

1 **Oxygen isotopes in orangutan teeth reveal recent and ancient climate variation**

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23 **Abstract**

24 Studies of climate variation commonly rely on chemical and isotopic changes
25 recorded in sequentially-produced growth layers, such as in corals, shells and tree rings, as
26 well as in accretionary deposits—ice and sediment cores, and speleothems. Oxygen isotopic
27 compositions ($\delta^{18}\text{O}$) of tooth enamel are a direct method of reconstructing environmental
28 variation experienced by an individual animal. Here we utilize long-forming orangutan
29 dentitions (*Pongo* spp.) to probe recent and ancient rainfall trends on a weekly basis over ~
30 3–11 years per individual. We first demonstrate the lack of any consistent isotopic
31 enrichment effect during exclusive nursing, supporting the use of primate first molar teeth as
32 environmental proxies. Comparisons of $\delta^{18}\text{O}$ values ($n = 2016$) in twelve molars from six
33 modern Bornean and Sumatran orangutans reveal a high degree of overlap, with more
34 consistent annual and bimodal rainfall patterns in the Sumatran individuals. Comparisons
35 with fossil orangutan $\delta^{18}\text{O}$ values ($n = 955$ measurements from six molars) reveal similarities
36 between modern and late Pleistocene fossil Sumatran individuals, but differences between
37 modern and late Pleistocene/early Holocene Bornean orangutans. These suggest drier and
38 more open environments with reduced monsoon intensity during this earlier period in
39 northern Borneo, consistent with other Niah Caves studies and long-term speleothem $\delta^{18}\text{O}$
40 records in the broader region. This approach can be extended to test hypotheses about the
41 paleoenvironments that early humans encountered in southeast Asia.

42

43 **Introduction**

44 Present-day rainfall patterns in Indonesia are controlled by the Asian and Australian
45 monsoon systems, yielding annual trends that vary considerably with geography, topography,
46 and the direction of monsoonal winds (Aldrian and Susanto, 2003; Moron et al., 2009; Qian
47 et al., 2013; Belgaman et al., 2017). Northern Sumatra and western Borneo experience high
48 annual rainfall and relatively stable annual temperatures, with a bimodal distribution of
49 rainfall governed by the Intertropical Convergence Zone (van Schaik, 1986; Aldrian and
50 Susanto, 2003; Belgaman et al., 2017). These islands are also under the influence of inter-
51 annual climate fluctuations driven by the El-Niño Southern Oscillation (ENSO); a periodic
52 coupling of atmospheric and oceanic temperature gradients that initiates in the tropical
53 Pacific, and influences global temperature and precipitation trends (McPhaden et al., 2006).

54 It is well understood that variation in rainfall patterns influences the fundamental
55 structure of primate habitats (Brockman and van Schaik, 2005; Wessling et al., 2018). Dense
56 tropical forests are sustained by fairly consistent rainfall and short, irregular dry seasons,
57 while woodland communities in more arid environments have smaller trees, less dense
58 canopies, and more deciduous trees (Vico et al., 2017; Archibald et al., 2019). In regions with
59 prolonged dry seasons, low annual rainfall and savannah landscapes abound, in addition to
60 disturbances such as wildfires (Pletcher et al., 2022).

61 Open woodland and savannah environments are unfavorable for slow-moving
62 orangutans, the largest mammal with an arboreal lifestyle, particularly in regions with
63 predators such as tigers or humans (Thorpe and Crompton, 2009; Ashbury et al., 2015;
64 Spehar et al., 2018). Supra-annual ENSO events may also impact orangutan energy balance,
65 reproduction, and social organization through the inducement of mast-fruiting, or dramatic
66 seed production events in dipterocarp forests (Knott 1998; Curran et al., 1999; Marshall et al.,
67 2009). Such climate fluctuations over the past several hundred years have been documented
68 in coral isotopes and tree-ring analyses, revealing especially marked changes during the past
69 few decades (Cole et al., 1993; Stahle et al., 1998; Hughen et al., 1999; Urban et al., 2000;
70 Tudhope et al., 2001; Pumijumnong et al., 2020).

71 Detailed climate records prior to the era of human-induced climate change are
72 somewhat limited for island southeast Asia, but they are directly relevant to understanding
73 the recent distribution of orangutans, and the arrival and dispersal of modern humans in the
74 region during the Late Pleistocene (e.g., Piper 2016; Bae et al., 2017; Spehar et al., 2018). A
75 small number of studies of fossil corals, molluscs, marine sediments, and speleothems has

76 provided insights into the last interglacial and glacial periods (e.g., Hughen et al., 1999;
77 Tudhope et al., 2001; Stephens et al., 2016; Yang et al., 2016; Buckingham et al. 2022). For
78 example, oxygen isotopes in fossil corals from seven periods during the last 130,000 years
79 suggest that ENSO activity in the western Pacific over that time was comparable to modern
80 records, although there was variation in the intensity of such activity at different timepoints
81 (Tudhope et al., 2001). This study was also able to resolve bimodal annual rainfall peaks in
82 modern corals, yet such detailed subannual records are extremely uncommon, particularly
83 from terrestrial environments where early humans once lived alongside orangutans and other
84 mammals.

85

86 *Oxygen isotope studies for paleoenvironmental reconstruction*

87 Oxygen isotope values ($\delta^{18}\text{O}$) in water vary with latitude, altitude, temperature and
88 precipitation cycles, and are also impacted by precipitation sources. In tropical regions the
89 primary determinant of rainfall isotope compositions is rainfall amount (Dansgaard, 1964;
90 Rozanski et al., 1993; Belgaman et al., 2017). During wet seasons, rainfall $\delta^{18}\text{O}$ values are
91 relatively low, while the opposite pattern is evident in periods with less rain, although other
92 meteorological factors can influence isotope values as well (Belgaman et al., 2016). This
93 primary tropical pattern influences isotopic variation in meteoric, surface, and leaf waters,
94 which may show further elevations in $\delta^{18}\text{O}$ values during dryer periods due to preferential
95 evaporative loss of the lighter isotope, ^{16}O (da Silveira et al., 1989; Bowen, 2010; Roberts et
96 al., 2017).

97 In addition to $\delta^{18}\text{O}$ values in fossil corals, tree rings and speleothems, other fine-
98 scaled oxygen isotopic climate proxies include otoliths (fish ear bones) and mollusc shells
99 (e.g., Aubert et al., 2012; Stephens et al., 2016; Prendergast et al., 2018)—although these are
100 rarely preserved in rainforest environments. Records of $\delta^{18}\text{O}$ values in mammalian tooth
101 enamel are a more direct means of studying seasonality (reviewed in Green et al., 2018,
102 2022), providing insight into the actual climates experienced by individuals, in contrast to
103 indirect proxies for which it can be difficult to establish concurrence. Unlike bone, teeth do
104 not remodel during life, and the phosphate component of the enamel mineral (hydroxyapatite)
105 is especially resistant to modification after burial (reviewed in Smith et al., 2018a; Pederzani
106 and Britton, 2019).

107 Tooth enamel is most commonly sampled with hand-held drills to recover the isotopic
108 composition of oxygen inputs from water and food preserved in the hydroxyapatite (e.g.,
109 Janssen et al., 2016; Roberts et al., 2020; Kubat et al., 2023). This coarse drilling method

110 yields spatially and temporally blurred powdered samples formed over a substantial and
111 unknown period of time, however, precluding the identification of precise seasonal
112 environmental patterns. To circumvent this limitation, we have employed the stable isotope
113 sensitive high-resolution ion microprobe (SHRIMP SI) to measure $\delta^{18}\text{O}$ values sequentially
114 from thin sections of teeth, relating these to daily increments and birth lines to determine
115 enamel formation times, and in some instances, calendar ages (Smith et al., 2018a, 2022;
116 Green et al., 2022; Vaiglova et al., in review).

117 It is well established that $\delta^{18}\text{O}$ values in tooth enamel are closely related to local water
118 oxygen isotope compositions (reviewed in Green et al., 2018, 2022). For teeth that form after
119 birth and during periods of milk consumption, $\delta^{18}\text{O}$ values are expected to be higher, as a
120 result of infant evaporative water loss while consuming ^{18}O -enriched mother's milk (Bryant
121 et al., 1996; Wright and Schwartz 1999; Britton et al., 2015). Studies of large-bodied
122 mammals report that milk $\delta^{18}\text{O}$ values are elevated by $\sim 1\text{--}6\text{‰}$ relative to local drinking
123 water $\delta^{18}\text{O}$ (Kornexl et al., 1997; Lin et al., 2003; Chesson et al., 2010; Green et al., 2018; but
124 see Cherney et al., 2010). Comparable data on human or nonhuman primate milk enrichment
125 appear to be lacking, save for a study of 44 British infants aged 5–16 weeks (Roberts et al.,
126 1988). The urine of infants who were breast-fed showed isotopic enrichment of 1–3 ‰
127 compared to infants who were fed formula prepared from sterile local tap water.

128 While such studies point to potential changes in infant body water during nursing, it is
129 unclear whether such differences prohibit the use of early-formed enamel in studies of
130 climate variation (Blumenthal et al., 2017; Luyt and Sealey 2018). Two studies of $\delta^{18}\text{O}$
131 values in the dentitions of modern sheep, horses, and zebras reported higher bulk values ($\sim 1\text{--}2\text{‰}$)
132 in five molars (M1) compared to the rest of the permanent dentition (Bryant et al.,
133 1996; Fricke and O'Neil, 1996). This led Fricke and O'Neil (1996) to suggest that M1s are
134 unlikely to reflect the values of local meteoric water due to the influence of maternal inputs in
135 utero and through lactation. However, near-weekly $\delta^{18}\text{O}$ values over the first 2.75 years of
136 life in a Neanderthal M1 measured with SHRIMP SI showed clear annual trends and
137 maximum $\delta^{18}\text{O}$ values corresponding to a period after nursing has ceased (Smith et al.,
138 2018a). An examination of longer continuous periods of enamel formation within and
139 between teeth will help to clarify whether early-formed primate teeth should be avoided for
140 studies of climate seasonality.

