

**Bromine soil/sediment enrichment in tidal salt marshes as a potential indicator of
climate changes driven by solar activity: new insights from W coast Portuguese
estuaries**

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Abstract

This paper aims at providing insight about bromine (Br) cycle in four Portuguese estuaries: Minho, Lima (in the NW coast) and Sado, Mira (in the SW coast). The focus is on their tidal marsh environments, quite distinct with regard to key biophysicochemical attributes. Regardless of the primary bromide (Br^-) common natural source, i.e. seawater, the NW marshes present relatively higher surface soil/sediment Br concentrations than the ones from SW coast. This happens in close relation with organic matter (OM) content, and is controlled by their main climatic context. Yet, the anthropogenic impact on Br concentrations cannot be discarded. Regarding [Br] spatial patterns across the marshes, the results show a general increase from tidal flat towards high marsh. Maxima [Br] occur in the upper driftline zone, at transition from highest low marsh to high marsh, recognized as a privileged setting for OM accumulation. Based on the discovery of OM ubiquitous bromination in marine and transitional environments, it is assumed that this Br occurs mainly as organobromine. Analysis of two dated sediment cores indicates that, despite having the same age (AD 1300), the Caminha salt marsh (Minho estuary) evidences higher Br enrichment than the Casa Branca salt marsh (Mira estuary). This is related to a greater Br storage ability, which is linked to OM build-up and rate dynamics under different climate scenarios. Both cores evidence a fairly similar temporal Br enrichment pattern, and may be interpreted in light of the sun–climate coupling. Thereby, most of the well-known Grand Solar Minima during the Little Ice Age appear to have left an imprint on these marshes, supported by higher [Br] in soils/sediments. Besides climate changes driven by solar activity and impacting marsh Br biogeodynamics, those [Br] positive peaks might also reflect inputs of enhanced volcanic activity covarying with Grand Solar Minima.

47 **Keywords:** Salt marshes; Br cycle; OM storage; Grand Solar Minima; Climate modelling;
48 Climate variability.

49

50 **1. Introduction**

51 Wetlands play an important role on the biogeochemical cycle of elements such as carbon,
52 nitrogen, phosphorus, sulphur and mercury at local, regional and even global scales (e.g.,
53 Marques et al., 2011; Neubauer et al., 2013). A considerable amount of research has
54 revealed that this is also true for bromine (Br) (Varner et al., 1999; Keppler et al., 2000;
55 Rhew et al., 2000, 2002, 2014; Dimmer et al., 2000; Drewer et al., 2006; Manley et al.,
56 2006; Hardacre et al., 2009, 2013; Blei et al., 2010; Martinez-Cortizas et al., 2007, 2016).
57 Specifically, coastal wetlands, in which tidal marshes are included, represent important
58 land-ocean-atmosphere interfaces that allow to capture spatiotemporal variability in
59 chemical fluxes. In these habitats, Br mainly supplied by seawater interacts with both
60 halophytes and the relatively large pool of soil/sediment organic matter (OM). This
61 connection occurs through, although still poorly understood, bromination processes that
62 contribute to the production of organobromine compounds, which have detrimental effects
63 on the atmosphere. For instance, salt marshes have been identified as globally significant
64 natural sources of methyl bromide (CH₃Br) (Rhew et al., 2014, and references therein), a
65 reactive trace gas contributing to ozone loss processes in the stratosphere (e.g.,
66 Chipperfield, 2015). On the other hand, significant widespread bromination of natural OM
67 may significantly impact the preservation and/or degradation of organic carbon (C_{org}) in
68 soils/sediments (Leri and Myneni, 2012), therefore affecting the recognized salt marsh
69 ecosystem's role on climate and carbon sequestration. It is also known that in coastal (and
70 open ocean) areas, bromine-radical chemistry provides alternative reaction pathways in the
71 marine atmospheric boundary layer for (i) sulphur cycling, with associated implications for

72 aerosol production (and growth), radiative heat transfer and climate (Keene et al., 2007),
73 and (ii) mercury (Hg) cycling (Obrist et al., 2011; Tas et al., 2012), with Br-induced mercury
74 oxidation as a likely important Hg source to world's oceans, which can contribute to human
75 mercury exposure by seafood consumption (Sunderland, 2007). Taking together the
76 previous findings about the Br influence on other key element biogeochemical cycles in the
77 marine domain, and the knowledge that natural CH₃Br emissions are contributing to
78 increase the stratospheric reactive Br budget (e.g., Carpenter et al., 2014), establishing the
79 foundations of the Br biogeochemical cycle in coastal areas and tidal marsh habitats has
80 gained a renewed significance.

81 Traditionally, Br has been used in conjunction with chlorine (Cl) as a geochemical proxy for
82 seawater intrusion in coastal areas (e.g., Jones et al., 1999; Alcalá and Custodio, 2008),
83 and alone as a paleosalinity indicator and a stratigraphic marker in brine cores (Adams,
84 1969; Ziegler et al., 2008). An alternative interpretation regarding Br concentrations and
85 fluxes has been proposed by Moreno et al. (2015) when studying the sedimentary record
86 recovered from the high marsh zone on a tidal salt marsh located in the NW coast of
87 Portugal (Caminha, in the Minho River estuary). They suggested that the most prominent Br
88 enrichment peaks between AD 1300 (tidal marsh settlement) and AD 1800 (considered as
89 the beginning of industrialization) were primarily driven by a series of biogeochemical
90 processes rather than an indication of seawater intrusion events in the Minho estuary.
91 Those processes responded to significantly prolonged environmental (e.g., temperature and
92 precipitation) shifts triggered by Grand Minima Episodes of solar activity (SA) during the
93 LIA, namely the Wolf and the Maunder Minima, as well as the Dalton Minimum. These
94 episodes corresponded to periods of solar minimal energy output, as demonstrated by long-
95 term records of SA proxies (e.g. Usoskin et al., 2007), affecting Earth's climate. Prolonged
96 changes in environmental conditions can lead to significant responses at all levels of
97 ecosystem organization, generating persistent alterations in its biogeochemical functioning

98 (e.g., Keller et al., 2006; Neubauer et al., 2013). Accordingly, and based on Moreno et al.
99 (2015) that linked Br biogeodynamics to past SA, the impact of the sun–climate coupling at
100 Grand Solar Minima resulted in cascading effects on Br cycling in the Caminha salt marsh,
101 in parallel with rate changes in OM bromination, which ultimately weakened the marsh’s role
102 as a source for CH₃Br.

103 Following the study conducted in Caminha, the current contribution expands and
104 generalises the analysis, including three other Portuguese estuaries (Figure 1): the Lima
105 estuary, also in the NW coast, and the Sado and Mira estuaries, both located in the SW
106 coast. Broadly, the Portuguese W coast is typically characterized by an Atlantic climate,
107 though two main climatic regions can be distinguished. Compared to the NW coast, climate
108 in SW coast has drier summers, lower annual precipitation as well as higher annual
109 temperatures and insolation. Therefore, the aim of this work is twofold: (i) acquire a wider
110 latitudinal range of Br measurements in waters (superficial and interstitial) and marsh
111 soils/sediments in order to infer trends associated to the bio-geomorphological settings and
112 climatic variability, and (ii) improve the understanding of bromine-climate relationships
113 driven by SA, also providing new independent data and source insight to the still ongoing
114 debate about the “missing source” for the CH₃Br global budget (Yvon-Lewis et al., 2009).

115

116 **2. Regional setting**

117 ***NW coast***

118 Taken together, the Minho and Lima watersheds cover an area of 19 550 km² under an
119 Atlantic wet climate, with relatively high exposition to maritime winds, high mean annual
120 precipitations (Minho: 1200–2400 mm; Lima: 1300–4200 mm), mild summers (summer

121 mean temperatures from ca. 18–22°C) and relatively low mean annual insolation (2200–
122 2500 hours) (APA, 2011).

123 The Minho estuary (23 km²) is oriented NE–SW and presents a semidiurnal, high-mesotidal
124 regime in which vertical stratification occurs during periods of large freshwater discharge.
125 The mean annual freshwater inflow is around 300 m³/s (Ferreira et al., 2003). The highest
126 high water spring (HHWS) is 4 m height, but this is often amplified by storm surges
127 (Taborda and Dias, 1991), which we observed during field work. The mean tidal range is of
128 about 2.0 m. The upstream limit of the tidal salt wedge in the Minho River is 9 km (Fatela et
129 al., 2009). Large tidal flat and tidal marsh surfaces with approximately 6 km² occur in the
130 Minho estuary's banks, with its largest expansion along the left bank – Caminha tidal marsh
131 (ca. 2.5 km²) – at the confluence with Coura tributary (Figure 1). Recently, Reis et al. (2014)
132 updated the estuary ecological quality to “moderately to remarkably polluted” based on
133 metal concentrations guidelines (SFT TA-1467/1997). Furthermore, the use of ethylene
134 dibromide in leaded gasoline and the vehicle emissions since 1930, a shared Br and Pb
135 anthropogenic source, appears to have had a significant impact on the Caminha salt marsh
136 (Moreno et al., 2015).

137 The Lima estuary (5 km²), located 20 km south of Minho, is oriented ENE–WSW and it is a
138 semidiurnal mesotidal estuary, with a mean tidal range of about 2.5 m and a HHWS of 3.7
139 m (Vale and Dias, 2011). The mean annual freshwater inflow is 50 m³/s (Vale and Dias,
140 2011). Here the tidal salt wedge effect is noticed to 3–5 km upstream in winter and no more
141 than 15 km in summer (Alves, 2003). Intertidal areas extend over more than 2 km² on the
142 banks of the Lima estuary, including the Nossa Senhora do Rosário salt marsh (NSR;
143 Figure 1). The Lima lower estuary's ecological status is considered moderate (Costa-Dias et
144 al., 2010). This results from the significant impact on the estuary of the harbour activities,
145 leading to continuous petrochemical contamination (Lima et al., 2007), and diffuse pollution

146 from agriculture, domestic, and industrial waste discharges, including a paper mill (Costa-
147 Dias et al., 2010).

148 The salt marshes of these two NW sites are classified as Eurosiberian, based on their plant
149 communities (Costa et al., 2009), with abundant reed meadows where the presence of
150 *Juncus maritimus* Lam. (C3 plant; non-succulent; perennial) is ubiquitous (Honrado et al.,
151 2004; Almeida et al., 2011). Although, other plant species can also be found such as the
152 non-native *Triglochin striatum* Ruiz & Pav. (C3 plant; succulent; perennial), the invasive
153 *Phragmites australis* (Cav.) Trin ex. Steud. (C3 plant) and the weed *Spartina patens* (Aiton)
154 Muhl (C4 plant; non-succulent; perennial) (Almeida et al., 2011).

155

156 **SW coast**

157 The SW coast of Portugal, where the Sado and Mira estuaries are located, is under a sub-
158 wet Mediterranean climate, with mean values of annual precipitation around 600–700 mm
159 (Bettencourt et al., 2003). Mean air surface temperatures are near 23°C in the hottest
160 months (July and August), with a yearly average number of sunshine hours ranging 2900–
161 3000 (APA, 2011), also showing almost permanent maritime moist winds (APA, 2011). The
162 size of Sado and Mira rivers watersheds is 6 700 km² and 1 576 km², respectively.

