344 $\mu\text{mol/mol}$ (340 m; June 2015). Ba/Ca in *T. clarkei* ‘big’ ranges from 0.39 $\mu\text{mol/mol}$ (120 m;
345 June 2015) to 246.54 $\mu\text{mol/mol}$ (450 m; March 2015). Ba/Ca in *T. clarkei* ‘encrusted’ ranges
346 from 0 $\mu\text{mol/mol}$ (April 2015) to 171.41 $\mu\text{mol/mol}$ (340 m; March 2015) (Fig. 7; Fig. S1). The
347 three morphotypes display varied ICV, although *T. clarkei* shows more prominent ICV during
348 spring months with SD values of 42.06 (340 m; April 2015, ‘encrusted’) and 98.98 (450 m;
349 March 2015, ‘big’) (Figs. 7i-7t and supplementary table S4) than *G. ruber albus* with SD values
350 of 19.14 (220 m; June-July 2014) (Figs. 7b-7f). Furthermore, RDA analyses exhibit for both *T.*
351 *clarkei* phenotypes, a clear RDA1 dominance with 93.6% and 95.8% for ‘big’ and ‘encrusted’
352 respectively. Additionally, for both types, sediment trap data is clustered with positive
353 correlation to MLD and negative to temperature, salinity and pH. For *G. ruber albus*, RDA1
354 value is 53.8% and RDA2 is 23.0%. The sediment traps data are scattered and while the
355 environmental parameters temperature and salinity point together negatively on the RDA2 axis,
356 MLD and pH are positioned on opposite directions on the RDA1 axis (positive and negative
357 respectively).

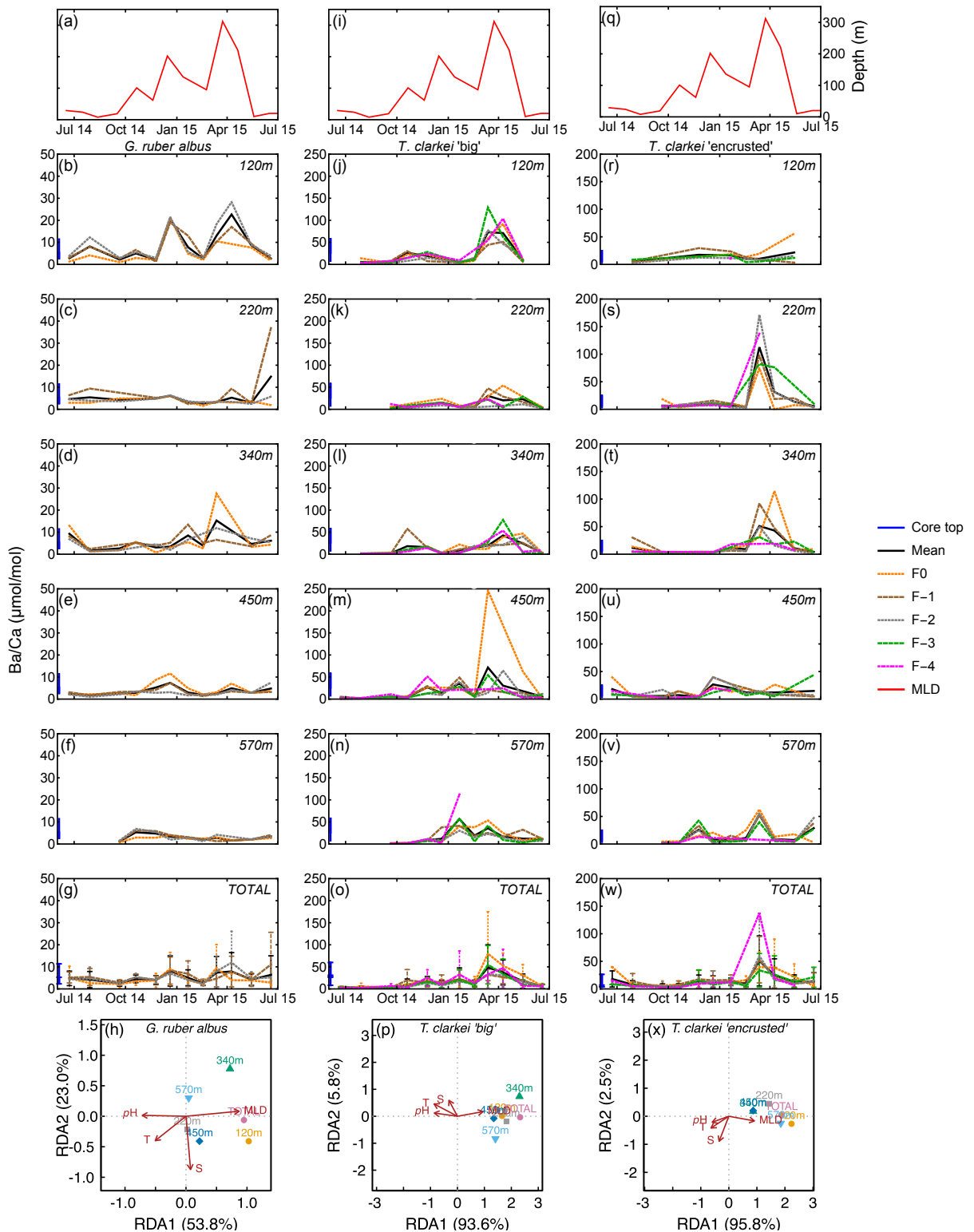


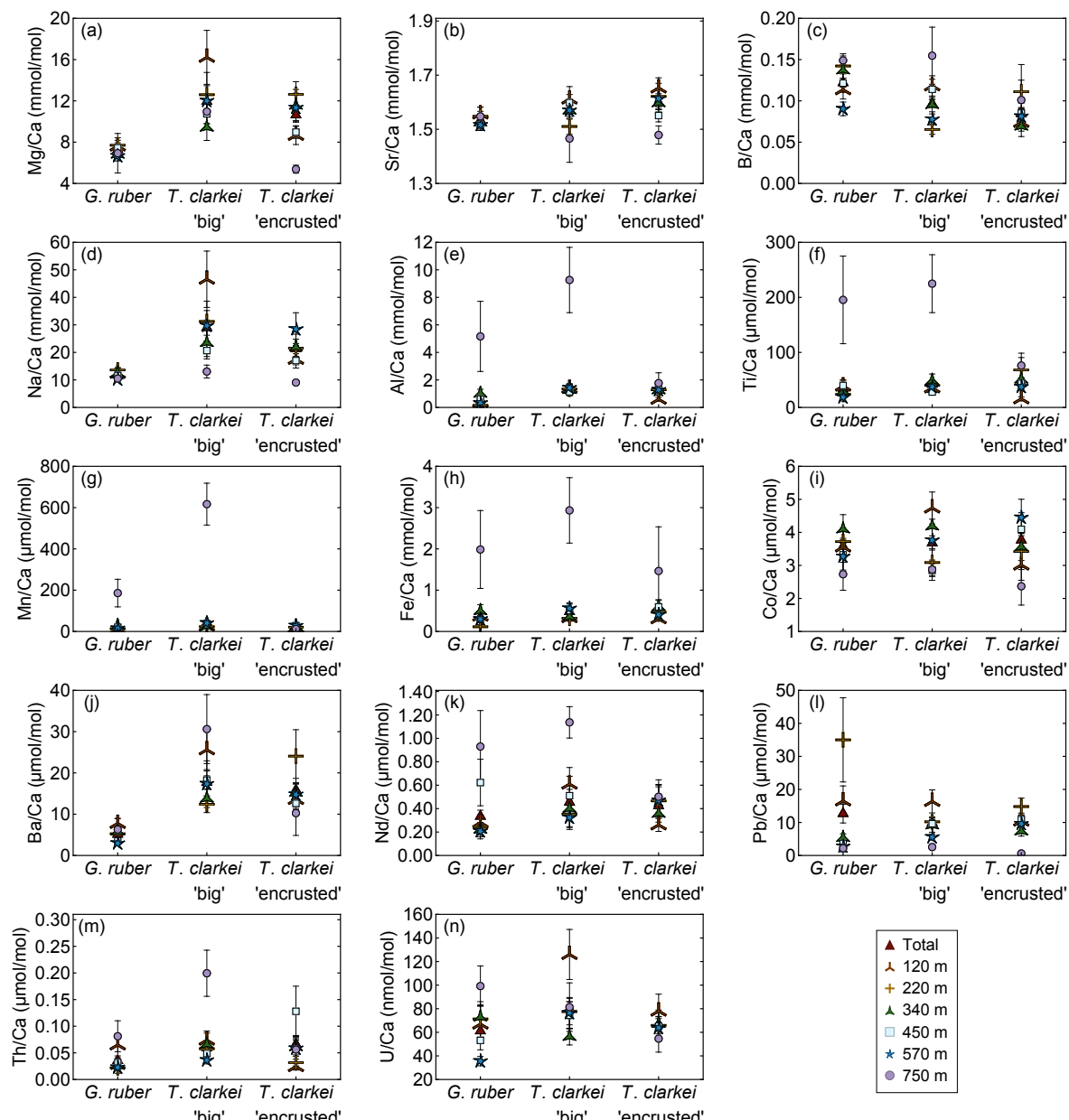
Figure 7: Time series of Ba/Ca values measured from the shells of *G. ruber albus* (b-f), *T. clarkei* 'big' (j-n) and *T. clarkei* 'encrusted' (r-v), derived from sediment traps located at different water depths (120-570 m). Ba/Ca of core top are marked by a blue bar along the left y-axes. Top panels a, i, and q show the depth (meters below surface layer) of the mixed layer depth (MLD). Panels g, o, and w represent the average values of all depths for each chamber