141 Here we first assess whether wild orangutans show elevated $\delta^{18}\text{O}$ values in early-
142 formed enamel, testing the suggestion that M1s are significantly affected by nursing ^{18}O -
143 enrichment, thereby precluding their use in climatological reconstructions. We then explore

144 approximately 30 years of weekly $\delta^{18}\text{O}$ values (n = 2016 measurements) to compare
145 orangutan individuals from the islands of Sumatra and Borneo. Finally, we contrast $\delta^{18}\text{O}$
146 values between modern and Pleistocene orangutans, including those from key regions of
147 early human occupation: Lida Ajer, Sumatra (Hooijer, 1948; Westaway et al., 2017) and Niah
148 Caves, Malaysia (Hooijer, 1961; Barker et al., 2007) (Figure 1, Table 1). Novel
149 understanding of climate patterns in these fossil assemblages may inform debates about the
150 likelihood of modern humans living in dense Asian rainforests, and the conditions that would
151 support savannah corridors for human dispersals throughout the region (e.g., de Vos, 1983;
152 Bird et al., 2005; Westaway et al., 2017; Louys and Roberts, 2020).

153

154 Insert Figure 1 and Table 1 here

155

156 **Results**

157 *Modern orangutans*

158 The $\delta^{18}\text{O}$ ranges of twelve modern and six fossil orangutan molars, representing 2971
159 near-weekly measurements spanning 57.6 years of tooth formation, are listed in Table 2.
160 Prior to making comparisons between individuals, geographic regions, or time periods, we
161 first consider the potential intra-individual effect of isotopic enrichment from maternal milk
162 on $\delta^{18}\text{O}$ values. Comparisons of $\delta^{18}\text{O}$ values during the first, second, and third years of life in
163 five modern orangutan first molars (M1) do not show consistently elevated values during
164 their first year (Figure 2). Mean yearly $\delta^{18}\text{O}$ values in the first year are elevated by only 0.3
165 ‰ compared to the second year. While three of the five M1s showed first year $\delta^{18}\text{O}$ values
166 higher than second year values ($p \leq 0.05$), only two individuals showed mean values that
167 were $\sim 1 - 2$ ‰ higher during year one; one individual showed no difference from the first to
168 the second year, and one individual showed lower values during the first year than during the
169 second year ($p \leq 0.05$) (Supplementary Table 1). A sixth individual was only sampled from
170 193 days of age, but maximum values from this point onwards were similar across more than
171 three years of life. Similarly variable patterns were observed for the six putative fossil
172 orangutan M1s (Supplementary Figure 1).

173

174 Insert Figure 2 and Table 2 here

175

176 Comparisons across serial molars in four modern orangutans show no consistent trend
177 of elevated $\delta^{18}\text{O}$ values in M1s relative to successive molars (Figure 3). Only two individuals

178 showed maximum $\delta^{18}\text{O}$ values in their M1s relative to M2s; in both instances M3s were
179 unavailable due to their lack of development prior to death. The other two individuals showed
180 higher $\delta^{18}\text{O}$ values in M2s or M3s than in their respective M1s. In the case of the oldest
181 individual (ZSM 1981/248), the highest $\delta^{18}\text{O}$ values appeared at approximately 5.8 years of
182 age, well past the age when exclusive nursing ends.

183

184 Insert Figure 3 here

185

186 Comparison of the $\delta^{18}\text{O}$ values in the full datasets of modern Bornean and Sumatran
187 orangutans reveals a high degree of overlap. Values from the three Bornean individuals
188 ranged from 12.7 to 20.0 ‰ (n = 955 near weekly measurements), while the three Sumatran
189 individuals ranged from 11.3 to 20.6 ‰ (n = 1061 measurements). Comparisons of periodic
190 trends via spectral power distribution analysis revealed more consistent bimodal patterns in
191 the Sumatran individuals; three of the six Bornean molars were aperiodic (statistical power of
192 0.1 or less), while all six of the Sumatran molars revealed annual or semiannual cycles with
193 greater power (Supplementary Figure 2). Rapid oxygen isotopic shifts on the order of ~ 6–8
194 ‰ are evident in the single Bornean and Sumatran individuals with $\delta^{18}\text{O}$ measurements
195 spanning M1 to M3, which may represent one or more supra-annual ENSO events captured
196 during the ~ 9–11 years these molars were forming.

197

198 *Fossil orangutans – oxygen isotopes*

199 Concurrently forming teeth (molar specimens 11594.12 and 11595.105) from same
200 individual at Lida Ajer, Sumatra are nearly isotopically identical; $\delta^{18}\text{O}$ values range from
201 15.1 to 19.9 ‰ and 15.7 to 20.0 ‰, respectively, supporting the biogenic fidelity of these
202 records. The $\delta^{18}\text{O}$ values of two individuals from the nearby Sibrabang site (15.3–20.4 ‰,
203 14.7–20.8 ‰) are very similar to those of the Lida Ajer individual. These Sumatran fossils all
204 fall at the upper end of the range of modern Sumatran orangutans (Figure 4), and reveal
205 approximately annual $\delta^{18}\text{O}$ periodicities (0.9–1.3 years), as well as strong bimodal
206 distribution patterns in one instance (11565.162).

207

208 Insert Figure 4 here

209

210 The two fossils from the Niah Caves were excavated from different regions and
211 stratigraphic depths; $\delta^{18}\text{O}$ values in the tooth from grid US/22 ranged from 15.9 to 24.8 ‰

212 and, unlike the three modern Bornean individuals, yielded an annual periodicity (1.0 years).
213 The $\delta^{18}\text{O}$ in the tooth from grid Y/F4 ranged from 14.2 to 22.9 ‰ and showed a stronger
214 bimodal trend than an annual one, although its short formation time may have prohibited
215 identification of longer trends. The range of values from these two fossil molars (14.2–24.8
216 ‰) markedly exceeds the range of modern Bornean orangutans (12.7–20.0 ‰) (Figure 4),
217 with the mean $\delta^{18}\text{O}$ value at least 2 ‰ heavier. This suggests possibly drier conditions with
218 greater seasonality during fossil molar formation (Supplementary Figure 3).

219

220 *Fossil orangutans – U-series age estimates*

221 The six fossil teeth have very low uranium concentrations in their enamel (<0.5 ppm),
222 regardless of their origin (Supplementary Table 2). These enamel values are very close to the
223 detection limit of the Nu Plasma II MC-ICP-MS, and thus are not useful for estimating
224 minimum ages. The dentine of Lida Ajer specimen 11595.105 shows a spatial gradient of
225 increasing uranium concentration from ~ 41 to 66 ppm, and decreasing age estimates from ~
226 51 to 40 ka (Supplementary Table 2). This trend might result from a preferential uranium
227 leaching overprint near the end of the root. Spot DE10, positioned near the EDJ, is less likely
228 to be impacted (Supplementary Figure 4), and is thus assumed to provide the most reliable
229 minimum age for the tooth, ~ 40 ka. Uranium values from Lida Ajer specimen 11594.12
230 show a similar trend of concentrations decreasing from ~ 31 to 24 ppm towards the root tip.
231 However, the U-series age estimates remain constant within the range 31–34 ka across the
232 dentine (Supplementary Table 2; Supplementary Figure 4). No evidence for a recent
233 overprint is observed, supporting a minimum age of 33 ka. In summary, this individual's age
234 is at least 33 ka, and possibly > 40 ka.

235 U-series analysis of the dentine of Sibrabang specimen 11565.162 shows a slight
236 decreasing trend of uranium concentration from the EDJ to the root tip (from > 60 ppm to <
237 60 ppm), and corresponding increasing age estimates (56–62 ka) (Supplementary Table 2;
238 Supplementary Figure 5). This might result from a slight uranium leaching overprint; a
239 minimum age of 60 ka is likely for this tooth. The U-series age estimates obtained for
240 Sibrabang specimen 11564.5 show a decreasing trend from the EDJ toward the
241 circumpulpal dentine from 75 to 65 ka (Supplementary Table 2; Supplementary Figure 5.)
242 However, given the associated uncertainties, this trend might not be meaningful. An average
243 dentine U-series age of 70.3 ± 5.5 ka (2σ) may be regarded as a minimum age for the fossil,
244 which is broadly consistent with the single age estimate obtained from the enamel (64 ka). In
245 summary, the two teeth from Sibrabang yield U-series apparent ages of ~ 60–70 ka.

246 The uranium concentration measured across the dentine of the Niah Caves specimen
247 from grid Y/F4 shows little variability, 4.2–4.9 ppm. The U-series age estimates are between
248 6.0 and 8.7 ka (Supplementary Table 2; Supplementary Figure 6). The average dentine U-
249 series minimum age is 7.6 ± 1.3 ka. Similarly, the Niah Cave specimen from grid US/22
250 shows a consistent uranium concentration through the dentine (1.3–1.4 ppm), with relatively
251 large uncertainties that nonetheless bracket individual age estimates (Supplementary Table 2;
252 Supplementary Figure 6). The average dentine U-series age is 8.8 ± 3.0 ka. In summary, the
253 two teeth from Niah Cave yield consistent apparent ages of ~ 8 –9 ka, which should be
254 regarded as a minimum age constraint for the fossils.

255

256 **Discussion**

257 *Primate oxygen isotope compositions do not reveal a clear milk enrichment effect*

258 Half of our modern sample, and potentially all of our fossil sample, are composed of
259 M1s. These begin forming around birth and continue growing for three or more years (Smith,
260 2016). Orangutan infants rely exclusively on maternal milk during their first year of life,
261 supplementing this with solid foods in the second year, which are increased until suckling
262 ceases prior to eight or nine years of age (van Noordwijk et al., 2013; Smith et al., 2017). Our
263 developmentally-guided sampling approach allows us to examine fine-scaled trends in $\delta^{18}\text{O}$
264 values during birth, exclusive nursing, supplemental feeding, and also after nursing ends (in
265 those individuals with available serial molar teeth).

266 We find that five modern orangutans show only minor and inconsistently elevated
267 $\delta^{18}\text{O}$ values during the first year of life when compared to the subsequent year. These data do
268 not support the hypothesis that primate infants have markedly elevated body water $\delta^{18}\text{O}$
269 values during exclusive nursing. Data from the majority of 12 human M1s studied by
270 Vaiglova et al. (in review) similarly reveal maximum $\delta^{18}\text{O}$ values after the first year of tooth
271 formation, well beyond the duration of exclusive milk intake. This is also evident in the M1
272 of a Neanderthal born in the spring (Smith et al., 2018a); $\delta^{18}\text{O}$ values mostly rose for the first
273 3.5 months of life, but did not reach a maximum for another two years — long after the infant
274 would have begun consuming supplemental foods and liquids. This final dataset points to the
275 influence of season of birth on initial postnatal $\delta^{18}\text{O}$ values, as inferred in other mammals
276 (Bryant et al., 1996; Frick and O’Neil, 1996).