163 The Sado estuary (170 km²), the second largest estuarine system in Portugal, is located
164 about 40 km south of Lisbon (Figure 1). It is a well-mixed estuary under normal river flow
165 conditions, however, high discharge in some winter months may cause moderate
166 stratification locally (Ferreira et al., 2003). It has a complex morphology generally oriented
167 NW–SE and presents wide tidal flats as well as narrow and discontinuous coastal salt
168 marshes covering around 7.2 km² (Moreira, 1992). The tidal pattern is semi-diurnal, with a
169 mean tidal range of about 2.7 m and a HHWS of 3.2 m (Martins et al., 2001). The maximum

170 salt-wedge limit is 70 km upstream. The mean annual freshwater input is ca. 40 m³/s,
171 exhibiting large interannual fluctuations. The lower estuary behaves as a coastal lagoon,
172 while the upper reaches present a greater fluvial influence (Martins et al., 2001). In general,
173 the Sado estuary can be classified as moderately contaminated, but the lower estuary and
174 some segments near industrial areas have revealed levels of concern for several
175 contaminants both organic and inorganic, with adverse toxicological effects to biota (e.g.,
176 Neuparth et al., 2005).

177 Finally, the vertically well-mixed Mira estuary (4.5 km²) is a narrow incised estuary oriented
178 NE–SW (e.g., Paula et al., 2006). It presents a semi-diurnal mesotidal regime with a mean
179 tidal range of about 2.4 m and a HHWS of 3.5 m (Amaral et al., 2007). The salt edge may
180 reach 32 km from the river mouth (Bettencourt et al., 2003). The lower section of the estuary
181 has a dominant marine influence due to low, seasonal and limited freshwater input by the
182 Mira River (2.9 m³/s). This characteristic has allowed the development in the lower 8 km of
183 large, intertidal, and homogenous seagrass meadows of *Zostera noltii* Hornemann, 1832
184 (e.g., Cunha et al., 2013). The estuarine area is also characterized by bare sandy areas and
185 muddy substrates, with a 2.9 km² area of fringing salt marshes occurring as far as 15–20
186 km upstream (Costa et al., 2001). These salt marshes have remained nearly unchanged
187 since 1958, with the entire ecosystem relatively undisturbed by anthropogenic activities
188 (Castro and Freitas, 2006).

189 As oppose to the NW sites, the SW salt marshes studied here belong to the Biogeographic
190 Mediterranean region (Costa et al., 2009). The halophytic community is mixed including
191 perennial succulent species such as *Halimione portulacoides* (L.) Aellen (C3 plant),
192 *Sarcocornia fruticosa* (L.) A.J. Scott (C3 plant), *Sarcocornia perennis* (Mill.) A.J. Scott (C3
193 plant) in the high marsh and the annuals *Spartina maritima* (Curtis) Fernald (C4 plant) and
194 *Salicornia fragilis* P. W. Ball & Tutin (C3 plant) in the low marsh (e.g., Costa, 2001).

195

196 **2. Materials and Methods**

197 **2.1. Water and sediment samples**

198 The methodologies used for sampling and analysis of water (superficial and interstitial) and
199 sediments (surface and cored) in the Lima, Sado and Mira estuaries follow Moreno et al.
200 (2015) and are fully described therein. Figure 1 and Tables 1 and 2 summarize the new
201 samples analysed here: eight interstitial water samples from three salt marsh transects
202 (NSR_L, TRO_S, PMF_M), ninety-one sediment surface samples from the intertidal domain
203 (tidal flat, devoid of vascular plants; low marsh and high marsh zones with typical halophytic
204 vegetation), along ten cross-shore transects, as well as an one-metre-long sediment core
205 (hereafter FWCB_r) recovered with a manual Auger sampler from the Casa Branca salt
206 marsh (1.74 m above mean sea level; 37°40'03.7" N and 8°43'12.7" W), located on the Mira
207 River estuary. A total of thirty sliced (1 cm thick) samples were analysed for Br and OM
208 contents. The FCPw1 core from Moreno et al. (2015), located in the Caminha high marsh
209 zone (1.55 m above mean sea level; 41°52'37.0" N and 8°49'28.0" W), is also indicated in
210 Figure 1.

211 In order to characterize the two possible end-members (fluvial and marine) of
212 biogeochemical sources to salt marshes, four marine seawater and four fluvial freshwater
213 samples were collected (Table 1).

214 Water sample analyses: The filtrate was analysed for bromide (Br⁻), amongst other anions,
215 by ion chromatography (IC) with suppressed conductivity detection (761 Compact IC
216 Metrohm), and raw data processed with Metrohm Metro data 1.1. The IC method no. S-73,
217 developed by Metrohm to determine anions in seawater, was used for the most saline
218 waters (see Valente et al., 2009 and Moreno et al., 2015 for detailed information). A set of

standards was prepared to make a 6-point calibration curve covering the range of Br⁻ (and Cl⁻) concentrations in water samples. The IC method not only allows an efficient separation of the Br⁻ and Cl⁻ peaks, but has also the advantage of measuring both anions in the same sample preventing the errors introduced by dilution. A standard (20 mg/L) was run independently of the calibration curve to check for accuracy (every two samples) and sample replicates were run to check for precision. The precision was within the relative standard deviation (RSD) of 5% for all determinations and results were accurate within precision.

Sediment sample analysis: Br concentrations in cored samples, along with the collected surface sediments, were determined by Energy-Dispersive X-Ray Fluorescence Spectrometry (EDXRF), using a KEVEX 771 spectrometer. To calibrate the spectrometer and verify the accuracy and precision of the overall procedure three certified reference materials were analysed: SGR1 (Green River Shale from the United States Geological Survey – USGS), SRM 2704 (Buffalo River Sediment) and SRM1646 (Estuarine Sediment), both from the National Institute for Standards and Technology (NIST). A complete description of the equipment, analytical conditions and spectral evaluation, along with the calibration and quantification techniques, is available in Araújo et al. (1998, 2003). Accuracy and precision on the Br determinations are better than 10% as previously fully described in Moreno et al. (2015). The OM content was determined as Loss-on-Ignition (LOI), with an aliquot of bulk sediment sample (2.0 g) dried and oven-heated at a temperature of 500 °C ± 50 °C for about 2 hours (Moreira et al., 2009). Quality control was checked by replicate analysis (40% of the total), with errors lying in the interval 0.1%–15.0% (average: 5.7%) of the measured value. In an attempt to test the reliability of LOI data for the estimation of C_{org} content, a regression analysis was performed for LOI vs. C_{org} for the FCPw1 core, with C_{org} data taken from de la Rosa et al. (2012). A strong statistically significant correlation ($r = 0.97$,

244 N= 19; $p < 0.001$) between LOI and C_{org} was achieved, ensuring that LOI results are
245 reflecting mostly C_{org} (%OM).

246 In order to characterize OM quality, i.e. the percentages of labile and recalcitrant OM, the
247 stepwise thermogravimetric procedure (STG) of Kristensen (1990) was applied to the cored
248 samples from the Casa Branca salt marsh. According to this method, these OM fractions
249 are defined as the percentage weight losses after ignition at 280°C and 520°C, respectively.
250 In short, samples of 0.5 g were grounded and pre-dried at 105°C for 6h. After cooling in a
251 desiccator, the sample weight was determined with a precision of 0.1 mg. Next, the samples
252 were combusted at precisely 280°C for 6h in a computer controlled Heraeus MR 170 muffle
253 furnace. After cooling in a desiccator and re-weighting, the samples returned to the muffle
254 furnace and combusted at 520°C for 6h. After cooling in a desiccator the final ash weight
255 was determined (Kristensen, 1990).

256

257 **2.2. Solar activity, temperature, and precipitation climatic modelled data**

258 Cosmogenic radionuclides are produced in the atmosphere through a nuclear cascade
259 mainly triggered by the high-energy galactic cosmic rays (GCR; Lal and Peters, 1967). As
260 GCR enter the heliosphere, they are subject to modulation processes due to variable solar
261 magnetic activity. This is the reason why during phases of low SA much higher particle
262 intensities occur inside the heliosphere than during solar maximum conditions (e.g., Herbst
263 et al., 2015; Adolphi and Muscheler, 2016). The two most noticeable cosmogenic
264 radionuclides suitable for reconstructing SA are ^{14}C and ^{10}Be . The production rate of both
265 isotopes reacts in a very similar way to changes in solar and geomagnetic shielding
266 (Masarik and Beer, 1999). The reconstructed dataset chosen for this work is the total solar
267 irradiance (TSI), considered as a proxy for SA, from Steinhilber et al. (2012). This

268 reconstruction is based on time series of ^{14}C stored in tree rings and of ^{10}Be extracted from
269 polar ice cores, and was downloaded from the NOAA web page (<http://www.noaa.gov/>).

270 Aiming to compare the Br and OM records with climate variables, this study also
271 incorporates the series of temperature and precipitation evolution in both study areas as
272 predicted by a high-resolution regional climate model (Gómez-Navarro et al., 2011). The
273 simulation implements a domain that encompasses the whole Iberian Peninsula (IP) and
274 spans the second millennium entirely. It was carried out with a climate version of the
275 mesoscale model MM5, and driven at the boundaries by a simulation with the global model
276 ECHO-G (see Gómez-Navarro et al., 2011 for details). Following Gómez-Navarro et al.
277 (2011), the use of a high resolution model aims to reduce the scale gap between the large-
278 scale correctly simulated by the GCM and the features of regional variability present in the
279 Br and OM records presented here. The model simulates coherently the evolution of most
280 relevant climate variables, and in particular reproduces the physically constrained co-
281 evolution of temperature and precipitation, as well as their relation with large-scale
282 dynamics (e.g., the North Atlantic Oscillation – NAO). It is jointly driven by reconstructions of
283 the variability of three external forcings: TSI, greenhouse gas concentrations and the effect
284 of volcanic activity.

285

286 **2.3. Chronology**

287 The geochronology of the top 15 cm of the FWCBBr core was calculated from the ^{210}Pb
288 profile using the constant rate of supply method (CRS) (Appleby and Oldfield, 1978)
289 supported by ^{137}Cs . Samples for ^{210}Pb and ^{137}Cs were analyzed following the methodology
290 described by Appleby (2001) at the University of Plymouth (UK) Consolidated Radioisotope
291 Facility, using an EG&G Ortec planar (GEM-FX8530-S N-type) HPGe gamma spectrometry

292 system built to ultra-low background specification for ^{210}Pb detection. Additional
293 information regarding the technique is provided in Appendix A. This core presented an
294 unsupported ^{210}Pb ($^{210}\text{Pb}_{\text{xs}}$) profile that suggested some changes in the sediment
295 accumulation rate in the upper section (ca. 150 years), although they could also reflect
296 sediment mixing or disruption of the sedimentation. While the available elemental data is
297 limited and presents generally low concentrations, the Pb profile shows slightly higher
298 values above 18 cm that could be coincident with the initial stages of the industrial
299 revolution (unclear date for this region but ca. AD 1800), which would be in agreement with
300 the CRS model used here. However, the inflexion indicated by the model for the two older
301 samples should be considered carefully (Leorri et al., 2010). In order to extend the
302 chronology down-core, two samples (69–70 cm, 90–91 cm depth) of total organic carbon
303 (TOC) were carbon-14 dated by accelerator mass spectrometry–AMS at Beta Analytic Inc.
304 (USA). The chronology for the FWCB core was created using a Bayesian age-depth model
305 (Bchron 4.1; Haslett and Parnell, 2008; Parnell et al., 2008) (Appendix A). The model
306 provides ages with an individual error for each sample averaging 73 years for a 95%
307 confidence interval. The obtained calendar ages are presented in years of *Anno Domini*
308 (years AD).