364 for each species. Panels h, p and x show the redundancy analysis (RDA) per species for each
 365 water depth with MLD depth, temperature, salinity and *pH* as explanatory variables.

366

367 **3.2. Depth-averaged values of element/Ca measured in *G. ruber albus* and *T. clarkei* shells**
 368 **using LA-ICP-MS**

369 Generally, the means of Mg/Ca, Sr/Ca, B/Ca, Na/Ca and Ba/Ca in *G. ruber albus* indicate that
 370 the composition of tests, from most water depths is similar to that of core-top samples (Figs.
 371 8a-8d, 8j). In contrast, Al/Ca, Ti/Ca, Mn/Ca, Fe/Ca, Nd/Ca, Th/Ca, and U/Ca (Figs. 8e-8i, 8k,
 372 8m, 8n) in the tests from sediment interface were higher than in the water column, and lower
 373 in case of Co/Ca and Pb/Ca (Figs. 8i, 8l).

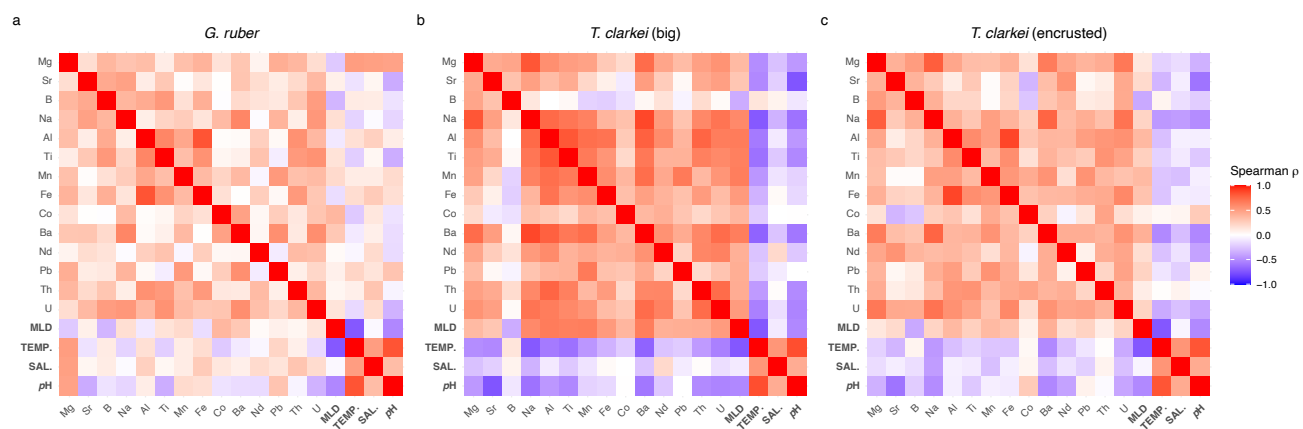


374

375 Figure 8: Pooled mean values of the calcium-normalized element ratios of *G. ruber albus*, *T.*
 376 *clarkei* ‘big’ and *T. clarkei* ‘encrusted’ shells, derived from sediment traps located at different
 377 water depths (120 m to 570 m) and a core top sample (750 m) from the Gulf of Aqaba. Error
 378 bars represent 1 sigma relative standard error (SD/\sqrt{n}).

379
 380 Furthermore, *T. clarkei* tends to demonstrate higher values and higher variability compared to
 381 *G. ruber albus* (e.g., Mg/Ca, Na/Ca, Ba/Ca, Nd/Ca). Compared to the core-top samples, *T.*
 382 *clarkei* from the water column also exhibit relative enrichment in Al/Ca, Ti/Ca, Mn/Ca, Fe/Ca,
 383 Nd/Ca, B/Ca, and Th/Ca (*T. clarkei* ‘big’), and depletion in Co/Ca, Pb/Ca, Sr/Ca, and Mg/Ca
 384 (*T. clarkei* ‘encrusted’) (Fig. 8).

385
 386 3.3. Relationships between element/Ca and environmental parameters
 387 A Spearman correlation matrix was applied to assess the relationships of the element/Ca and
 388 environmental variables in the three analyzed PF phenotypes (Fig. 9; Tab. S2; Fig S12). The
 389 *T. clarkei* types exhibit similar pattern of relationships, with minor differences mainly in
 390 correlation strength (Fig. 9a, 9b). In general, *T. clarkei* shows more significant relationships
 391 than *G. ruber albus*, while, *G. ruber albus*, display different relationships to those of the two
 392 *T. clarkei* types. In *T. clarkei*, Mg/Ca displays relatively strong relationships with Na/Ca,
 393 Ba/Ca, and Al/Ca (Fig. 9b, 9c). Sr/Ca, B/Ca, Co/Ca and Nd/Ca do not display significant
 394 relationships to other elements in *G. ruber albus* as well as in *T. clarkei* ‘big’ and *T. clarkei*
 395 ‘encrusted’.



396
 397 Figure 9: Spearman correlation Matrix of element-Ca means in *G. ruber albus* (a), *T. clarkei*
 398 “big” (b) and, *T. clarkei* “encrusted” (c) and environmental variables.

399 For both *T. clarkei* ‘big’ and *T. clarkei* ‘encrusted’, Na/Ca significantly correlates with
400 Al/Ca, Mn/Ca, and Ba/Ca, the later showing the strongest relationships in *T. clarkei* ‘big’ ($r =$
401 0.82, Fig. 9b; Tab. S2). Relationships between Al/Ca, Ti/Ca, Mn/Ca, Fe/Ca, Ba/Ca and Th/Ca
402 are generally stronger in *T. clarkei* ‘big’ than in *T. clarkei* ‘encrusted’, except for Al/Ca and
403 Fe/Ca, which are stronger related in *T. clarkei* ‘encrusted’ ($r = 0.85$; Tab. S2) than in *T. clarkei*
404 ‘big’ ($r = 0.74$; Tab. S2). Unlike *G. ruber albus*, the U/Ca in *T. clarkei* exhibit relatively strong
405 relationships with Ba/Ca, Na/Ca and Al/Ca (in *T. clarkei* ‘big’) and Mg/Ca (in *T. clarkei*
406 ‘encrusted’, Fig. 9c) while in *G. ruber albus*, U/Ca is poorly related to the other elements (Fig.
407 9a). Mg/Ca of *G. ruber albus*, has a positive correlation with temperature, salinity, and pH, and
408 a negative correlation with MLD; other element/Ca in *G. ruber albus* do not exhibit significant
409 correlations with these environmental variables (Fig. 9a). By contrast, in *T. clarkei* (‘big’ and
410 ‘encrusted’), element/Ca show strong correlations with MLD but not with temperature, salinity,
411 or pH (Figs. 9b, 9c). In the *T. clarkei* ‘big’ morphotype, SD (ICV) shows strong correlations
412 between multiple element/Ca and with MLD. On the other hand, in *T. clarkei* ‘encrusted’ and
413 *G. ruber albus*, correlations are weaker and/or inconsistent across variables, and no uniform
414 pattern emerges.