277 Comparisons of serially-forming teeth in four wild orangutans also fails to show a
278 consistent elevation of $\delta^{18}\text{O}$ values in M1s versus M2s (or M3s in two cases). Comparisons of

279 M1 $\delta^{18}\text{O}$ values with subsequent-forming teeth in four baboons, two tantalus monkeys, and
280 one mona monkey (from Green et al., 2022: SI Dataset S1) also largely fail to support the
281 enriched “Pattern 1” trend modelled by Bryant et al. (1996: Figure 4, p. 401). This is also the
282 case in comparisons of $\delta^{18}\text{O}$ values from bulk samples of human teeth—Wright and Schwartz
283 (1999) demonstrated that M1s have higher $\delta^{18}\text{O}$ values than later-forming teeth in only four
284 of seven individuals. In summary, the data from a range of primates, humans included, do not
285 support the exclusion of early-forming primate teeth from the assessment of environmental
286 seasonality.

287

288 *Modern orangutans show similar isotopic values across the islands of Borneo and Sumatra*

289 The two Bornean juveniles from the Munich collection (ZSM 1981/48, ZSM 1981/87)
290 reflect the environmental conditions of the late 1880s and early 1890s in Skalau—a region
291 where orangutans might now be locally extinct. Similarly, the teeth from the two Sumatran
292 individuals from the Munich collection (ZSM 1981/246, ZSM 1981/248) were collected prior
293 to 1939 in northern Aceh, from where orangutans also have since disappeared (Spehar et al.,
294 2018). While the individuals from northernmost Sumatra might have inhabited somewhat
295 higher elevations than those from western Borneo, there does not appear to be an evident
296 altitude effect (lower isotopic values at higher altitudes), as these four individuals show
297 similar isotopic values, save for a single brief excursion below 12 ‰ in ZSM 1981/248
298 (Table 1, Figure 3). It is unknown to what extent local rainfall may have been isotopically
299 distinct at the time the teeth were forming.

300 The $\delta^{18}\text{O}$ values shown in Figure 1 reflect estimates of monthly and annual average
301 precipitation from the Online Isotopes in Precipitation Calculator (3.0) compiled for
302 www.waterisotopes.org. Actual measurements of precipitation $\delta^{18}\text{O}$ from the islands of
303 Borneo and Sumatra are extremely limited. The closest observation facilities to the ZMS
304 orangutan locations yield similar patterns of modern annual rainfall $\delta^{18}\text{O}$ variability
305 (Belgaman et al., 2017), yet specific measurements from the six facilities that make up
306 “Cluster 3” in this reference are not available for comparison.

307 Other studies underscore the complexity of water transport in this region—multiple
308 factors such as the oceanic origin of water vapor, cloud cover and type, and the post-
309 condensation process influence the short-term variability of $\delta^{18}\text{O}$ values in rainfall (Moerman
310 et al., 2013; Suwarrman et al., 2013; Belgaman et al., 2016). For example, Moerman et al.
311 (2013) provided five years of daily rainfall $\delta^{18}\text{O}$ measurements from Northern Borneo
312 (Gunung Mulu National Park, Malaysia); daily rainfall $\delta^{18}\text{O}$ values ranged from +0.7 ‰ to

313 –18.5 ‰ and showed 1 – 3 month, annual, and supra-annual cycle frequencies. Interannual
314 rainfall $\delta^{18}\text{O}$ fluctuations of 6–8 ‰ were significantly correlated with ENSO events; these are
315 similar in scale to the large fluctuations in our serial tooth datasets (Figure 3).

316 Another potential source of isotopic variability derives from dietary variation, as
317 orangutans obtain the majority of their body water from plants (MacKinnon, 1974). Plant
318 oxygen isotope compositions can be stratified within tropical forest canopies (Sternberg et al.,
319 1989; Roberts et al., 2017; Lowry et al. 2021)—potentially leading to offset values amongst
320 various animals, including primates, that consume different resources in the same forest
321 (Krigbaum et al., 2013; Nelson, 2013; Fannin and McGraw, 2020). Orangutans forage at
322 different canopy heights ranging from the ground to high in the canopy (MacKinnon, 1974;
323 Ungar, 1996; Thorpe and Crompton, 2005; Ashbury et al. 2015). MacKinnon (1974) reported
324 that Bornean and Sumatran orangutans obtain 95 % of their food from the middle and upper
325 levels of the canopy, where preferred foods are most abundant. In contrast, Ungar (1996)
326 reported that Sumatran orangutans were quite variable in feeding heights, with a mean of
327 approximately 19 meters; lower than gibbons who fed preferentially in the high canopy.
328 Thorpe and Crompton (2005) reported stratification in Sumatran orangutans, with immature
329 individuals feeding below 20 meters, females feeding both below and above this height, and
330 adult/subadult males preferring to feed high in the canopy.

331 While differences in enamel $\delta^{18}\text{O}$ values are apparent in comparisons of sympatric
332 arboreal and terrestrial mammals (reviewed in Lowry et al. 2021; Green et al., 2022), it
333 remains to be seen whether primates with broadly similar diets and habitats show meaningful
334 differences in $\delta^{18}\text{O}$ values, and to what degree plant physiology influences the pattern and
335 amplitude of seasonality relative to rainfall. Oxygen isotope compositions in the six modern
336 individuals from the islands of Borneo and Sumatra are very similar. Orangutans from both
337 islands prefer ripe fruit when available, with some differences in the consumption of bark,
338 leaves, unripe fruits, and insects—which varies between sites and across seasons (reviewed in
339 Smith et al., 2012). Seasonal variation in diets and the stratification of food within the canopy
340 may also contribute to enamel oxygen isotope variation within individuals, in addition to the
341 seasonal rainfall trends we observe in our datasets. Orangutan $\delta^{18}\text{O}$ values are also quite
342 similar to the $\delta^{18}\text{O}$ values from five humans from Flores, Indonesia (14.8–21.0 ‰) dated at ~
343 2.2–3.0 ka (Vaiglova et al., in review). This is remarkable given the major dietary differences
344 between frugivorous orangutans and omnivorous coastal-dwelling humans, and suggests that
345 their enamel $\delta^{18}\text{O}$ values are predominantly influenced by regional precipitation.

346

347 *Fossil orangutan isotope values suggest different ancient climates in Sumatra and Borneo*
348 Dating studies at Lida Ajer have established the presence of the oldest human remains
349 in insular Southeast Asia, ~ 63–73 ka (Westaway et al., 2017), and a broad survey of the cave
350 has reconfirmed an age of MIS 4 (59–76 ka) for the mammalian fauna (Louys et al., 2022).
351 This is consistent with the minimum age of ~ 33–40 ka estimated for the two molars
352 examined in the current study. The Sumatran Sibrambang Cave has been regarded as roughly
353 contemporaneous to Lida Ajer given broad faunal similarities (de Vos, 1983). Recent U-
354 series dating of two fossil orangutans from the Sibrambang assemblage yielded minimum
355 ages of >56 ka and >85 ka (Louys et al. in press), which bracket the apparent U-series
356 minimum ages of ~ 60–70 ka in the current study. Sibrambang primates appear similar to, or
357 slightly older than, those from Lida Ajer, given the minimum U-series age estimates for teeth
358 from both sites, but this is not definitive given the absence of finite numerical ages for the
359 fossils. Our analysis of $\delta^{18}\text{O}$ values in Sumatran orangutan fossil molars reveals a close
360 similarity across sites and with modern Sumatran individuals, although the fossil
361 compositions fall at the upper end of the modern range. This may indicate a slightly dryer and
362 less variable climate during the late Pleistocene; elevated tooth $\delta^{18}\text{O}$ values are also indicative
363 of elevated values in hydrological systems globally, resulting from increased ice volumes at
364 in glaciers and at the poles.

365 Pollen records from the Niah Caves archaeological site indicate that there were a
366 number of local ecological shifts from lowland rainforest to more open environments during
367 the Late Pleistocene and into the Holocene (Hunt et al., 2012), where humans may have
368 begun hunting orangutans at ~ 45 ka (Spehar et al., 2018). While it is not possible to locate
369 the two fossil orangutan molars in these pollen records, Piper and Rabett (2016) considered
370 that the large animal bone assemblages accumulated within the Lobang Hangus entrance and
371 defined by the Harrisson spit depths of 12"- 42" were of terminal Pleistocene age. More
372 broadly, the orangutan specimen from grid US/22 (32"-36") is stratigraphically positioned
373 between radiocarbon ages of 14,206–15,061 cal. BP (OxA-13936) and 36,583–38,059 cal.
374 BP (OxA-13938), and this provides plausible minimum and maximum age constraints that
375 are not incompatible with the apparent minimum U-series age of ~ 9 ka. Based on these
376 results, the tooth is likely to date from the latest part of the Late Pleistocene. The specimen
377 from grid Y/F4 might date from the latest part of the Late Pleistocene to the early Holocene,
378 by comparison with the shell and fauna assemblage from other excavated areas (Piper et al.,
379 2016.)

380 Both orangutan molars from the Niah Caves yield wide ranges of $\delta^{18}\text{O}$, which is
381 particularly notable given the short periods of time sampled compared to the other fossils and
382 most modern orangutan molars. Given the similar offsets in $\delta^{18}\text{O}$ values between modern
383 baboons living in Ugandan forests and the Ethiopian rift region (Green et al., 2022) and
384 modern and prehistoric Bornean orangutans, we regard the higher $\delta^{18}\text{O}$ values in the Niah
385 Cave orangutans as possibly indicative of reduced rainfall when compared to recent
386 conditions. This is consistent with paleoclimate reconstructions for Borneo and Flores during
387 the late Pleistocene and early Holocene (Griffith et al. 2009; Buckingham et al., 2022), when
388 the environment around the Niah Caves is believed to have been a drier, more open seasonal
389 forest (Harrison, 1996; Hunt et al., 2012). A study of $\delta^{18}\text{O}$ values in Niah Caves shell
390 middens dating from the early to mid- Holocene indicates a shift to periods of high rainfall
391 with less variation than modern conditions (Stephens et al., 2016). The transition from a drier
392 environment to moist tropical rainforest is also reflected in the increasing number and higher
393 frequencies of canopy-adapted mammalian taxa in excavated layers of the Pleistocene-
394 Holocene transition (Piper and Lim, 2021).