309 The chronology of the FCPw1 core can be found in Moreno et al. (2015), but also relies on
310 the combination of ^{210}Pb and carbon-14 data from TOC.

311

312 **3. Results and discussion**

313 **3.1. Br^- in surface and interstitial water samples**

314 The results of Br^- concentrations in surface and interstitial waters are presented in Table 1.
315 Freshwater samples show Br^- contents between less than 0.01 mg/L (Minho River) and a

316 maximum of 0.8 mg/L (Sado River). On the other hand, marine surface water samples
317 throughout the W coast have Br^- contents ranging from 185 to 197 mg/L (Table 1). Such
318 high values, outside the typical marine $[\text{Br}^-]$ range: 60–80 mg/L, are somewhat expected
319 and connected or linked to the known supersaturation in brominated organic compounds in
320 the Portuguese offshore (Raimund et al., 2011). This is related to the presence of strong
321 macroalgal sources and to the Iberian Peninsula coastal upwelling, as discussed previously
322 by Moreno et al. (2015). Brominated organic compounds (also including CH_3Br) are
323 produced and degraded at relatively fast rates in the coastal ocean, with their degradation
324 mechanisms (e.g., hydrolysis and chloride exchange reactions) in the water column as
325 major suppliers of bromide anions to seawater. Our hypothesis – linking the Br^- enrichment
326 of western Portuguese superficial coastal waters to loss reactions of brominated organic
327 compounds in seawater column – may be supported by other authors' findings. Namely, Hu
328 et al. (2010) discovered for the east coast of United States evidence of a vertical distribution
329 in the CH_3Br saturation anomalies, with highest concentrations in the subsurface seawater
330 below the mixed layer, due to high degradation rates near the surface. An analogous
331 subsurface seawater enhancement in depth profiles of two of the most important short-lived
332 carriers of atmospheric Br, i.e. dibromomethane (CH_2Br_2) and bromoform (CHBr_3), was
333 described by Raimund et al. (2011) when sampling the Iberian Peninsula upwelling system
334 off the coast of Portugal.

335 The detected Br^- enrichment in the nearshore surface water samples also implies that
336 marine water would have Cl^-/Br^- mass ratios lower than the reported value for the average
337 seawater (typically 290 ± 4 ; e.g., Katz et al., 2011). This chemical signature in our water
338 samples is presented in the plot of Cl^-/Br^- mass ratios versus Cl^- in Figure 2, along with a
339 typical marine water sample (TA_SW: $[\text{Cl}^-] = 19,353 \text{ mg/L}$ and $[\text{Br}^-] = 67 \text{ mg/L}$; Millero,
340 2013). The clusters displayed by the Cl^-/Br^- vs. Cl^- plot allow to identify freshwater (with
341 consistently lower levels of Cl^- and Br^-) from marine and brackish waters reflecting an

342 increase in Cl^- (Figure 2A). In addition, it is possible to differentiate between the interstitial
343 water samples of the NW coast salt marshes (Minho and Lima) and the SW coast samples
344 (Sado and Mira) (Figure 2B). This suggests that beyond the inferred common main source
345 of Br^- (and Cl^-), i.e., seawater (theoretical ranges after Panno et al., 2006), a clear N–S
346 differentiation can be established based on the relationship between the chemical indicators
347 chloride and bromide. As shown in Figure 2B, the Sado and Mira interstitial waters, ranging
348 from polyhaline (18–30‰) to euhaline (30–40‰), can be considered closer, based on these
349 anions, to marine water samples (all clustering together; Figure 2B) than the ones from the
350 Caminha and Lima salt marshes (mostly mesohaline: 5–18‰). These results are
351 symptomatic of a stronger mixing with freshwater, originated by higher inputs of rainfall-land
352 runoff production to the NW coast salt marshes, leading to salt dilution and lower ionic
353 concentrations.

354 At this point, it must be emphasized that a rising body of evidence suggests that Br^- does
355 not act conservatively in soils or water. Br^- can be (re)actively involved in OM cycling (e.g.,
356 Gerritse and George, 1988; Mahn and Gieskes, 2001; Biester et al., 2004, 2006; Leri et al.,
357 2010, 2014), and it is frequently related to Fe and Mn cycling (e.g., Leri et al. 2010). This
358 could diminish the “sensitivity” of the ratio Cl^-/Br^- , and limits its applicability as an inorganic
359 tracer of marine intrusion. However, plotting Cl^-/Br^- ratios provides insight into the Br^- (and
360 Cl^-) major sources in the sampled salt marshes. Panno et al. (2006) proposed Cl^-/Br^-
361 fingerprinting as a valuable diagnostic method in the identification of anthropogenic sources
362 of salinization; though additional analysis should be required to complement this technique,
363 and thus determine more accurately the source of the contaminant present in water
364 samples. For instance, Figure 2B depicts a group of samples that evidence an
365 anthropogenic impact based on this methodology – NSR_P1 (tidal flat; Lima), TRO_P7
366 (high marsh zone; Sado) and PMF_P6 (high marsh zone; Mira). The three fit in the “basin
367 brines and animal waste” water-type (Panno et al., 2006). In addition, the sample TRO_P7

368 presents a measurable $[\text{NO}_3^-]$ of 54 mg/L, by opposition to the other two, both with $[\text{NO}_3^-] <$
369 0.01 mg/L (results not shown). Historical Br anthropogenic sources such as emissions from
370 an antiknock additive in leaded gasoline, flame retardants, dyes, pharmaceuticals or
371 pesticides in agriculture are well-known (e.g., Flury and Papritz 1993), and while some were
372 phased out, others are still in use (e.g., Shaw et al., 2010). As briefly pointed in Section 2,
373 the Lima, Sado and Mira estuaries have their specific histories of anthropogenic
374 disturbances, but the effort to better distinguish the likely detected Br^- contamination in
375 water samples (Figure 2B) is beyond the scope of the present study.

376

377 **3.2. Br–OM relationships in marsh surface environments**

378 The data gathered in the newly investigated salt marshes regarding interstitial waters (Table
379 1) are scarce due to several sampling and analytical constraints. This, together with the
380 presence of samples disturbed by anthropogenic activities frustrated our goal of broadening
381 the “evidence base” for natural $[\text{Br}^-]$ interstitial water patterns in salt marshes. It was earlier
382 detected in the Caminha salt marsh that a pattern appears to be mainly dictated by marine
383 influence (periodic tidal flooding), i.e., higher $[\text{Br}^-]$ in low marsh zone (average 71 mg/L;
384 Table 1), exposed to longer inundation periods by seawater and, consequently, prone to
385 greater inflow of bromide (Moreno et al., 2015). This is correlated with the relative
386 submersion times (annual basis) in the Caminha salt marsh, which is ca. 76–53% for tidal
387 flat, ca. 53–10% for low marsh and ca. 10–2% for the high marsh (Fatela et al., 2009).

388 Br^- entering salt marshes can be cycled by several biotic and abiotic OM bromination
389 mechanisms (e.g., Keppler et al., 2000, 2004; Hamilton, 2003; Saito and Yokouchi, 2006;
390 Wishkerman et al., 2008; Leri et al., 2010; Leri and Myneni, 2012; Leri and Ravel, 2015). All
391 include the production of an oxidized form of Br that reacts with electron-rich organic

392 molecules, with the subsequent formation of organobromine by-products (Leri et al., 2014).
393 Among them, it is the highly volatile gas CH_3Br (e.g., Wuosmaa and Hager, 1990; Hamilton
394 et al., 2003; Keppler et al., 2000, 2004; Saito and Yokouchi, 2006). Therefore, both the
395 halophytic vegetation cover and the soil/sediment organic fraction of tidal salt marshes are
396 the main settings and substrates for the conversion of inorganic Br (Br_{inorg}) into organic Br
397 (Br_{org}), which can largely contribute to their Br pool (Moreno et al., 2015). It also explains
398 why the total [Br] in (coastal) soils/sediments might not necessarily correlate with $[\text{Br}^-]$ in
399 water (Leri and Ravel, 2015). This lack of correlation is clear in the investigated salt
400 marshes, where some mismatch is observed between $[\text{Br}^-]$ in interstitial waters and the total
401 [Br] gradient in surface soils/sediments (Tables 1 and 2; Figure 3). The highest total Br
402 concentrations were found in the high marsh zones from Minho (average: 389 mg/kg) and
403 Mira (avg.: 233 mg/kg) estuaries, while in their respective low marshes these contents did
404 not exceed, on average, 133 mg/kg and 152 mg/kg (Table 2). Salt marshes from the Sado
405 are the most depleted in both Br and OM, with average (median) values of 53 mg/kg and
406 8.2% (low marsh), and 81 mg/kg and 11.7% (high marsh), respectively. In the Lima estuary,
407 Br trends are quite distinct (Table 2). This might be explained by the dominance of a coarser
408 soil/sediment fraction (mainly sands) in the samples from these salt marshes transects
409 (results not shown). Since a direct correlation has been well-established and widely
410 accepted between mud (clay and/or silt) grain size fractions and both Br (e.g., Correns,
411 1956; Vinogradov, 1959) and OM (e.g., Buchanan and Longbottom, 1970; Mayer, 1994)
412 contents, Br concentrations can be diluted in coarse-grained samples. The NSR_L transect
413 is additionally impacted by a dredged sand processing facility located in the marsh
414 surrounding area, causing resuspension of large quantities of sediment and disturbing this
415 salt marsh (Cardoso et al., 2008).

416 The spatial pattern across soil/sediment surface transects (Figure 3) presents a strong
417 direct correlation between Br and %OM (Lima: $r = 0.89$; $N = 28$; $p < 0.001$; Minho: $r = 0.86$;

418 N= 21; $p < 0.001$; Mira: $r = 0.79$; N= 29; $p < 0.001$, and Sado $r = 0.67$; N= 34; $p < 0.001$; tidal
419 flat samples are also included in the computed correlation coefficients). These results are in
420 good agreement with other studies (e.g., Cundy et al., 2005), suggesting that elevation is a
421 key factor controlling the soil/sediment C_{org} pools of tidal salt marshes, as suggested by
422 Spohn and Giani (2012, 2013). These authors not only have found higher C_{org} stocks in the
423 high marsh zones with limited flooding, but a major contribution of autochthonous (in situ)
424 OM inputs of the halophytic plant cover to soils and sediments at higher elevation within the
425 tidal frame. Such findings could likewise enlighten the preferential total Br enrichment in the
426 surface soils/sediments from the studied high marshes. Assuming that this Br is present
427 mainly as organobromine, the higher concentrations found in the high marsh zones,
428 typically characterized by greater density of stems and litter, can be associated to the
429 relatively fast oxidation of part of Br^- , which seems to lead to a rapid conversion to Br_{org}
430 (Leri and Myneni, 2012).