415

416 4. Interpretation

417 4.1 Inter chamber variability (ICV)

418 Shell-bound element/Ca display varying trends across different chambers depending on the
419 specific element ratios, and varying over water depth and time (Figs. 3-7). In most element/Ca
420 ICV is higher during water column mixing months (March-May; e.g., Al/Ca, B/Ca, Ba/Ca,
421 Co/Ca, Fe/Ca, Mg/Ca) in all water depth horizons for *T. clarkei* ‘big’ and *T. clarkei* ‘encrusted’,
422 while for *G. ruber albus* it increases in the two upper water depth horizons (i.e., 120 m and 220
423 m). Elevated element/Ca values and high ICV may reflect the changes in the water properties
424 like the temperature, salinity, pH and nutrient availability derived from the mixing of the water
425 column (Fig. S12, Figs 3-7 panels h, p, and x). For some element/Ca ratios (e.g., Na/Ca, Fig.
426 6/panels g, o, and w; Ba/Ca, Fig. 7/panels g, o, and w) ICV varies with depth and shows
427 seasonal differences (i.e., less variation with depth during water column stratification and more
428 variation with depth during water column mixing), whereas for others (e.g., B/Ca, Fig. 5/panels
429 g, o, and w; Sr/Ca, Fig. 4/panels g, o, and w) it remains relatively constant with depth. In the
430 *T. clarkei* ‘big’ morphotype, ICV shows strong correlations across most element/Ca and with
431 MLD, indicating that increases in ICV reflects a common driver, most likely the mixed-layer
432 depth (Fig. S12).

433 Typically, most PF reproduction-cycles span about a month with individual chambers forming
434 within several hours (Bé et al., 1977), while the time interval between chamber formation can
435 range from hours to weeks (Schiebel & Hemleben, 2017, and references therein). Setting aside
436 the March-May time-interval where PF shells show exceptionally high ICV, *G. ruber albus*
437 generally exhibits lower values (e.g., Mg/Ca, B/Ca), and less ICV compared (e.g., Mg/Ca
438 SD=1.67) to *T. clarkei* 'big' and 'encrusted' (e.g., Mg/Ca SD=5.24 for 'big' type and Mg/Ca
439 SD=3.55 for 'encrusted' type). The residence of *G. ruber albus* in the relatively homogenous
440 and narrow living environment in the surface mixed layer (Schiebel & Hemleben, 2017;
441 Thirumalai et al., 2014; and others), could explain relatively lower ICV. In contrast, *T. clarkei*
442 dwell in the dynamic region near/under the thermocline (Schiebel & Hemleben, 2017; Levy et
443 al., 2023) over a wider dwelling depth horizon, and may experience more heterogeneous
444 environmental conditions which may result in higher ICV.

445 The secondary crust observed on *T. clarkei* 'encrusted' morphotypes, which covers all
446 chambers of the tests analyzed here, does not significantly alter element/Ca values when
447 compared to *T. clarkei* 'big', unlike the crust of *Neogloboquadrina dutertrei* (Jonkers et al.,
448 2012). Nonetheless, the RDA (Figs. 3-7) and the Spearman correlation analyses (Fig. 9) of both
449 *T. clarkei* types indicate that while *T. clarkei* 'big' show a positive correlation to MLD and a
450 weaker or negative correlation to other environmental parameters (T, S and pH), the *T. clarkei*
451 'encrusted' type calcification mechanism is more complex and could be influenced by
452 additional factors / environmental variables. Therefore, while both types are eligible for
453 paleoceanography and paleoclimate reconstructions, the interpretation of their measurements
454 might be different.

455 The ultimate chamber (F0) presents different systematics compared to the preceding
456 chambers in both *T. clarkei* and *G. ruber albus* (Fig. S11). In *T. clarkei* (both 'big' and
457 'encrusted'), the F0 typically exhibits higher values of B/Ca, Na/Ca, Mg/Ca, and Al/Ca
458 compared to the previous chambers. In contrast, *G. ruber albus* displays relatively lower values
459 in F0 for the same ratios highlighting species-specific differences in chamber formation (Fig.
460 S11). Interestingly, Sr/Ca does not follow the same pattern. In *T. clarkei* 'big' the Sr/Ca
461 distribution mirrors the trends of other elements, while F0 in *G. ruber albus* and *T. clarkei*
462 'encrusted' shows an even distribution of Sr/Ca, likely reflecting the relatively constant Sr/Ca
463 values in the water column during the lifespan of a single test. These observations in *G. ruber*
464 *albus* are consistent with previous studies that measured Mg/Ca in individual chambers (Bolton
465 et al., 2011; Davis et al., 2020; Fischer et al., 2024). The contrasting systematics of F0 leading
466 to elevated ICV in the ultimate chamber compared to the previous chambers was previously

467 suggested to be associated with a chamber wall that is not fully calcified (Schiebel &
468 Hemleben, 2017; Bolton et al., 2011; Fischer et al., 2024). Differences in F0 systematics
469 between *T. clarkei* and *G. ruber albus* could be driven by species-specific calcification
470 processes, though further research is needed to clarify this issue. Additionally, it is important
471 to consider potential biases in small chambers such as F-4 in *T. clarkei* morpho-species, where
472 methodological challenges (e.g., laser spots hitting sutures) may skew element/Ca
473 measurements. Consequently, we conclude that the exclusion of F0 and F-4 may enhance the
474 robustness of reconstructions of the marine environment in studies of downcore records.

475

476 4.2 Relationships of element ratios of the three PF morphotypes

477 The contrasting results of the correlation matrixes of the three morpho-species, suggests
478 species-specific mechanisms while calcifying their shells. The Mg/Ca in *T. clarkei* which
479 strongly correlates with Na/Ca, Ba/Ca, and Al/Ca (Fig. 9b, 9c), suggests more than one
480 environmental process affects Mg/Ca in the tests as the other element/Ca are considered proxies
481 to different environmental characteristics such as salinity, productivity, and terrigenous input
482 (Chang et al., 2015; Mesa-Fernández et al., 2022; Beasley et al., 2021). This is also reinforced
483 by the positive correlation to MLD and the negative correlation to temperature, salinity and pH
484 (Fig. 9 and RDA panels in figs. 3-7). Additionally, Mg/Ca in *G. ruber albus* show a relative
485 strong positive correlation to temperature, salinity and pH and a negative correlation to MLD
486 (Fig. 9a and Figs. 3h, 3p and 3x). Similar to *G. ruber albus*, in the *T. clarkei* types Sr/Ca, B/Ca,
487 Co/Ca and Nd/Ca display a much weaker relationship to other elements making them more
488 suitable proxies for distinct and independent environmental properties.

489 In *G. ruber albus*, Mg/Ca, Sr/Ca and B/Ca show no significant relationships with other element
490 ratios, indicating that independent processes likely govern their proxy systematics (Fig. 9c).
491 Similarly, Co/Ca, Nd/Ca and U/Ca also do not correlate with other element/Ca. While Na/Ca
492 and Ba/Ca exhibit some degree of correlation, as do Mn/Ca and Pb/Ca, the lithophilic elements,
493 Al/Ca, Ti/Ca, which are considered proxies for terrigenous dust input (Chang et al., 2015;
494 Mesa-Fernández et al., 2022; Beasley et al., 2021), as well as, Fe/Ca, and Th/Ca, all show a
495 relative strong correlation. Their correlation implies they can be used together for
496 reconstructing terrigenous input to the water column. Among the lithophilic elements, Th/Ca
497 display a relatively weaker relationship, suggesting a potential effect of additional processes
498 such as scavenging (Anderson et al., 1983; Francois et al., 2004; Costa et al., 2020).