395 Our approach has the potential to contribute to reconstructions of ancient
396 paleoenvironments in SE Asia based on studies of pollen, molluscs, faunal community
397 compositions, guano records, and stable isotopes of teeth (e.g., Jablonski et al., 2000; Bird et
398 al., 2005; Louys and Meijaard, 2010; Wurster et al., 2010; Hunt et al., 2012; Janssen et al.,
399 2016; Stephens et al., 2016; Louys and Roberts, 2020; Bacon et al., 2021; Louys et al., 2022,
400 in press). This may be especially timely given that recent work examining modern fauna
401 compositions in African landscapes has cautioned that fossil herbivore assemblages tend to
402 overestimate the extent of ancient grasslands in comparison to woodlands (Negash and Barr,
403 2023; also see Sokolowski et al. 2023). Fine-scaled tooth sampling may also allow an
404 expansion of inferences from $\delta^{18}\text{O}$ values of bulk-sampled Asian hominin remains (Janssen et
405 al., 2016; Roberts et al., 2020; Kubat et al., 2023), which are difficult to interpret for
406 understanding seasonal rainfall dynamics in tropic environments (Green et al., 2022). Such
407 information could better inform debates about whether humans employed arid savannah
408 corridors to avoid dense tropical forests, or whether humans were adept at colonizing such
409 environments during their consequential migration throughout island Southeast Asia.

410
411 **Materials and Methods**

412 *Orangutan samples*

413 Thin (histological) sections of twelve molar teeth from six modern orangutans and six
414 molar teeth from five fossil orangutans were employed (Table 1). These sections were
415 previously prepared for studies of incremental tooth development, enamel thickness,
416 elemental chemistry, and Asian hominoid taxonomy (Smith, 2016; Smith et al., 2011, 2012,
417 2017, 2018b). Four modern individuals were sourced from the Munich State Anthropological
418 Collection (ZSM); two were collected in 1893-1894 from Skalau (north of the Kapuas River
419 and south of the Klingkang Mountains in eastern West Borneo), and two were collected prior
420 to 1939 from Aceh (northwest Sumatra) (Röhrer-Ertl, 1988: Figure 3, p. 14) (Figure 1). It
421 was not possible to determine from which specific regions or time periods the two other
422 modern individuals derive—collection notes were not available for these specimens from the
423 Harvard Museum of Natural History (MCZ) or the Humboldt Museum (ZMB). Ages at death
424 were determined for five of six individuals from assessments of incremental features and
425 elemental registration of serially forming molars (detailed in Smith, 2016; Smith et al., 2017).

426 We also studied four Sumatran fossil orangutan teeth that were collected more than a
427 century ago from the Lida Ajer and Sibrambah Caves in the Padang Highlands by Eugene
428 Dubois (de Vos, 1983). Right and left lower molars from Lida Ajer (11594.12, 11595.105)
429 show identical trace element patterns in their dentine (Supplementary Figure 7), as well as
430 similar occlusal fissure patterns and light wear, consistent with their attribution to the same
431 individual. Two Bornean fossil orangutan teeth from Niah Caves (Malaysia) were also
432 included in this study. The caves have yielded significant late Pleistocene and early Holocene
433 human remains since the Harrissons began excavations in the 1950s (Barker et al., 2007).
434 These lower molars were derived from two different entrances to the cave system, Gan Kira
435 (grid square Y/F4) and Lobang Angus/Hangus (grid square US/22), with burial depths of 12–
436 18 inches and 30–36 inches, respectively (Hooijer, 1961). Although Hooijer (1948, 1961)
437 identified all six of these fossil teeth as M1s, we regard this as tentative, given that isolated
438 orangutan molars are notoriously difficult to seriate (Grine and Franzen, 1994).

439

440 *Dating of fossil samples*

441 Preliminary assessments at the Australian National University Radiocarbon Dating
442 Laboratory confirmed that collagen preservation in the six fossil orangutans was insufficient
443 for radiocarbon dating, a common limitation in tropical environments (e.g., Wood et al.,
444 2016). Laser ablation uranium series (U-series) analyses were carried out on longitudinal
445 sections of teeth at the Radiogenic Isotope Facility of the University of Queensland using an
446 ASI RESolution SE laser ablation system connected to a Nu Plasma II MC-ICP-MS. A

447 succession of several rasters (< 2-minute linear ablations) was made in a transect across the
448 dentine and enamel of each tooth (Supplementary Figures 4-6) following Grün et al. (2014).
449 The $^{230}\text{Th}/^{238}\text{U}$ and $^{234}\text{U}/^{238}\text{U}$ activity ratios of the samples were normalized to bracketing
450 analyses of a homogeneous rhino tooth standard that has been precisely calibrated by isotope
451 dilution (Grün et al., 2014). Importantly, dental tissues are known to behave as open systems
452 for U-series elements; provided there is no occurrence of uranium leaching, age estimates
453 should therefore be regarded as minimum age constraints since uranium uptake into dental
454 tissues may be significantly delayed after death.

455

456 *Tooth formation and oxygen isotope analyses*

457 Thin sections were first imaged with transmitted light microscopy. Enamel daily
458 secretion rates were measured between sequential accentuated growth lines to yield the time
459 of formation (see Smith, 2016: Fig. 1, p. 94), and enamel extension rates were calculated
460 between accentuated lines to guide placement of the analysed spots at approximately weekly
461 intervals of growth from the dentine horn tip to the enamel cervix (Smith et al., 2018a; Green
462 et al., 2022). Following the removal of cover slips by immersion in xylene, each thin section
463 was analysed for $\delta^{18}\text{O}$ at the SHRIMP Laboratory at the Australian National University
464 according to methods detailed in Vaiglova et al. (in review).

465 In brief, a 15 kV Cs primary ion beam focused to a spot $\sim 15 \times 20 \mu\text{m}$ diameter was
466 used to sequentially sample the enamel as close as possible to the enamel-dentine junction
467 (EDJ). Oxygen secondary ions were extracted at 10 kV and analysed isotopically by a
468 multiple collector equipped with dual electrometers operated in resistor mode. The $\delta^{18}\text{O}$
469 values were calculated relative to reference apatite (Durango 3) measured every 10–15
470 sample analyses. Distances of SHRIMP $\delta^{18}\text{O}$ measurements along the innermost enamel from
471 the cusp to cervix were converted to secretory time in days following Green et al. (2022). A
472 polynomial regression relating distances to days was created using the enamel extension
473 rates, and this regression was applied to estimate the timing of secretory deposition at every
474 SHRIMP spot location. The Lomb–Scargle periodogram was used to assess time-dependent
475 patterns of $\delta^{18}\text{O}$ values, which estimates the power of sine wave periods within a given range
476 to produce the temporal patterns present within those measurements.

477 The probability that differences between first and second year $\delta^{18}\text{O}$ values in modern
478 first molars might have arisen by chance was assessed by one-way paired t-tests, with alpha =
479 0.05 adjusted by Bonferroni correction due to repeated comparisons across multiple teeth.

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482

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506

507

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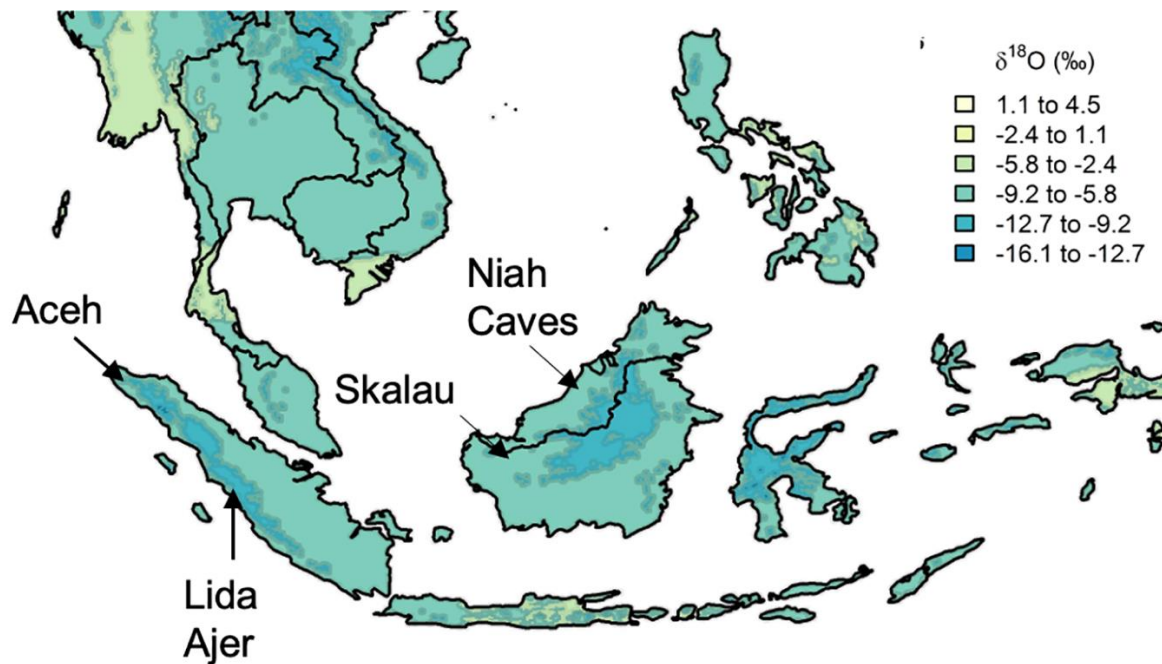
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942 **Figures**

943

944 Figure 1. Approximate location of select modern and fossil orangutans superimposed on
945 modeled isotopic variation.