431 Similarly, the role of vegetation should be highlighted. The ubiquity of non-succulent
432 perennial species (*Juncus maritimus*) in the Caminha and Lima salt marshes contrasts with
433 the dominance of succulent species (e.g., from *Sarcocornia/Salicornia* genera) in Sado and
434 Mira salt marshes. Manley et al. (2006) and Blei et al. (2010) found higher relative Br
435 contents in succulent species, which (succulence) results from their strategy to survive in
436 saline soils by maintaining a large amount of tissue water. According to Manley et al.
437 (2006), high Br tissue levels would make of these succulent halophytes prolific CH_3Br
438 producers, although suggesting that each plant species has very different intrinsic abilities
439 to produce CH_3Br . Blei et al. (2010) showed that the variations in the Br content found on
440 salt marsh vegetation do not explain the spatial differences in CH_3Br flux magnitudes,
441 concluding instead that the limiting factor lies on the plants conversion mechanism (abiotic
442 and/or biotic). Wishkerman et al. (2008) reported that the abiotic reaction (occurring
443 between plant pectin and Br followed by CH_3Br emission) is strongly influenced by both air

444 temperatures, increasing by a factor of two for every 5°C increase, and plants succulence,
445 becoming more efficient as plants dry out. Rhew et al. (2014) estimated that only
446 approximately 0.17% Br in the leaf tissue of *Batis maritima* L. (known as one of the CH₃Br
447 greatest producers in salt marshes) is daily removed via CH₃Br emissions, indicating that to
448 impact Br availability, a small separated subset of “active Br” at the enzyme site would be
449 needed. According to them, if this *active* Br pool was 0.5% of the overall tissue content, then
450 CH₃Br emissions could reduce daily that pool by 34% for Br. This ongoing discussion is of
451 major importance as it holds the power for unbalancing the total [Br⁻] (either reducing or
452 increased it) available to ultimately be converted in Br_{org} in a given marsh. As a final remark
453 we would like to draw attention to the extensive intertidal habitat of *Zostera noltii* in Sado
454 and Mira estuaries (Cunha et al., 2013), and its possible impact on the Br estuarine cycling.
455 This seagrass along with *Zostera marina* Linnaeus 1753, which though very rare occurs as
456 well in the Mira estuary (Cunha et al., 2013), also have been identified as
457 producers/emitters of volatile Br compounds, such as CH₃Br and CHBr₃ (Weinberg et al.,
458 2013, 2015).

459 Interestingly, maximum Br soil/sediment enrichment in all studied salt marshes occurs in the
460 highest low marsh transitioning to high marsh, in the so-called upper driftline zone (Adam,
461 1990; Gerlach 1999; Persicke et al. 1999; Lefeuve et al., 2000; Gettner, 2003). These
462 areas represent a tidal-terrestrial/freshwater transition interface, where most drift litter
463 accumulates, usually containing a high concentration of seeds and vegetative material
464 (Mineke and Bakker 2002). While this litter is effectively taken out of the estuarine
465 circulation (e.g., Boorman, 2003), it becomes potentially accessible for promoting the
466 magnification of the local C_{org} pool of soils/sediments. As a result, and even if the
467 understanding of the internal marsh processes affecting OM accumulation and turnover is
468 limited (Fagherazzi et al., 2013) – identical to the mechanisms regulating Br⁻ fluxes –, it
469 seems plausible to consider the upper driftline zones as promising Br_{org} sink areas, from the

standpoint of the current knowledge about OM bromination. Driftline zones might also be natural laboratories for studying the short and longer-term impacts of counterbalancing controls, like temperature, moisture and inundation (e.g., Lewis et al., 2014) as well as priming (e.g., Gontikaki et al., 2013) on OM mineralization, distressing Br sequestration and its fate in coastal environments.

Finally, the identified Br reduction in the soils/sediments collected at the highest high marsh (Figure 3) can be attributed to the increased influence of adjacent terrestrial uplands, taking into account the principle that the terrestrial environment, and thus terrestrial OM, is relatively poor in bromine (Mayer et al., 2007).

3.2. Br temporal trends in SW (Mira estuary) and NW (Minho estuary) coasts

3.2.1. Comparing Br enrichment in relation to the long-term OM storage ability from salt marshes in their soils/sediments

Casa Branca salt marsh (FWCBr core; Mira estuary) down-core profiles of Br and OM up to 89 cm depth (AD 1190) are presented in Figure 4, along with the profiles previously obtained from the Caminha salt marsh up to 62 cm depth (AD 1143) (FCPw1 core; Fig. 3 of Moreno et al., 2015). Also shown in Figure 4 are the corresponding computed Br/OM ratio trends for both cores.

The FWCBr core presents Br concentrations in the range 129–560 mg/kg while the OM content varies between 4.9% and 23.6% (Appendix B), with values uniform from the base (AD 1190) until around AD 1920 (Br: 129–215 mg/kg; average 161 ± 5 mg/kg; OM: 4.9–8.3%; average $6.7 \pm 0.2\%$). Then, in the core's uppermost part, both Br and OM increase significantly, with two peaks at AD 1984 and AD 2010. This depth profile contrasts with the wider range and higher average Br content recorded in the FCPw1 core (Caminha):

average concentration of 747 mg/kg after the tidal marsh set up in AD 1330; minimum of 68 mg/kg, AD 1143 and maximum of 1300 mg/kg by AD 1700 (Moreno et al., 2015). It seems important to mention the concomitant tidal marsh build-up in both places, as indicated by the core's analysis of preserved benthic foraminifera associations: AD 1330 in Caminha (Moreno et al., 2014) and AD 1323 in Casa Branca (unpublished data). This event occurs during the transition from the Medieval Climatic Anomaly (MCA; 900–1300) to the LIA (1350–1900) and it is likely related to the main MCA–LIA shifts in local-to-regional hydroclimatic conditions in Iberian Peninsula (e.g., Lebreiro et al., 2006; Moreno et al., 2012).

The strong direct Br–OM correlation identified in surface marsh environments is preserved in the cored sediment samples with depth ($r = 0.91$; $N = 30$; $p < 0.001$), like previously described for the Caminha salt marsh ($r = 0.83$; $N = 49$; $p < 0.001$). This Br–OM correlation occurs independently of the large differences observed between the two cores regarding their Br and OM inventories (Figure 4). Indeed, and despite the evidence provided herein showing that the Mira estuary is subject to greater influence from Br^- enriched coastal waters than Minho, the FWCBBr core is relatively depleted in total Br (Appendix B). This depletion is also true for the amounts of long-term OM storage in both salt marshes soils/sediments. While in the FCPw1 core near 45% of the samples can be labelled as highly organic (OM > 30%) and *ca.* 41% as organic (organic content in the 15%–30% range), in the FWCBBr core almost 94% of the samples can be classified as mineral soils with organics (organic content > 3% and \leq to 15%) (Huang et al., 2009). Therefore, and consistent with the Br–OM relationship (leading to the production of organobromine compounds) found in the surface/modern marsh habitats, it can be hypothesized that the primary driver of the whole dissimilar Br pool size of these two coastal tidal marshes is their technical C_{org} sink capacity, constraining the amount of C_{org} that is sequestered in each. Generally, decomposition rates in salt marshes are lower than OM inputs (allochthonous

520 and autochthonous), the reason why they are recognized as one of the most powerful C_{org}
521 sinks on the planet (e.g., Macreadie et al., 2013). However, it is expected that at a regional
522 scale, soil/sediment C_{org} pools are dependent upon several decomposition rate modifiers
523 (e.g., litter chemical composition, climate, nutrient availability, communities of soil/sediment
524 organisms, and site-specific factors), creating diverse geographic patterns as regards C_{org}
525 sequestration. Among those controls, salinity seems to be a major factor, and it is
526 suggested that on tidal marshes soil C_{org} sequestration increases with decreasing salinity
527 (e.g., Poffenbarger et al., 2011; Van de Broek et al., 2016). Actually, salinity seems to have
528 an even stronger impact than elevation on the soil/sediment OM pools of tidal marshes,
529 inhibiting above-ground biomass and by enhancing OM mineralization (Hansen, 2015). In
530 the climatic context of the Mira estuary, the evapotranspiration rates usually exceed
531 precipitation, with tidal seawater supplying most of the moisture to the Casa Branca salt
532 marsh soils/sediments. This induces high salinity even on the high marsh (Table 1), with
533 Fatela et al. (2016) referring the occurrence of modern hypersaline conditions, with maxima
534 records of 48‰, in the Mira lower estuary. Foraminiferal evidence (that will be discussed
535 elsewhere) supports the idea that the (higher) Mira salinity baseline has been dominant
536 across the timeframe investigated, with assemblages dominated by *Jadammina*
537 *macrescens* (Brady, 1870) and *Trochammina inflata* (Montagu, 1808) (average 92%)
538 (unpublished data). Such high salinity baseline is, in turn, a possible explanation to the
539 lower OM in-depth concentrations from the FWCB_r profile, which agrees with results from
540 other studies (e.g., Van de Broek et al., 2016 and references therein).

541 Moreover, climate has a fundamental influence on the quantity (and quality) of inputs to the
542 soil OM pool, with C_{org} stocks being largest toward cooler and wetter locations, and smallest
543 at hotter and drier regions, as established by other studies of terrestrial ecosystems (e.g.,
544 Jenny, 1941; Meentemeyer, 1978; Liu et al., 2012, and references therein). These climatic
545 gradients can have left their signature on the temporal evolution of the C_{org} storage in the

546 Caminha and Casa Branca study sites. This is revealed by their individual temporal
547 soil/sediment OM patterns, developed in response to the long-term climatic gradient
548 between the NW and SW coasts of rising temperature and decreasing precipitation (as can
549 be observed in Figure 4), with plausible direct implications on Br longer-time-scale trend, as
550 explained before.

551 The strong direct Br–OM correlation holds for both the labile ($r = 0.92$, $N = 30$; $p < 0.001$) and
552 the relatively more recalcitrant OM ($r = 0.88$; $N = 30$; $p < 0.001$) fractions from the Casa
553 Branca salt marsh soils/sediments (results not shown). This result is consistent with recent
554 investigation, in which a series of model experiments allowed to establish the existence of a
555 natural, abiotic mechanistic source both of aliphatic (more labile) and aromatic (more
556 recalcitrant) forms of Br_{org} in plant debris and humic substances in soil environment (Leri et
557 al., 2014), and marine particulate OM (Leri and Ravel, 2015). The soil humic substances
558 showed a recalcitrant aromatic Br_{org} speciation, leading Leri et al. (2014) to suggest that this
559 might provide a useful proxy for evaluating the rate of OM burial in sediments. In this
560 direction, is worth mentioning the work by Biester et al. (2004, 2006) and Martínez-Cortizas
561 et al. (2007, 2016) in peatland soils, where they have started to study the temporal trends of
562 the more stable Br_{org} compounds and the role of the main pedogenetic processes on Br
563 accumulation. Altogether, this analysis can help future research on the Br–OM link in tidal
564 marshes, needed to support the use of Br in marsh soils/sediments as a paleoclimatic
565 indicator.