499

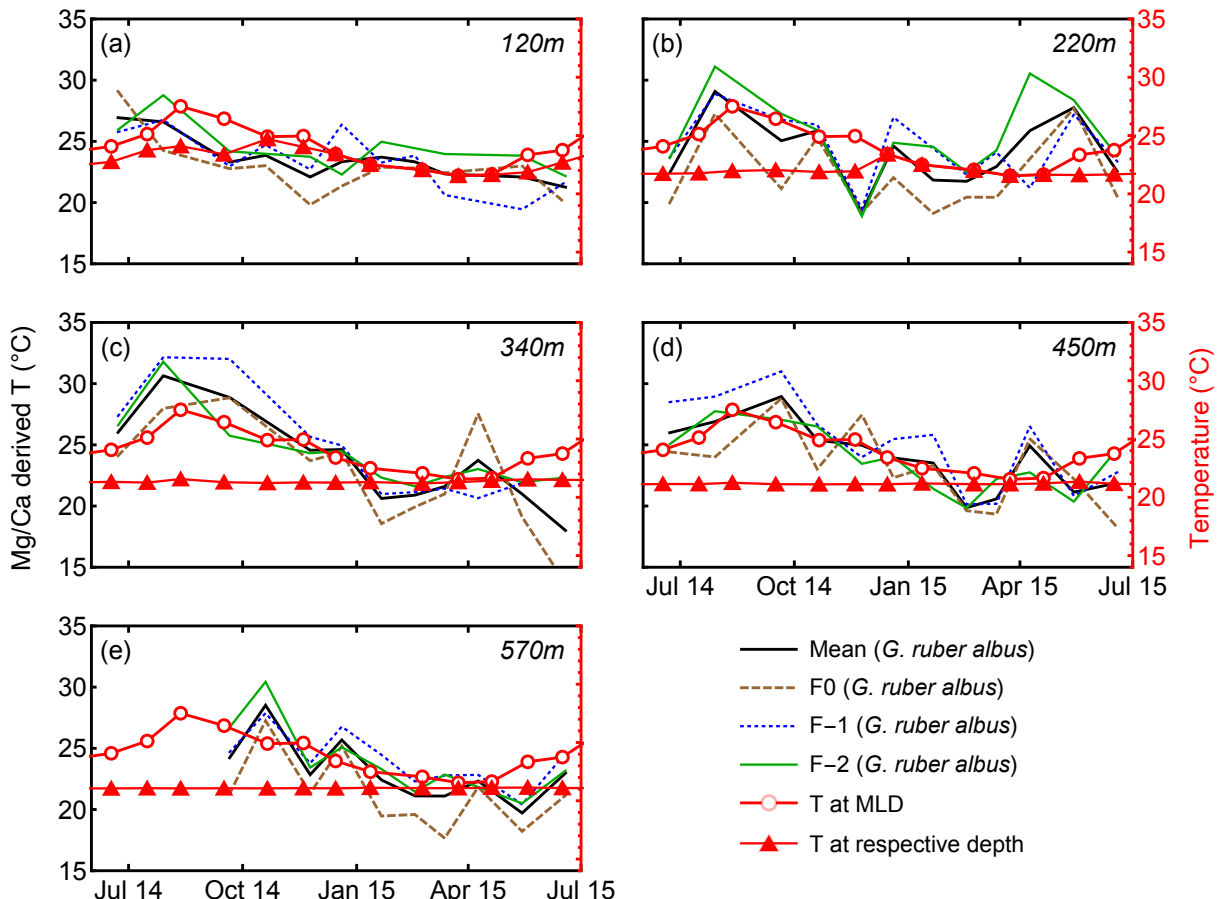
500 4.3 Mg/Ca as a proxy for sea surface temperature

501 Shell-bound Mg/Ca of calcareous foraminifera have been extensively utilized as a paleo-
502 thermometer (e.g., Nürnberg et al., 1996; Sadekov et al., 2009). Many of these Mg-temperature
503 calibrations rely on whole-test or pooled-mean Mg/Ca values to reconstruct past sea surface
504 temperatures (Spero et al., 2003; Ganssen et al., 2010; and others). Several studies have
505 measured intra-test and inter-test Mg/Ca in an effort to produce Mg-temperature calibrations
506 using single chamber measurements of *G. ruber* (Sadekov et al., 2008; Bolton et al., 2011;
507 Davis et al., 2020; Levy et al., 2023; Fischer et al., 2024). Previous work on sediment trap-
508 derived specimens of *T. clarkei* and *G. ruber albus* from the GOA indicated that *T. clarkei* is
509 not suitable for temperature reconstructions, due to its presumed deep dwelling-depth below
510 the thermocline together with its high sensitivity to water column mixing events. However,
511 while *G. ruber albus* shows exceptionally high pooled mean Mg/Ca values in the GOA in
512 comparison to other ocean regions, it also exhibits seasonal variations that indicate effective
513 applicability as a paleothermometer (Levy et al., 2023). Due to the high seawater salinity of
514 the GOA, a local calibration curve was proposed (Eq. 1; Levy et al., 2023).

515

516
$$\frac{\text{Mg}}{\text{Ca}} = 0.39(\pm 0.30) \cdot e^{0.12(\pm 0.03)T} \quad (1)$$

517



518

519 Figure 10: *G. ruber albus* Mg/Ca-derived temperatures versus measured temperatures (red).
 520 The calculated temperatures were derived from Eq. 1 for individual chambers. See also Levy
 521 et al. (2023).

522

523 Pooled mean values of Mg/Ca in *G. ruber albus* taken from all water column depths in the
 524 GOA reflect MLD temperatures (Fig. 10, Fig 9). Inter-chamber variability (ICV) has been
 525 shown to affect the local Mg/Ca temperature calibration (Eq. 1) of *G. ruber albus* (Levy et al.,
 526 2023; Fig. 10). Generally, Mg/Ca-derived temperatures from chambers F-1 and F-2 correspond
 527 closely with mixed layer depth (MLD) temperatures. However, beneath the photic zone, Mg/Ca
 528 of all three chambers F0, F-1, and F-2 of the *G. ruber albus* specimens exhibit poor fits with
 529 measured temperatures (Fig. 10). Given that *G. ruber albus* calcifies its shell in the photic zone
 530 (Schiebel and Hemleben, 2017), these findings support the use of Mg/Ca as a
 531 paleothermometer for the mixed layer (Nürnberg et al., 1996). Sadekov et al. (2009) measured
 532 F0-Mg/Ca in core top individuals of *G. ruber albus* from various latitudes and have reported
 533 an agreement with sea surface temperature. Hupp and Fehrenbacher (2024) measured intra-test
 534 variability in the polar and sub-polar species *Neogloboquadrina incompta*, *N. pachyderma*, and
 535 *Turborotalita quinqueloba* and have not reported any issues regarding the use of F0 for

536 temperature reconstruction. In the GOA however, the Mg/Ca-derived temperatures from
537 chamber F0 calculate lower Mg/Ca temperatures of the MLD than chambers F-1 and F-2 (Fig.
538 10). Although Mg/Ca data from chambers F-1 and F-2 appear suitable for reconstructing
539 temperatures and demonstrate agreement with MLD temperature trends, the high ICV in *G.*
540 *ruber albus* is evidently too great to accurately reflect ambient temperatures using this
541 calibration. Therefore, and based on these new observations, we suggest that optimal Mg/Ca-
542 temperature calibration (Eq. 1) should be based on the pooled mean of the F-1 and F-2
543 chambers at all depths as the final chamber might lead to different results while reconstructing
544 temperature.