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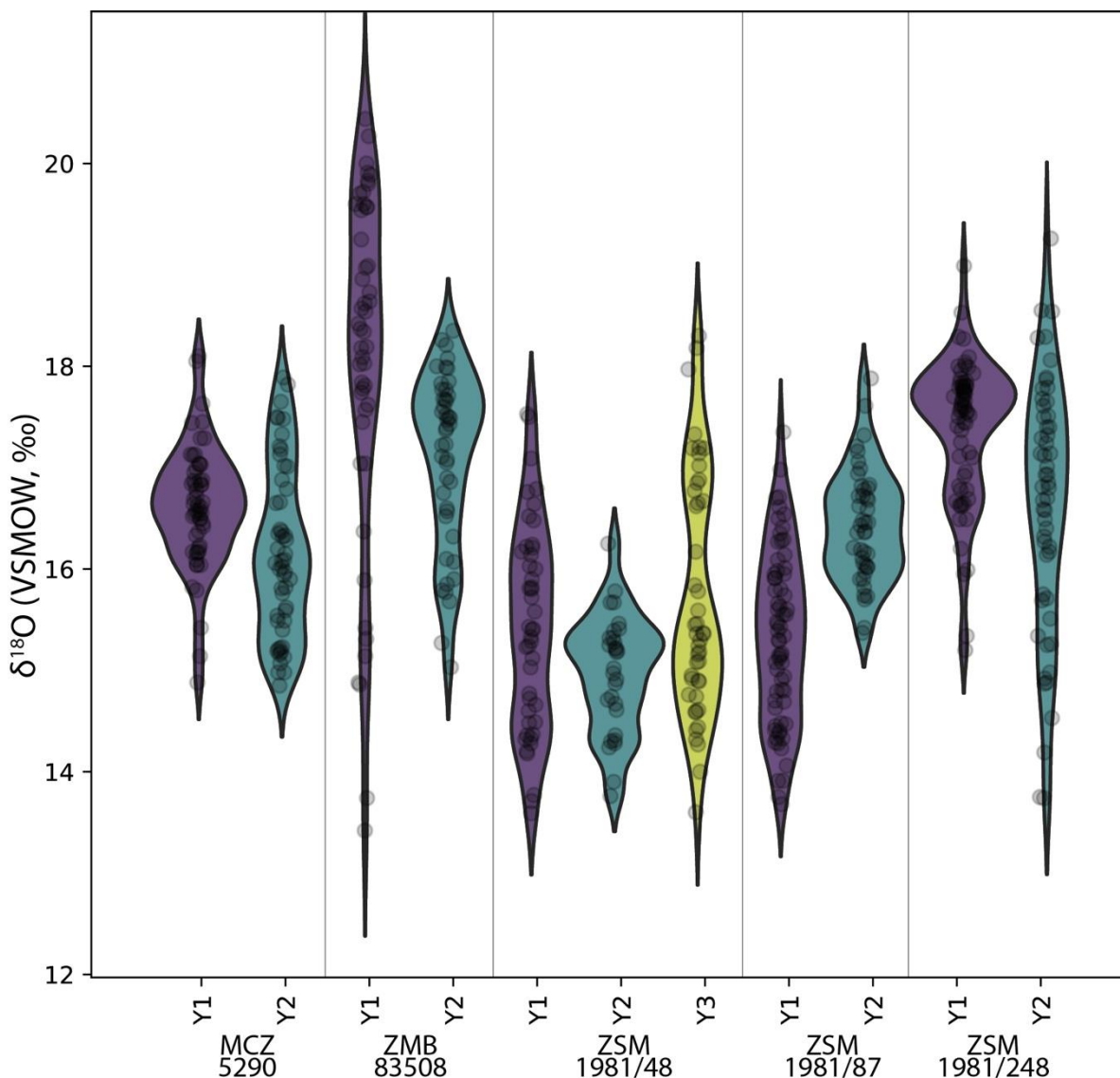
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950 Figure modified from www.waterisotopes.org based on data from the Online Isotopes in
951 Precipitation Calculator (3.0). See Table 1 for the location of particular individuals.

952 Sibrabang Cave has yet to be relocated since Eugene Dubois' original excavations, but it is
953 known to be in the general vicinity of Lida Ajer in the Padang Highlands, possibly near to the
954 modern village of a similar name (Louys et al. in press).

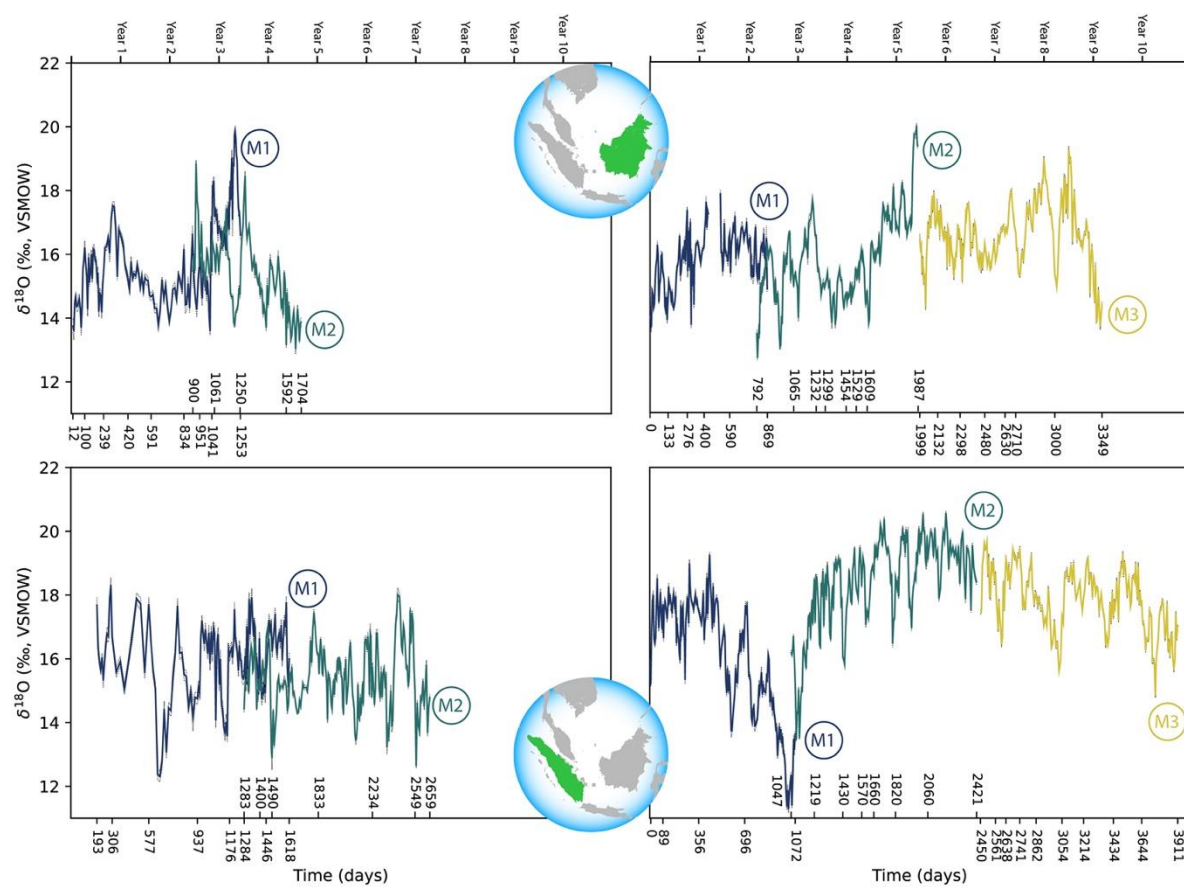
955

956 Figure 2. Comparison of sequential $\delta^{18}\text{O}$ values across multiple years of first molar formation
957 in five modern orangutans from Borneo and Sumatra.
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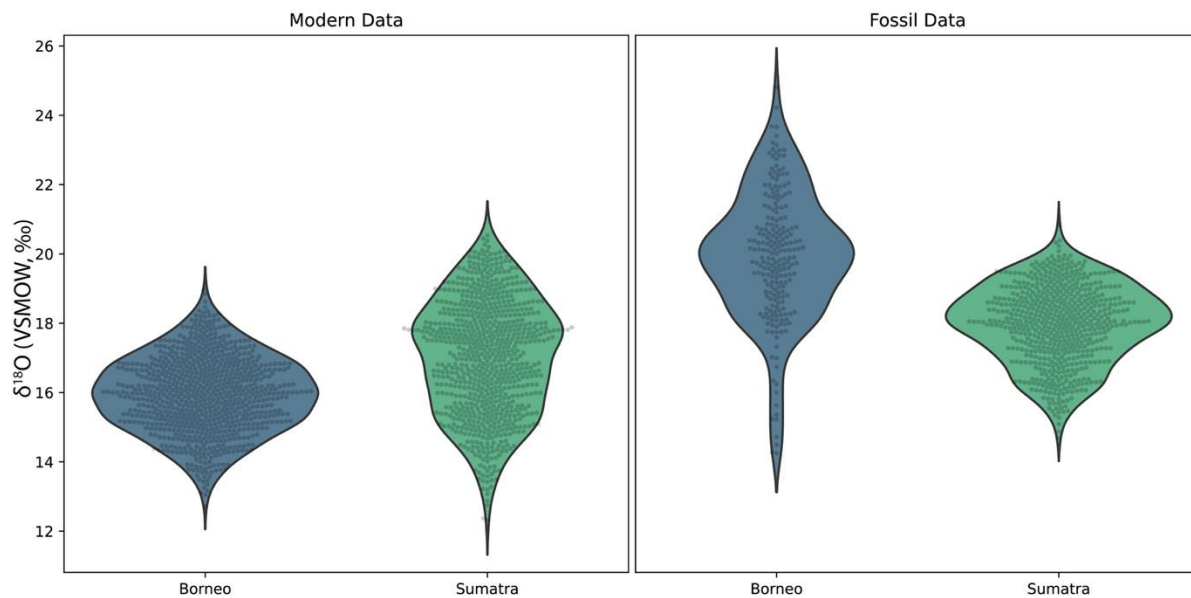
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961 Bornean individuals: MCZ 5290, ZSM 1981/48 ZSM 1981/87; Sumatran individuals: ZMB
962 83508, ZSM 1981/248. The width of each curve is a kernel density estimate (KDE)
963 corresponding to the distribution of $\delta^{18}\text{O}$ values. First year data (Y1) is shown with a purple
964 violin plot, second year data (Y2) with a green plot, and third year data (Y3) with a yellow
965 plot where complete/available. Actual data are plotted as black circles.
966

967 Figure 3. Comparison of sequential $\delta^{18}\text{O}$ values across multiple years of serial molar
968 formation in two modern orangutans from Borneo (top) and two from Sumatra (bottom).
969



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974 Individual in upper left: ZSM 1981/48; upper right: ZSM 1981/87; lower left: ZSM
975 1981/246; lower right: ZSM 1981/248. Developmental overlap was determined through
976 registration of trace elements as in Smith et al. (2017).
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982 Figure 4. Comparison of $\delta^{18}\text{O}$ values in fossil and modern orangutans from Borneo (blue) and
983 Sumatra (green).
984



985
986 Violin plots show kernel density estimates representing the distribution of $\delta^{18}\text{O}$ values in
987 modern individuals (left plot), and in fossil individuals (right plot). Actual $\delta^{18}\text{O}$
988 measurements are shown as black circles.
989

990 Table 1. Modern and fossil orangutan teeth employed in the current study.

991

Taxon	Accession	Origin	Sex	Age (Years)	Teeth
<i>Pongo pygmaeus</i>	ZSM 1981/48	Skalau, Borneo	F	~8.4	RUM1, LLM2
	ZSM 1981/87	Skalau, Borneo	F	> 9	LUM1, RUM2, RLM3
	MCZ 5290	Borneo (location unspecified)	n/a	4.5	RUM1
<i>Pongo abelii</i>	ZSM 1981/246	Aceh, Sumatra	M	~8.5	LLM1, LUM2
	ZSM 1981/248	Aceh, Sumatra	F	adult	LUM1, LUM2, LLM3
	ZMB 83508	Sumatra (location unspecified)	n/a	8.8	RLM1
Fossil <i>Pongo</i> spp.	11564.5	Sibrabang, Sumatra	n/a	n/a	RUM
	11565.162	Sibrabang, Sumatra	n/a	n/a	LUM
	11594.12	Lida Ajer, Sumatra	n/a	n/a	RLM
	11595.105	Lida Ajer, Sumatra	n/a	n/a	LLM
	US/22	Niah Caves, Malaysia	n/a	n/a	RLM
	Y/F4	Niah Caves, Malaysia	n/a	n/a	LLM

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994

995 Numerous taxonomic assignments have been made for fossil orangutans (*Pongo* spp.), some
996 of which have not been based on clear morphological characteristics (Lim, 2016), and are not
997 relevant for the focus of this paper.