566 Finally, we propose a conceptual model for the salt marshes Br cycle in our case study
567 (Caminha and Casa Branca) in order to summarize the interplay between the forcing factors
568 analysed along the lines of the previous discussion. The interactions illustrated in Figure 5
569 are, in our view, the most likely to have a strong influence on salt marshes short and even
570 longer-term role to act as a source and/or as a sink for Br. Accordingly, NW coast high

marshes under the sustained influence of concomitant lower photosynthetically active radiation (PAR), corresponding to the spectral range of solar radiation from 400 to 700 nm that is used in photosynthesis reactions (e.g., Mariscal et al., 2000), and colder wetter conditions developed lower salinity baselines where the vegetation cover largely consists of non-succulents. These lower salinity (mesohaline) tidal marshes have typically higher rates of plant productivity and lower decomposition rates of dead and senescing plant material, leading to higher accumulation of soil/sediment OM (e.g., Morrissey et al., 2014). Collectively, these processes culminate in lesser CH₃Br atmospheric emissions from marshes and higher Br_{org} concentration in their soils/sediments. On the other hand, the SW coast high marshes, settled under higher available PAR conditions and a hotter and dryer climate, show a higher (polyhaline to euhaline) salinity baselines and consequently lower productivity, being mostly colonized by succulent plants more adapted to saline conditions and greater emitters of CH₃Br. These conditions finally lead to a lower long-term storage of OM and Br_{org} in soils/sediments.

The afore-described model establishes a connection with the analysis of the Br temporal variability in light of the past SA–climate link made in the next section.

3.2.2. Br enrichment peaks in association with Grand Minima of solar activity

The manifestation of a solar activity (SA) Grand Minima in terrestrial climate has been well established after the pioneering work of Eddy (1976). The most recent Grand Minima in reconstructed SA, as expressed by the TSI variation (Steinhilber et al., 2012), are presented in Figure 4 (Maunder Minimum – MM; 1645–1715; Spörer Minimum – SM; 1450–1550; and Wolf Minimum – WM; 1282–1342). Note that the Dalton Minimum (DM; 1790–1820) has a

594 distinct physical origin (Duhau and de Jager (2010), and it is therefore not regarded today
595 as a Grand Episode of SA.

596 In the last few decades, extensive research work has been done towards a better
597 understanding of the Sun-Earth's climate coupling system, with great progress being
598 achieved (e.g., Haigh, 2007; Soon et al., 2014 for a review). Recently, Brugnara et al.
599 (2013) referred that the Euro–Atlantic sector, in which Portugal is located, seems to be a
600 region with a particularly strong solar influence on the troposphere, finding a weak but
601 significant change in the mean late winter circulation over Europe, which culminates in
602 detectable impacts on the near-surface climate. The results obtained by Jiang et al. (2015)
603 suggest not only that climate in the northern North Atlantic regions follows SA fluctuations
604 on multidecadal to centennial time scales, but also that it is more susceptible to the
605 influence of those fluctuations throughout cool periods with, for instance, less vigorous
606 ocean circulation. Similar results were found by Gómez-Navarro et al. (2012) in the context
607 of climate simulations for the second millennium over the Iberian Peninsula. These
608 researchers studied the impact of natural forcing and internal variability on climate, and
609 found that temperature and precipitation variability are significantly affected at centennial
610 scales by variations in the SA.

611 Grand Minima and Dalton-type Minimum scenarios are broadly characterized by (i) lower
612 TSI (i.e., lower available PAR) (Lean, 1991, and references therein), (ii) development of
613 cloudiness (e.g., Usoskin and Kovaltsov, 2008), and (iii) decreased global/regional air
614 surface temperatures (e.g., Neukom et al., 2014) in tandem with greater regional
615 precipitation variability. In the Iberian Peninsula, according with the modelled results from
616 Gómez-Navarro et al. (2011), precipitation could have increased in response to reduced
617 solar forcing (Figure 4), prompting greater river discharges. Also Cruz et al. (2015) related
618 maxima rainfall episodes, as recognized in their stalagmite record, with other Grand Solar

619 Minima, suggesting a strong coupling between SA and precipitation over northern Iberia,
620 which agrees with Gómez-Navarro et al. (2012) outcomes in the context of climate
621 simulations.

622 Up to now, and in the absence of a “unified hypothesis”, the explanations for the
623 connections between solar phenomena and the lower atmosphere processes can be
624 summarized in two types of mechanisms: (i) “top-to-down”, influencing the pole-to-equator
625 temperature gradient and exerting an impact on the modulation of the atmospheric
626 circulation cells, weakening or strengthening the zonal winds, and (ii) “bottom-to-up” that
627 directly impact on the radiation fluxes, energy balance and temperatures on the ground.
628 Depending on the surface albedo a part of this radiation is absorbed and transformed into
629 latent or sensible heat. During periods of lower SA, less radiation is available in the tropics
630 for conversion to latent heat, which is thought to lead to a weakening of the Hadley and
631 Ferrel cells (Labitzke et al., 2002). Other than intensity, the position and extent of those cells
632 are also affected, inducing latitudinal shifts. Hence both mechanisms finally impact the
633 atmospheric circulation modes responsible for the global/regional precipitation and
634 temperature patterns (e.g., Gray et al., 2010; Martin-Puertas et al., 2012; Thiéblemont et al.,
635 2015).

636 Therefore, if solar variations are an important source of regional climate variability, we might
637 expect that paleoclimate proxies reproduce somewhat the climatic response to SA changes.
638 In line with this, Moreno et al. (2015) suggested that Br soil/sediment enrichment in the
639 Caminha salt marsh (NW coast) is, at least partially, related to the SA pattern over the last
640 almost 800 years. This can be extended to Casa Branca salt marsh (SW coast), essentially
641 through a SA control on (i) available PAR fluctuations and (ii) regional temperature and
642 precipitation regimes, affecting evapotranspiration rates and, as a result, interstitial water
643 salinity at the upper elevations within the marsh. Indeed, the relationship between the

644 curves presented in Figure 4 is clear, with periods of highest Br enrichment (FCPw1 and
645 FWCBBr) agreeing with major excursions in SA. This means that the TSI negative anomalies
646 (Steinhilber et al., 2012) from the Dalton, Maunder, Spörer and Wolf Solar Minima, to which
647 correspond periods of both modelled lower temperature and increased precipitation in the
648 NW and SW of Portugal (Gómez-Navarro et al., 2011), are marked by Br positive peaks in
649 marshes soils/sediments, more clearly observed in Br/OM ratio curves to account for
650 changes in OM content (see black arrows). Considering only the group of samples from the
651 FWCBBr core falling within the LIA (with stronger impacts on the climate of Europe and other
652 regions neighbouring the North Atlantic during the 16th–19th centuries; e.g., Mann, 2002),
653 three display a more pronounced Br excess relative to OM, lying outside the upper 95%
654 confidence limit of their linear regression interval: AD 1451 (Spörer Minimum), and AD 1694
655 and AD 1660 (both in the Maunder Minimum). These sedimentary records, considered
656 altogether with other climatic proxies for the NW of Portugal (Moreno et al., submitted),
657 strongly suggest that the LIA resulted in a wetter and cool climate in this south-western
658 European region, triggering major hydrological changes present in paleoecological records,
659 namely from high marsh benthic foraminifera (Moreno et al., 2014). Specifically, climatic
660 shifts driven by Grand Minima on western Portuguese coast could have forced a
661 deceleration of the whole dynamics involving the net CH₃Br phytogenic emissions to the
662 atmosphere, thus favouring Br sequestration and storage (as Br_{org}) in marsh
663 soils/sediments. Throughout SA Grand Minima (↓TSI/PAR), the climate controls departed
664 from normal values (↓T; ↓ET; ↑P; see Figure 5), inducing a decrease in marsh interstitial
665 salinity, certainly ended in higher plant productivity peaks (e.g., De Leeuw et al., 1990),
666 thereby causing a rise in salt marsh sediment C_{org} accumulation over time, with plant debris
667 more enriched in Br and liable to further bromination during the humification process.

668 Rhew et al. (2014) emphasized that CH₃Br phytogenic emissions from coastal salt marshes,
669 present a dramatic inter- and intra-marshes variability, namely in relation to magnitude

670 (subtropical salt marshes showing much higher emission rates than temperate salt
671 marshes) and seasonality. They recorded maxima CH₃Br emission fluxes in peak summer
672 growing season (July) and lowest at the end of the growing season (November). The latter
673 were registered during the morning and coincident with the high tide. They also found a
674 pronounced mid-day peak, coinciding with the time of highest ambient air and surface soil
675 temperatures, in the diurnal CH₃Br emission trend, with this one mirroring the variation of
676 PAR.

677 Considering both CH₃Br phytogenic emissions and the ubiquity of soil/sediment OM
678 bromination, and applying them to a Grand Minima scenario (↓TSI/available PAR; ↓air
679 surface temperatures; ↓growing seasons, ↑rainfall leading to ↑soil/sediment saturation), a
680 reliable framework for the Br enrichment in temperate marsh habitats triggered by climatic
681 shifts driven by SA can be proposed (see Figure 5), recognizing that further work would be
682 required to completely prove this assumption.

683 Volcanic eruptions may also represent an alternative source of Br to salt marshes during
684 Grand Minima Episodes (see Figure 4). The first discovery of volcanic BrO (Bobrowski et
685 al., 2003), and its subsequent measurement in many volcanic plumes around the globe
686 (e.g., Roberts et al., 2014) demonstrates the formation of reactive bromine (firstly as Br₂,
687 which then converts into other forms including Br, BrO, HOBr, BrONO₂) during these
688 events, which can be removed (at least partially) from gaseous phase by aerosol, water and
689 ice-uptake (followed by particle sedimentation) (Fernandez et al., 2014; Jourdain et al.,
690 2015). Reactive bromine acts as a catalyst to its own formation, leading to an exponential
691 growth called “bromine explosion”. The LIA Grand Minima have been punctuated by
692 considerable volcanic activity (e.g., D’Arrigo et al., 2013; Figure 4). D’Arrigo et al. (2013)
693 highlighted, as two of the major volcanic events over the past millennium, the eruptions of
694 AD 1453 (Kuwae Volcano, Vanuatu; Spörer Minimum) and AD 1815 (Mount Tambora,

695 Indonesia; Dalton Minimum). Such episodes of great volcanic activity and their worldwide
696 effects (e.g., Trigo et al., 2009; Koffman et al., 2013) might also have had the potential for
697 causing disruption on the Br cycling in Caminha and Casa Branca salt marshes, contributing
698 somewhat for their soils/sediments Br enrichment throughout Grand Minima.

699

700 **4. Conclusions**

701 The present study addresses major topics concerning the Br cycling in contrasting salt
702 marshes environments of four western Portuguese estuaries – Minho, Lima, Sado and Mira,
703 drawing attention for its complexity and linkages with OM dynamics. We provided evidence
704 that besides the marine influence, Br enrichment of these marshes is ultimately connected
705 to their ability for long-term C_{org} storage. A clear difference between the marshes from NW
706 and SW coasts stands out, with the former being more enriched in both OM and total Br.
707 This contrasting behaviour is driven by different climatic conditions between the two regions
708 that favour more strongly the mechanisms and processes of OM production, burial, and
709 preservation (with concomitant incorporation of Br) in the north-western coastal salt
710 marshes, in contrast with the SW coast, most probably as a result of the higher salinity
711 (lesser productivity) of the latter. Seemingly, this same NW climatic setting inhibits the
712 emission of comparatively larger phytogenic CH_3Br fluxes to atmosphere. This might be
713 intensified as a result of the marshes colonization by non-succulent species (less efficient in
714 the CH_3Br production), further promoting the Br enrichment of soil/sediment OM.