545

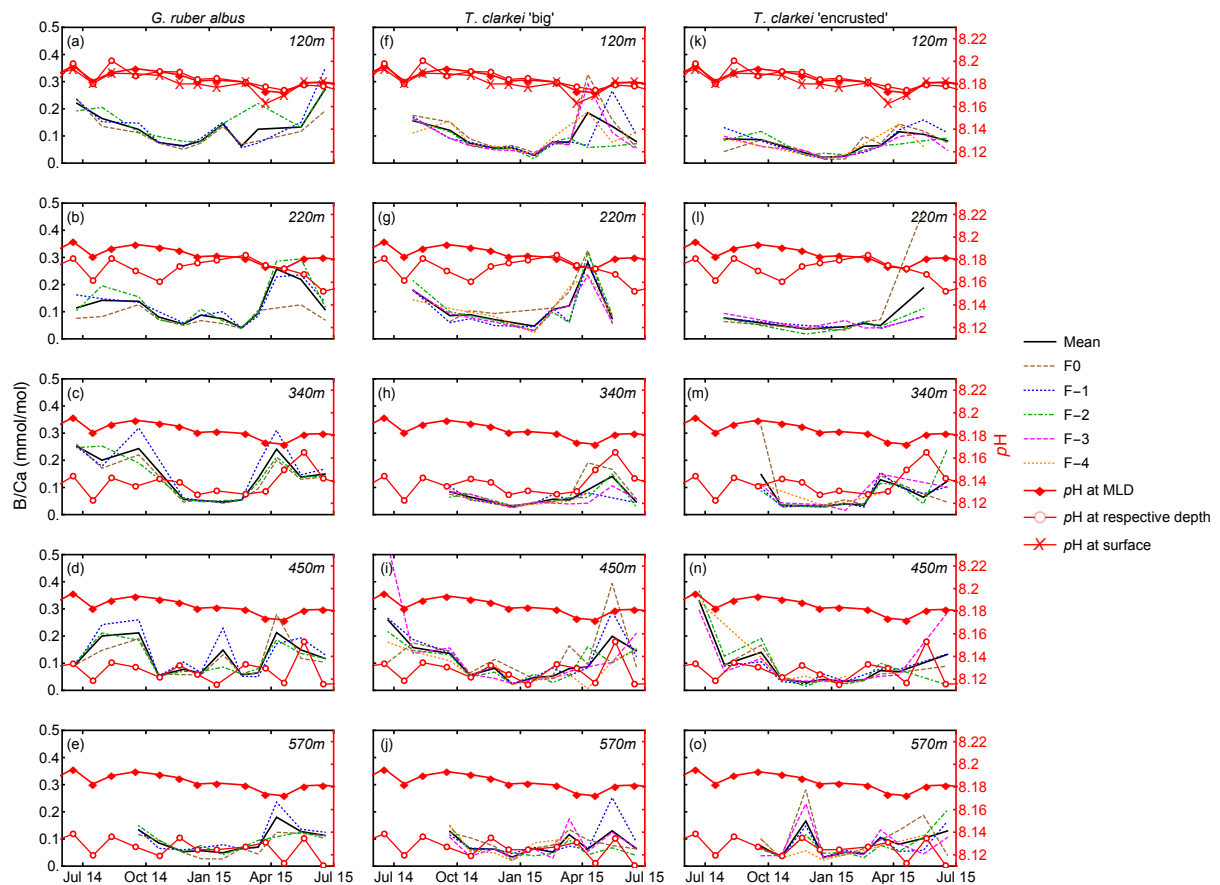
546 4.4 B/Ca as a proxy for pH

547 B/Ca in some PF species has been suggested to be a proxy for pH (Yu et al., 2007; Allen et al.,
548 2011). Comparing chamber B/Ca of both *G. ruber albus* and *T. clarkei* (both ‘big’ and
549 ‘encrusted’) alongside pH at various water column depths in the GOA reveals contrasting
550 results. While B/Ca in *G. ruber albus* exhibits seasonality (Fig. 5), with lower values during
551 winter months, it does not appear to be consistent with the pH of respective water depth nor
552 the MLD (Fig. 11). This inconsistency suggests that B/Ca in *G. ruber albus* from the GOA is
553 not a reliable recorder of ambient water pH. Similarly, Henehan et al. (2015) and Naik & Naidu
554 (2014) reported that B/Ca of open ocean core-top samples and down-core sediment samples do
555 not display a pH relationship.

556 Alternatively, B/Ca in *G. ruber albus* may be sensitive to salinity and micro-environments
557 produced by PF symbionts with pH levels which are distinct from the ambient water column.
558 Culture experiments have shown that B/Ca is affected by salinity and increases with increasing
559 salinity (Allen et al., 2012). However, only small salinity changes occur in the GOA (Fig. 1),
560 which argue against a strong B/Ca-salinity relationship that would result in a B/Ca seasonal
561 trend. It was suggested that photo-symbionts such as dinoflagellates in *G. ruber albus* create
562 micro-environments with pH levels, which are distinct from ambient seawater, to accommodate
563 for their photosynthetic activity, and indicate that B/Ca is more affected by pH in those micro-
564 environments than the water column pH (Hönisch et al., 2021; Babila et al., 2014). An
565 additional observation for the *G. ruber albus* B/Ca values is that they are relatively high in
566 comparison to values from other studies. The relatively high salinity in the GOA (~41),
567 combined with the photosymbiont activity in *G. ruber albus* may explain the elevated B/Ca
568 values (Henehan et al., 2015; Hönisch et al., 2021; Babila et al., 2014).

569 In contrast to *G. ruber albus*, B/Ca in the photosymbiont barren *T. clarkei* may possibly record
 570 the changes in pH (Fig. 11) of seawater at its assumed ambient dwelling depth (i.e., 340 m-570
 571 m), indicative of shifting between the deeper water column depth horizons where pH changes
 572 are evident. Based on the fluxes of *T. clarkei* (Chernihovsky et al., 2018; Fig. 2), the B/Ca of
 573 *T. clarkei* in the sediment record likely represent the pH beneath the thermocline and within
 574 the deep-water column horizons for specimens that lived from early winter through spring. In
 575 particular, pH at 340 m trends appear to follow the B/Ca trends of *T. clarkei* types. For B/Ca-
 576 pH calibrations utilizing the pooled mean of data from the chambers F-1, F-2, and F-3 may be
 577 used, while excluding the F0 and F-4 chambers where more ICV is visually apparent (Fig. 11).

578



579
 580 Figure 11: Single chamber B/Ca and in situ pH measured at MLD depth (empty red circles)
 581 and 120 m water depth for *G. ruber albus*, *T. clarkei* 'big' and *T. clarkei* 'encrusted'.

582

583 4.5 Na/Ca as a proxy for salinity and Ba/Ca as an indicator for productivity

584 Cultured individuals and samples from the surface Caribbean and the Gulf of Guinea of live *T.*
 585 *sacculifer* indicate that Na/Ca can be used as a proxy for salinity, without temperature
 586 dependence, however, a species-specific calibration might be required (Bertlich et al., 2018).

587 Despite the high variability of PF Na/Ca values in the GOA during water column mixing (Fig.
588 6), salinity remains high and relatively constant, ranging between 40.4-40.7. Consequently, a
589 local Na/Ca-salinity calibration shows no significant relationship for any of the three PF
590 morphotypes.

591 Na/Ca values in PF from the GOA are notably higher compared to other regions. Gray *et*
592 *al.* (2023) explored the relationship between Na/Ca and salinity in *G. ruber albus* collected
593 from sediment traps, plankton tows, culture samples, and core top samples, contributing to the
594 ongoing discussions regarding the reliability of Na/Ca as a proxy for salinity in both planktic
595 and benthic foraminifera (Allen *et al.*, 2016; Geerken *et al.*, 2018; Mezger *et al.*, 2016, 2018;
596 Gray *et al.*, 2023, and references therein). They concluded that the measurement method (i.e.,
597 ‘solution’ ICP-MS vs. LA-ICP-MS) influences the values of Na/Ca and in turn the relationship
598 with salinity, i.e., weak in solution-based compared to significant in laser ablation-based, at
599 salinity over 36.