998

999 Table 2. Modern and fossil orangutan molar $\delta^{18}\text{O}$ values.

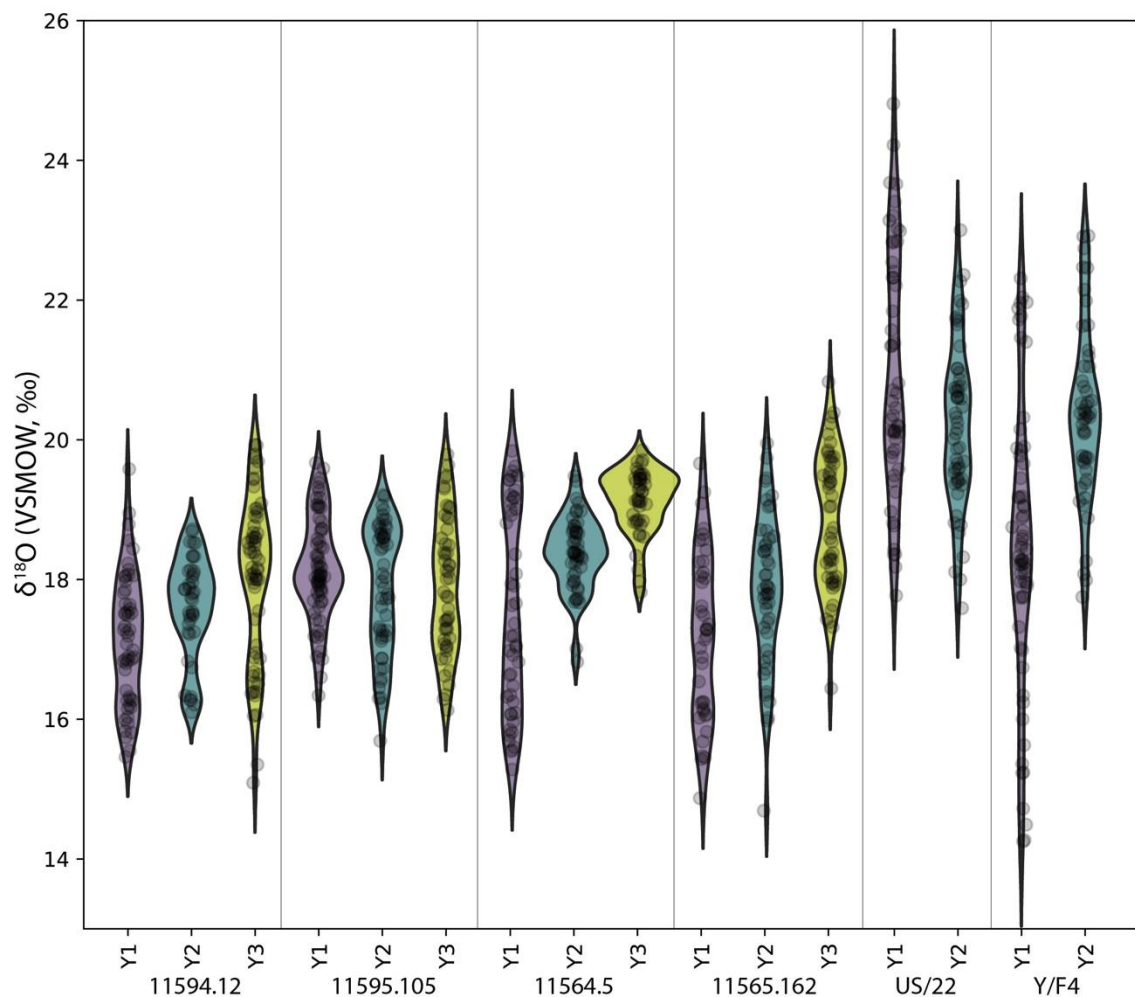
1000

Taxon	Accession	Tooth	Cusp	Spots	Time (Days)	dO18 Range
<i>Pongo pygmaeus</i>	ZSM 1981/48	RUM1	dl	151	1241	13.6-19.9
	ZSM 1981/48	LLM2	mb	107	804	13.0-18.8
	ZSM 1981/87	LUM1	ml	131	869	13.7-18.2
	ZSM 1981/87	RUM2	ml	196	1195	12.7-20.0
	ZSM 1981/87	RLM3	mb	220	1350	13.7-19.2
	MCZ 5290	RUM1	ml	150	1002	13.8-18.1
<i>Pongo abelii</i>	ZSM 1981/246	LLM1	mb	136	1425	12.3-18.3
	ZSM 1981/246	LUM2	ml	229	1376	12.6-18.0
	ZSM 1981/248	LUM1	db	177	1072	11.3-19.3
	ZSM 1981/248	LUM2	db	193	1374	13.5-20.6
	ZSM 1981/248	LLM3	db	191	1461	15.2-21.2
	ZMB 83508	RLM1	db	135	1029	13.4-20.4
Fossil <i>Pongo</i> spp.	11564.5	RUM	mb	178	1387	15.3-20.4
	11565.162	LUM	ml	143	1144	14.7-20.8
	11594.12	RLM	ml	154	1081	15.1-19.9
	11595.105	LLM	mb	197	1312	15.7-20.0
	US/22	RLM	mb	149	1023	15.9-24.8
	Y/F4	LLM	db	134	869	14.2-22.9

1001

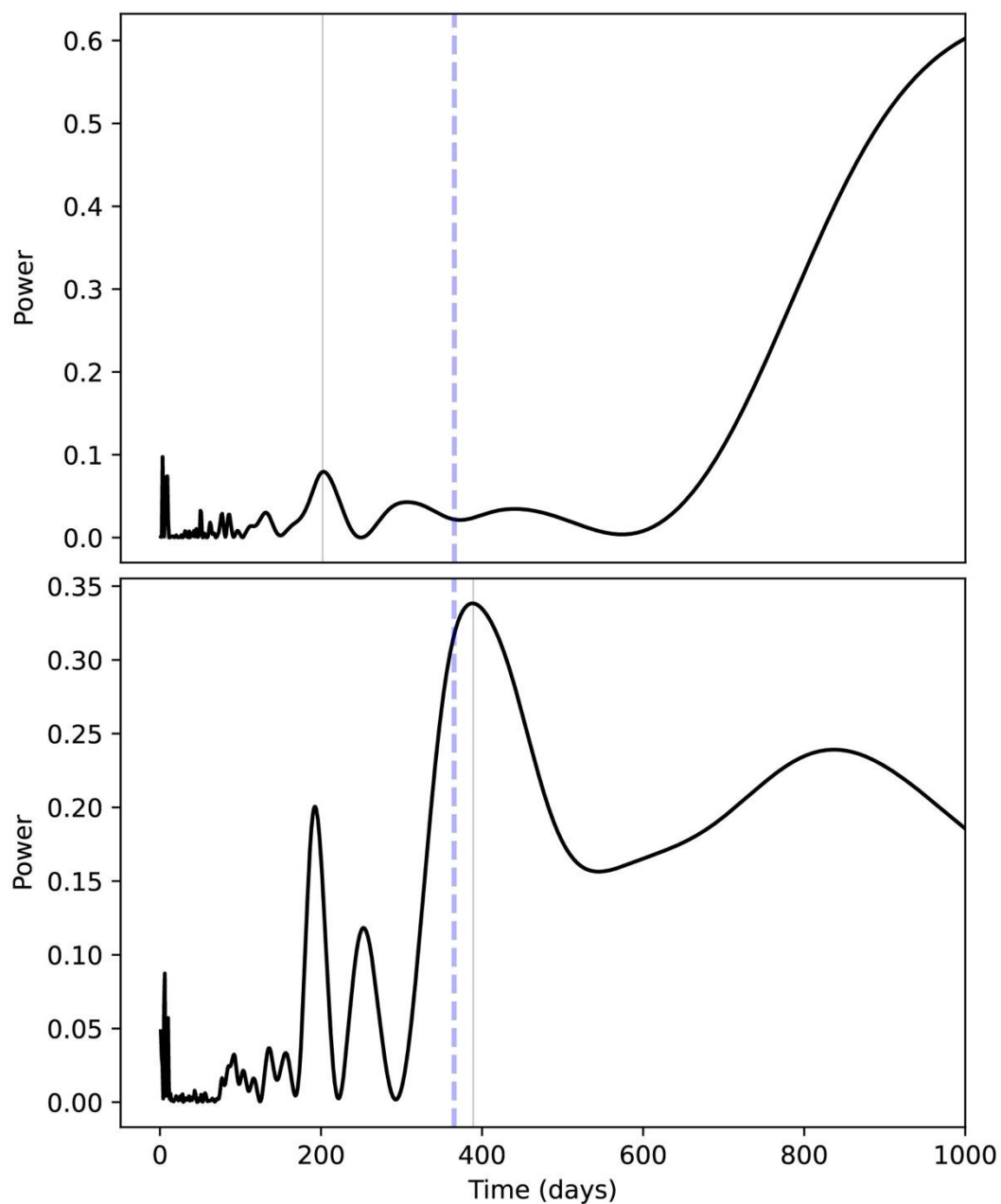
1002

1003 Supplementary Figure 1. Comparison of sequential $\delta^{18}\text{O}$ values across multiple years of
1004 molar formation in six putative fossil orangutan M1s from Borneo and Sumatra.
1005