715 Although the applied approach is constrained by chronological uncertainties, cores sampling
716 resolution or the (relatively short) time series lengths, preventing the application of
717 enhanced methodologies in the time-frequency domain, the Br temporal variability in the
718 Caminha and Casa Branca salt marshes can be related to SA oscillations, showing greater

719 Br enrichment during Grand Minima or Minima-like Episodes. This can be explained in
720 connection with the changing temperature (decrease) and precipitation (increase) regimes
721 in the NW and SW coasts of Portugal induced by lower TSI (available PAR), as pointed out
722 by previous studies and supported by climate simulations. The contribution of major tropical
723 volcanic explosions at Grand Solar Minima during the registered higher Br enrichment in the
724 two Portuguese salt marshes is also considered.

725 Finally, we expect that the issues encompassed here can be deepened in future research
726 about the Br biogeochemical cycle in salt marshes worldwide. The proposed conceptual
727 framework identifies several influences capable of imbalance the Br–OM interconnections
728 and helps to prioritize which are likely to play key roles on salt marshes Br recycling, in
729 order to improve, henceforward, its reliability as a marker of climate change driven by past
730 SA. In line with this, it is important to bear in mind that for robustly test solar-climate signals
731 in Br tidal salt marsh records, large and widespread ensembles of well-dated data are
732 required along with high-resolution sampling.

733

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743

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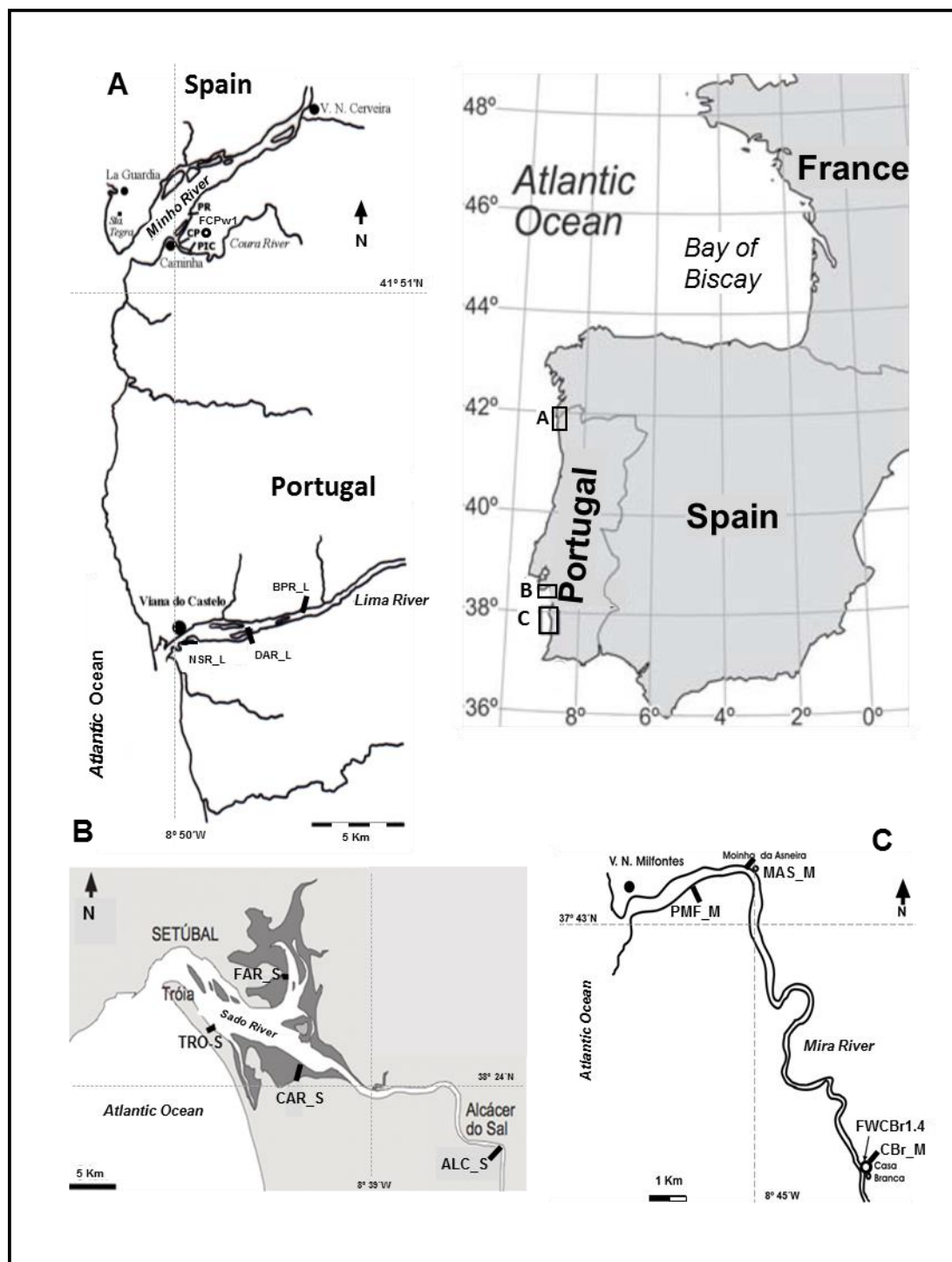
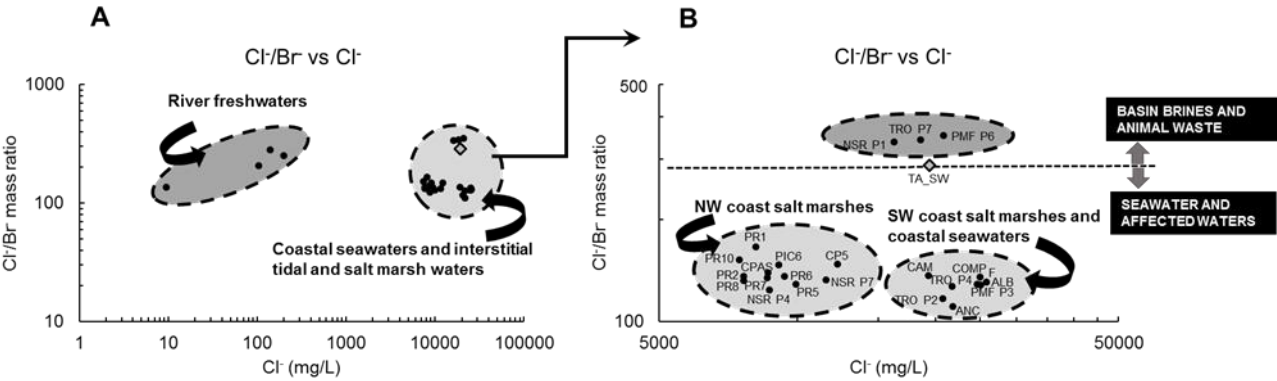


Figure 1. A. Study areas general location; B. Minho and Lima estuaries (NW coast); C. Sado estuary (SW coast); D. Mira estuary (SW coast). The location of the complete sampling set is also signalled. This includes thirteen surface sampling transects across the four tidal salt marshes (where both interstitial waters and sediments have been collected), and the two sediment cores obtained in the high marsh zones of the Caminha (FCPw1; Minho estuary; 1.55 m above mean sea level; 41°52'37.0" N and 8°49'28.0" W) and Casa Branca (FWCB1.4; Mira estuary; 1.74 m above mean sea level; 37°40'03.7" N and 8°43'12.7" W) salt marshes.

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1355 **Figure 2.** Cl⁻/Br⁻ mass ratios vs. Cl⁻ concentrations of (surface and interstitial) water samples and potential
1356 sources (zones and theoretical limits from Panno et al., 2006). **A.** Clusters of freshwater (riverine), marine, and
1357 impacted water samples are indicated (i.e., tidal and salt marsh interstitial waters). **B.** Clusters of interstitial
1358 waters from NW and SW coastal salt marshes. Marine water samples are in the same group as the SW coast
1359 salt marshes interstitial water. A group of three anomalous samples (TRO_P7, NSR_P1 and PMF_P6) is also
1360 identified. The diamond represents a typical marine sample.

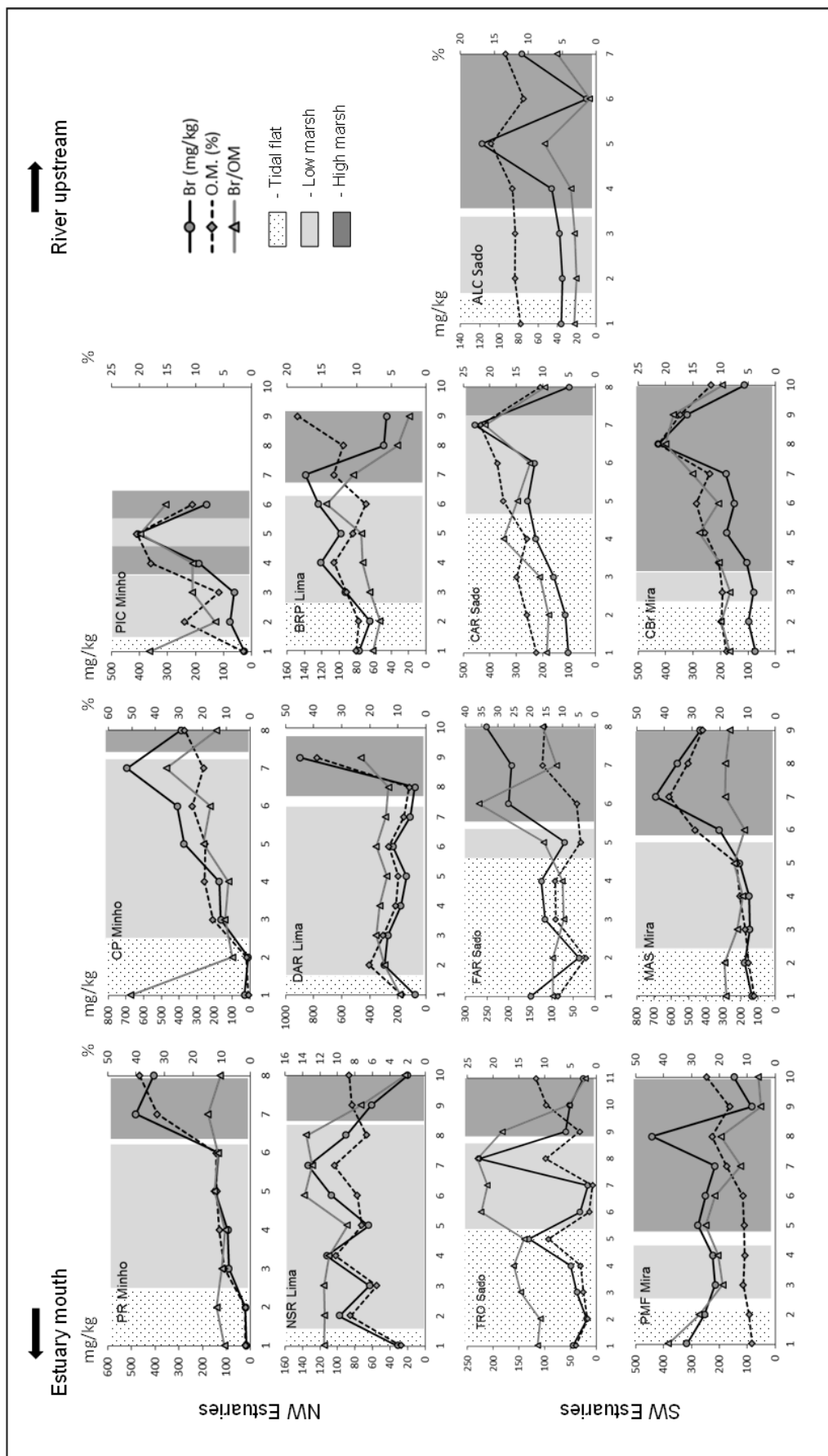


Figure 3. Bromine and organic matter contents (OM) of superficial tidal salt marsh sediment transects from Minho, Lima, Sado and Mira estuaries.