600 Comparing Na/Ca of *G. ruber albus* from the shallowest sediment trap (120 m) in the GOA
601 with the Na/Ca of *G. ruber albus* plankton tows-samples from the GOA deployed and collected
602 in January 2010 and October 2013 (Gray *et al.*, 2023), both measured using LA-ICP-MS,
603 generally reveals similar results, excluding the high-value excursions observed in some single
604 chamber measurements (Fig. 6). *Turborotalita clarkei* in the GOA exhibits elevated Na/Ca
605 values in both ‘big’ and ‘encrusted’ compared to *G. ruber albus*. Unlike *G. ruber albus*, there
606 is relatively higher variability between water depths as well as significantly higher values in
607 March, April, and May associated with water column mixing (Fig. 6). During these mixing
608 events, nutrient-rich, high salinity (~40.7) water ascend upward. Therefore, the Na/Ca of *T.*
609 *clarkei* may serve as a proxy for water column stability, i.e., stratification vs. mixing.

610 The Ba/Ca in the three morpho-species show a relatively strong correlation with Na/Ca
611 (0.74 and 0.82 in *T. clarkei* ‘big’ and ‘encrusted’ respectively, and 0.54 in *G. ruber albus*, the
612 second highest ratio and exceeded only by the 0.57 of Pb/Ca). Ba/Ca is presumably unaffected
613 by temperature, salinity, and pH (Hönisch *et al.*, 2011). In non-spinose species, Ba/Ca typically
614 shows positive relationships with productivity and potentially can be used as an indicator of
615 river run-off (Fritz-Endres *et al.*, 2022; Hönisch *et al.*, 2011; Weldeab *et al.*, 2014). Although
616 floods in the catchment area of the GOA are brief and occur only few times each year (Katz *et*
617 *al.*, 2015), significant Ba/Ca perturbations during water column mixing may reflect nutrient-
618 rich water admixing to the surface water (Fig. 7).

619

620 5. Discussion:

621

622 5.1 Temporal and vertical dynamics of element/Ca in the GOA

623 Trace element incorporation into the calcium carbonate shells of planktic foraminifera during
624 calcification is controlled by environmental and ecological factors in the water column such as
625 temperature, salinity, pH, the carbonate system, dust and terrigenous inputs, as well as whether
626 a species harbor photosymbionts (Schiebel & Hemleben, 2017; and others). Shells of *G. ruber*
627 *albus*, *T. clarkei* ‘big’ and *T. clarkei* ‘encrusted’ from the GOA show species-specific behavior
628 and offer new insights into how these species respond to the vertical and temporal variations
629 in the water column. For most elements, the smaller *T. clarkei* specimens display higher trace
630 element ratios than the larger *G. ruber albus*, suggesting more efficient trace element
631 incorporation to the shell or implying that its habitat deeper in the water column has conditions
632 which result in higher trace element incorporation (Fig. 8). Some element ratios such as Mg/Ca,
633 Sr/Ca, B/Ca, Na/Ca (for *G. ruber albus*) and Ba/Ca for both *G. ruber albus* and *T. clarkei*
634 ‘encrusted’, show overlap between specimens from the water column and from core-tops (Fig.
635 8), confirming the robustness of downcore-based records allowing to further consider these
636 element/Ca recorders of the water column as paleo-proxies.

637 While water depth likely influences element/Ca through variations in physical and
638 chemical conditions, the observed inter-chamber variability (ICV) and element/Ca differences
639 between species cannot be attributed to any single environmental parameter. Nonetheless,
640 elements such as Al/Ca, Ti/Ca, Mn/Ca, and Fe/Ca for all species, and Mg/Ca, Sr/Ca, Na/Ca,
641 and Ba/Ca for *G. ruber albus* alone, demonstrate consistent behavior across the water column,
642 suggesting that depth-related factors do not significantly alter calcification mechanisms. This
643 supports the use of pooled mean values for specimens over multiple sediment traps spread over
644 depths (Levy et al., 2023). Interestingly, most element/Ca peak during water column mixing in
645 March-April 2015 for all three morphotypes analyzed here, accompanied by larger ICV (Figs.
646 3-7). Mg/Ca in *G. ruber albus* and Sr/Ca in all three morpho-species show less pronounced
647 excursions, while other trace element ratios (e.g., Co/Ca, U/Ca) exhibit more variability and
648 more extreme values (Figs. S5 and S10). These observations can reflect: i) primary calcite
649 structure alterations driven by environmental shifts and life cycle changes, ii) secondary
650 mineralization (e.g., barite, Amorphous Calcium Carbonate, ACC) (Torres et al., 2010; Evans
651 et al., 2020 and references therein), and iii) fluid inclusions within the shell structure (Gray et
652 al., 2023).

653 All of these relationships do possibly concern the ontogenetic PF calcite, since SEM
654 imaging of GOA specimens did not reveal secondary minerals or overgrowth on shell calcite

655 (Levy et al., 2023). Moreover, the enrichment of multiple trace elements across species
656 suggests that secondary minerals are unlikely to be responsible for these trends. Discrepancies
657 between Na/Ca in plankton tow versus core-top samples in the Red Sea (Mezger et al., 2018),
658 as well as higher Na/Ca values measured by LA-ICP-MS compared to solution ICP-MS, have
659 been linked to early diagenesis of Na-enriched phases like spines, ACC, or fluid inclusions
660 (Gray et al., 2023). However, spines and ACC were ruled out for GOA samples, as all of the
661 specimens had lost their spines before analysis and ACC was not detected via SEM. Given that
662 most element/Ca in GOA shells are elevated relative to PF data from elsewhere, fluid inclusions
663 may be a contributing factor (Gray et al., 2023). However, more research is required to
664 investigate whether fluid inclusions are evident in PF shells from the GOA. In the absence of
665 fluid inclusions, environmental changes, particularly during water column mixing, are
666 considered to be the primary drivers of the observed trace element/Ca enrichments in the GOA.
667

668 5.2 Water column and sediment signal correlation: Implications to Paleoceanographic 669 studies

670 Several element ratios (e.g., Al/Ca, Ti/Ca, Mn/Ca, Fe/Ca, Nd/Ca, U/Ca, Co/Ca, and Th/Ca)
671 exhibit discrepancies between water column and core-top specimens (Fig. 8). Some, like
672 Co/Ca, have lower values in surface sediment than the water column, while others, like Fe/Ca
673 show higher values. Differences between sediment trap samples and core-top samples may
674 stem from differential diagenetic processes that affect element/Ca in specimens taken from the
675 water column and the sea floor. For example, diagenetic processes can lead to Mn accumulation
676 and higher Mn/Ca in PF from the core top (McKenzie, 1980; Steiner et al., 2017). Conversely,
677 core-top PF samples may show lower ratios due to the release of these metals into pore water
678 over time (e.g., Co/Ca, Fig. 8i). This release can alter the elemental composition, potentially
679 skewing paleoenvironmental reconstructions. Understanding these processes is crucial for
680 accurately interpreting geochemical data from both sample types.

681 Despite the offsets of Al/Ca and Ti/Ca between core top and water column specimens, they
682 nevertheless may be utilized to trace the origins of terrigenous inputs and identify periods of
683 dust deposition in the geological record (Torfstein et al., 2017; Martinez-Garcia et al., 2011).
684 Our data reveal significant seasonal excursions in Al/Ca and may demonstrate the use of Al/Ca
685 and Ti/Ca in PF tests as proxies for dust or terrigenous input to the ocean (Fig. S3).