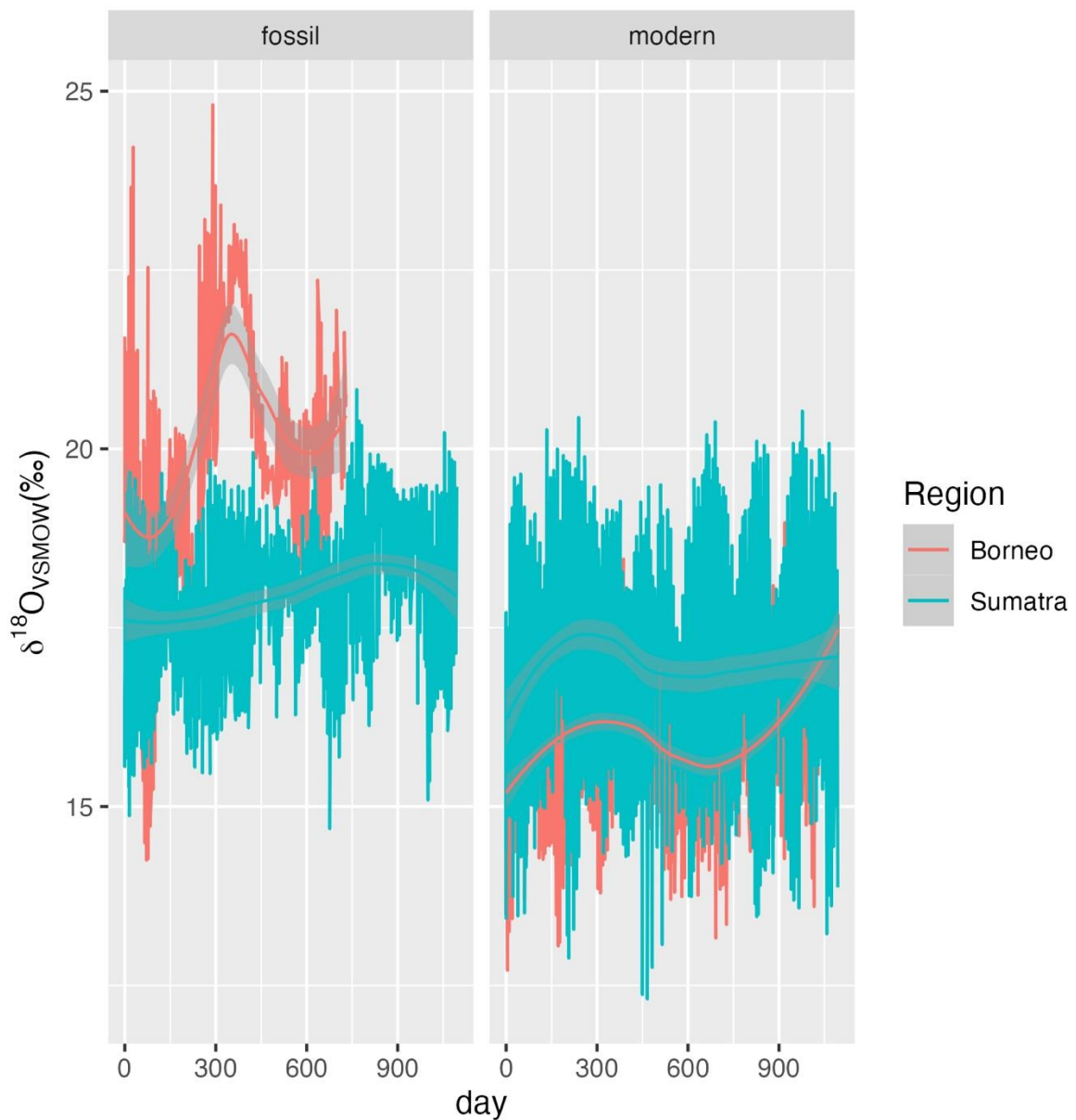


1006
1007
1008 The width of each curve is a kernel density estimate (KDE) corresponding to the distribution
1009 of oxygen isotope values measured from different teeth. From each tooth, first year data (Y1)
1010 is shown with a purple violin plot, second year data (Y2) with a green plot, and third year
1011 data (Y3, if present) with a yellow plot. Actual data are plotted as black circles.

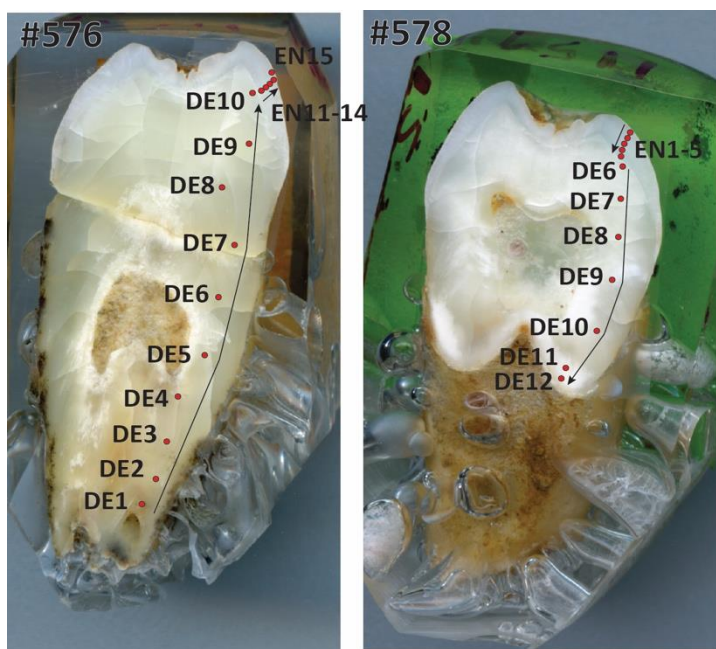
1012 Supplementary Figure 2. Inferred seasonality of $\delta^{18}\text{O}$ values from a Bornean (top) and
1013 Sumatran (bottom) M1.



1025 Supplementary Figure 3. Comparison of $\delta^{18}\text{O}$ values in fossil and modern orangutans from
1026 Borneo and Sumatra.



1029 Supplementary Figure 4. Laser ablation profiles performed across the two teeth from Lida
1030 Ajer.

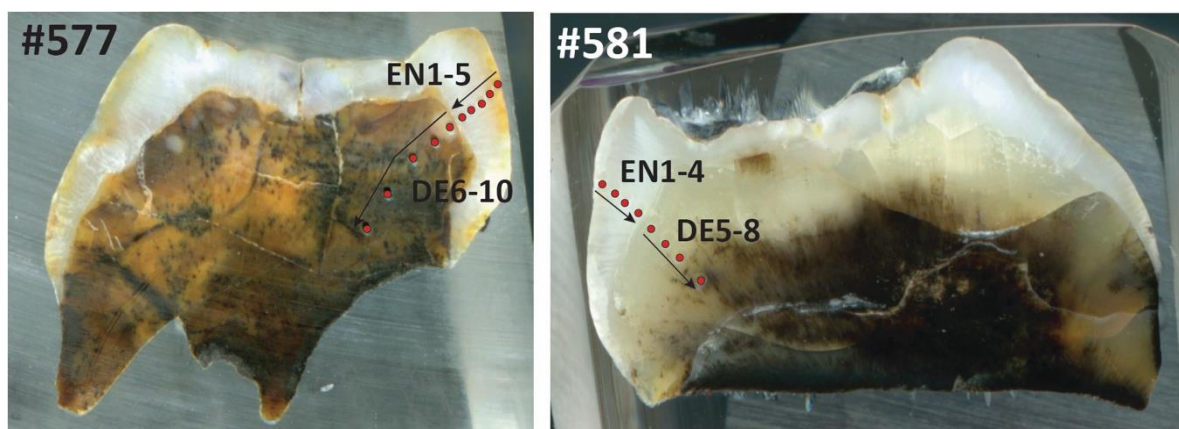


1031

1032 #576 (left) refers to specimen 11595.105; #578 (right) refers to specimen 11594.12. The red
1033 dots represent the position of the rasters, and arrows indicate the sequence of the analyses.
1034 EN = enamel. DE = dentine.

1035

1036 Supplementary Figure 5. Laser ablation profiles across the two teeth from Sibrambah Cave.

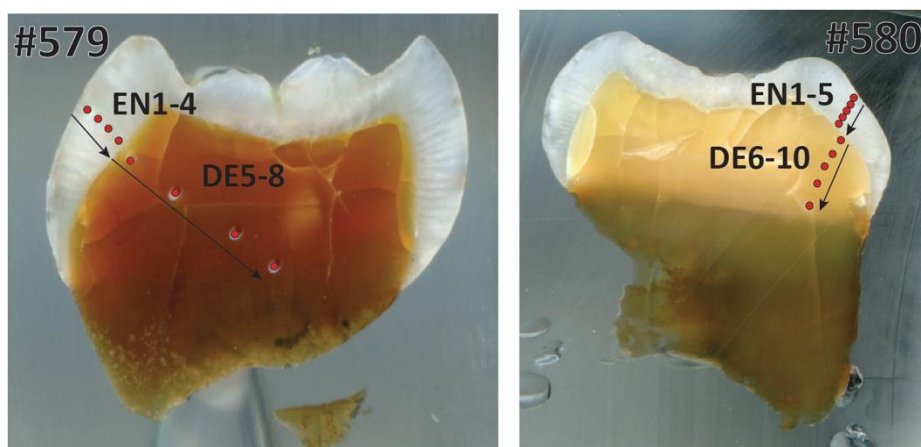


1037

1038 #577 (left) refers to specimen 11565.162; #581 (right) refers to specimen 11564.5. The red
1039 dots represent the position of the rasters, and arrows indicate the sequence of the analyses.
1040 EN = enamel. DE = dentine.

1041

1042 Supplementary Figure 6. Laser ablation profiles across the two teeth from Niah Caves.



1043

1044 #579 (left) refers to specimen from grid Y/F4; #580 (right) refers to specimen from grid
1045 US/22. The red dots represent the position of the rasters, and arrows indicate the sequence of
1046 the analyses. EN = enamel. DE = dentine.