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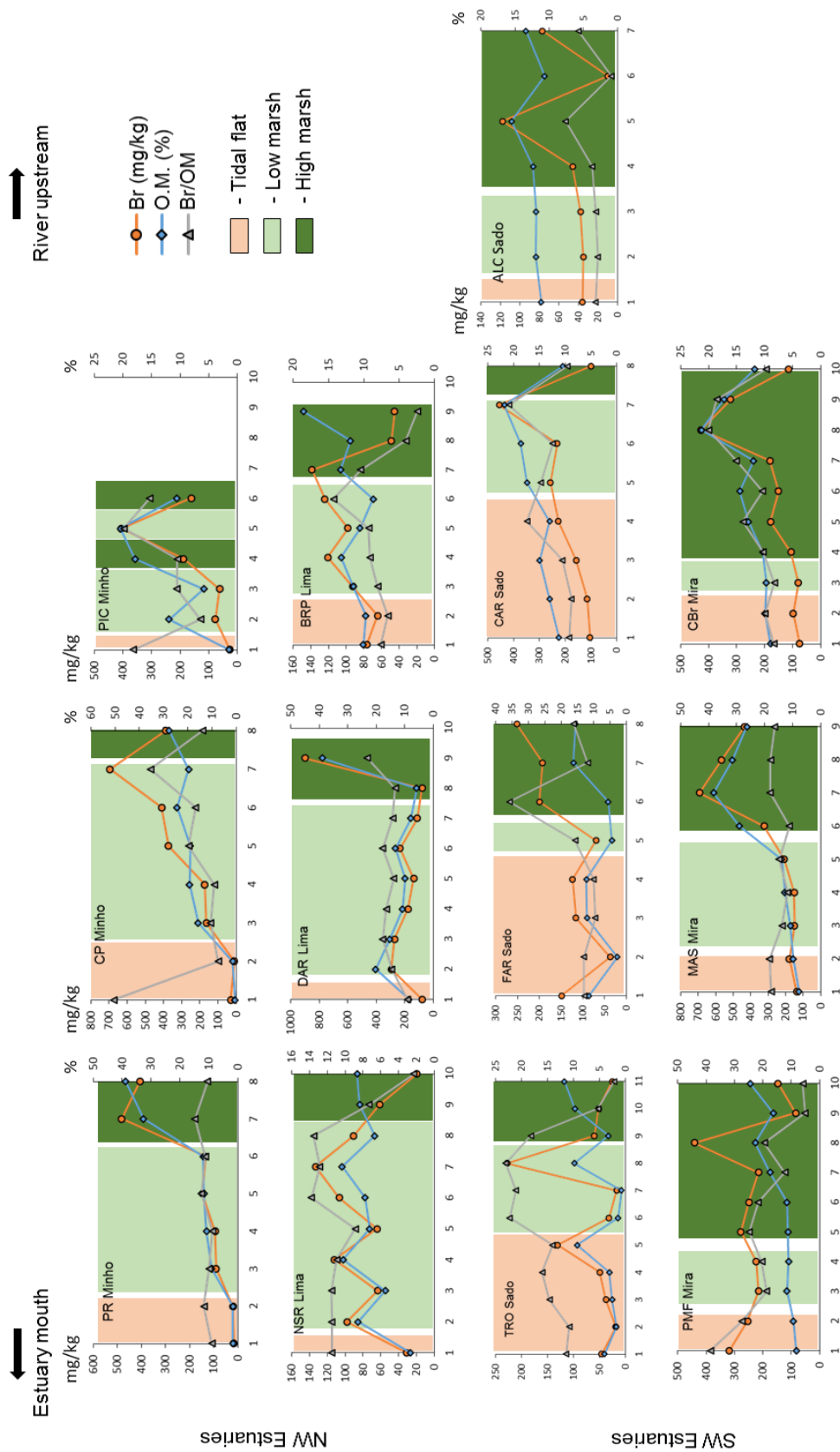


Fig 3. Bromine and organic matter (OM) contents of superficial tidal salt marsh sediment transects from Minho, Lima, Sado and Mira estuaries.

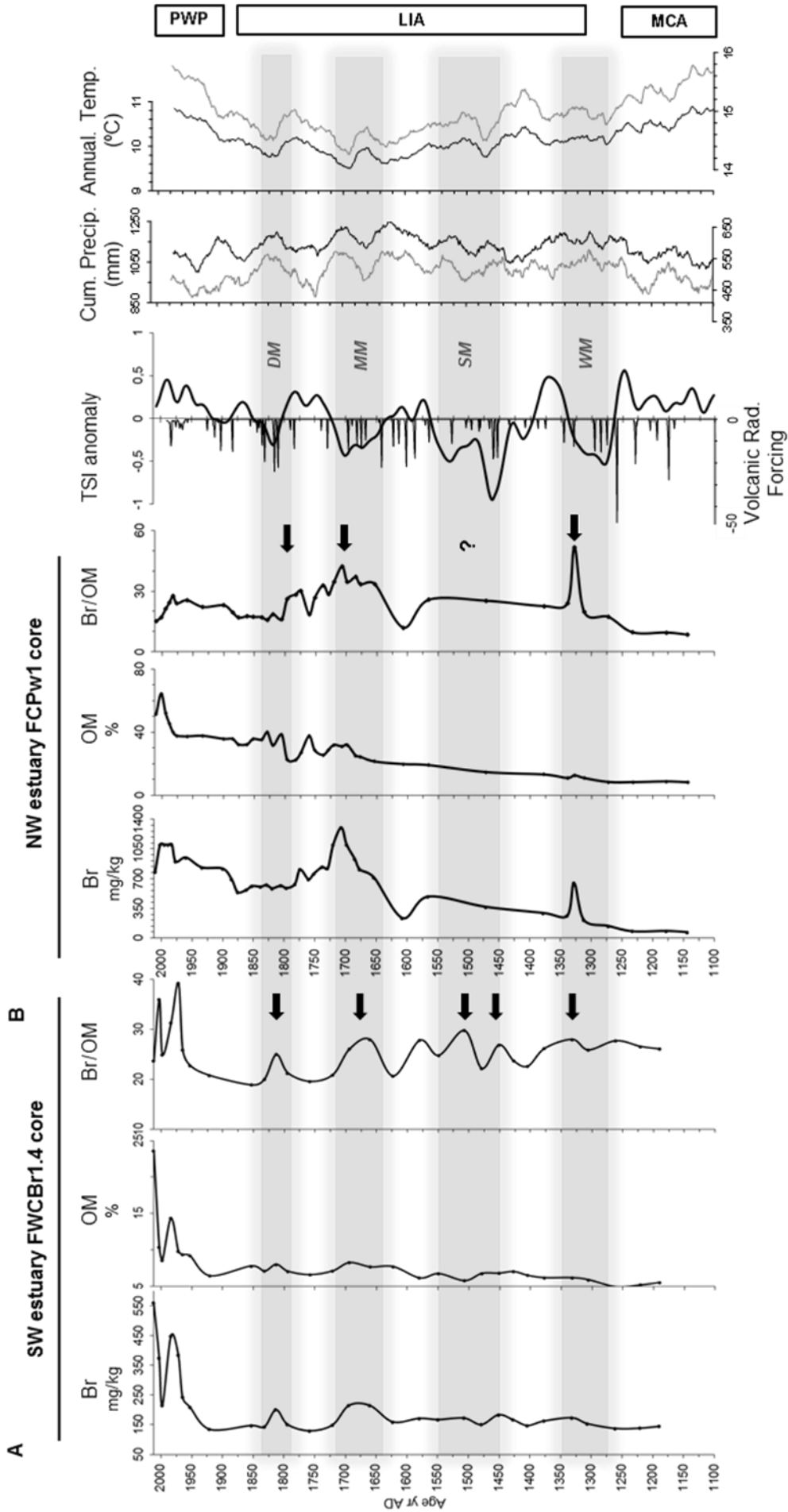


Figure 4. FCPw1 and FWCB1.4 cores geochemical data: Br (mg/kg), OM (%) and Br/OM ratios. Other data represented are: TSI anomaly using a 21-year running average reconstruction (Steinhilber et al., 2012); volcanic radiative forcing (black bars) after Crowley (2000); cumulative precipitation and annual temperature (31-year running averages) for NW coast (black lines and top axis) and SW coast (grey lines and bottom axis) of Portugal (Gómez-Navarro et al., 2011). DM – Dalton Minimum; MM – Maunder Minimum; SM – Spörer Minimum; WM – Wolf Minimum.