686 Core top element/Ca values that fall within the same range of values of the sediment trap
687 specimens (Mg/Ca, Sr/Ca, B/Ca, Na/Ca, and Ba/Ca; Fig. 8) suggest that they could reflect
688 water column conditions. The high temporal variability in many of these element/Ca data,

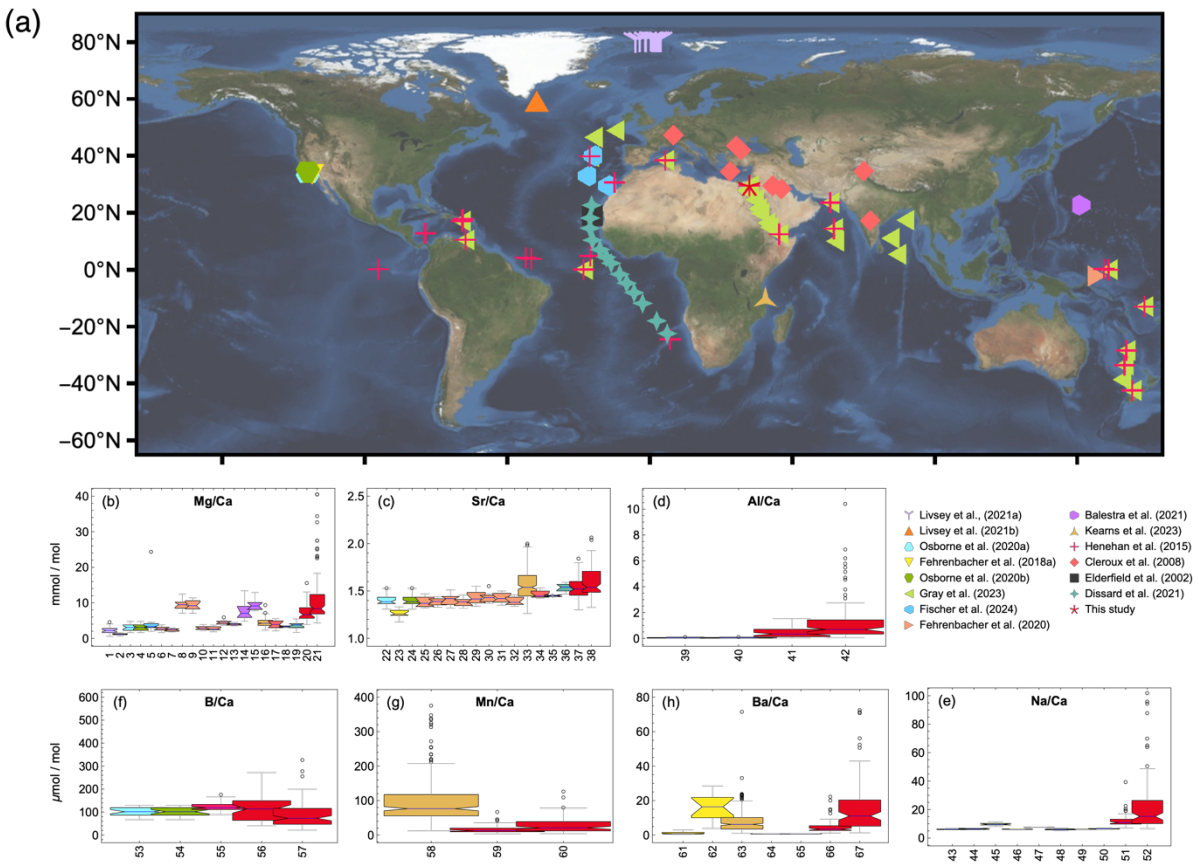
689 together with the varying PF population dynamics throughout the year (Fig. 2) may be
690 considered when approaching PF from sediment cores. Seasonal trends in element/Ca are often
691 obscured by the spring mixing event. However, exceptions to this are observed in Mg/Ca for
692 *G. ruber albus* (Fig. 4; Levy et al., 2023) and B/Ca for *T. clarkei* (Fig. 5), where clear seasonal
693 patterns emerge. A key limitation of reconstructing past environments from element/Ca in PF
694 shells is the challenge of disentangling seasonal effects from other more episodic
695 environmental signals. However, by identifying water column mixing events through positive
696 element/Ca excursions and elevated ICV, which are evident across all species (Figs. 3-7), it
697 may be possible to identify the time intervals over which environmental changes are
698 reconstructed. This could allow for more accurate reconstructions of shifts in temperature,
699 carbonate chemistry, and nutrient availability during specific mixing events, improving our
700 understanding of past ocean conditions.

701

702 5.3 Regional comparison of geochemical conditions and PF element/Ca

703 The Mg/Ca, Al/Ca, and Na/Ca in PF from the GOA generally exceed those reported from other
704 regions (Fig. 12b – 12e). Sr/Ca values, while reaching up to 2.2 mmol/mol during spring, have
705 an average of 1.5 mmol/mol, consistent with previous studies (Fig. 12c; Kisakürek et al., 2008;
706 Cleroux et al., 2008; Elderfield et al., 2002; Brown & Elderfield, 1996; Dissard et al., 2021).
707 The high Mg/Ca range in the GOA versus typical open-ocean levels (0.5-5 μ mol/mol) is
708 attributed to elevated salinity (~ 41 compared to mean ocean values of 34.7), which is also
709 evident by the high Na/Ca. The high Al/Ca values and their large variation may be attributed
710 to the close proximity of GOA to terrestrial input. Ba/Ca in the GOA are significantly higher
711 than the values reported in prior studies from Atlantic Ocean core samples and culture
712 experiments (Hönisch et al., 2011; Lea & Boyle, 1991), representing a roughly ten-fold
713 difference. These discrepancies likely stem from two factors: (1) higher salinity in the GOA
714 increases the availability of cations and trace element incorporation into foraminifera shells,
715 and (2) higher-resolution measurements here which reveal chamber-specific elemental ratios,
716 where early chambers (F-1 and F-2) exhibit higher values than final chambers, leading to more
717 accurate, chamber-level data compared to bulk measurements. Combined, these factors explain
718 the elevated values relative to global reports.

719



720

721 Figure 12. Global comparison of major and trace element-to-calcium ratios. (a) sample global
 722 map, (b) Mg/Ca of *N. pachyderma*, *G. bulloides*, *G. ruber white*, *N. dutertrei*, *O. universa*, *P.*
 723 *obliquiloculata*, *T. sacculifer* and *T. clarkei* derived from various sources (plankton tows/nets,
 724 sediment traps, cores) and measured by Laser Ablation (LA)-ICP-MS, solution-ICP-MS (SOL)
 725 and Electron micro-probe analyses (EPMA). (c) Sr/Ca of *G. bulloides*, *G. ruber white*, *N.*
 726 *dutertrei*, *O. universa*, *P. obliquiloculata*, *T. sacculifer* and *T. clarkei* derived from various
 727 sources (plankton tows/nets, sediment traps, cores) and measured by LA-ICP-MS and solution-
 728 ICP-MS. (d) Al/Ca of *G. bulloides*, *G. ruber white* and *T. clarkei* derived from sediment traps
 729 and measured by LA-ICP-MS. (e) Na/Ca of *G. ruber white* and *T. clarkei* from various sources
 730 (plankton tows/nets, sediment traps, cores and cultured samples) and measured by LA-ICP-MS
 731 and solution-ICP-MS. (f) B/Ca of *G. bulloides*, *G. ruber white* and *T. clarkei* derived from
 732 sediment traps and measured by LA-ICP-MS. (g) Mn/Ca of *G. ruber white* and *T. clarkei*
 733 derived from cores and sediment traps and measured by LA-ICP-MS. (h) Ba/Ca of *G. ruber*
 734 *white*, *N. dutertrei* and *T. clarkei* derived from various sources (plankton tows/nets, sediment
 735 traps, cores and cultured samples) and measured by LA-ICP-MS. See table 1 for detailed
 736 description of methods.