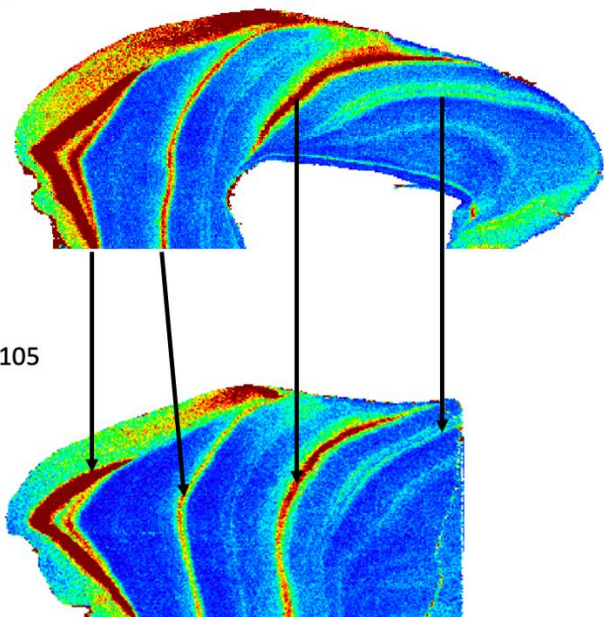
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1048

1049 Supplementary Figure 7. Matching trace element patterns in cross-sections of two isolated
1050 molars from the Dubois collection of fossil orangutan teeth from Lida Ajer.
1051

11594.12

11595.105



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1054 High concentrations are shown in warm colors, low concentration are in cool colors; here
1055 Li/Ca is shown, but identical corresponding patterns were also observed for Ba/Ca and Sr/Ca
1056 (not shown). The enamel cap of each tooth is to the left, and root dentine is to the right. Trace
1057 elements were measured according to LA-ICP-MS methods detailed in Smith et al. (2017).
1058

1059 Supplementary Table 1. Comparisons of first and second year $\delta^{18}\text{O}$ values in five first molars.
1060

Specimen	Adjusted p-values	Higher $\delta^{18}\text{O}$ values
MCZ 5290	p = 0.010	Year 1
ZMB 83508	p = 0.006	Year 1
ZSM 1981/48	p = 0.161 (N.S.)	Year 1
ZSM 1981/87	p < 0.001	Year 2
ZSM 1981/248	p < 0.001	Year 1

1061
1062

1063 Supplementary Table 2. U-series dates for fossil orangutan material.

1064

LA raster	Tissue	U(ppm)	$^{230}\text{Th}/^{238}\text{U}$	\pm	$^{234}\text{U}/^{238}\text{U}$	\pm	Age (ka)	\pm
11595.105-EN11	enamel	0.02	-1.230	0.090	0.459	0.037	n.c.	n.c.
11595.105-EN12	enamel	0.00	-12.82	0.084	-4.590	0.018	n.c.	n.c.
11595.105-EN13	enamel	0.00	-31.04	0.082	-17.11	0.018	n.c.	n.c.
11595.105-EN14	enamel	0.10	0.623	0.168	1.155	0.105	n.c.	n.c.
11595.105-EN15	enamel	0.04	-0.826	0.082	0.752	0.051	n.c.	n.c.
11595.105-DE1	dentine	41.4	0.516	0.020	1.360	0.047	50.6	3.40
11595.105-DE2	dentine	44.2	0.470	0.020	1.359	0.055	45.2	3.33
11595.105-DE3	dentine	48.6	0.471	0.015	1.344	0.045	46.0	2.65
11595.105-DE4	dentine	58.9	0.481	0.015	1.356	0.038	46.6	2.50
11595.105-DE5	dentine	61.1	0.482	0.019	1.341	0.045	47.5	3.10
11595.105-DE6	dentine	65.8	0.446	0.019	1.346	0.051	43.0	3.04
11595.105-DE7	dentine	64.0	0.440	0.016	1.346	0.044	42.3	2.55
11595.105-DE8	dentine	60.0	0.430	0.016	1.338	0.046	41.4	2.56
11595.105-DE9	dentine	64.6	0.433	0.018	1.336	0.049	41.8	2.81
11595.105-DE10	dentine	65.9	0.411	0.016	1.325	0.047	39.7	2.53

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LA raster	Tissue	U(ppm)	$^{230}\text{Th}/^{238}\text{U}$	\pm	$^{234}\text{U}/^{238}\text{U}$	\pm	Age (ka)	\pm
11594.12-EN1	enamel	0.00	-13.08	0.082	-9.394	0.019	n.c.	n.c.
11594.12-EN2	enamel	0.00	-23.91	0.104	-11.05	0.016	n.c.	n.c.
11594.12-EN3	enamel	0.00	-22.35	0.078	-4.621	0.027	n.c.	n.c.
11594.12-EN4	enamel	0.00	-16.49	0.089	-6.110	0.015	n.c.	n.c.
11594.12-EN5	enamel	0.01	17.58	0.103	16.21	0.031	224	3.12
11594.12-DE6	dentine	30.9	0.337	0.016	1.335	0.041	31.2	2.02
11594.12-DE7	dentine	30.6	0.347	0.013	1.337	0.036	32.2	1.68
11594.12-DE8	dentine	31.9	0.355	0.017	1.332	0.049	33.3	2.31
11594.12-DE9	dentine	31.4	0.366	0.013	1.335	0.035	34.4	1.78
11594.12-DE10	dentine	32.1	0.359	0.011	1.329	0.038	33.8	1.71
11594.12-DE11	dentine	24.6	0.344	0.014	1.357	0.031	31.4	1.67
11594.12-DE12	dentine	24.9	0.352	0.015	1.351	0.032	32.3	1.85

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LA raster	tissue	U(ppm)	$^{230}\text{Th}/^{238}\text{U}$	\pm	$^{234}\text{U}/^{238}\text{U}$	\pm	Age (ka)	\pm
11565.162-EN1	enamel	0.58	0.434	2.630	1.025	1.495	59.7	496
11565.162-EN2	enamel	0.00	-4.899	0.132	-81.42	0.020	n.c.	n.c.
11565.162-EN3	enamel	0.00	-23.35	0.075	-16.84	0.018	n.c.	n.c.
11565.162-EN4	enamel	0.00	-37.09	0.082	-22.74	0.021	n.c.	n.c.
11565.162-EN5	enamel	0.01	-5.283	0.068	-0.022	0.029	n.c.	n.c.
11565.162-DE6	dentine	62.2	0.438	0.287	1.073	0.470	56.5	59.4
11565.162-DE7	dentine	63.6	0.461	0.023	1.066	0.048	61.2	5.73
11565.162-DE8	dentine	61.4	0.467	0.021	1.072	0.044	61.7	5.23
11565.162-DE9	dentine	59.8	0.470	0.028	1.073	0.060	62.2	7.01
11565.162-DE10	dentine	57.4	0.480	0.026	1.072	0.051	64.2	6.51

LA raster	tissue	U(ppm)	$^{230}\text{Th}/^{238}\text{U}$	\pm	$^{234}\text{U}/^{238}\text{U}$	\pm	Age (ka)	\pm
11564.5-EN1	enamel	0.11	0.304	0.181	0.959	0.086	41.6	30.6
11564.5-EN2	enamel	<i>0.00</i>	-2215	0.086	-486.53	0.018	n.c.	n.c.
11564.5-EN3	enamel	<i>0.00</i>	-97.95	0.065	-4.236	0.031	n.c.	n.c.
11564.5-EN4	enamel	9.86	0.502	0.211	1.115	0.162	64.3	38.8
11564.5-DE5	dentine	51.6	0.564	0.055	1.116	0.052	75.4	11.8
11564.5-DE6	dentine	45.8	0.541	0.057	1.111	0.047	71.6	11.6
11564.5-DE7	dentine	49.1	0.530	0.039	1.117	0.079	69.0	10.2
11564.5-DE8	dentine	47.8	0.512	0.010	1.128	0.016	65.0	2.13

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LA raster	tissue	U(ppm)	$^{230}\text{Th}/^{238}\text{U}$	\pm	$^{234}\text{U}/^{238}\text{U}$	\pm	Age (ka)	\pm
Y/F4-EN1	enamel	<i>0.01</i>	-4.737	0.084	-0.283	0.025	n.c.	n.c.
Y/F4-EN2	enamel	<i>0.00</i>	-42.98	0.070	-2.119	0.024	n.c.	n.c.
Y/F4-EN3	enamel	<i>0.00</i>	-137.8	0.078	-18.50	0.017	n.c.	n.c.
Y/F4-EN4	enamel	<i>0.00</i>	-49.66	0.079	3.398	0.033	n.c.	n.c.
Y/F4-DE5	dentine	4.41	0.058	0.016	1.082	0.031	5.97	1.74
Y/F4-DE6	dentine	4.93	0.072	0.018	1.057	0.034	7.69	1.99
Y/F4-DE7	dentine	4.80	0.077	0.018	1.088	0.031	7.95	1.93
Y/F4-DE8	dentine	4.15	0.085	0.041	1.102	0.065	8.74	4.43

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LA raster	tissue	U(ppm)	$^{230}\text{Th}/^{238}\text{U}$	\pm	$^{234}\text{U}/^{238}\text{U}$	\pm	Age (ka)	\pm
US/22-EN1	enamel	<i>0.00</i>	-7.442	0.102	-6.047	0.023	n.c.	n.c.
US/22-EN2	enamel	<i>0.00</i>	-53.99	0.082	-14.98	0.020	n.c.	n.c.
US/22-EN3	enamel	<i>0.00</i>	215.0	0.103	78.83	0.019	n.c.	n.c.
US/22-EN4	enamel	<i>0.00</i>	2.054	0.109	-12.69	0.020	n.c.	n.c.
US/22-EN5	enamel	0.02	-2.955	0.109	1.044	0.052	n.c.	n.c.
US/22-DE6	dentine	1.28	0.066	0.070	1.205	0.100	6.09	6.66
US/22-DE7	dentine	1.36	0.085	0.073	1.179	0.086	8.10	7.22
US/22-DE8	dentine	1.38	0.094	0.060	1.183	0.071	8.94	5.98
US/22-DE9	dentine	1.41	0.126	0.062	1.183	0.087	12.27	6.38
US/22-DE10	dentine	1.41	0.090	0.050	1.234	0.080	8.17	4.77

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1070 $^{230}\text{Th}/^{238}\text{U}$ and $^{234}\text{U}/^{238}\text{U}$ are activity ratios. It is worth noting that, for most transect analyses, 1071 the ^{232}Th signal, which was measured on a Faraday collector, was indistinguishable from 1072 background noise. In this regard, the corresponding $^{230}\text{Th}/^{232}\text{Th}$ activity ratio of each transect 1073 should be $>>100$, and thus non-radiogenic or detrital ^{230}Th correction would have negligible 1074 impact on the age. U-series data in italics should be viewed with caution due to U 1075 concentrations of ≤ 0.5 ppm. All errors are 2σ . Key: EN= enamel; DE = dentine; n.c. = not 1076 calculable. Negative values were caused by background extraction from their measured peaks 1077 with intensities at detection levels.