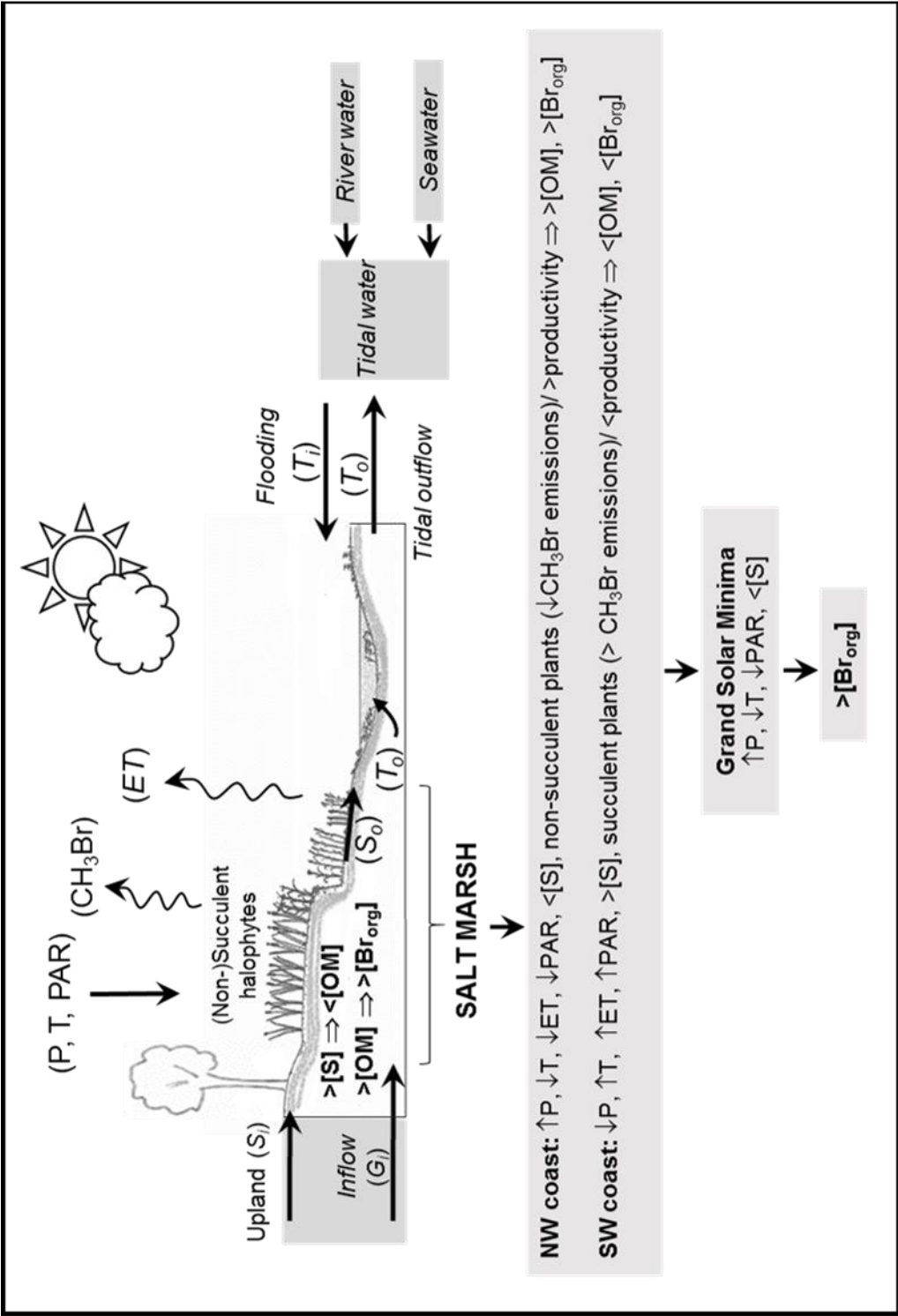


Table 1. Bromide concentrations in surface water (river freshwater, estuarine and coastal seawater) samples and in marsh interstitial water samples (Minho, Lima, Sado and Mira tidal salt marshes).

Surface and interstitial waters	NW coast				SW coast			
	Minho		Lima		Sado		Mira	
Zone	Br ⁻ (mg/L)	Salinity (‰)	Br ⁻ (mg/L)	Salinity (‰)	Br ⁻ (mg/L)	Salinity (‰)	Br ⁻ (mg/L)	Salinity (‰)
River	<d.l.	0.0	0.07	0.0	0.8	0.3	0.5	0.2
Estuary mouth	141	32.5	—	—	—	—	—	—
Coastal seawater	197	35.2	197	35.2	185	35.7	197	35.4
Tidal flat	52 (48-55)*	9.5 (2-17)*	48	24.6	177	31.8	—	—
Low marsh	71 (64-83)*	14.5 (10-18)*	70	14.4	171	31.4	195	37.0
High marsh	56 (49-62)*	12.6 (11-15)*	87	13.5	54	26.6	59	30.6
Marsh transect references	PR; CP; PIC		NSR-L		TRO-S		PMF-M	
Distance to river mouth (km)	3.8 - 4.8		2.0		14.0		2.0	

*Average from three transects and range

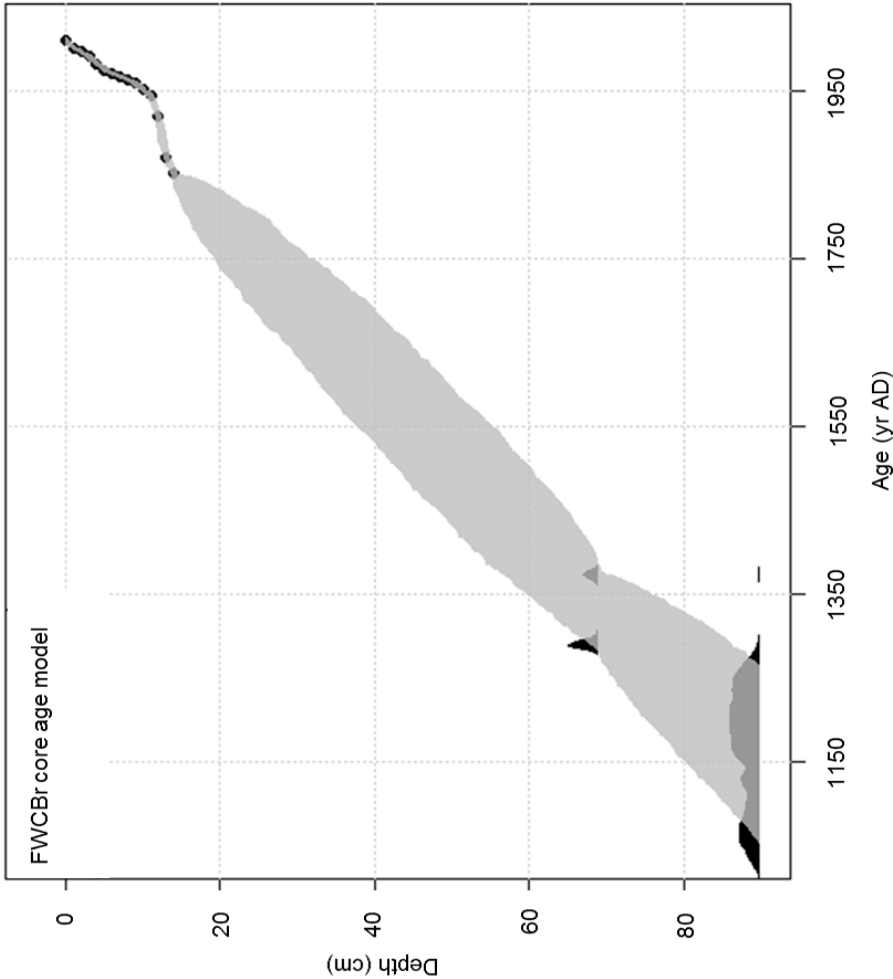
d.l. detection limit (0.01 mg/L Br⁻)

Table 2. Bromine content and OM contents in the surface sediment cross shore transects from the Minho, Lima, Sado and Mira intertidal domains (tidal flat, low marsh and high marsh).

Surface sediment samples / Estuaries	NW coast				SW coast			
	Minho		Lima		Sado		Mira	
Zone	Br (mg/kg)	OM (%)	Br (mg/kg)	OM (%)	Br (mg/kg)	OM (%)	Br (mg/kg)	OM (%)
Tidal Flat	21 (11-30)*	1.4 (0.6-1.7)*	71 (31-79)*	8.6 (2.7-10.1)*	83 (19-226)**	11.5 (1.7-15.0)**	158 (75-318)*	8.8 (7.7-9.9)*
Low Marsh	133 (62-695)*	15.0 (5.9-24.4)*	110 (64-296)*	11.0 (5.5-20.2)*	53 (17-456)**	8.2 (0.8-21.8)**	152 (80-224)*	11.2 (9.7-12.8)*
High Marsh	389 (162-482)*	27.7 (10.6-38.8)*	49 (45-900)*	13.3 (5.9-38.9)*	81 (10-252)**	11.7 (3.3-16.3)**	233 (85-693)*	16.9 (11.2-38.3)*
Marsh transect/cores surface references	PR; CP; PIC		NSR_L; DAR_L; BPR_L		TRO_S; CAR_S; FAR_S; ALC_S		PMF_M; MAS_M; CBr_M	
Distance to river mouth (km)	3.8 - 4.8		2; 5; 8.5		14; 19; 24; 43.5		2; 3.5; 13	
Correlation coefficient Br vs.OM (p < 0.001)	0.86 (N= 21)		0.89 (N= 28)		0.67 (N= 34)		0.79 (N= 29)	

*Average (median) from three transects and range

**Average (median) from four transects and range



Core/Sample	Depth (cm)	$\delta^{13}\text{C}$ ‰	Conventional radiocarbon age	$\Delta^{14}\text{C}$ ‰	Calibrated years AD (2 Sigma range)
FW CBr 2.0 CM 70	69-70	-25.0	720 +/- 30 BP	-85.7±3.4	1260 to 1295
FW CBr 2.0 CM 91	90-91	-25.6	920 +/- 30 BP	-108.2±3.3	1025 to 1190

Depth (cm)	^{137}Cs	2-sigma	^{210}Pb	2-sigma	^{214}Pb	2-sigma	$^{210}\text{Pb}_{\text{excess}}$
0.5	5.56	2.27	152.5	30.6	37.5	6.76	115
1.5	5.44	1.6	84.52	20.11	40.02	3.78	44.5
2.5	6.95	1.63	72.48	20.36	38.38	3.9	34.1
3.5	8.78	2.13	98.24	21.85	37.22	5.58	61.02
4.5	10.64	2.18	70.55	21.47	34.65	5.25	35.9
5.5	13.23	2.23	53.13	21.05	32.45	4.79	20.68
6.5	19.19	2.32	48.51	25.88	34.29	3.97	14.22
7.5	23.69	2.73	46.97	17.43	36.53	6.44	10.44
8.5	14.41	2.04	54.26	18.92	42.94	5.77	11.32
9.5	6.74	1.81	58.57	20.37	41.42	4.24	17.15
10.5	6.4	1.72	52.57	18.23	40.33	4.8	12.24
11.5	5.45	1.65	65.43	19.23	41.41	3.99	24.02
12.5	2.92		65.13	22.17	47.18	4.21	17.95
13.5	2.37		53.57	21.12	51.31	5.19	2.26
14.5	2.58		56.08	21.57	52.17	5.83	3.91
15.5	2.5		41.48	19.35	43.05	4.29	-1.57

Pb and Cs isotopes contents (Bq kg⁻¹)

Appendix A. Age model for the core FWCBr and estimated 2 σ errors, based on two AMS ^{14}C dates, performed on total organic sediment, and ^{210}Pb and ^{137}Cs chronology. The data interpolation was obtained with Bchron 4.1 software.

1385 **Appendix B.** Geochemical data from the sediment core FWCB (Casa Branca salt marsh;
 1386 Mira estuary).

Depth (cm)	Br (ppm)	OM (%)	Br/OM	Age (AD years)
0-1	560	23.6	23.7	2012
1-2	374	10.4	36.0	2003
2-3	214	8.6	24.9	1998
4-5	448	14.3	31.2	1984
6-7	385	9.8	39.3	1972
8-9	242	9.4	25.8	1965
10-11	209	9.2	22.7	1953
12-13	134	6.5	20.7	1921
14-15	147	7.8	18.8	1853
16-17	142	7.1	20.0	1832
18-19	200	8.0	24.9	1813
20-21	150	7.1	21.2	1794
24-25	129	6.6	19.5	1758
28-29	149	7.1	20.9	1721
31-32	215	8.3	26.0	1694
35-36	214	7.7	27.9	1660
39-40	159	7.7	20.6	1623
43-44	171	6.2	27.8	1580
47-48	167	6.7	24.8	1549
52-53	172	5.8	29.7	1507
55-56	150	6.8	22.1	1479
58-59	183	6.8	26.9	1451
61-62	166	7.0	23.6	1427
64-65	147	6.5	22.6	1404
68-69	162	6.2	26.2	1377
72-73	173	6.2	28.0	1332
75-76	153	5.9	25.9	1306
80-81	137	4.9	27.7	1261
85-86	138	5.2	26.5	1221
89-90	144	5.5	26.0	1190

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