#	Element/Ca	Reference	Species	Collecting method	Measuring method
1	Mg/Ca	Livsey et al. (2021a)	<i>N. pachyderma</i>	Plankton tows / nets	LA
2	Mg/Ca	Livsey et al. (2021b)	<i>N. pachyderma</i>	Sediment traps	LA
3	Mg/Ca	Osborne et al. (2020)	<i>G. bulloides</i>	Sediment trap	LA
4	Mg/Ca	Osborne et al. (2020b)	<i>G. bulloides</i>	Sediment trap	LA
5	Mg/Ca	Fischer et al. (2024)	<i>G. ruber</i>	Plankton tows / nets	LA
6	Mg/Ca	Fehrenbacher et al. (2020)	<i>N. dutertrei</i>	Core	LA
7	Mg/Ca	Fehrenbacher et al. 2020	<i>N. dutertrei</i>	Core	SOL
8	Mg/Ca	Fehrenbacher et al. (2020)	<i>O. universa</i>	Core	LA
9	Mg/Ca	Fehrenbacher et al. (2020)	<i>O. universa</i>	Core	SOL
10	Mg/Ca	Fehrenbacher et al. (2020)	<i>P. obliquiloculata</i>	Core	LA
11	Mg/Ca	Fehrenbacher et al. (2020)	<i>P. obliquiloculata</i>	Core	SOL
12	Mg/Ca	Fehrenbacher et al. (2020)	<i>T. sacculifer</i>	Core	LA
13	Mg/Ca	Fehrenbacher et al. (2020)	<i>T. sacculifer</i>	Core	SOL
14	Mg/Ca	Balestra et al. (2021)	<i>O. universa</i>	Plankton tows / nets	EPMA

15	Mg/Ca	Balestra et al. (2022)	<i>O. universa</i>	Plankton tows / nets	EPMA
16	Mg/Ca	Kearns et al. (2023)	<i>G. ruber</i>	Core	LA
17	Mg/Ca	Cleroux et al. (2008)	<i>G. ruber</i>	Core	SOL
18	Mg/Ca	Elderfield et al. (2002)	<i>G. ruber</i>	Core	SOL
19	Mg/Ca	Dissard et al. (2021)	<i>T. sacculifer</i>	Plankton tows / nets	LA
20	Mg/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
21	Mg/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA
22	Sr/Ca	Osborne et al. (2020)	<i>G. Bulloides</i>	Sediment trap	LA
23	Sr/Ca	Fehrenbacher et al. (2018a)	<i>N. dutertrei</i>	Plankton tows / nets	LA
24	Sr/Ca	Osborne et al. (2020b)	<i>G. bulloides</i>	Sediment trap	LA
25	Sr/Ca	Fehrenbacher et al. (2020)	<i>N. dutertrei</i>	Core	LA
26	Sr/Ca	Fehrenbacher et al. (2020)	<i>N. dutertrei</i>	Core	SOL
27	Sr/Ca	Fehrenbacher et al. (2020)	<i>O. universa</i>	Core	LA
28	Sr/Ca	Fehrenbacher et al. (2020)	<i>O. universa</i>	Core	SOL
29	Sr/Ca	Fehrenbacher et al. (2020)	<i>P. obliquiloculata</i>	Core	LA
30	Sr/Ca	Fehrenbacher et al. (2020)	<i>P. obliquiloculata</i>	Core	SOL
31	Sr/Ca	Fehrenbacher et al. (2020)	<i>T. sacculifer</i>	Core	LA

32	Sr/Ca	Fehrenbacher et al. (2020)	<i>T. sacculifer</i>	Core	SOL
33	Sr/Ca	Kearns et al. (2023)	<i>G. ruber</i>	Core	LA
34	Sr/Ca	Cleroux et al. (2008)	<i>G. ruber</i>	Core	SOL
35	Sr/Ca	Elderfield et al. (2002)	<i>G. ruber</i>	Core	SOL
36	Sr/Ca	Dissard et al. (2021)	<i>T. sacculifer</i>	Plankton tows / nets	LA
37	Sr/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
38	Sr/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA
39	Al/Ca	Osborne et al. (2020)	<i>G. Bulloides</i>	Sediment trap	LA
40	Al/Ca	Osborne et al. (2020b)	<i>G. bulloides</i>	Sediment trap	LA
41	Al/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
42	Al/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA
43	Na/Ca	Gray et al. (2023)	<i>G. ruber</i>	Core	SOL
44	Na/Ca	Gray et al. (2023)	<i>G. ruber</i>	Cultured	SOL
45	Na/Ca	Gray et al. (2023)	<i>G. ruber</i>	Plankton tows / nets	LA
46	Na/Ca	Gray et al. (2023)	<i>G. ruber</i>	Plankton tows / nets	SOL
47	Na/Ca	Gray et al. (2023)	<i>G. ruber</i>	Sediment trap	LA
48	Na/Ca	Gray et al. (2023)	<i>G. ruber</i>	Sediment trap	SOL
49	Na/Ca	Gray et al. (2023)	<i>G. ruber</i> mixed	Core	SOL
50	Na/Ca	Gray et al. (2023)	<i>G. ruber sl</i>	Core	SOL

51	Na/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
52	Na/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA
53	B/Ca	Osborne et al. (2020)	<i>G. Bulloides</i>	Sediment trap	LA
54	B/Ca	Osborne et al. (2020b)	<i>G. Bulloides</i>	Sediment trap	LA
55	B/Ca	Henehan et al. (2015)	<i>G. ruber</i>	Core	SOL
56	B/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
57	B/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA
58	Mn/Ca	Kearns et al. (2023)	<i>G. ruber</i>	Core	LA
59	Mn/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
60	Mn/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA
61	Ba/Ca	Fehrenbacher et al. (2018a)	<i>N. dutertrei</i>	Cultured	LA
62	Ba/Ca	Fehrenbacher et al. (2018a)	<i>N. dutertrei</i>	Plankton tows / nets	LA
63	Ba/Ca	Kearns et al. (2023)	<i>G. ruber</i>	Core	LA
64	Ba/Ca	Hönisch et al. (2011)	<i>G. bulloides</i>	Cultured	SOL
65	Ba/Ca	Hönisch et al. (2011)	<i>O. universa</i>	Cultured	SOL
66	Ba/Ca	This study	<i>G. ruber</i>	Sediment trap	LA
67	Ba/Ca	This study	<i>T. clarkei</i>	Sediment trap	LA

739

740 Table 1: detailed description of the different species, measurement methods and sample
 741 origin used for the compilation in figure 12. LA stands for Laser Ablation (LA)-ICP-MS,
 742 SOL is solution-ICP-MS and EPMA is Electron micro-probe analyses.

743

744 6. Summary and conclusions:

745 We investigated the effects of inter-chamber variability on the proxy systematics in the hyper
746 saline oligotrophic GOA using single chamber LA ICP-MS analysis measured on two flux-
747 dominating planktic foraminifer (PF) species *G. ruber albus* and *T. clarkei* with its two
748 phenotypes ‘big’ and ‘encrusted’. We observed how element/Ca varies in PF chambers as a
749 function of environmental changes in order to then be used as proxies for past oceanic and
750 climatic reconstruction. The results show that some element/Ca exhibit temporal and seasonal
751 variations related to environmental conditions in the water column such as Mg/Ca in *G. ruber*
752 *albus* as a temperature proxy, and B/Ca in *T. clarkei* as a proxy of pH. Although other
753 element/Ca values display more limited variability (e.g., Na/Ca) they may still be of use as
754 paleo-proxies when combined in global calibration studies.

755 Water column mixing has been shown to have a significant effect of element/Ca positive
756 excursions in the analyzed *G. ruber albus*, and two *T. clarkei* morphotypes, which may limit
757 the use of some element ratios as proxies, or alternatively, be used as a proxy for water column
758 mixing. Generally, pooled-mean values of element/Ca in the PF tests in the GOA are species-
759 specific and element-specific, and are elevated compared to other regions (e.g., Mg/Ca, Al/Ca,
760 Na/Ca). However, the final chamber F0 is different in comparison to the preceding chambers
761 F-1 and F-2, suggesting that the element composition of F0 may be biased and unreliable in
762 terms of recording environmental conditions.

763 Our findings indicate that high-resolution analytical techniques, such as LA ICP-MS
764 enable studying single chamber compositions and variations. Although pooled mean values of
765 specimens over various water depths are recommended for their incorporation as proxies, ICV
766 can also be used as a tracer of environmental factors. Exploring different biochemical or
767 physiological mechanisms which are responsible for the element/Ca variations between species
768 and chambers are critical to shed light on how element/Ca are incorporated to the PF shells.
769 Despite these limitations, the results provide valuable insights into the complex behavior of
770 element/Ca in PF shells.

771 Data availability

772 Tabular supplementary data generated in this study can be found in the supplementary
773 material.

774

775 Author contributions

776 NL, AT, and RS designed the study; NL, BS, UW, and KPJ, performed the measurements;
777 NL, NC, AT, and RS analyzed the data; NL, RS and AT wrote the manuscript draft; NL, RS,
778 AT and GH reviewed and edited the manuscript.

779

780 The authors declare that they have no conflict of interest.

781

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