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Astronomically controlled aridity in the Sahara since at least

2 11 million years ago

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- 15

16 Abstract

The Sahara is the largest hot desert on Earth. Yet, the timing of its inception and 17 its response to climatic forcing is debated, leading to uncertainty over the causes 18 19 and consequences of regional aridity. Here we present detailed records of 20 terrestrial inputs from Africa to North Atlantic deep-sea sediments, documenting a long and sustained history of astronomically-paced oscillations between a 21 humid and arid Sahara from over 11 million years (Myr) ago. We show that 22 intervals of strong dust emissions from the heart of the continent predate both 23 24 the intensification of northern hemisphere glaciation and the oldest land-based evidence for a Saharan Desert by millions of years. We find no simple long-term 25 gradational transition towards an increasingly arid climate state in northern 26 27 Africa, suggesting that aridity was not the primary driver of gradual Neogene 28 expansion of African savanna C₄ grasslands. Instead, insolation-driven wet-dry shifts in Saharan climate were common over the last 11 Myr and we identify 29 30 three distinct stages in the sensitivity of this relationship. Our data provide context for evolutionary outcomes on Africa, for example, we find that 31 32 astronomically-paced arid intervals predate the oldest fossil evidence of hominid bipedalism by at least 4 Myr. 33

34

35 Main text

Prominent from space, the Saharan desert is a vast, bare, intensely arid, dust-exporting landscape that grew 10% in response to climatic forcing during the 20th century¹. A recognisable Saharan desert is widely suggested to originate in the late Pliocene, ~2.6 Myr ago, coupled to the development of major continental ice sheets in the Northern

Hemisphere and their expansion during the mid-Pleistocene transition (ca. 1.2–0.6 40 Myr ago)^{2,3}. Yet, this interpretation has been challenged⁴, and evidence of Saharan 41 dust in ~4.8 Myr old Canary Island palaeosols⁵ and proposed aeolian dune deposits 42 from the Djurab of Chad dated to \sim 7 Myr ago⁶ hint at an earlier inception of desert 43 conditions. Weakening of the summer monsoon in climate model simulations of the 44 response to Tethys Sea closure is hypothesized⁷ to account for earlier desertification. 45 However, the significance of the fossil dunes is highly controversial⁸, estimated dates 46 of Tethys closure span more than 20 Myr^{9,10} and existing climate records are too 47 48 sparse to test these competing hypotheses for desert inception. Thus, the geological history of the world's largest hot desert, which exerts a major influence on the Earth's 49 energy balance and global biogeochemical cycles, remains poorly constrained. 50

51

52 Unearthing terrestrial climate history in marine sediments

We investigated the evolution of African aridity and atmospheric dust export over the 53 last 11 Myr by developing continuous, high resolution and well-dated geochemical 54 records of terrigenous inputs from northern Africa to marine sediment archives. We 55 studied Ocean Drilling Project (ODP) Site 659 off Mauritania (Supplementary Figure 56 1), situated underneath the main modern Saharan dust plume. Pronounced cycles in 57 colour and lithology of Site 659 sediments record changes in terrigenous content and, 58 in the seminal study of Tiedemann et al.¹¹, these changes were quantified and used to 59 infer dust inputs and aridity change on Africa over the past 5 million years. Yet, that 60 hydroclimate reconstruction is now called into question for three reasons: First, distal, 61 fine-grained material, likely of riverine origin, also contributes to the lithogenic 62 fraction of sediment accumulating on the northwest African margin including Site 63

64 659^{12,13} (see Supplementary Information). Second, the proportion of terrigenous
65 material in marine sediments is also influenced by marine carbonate deposition and
66 dissolution, which may exert a significant control, particularly in Pliocene and
67 Miocene-aged sediments at Site 659^{14,15}. Third, dust supply to the atmosphere and
68 marine sediments is controlled not only by aridity but also wind strength¹⁶.

69

We used X-ray fluorescence (XRF) core scanning to both document dust 70 accumulation in the North Atlantic Ocean and infer African hydroclimate over the last 71 11 Myrs geochemically (Figure 1). Because dust flux estimates show a strong wind 72 strength dependence and can be biased by carbonate dissolution and rapid fluctuations 73 in sediment accumulation¹⁵, we use two geochemical ratios, [Al+Fe]/[Si+K+Ti] and 74 ln[Zr/Rb], to reconstruct hydroclimate variability. High [Al+Fe]/[Si+K+Ti] values are 75 associated with low inputs of Si-rich dust and more intense chemical weathering 76 under humid climates¹⁷ while high ln[Zr/Rb] values indicate dominance of coarse dust 77 over fine river clays¹². Crucially, these two ratios employ only lithophile elements and 78 so, unlike dust fluxes, they are independent of sea-floor carbonate dissolution and 79 dilution and show strong agreement with hydroclimate reconstructions based upon 80 deuterium isotopic signatures of plant waxes (Figure 2, Extended Data Figures 1–2 81 and Supplementary Information). Our dust flux reconstructions are similar to thorium-82 normalization based estimates from nearby site MD03-2705¹⁵ (Extended Data Figure 83 3 and Supplementary Information) suggesting that, although the influence of dilution, 84 85 dissolution, rapid sedimentation rate fluctuations and sediment remobilisation cannot be entirely ruled out, they are not likely main drivers of reconstructed dust fluxes at 86 Site 659. 87

88

Our high-resolution records (Figure 1) reveal a long history of repeated dramatic 89 shifts in hydroclimate between dry dusty conditions and more humid intervals 90 91 reaching back 11 Myrs. The most recent of these is the African Humid Period (AHP1) in the early Holocene, when the Saharan landscape was well-vegetated and cross-cut 92 by a network of rivers and lakes^{13, 18-20} with abundant archaeological sites, some 93 adorned with rock art depictions of giraffe, elephant, hippopotamus and domesticated 94 cattle²¹. Throughout our record, dark terrigenous-rich sediment layers at Site 659 have 95 significantly higher ln[Zr/Rb] and lower [Al+Fe]/[Si+K+Ti] values than pale layers 96 97 indicating that they contain a greater proportion of coarser grains, and have a chemical signature approaching dust end-member values (Figure 3, Extended Data 98 Figure 4 and Supplementary Information). Thus, the dark layers consistently signify 99 high dust deposition linked to aridity over northern Africa rather than high riverine 100 101 sediment input or merely strong sea floor carbonate dissolution. Intervening paler layers consistently record low ln[Zr/Rb] and high [Al+Fe]/[Si+K+Ti], indicating far 102 less dusty conditions on Africa. Spectral analysis reveals that this climatic variability 103 is primarily paced by precession and obliquity, with a strong signature of eccentricity 104 only documented in the oldest (Miocene) part of our records (Extended Data Figures 105 5–6 and Supplementary Information). 106

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108 Long-lived pulsing of Saharan dust export

We identify three distinct stages in the evolution of Saharan hydroclimate state over the past 11 Myrs, separated by transitions with statistically significant shifts in central tendency and dispersion in our datasets (Figure 1, Extended Data Figure 7 and Supplementary Information). Stage I (>6.9 Myr ago) reveals astronomically-paced

development of arid and dust-producing regions on Africa from at least 11 Myr ago. 113 Pronounced dark sediment layers consistently show high ln[Zr/Rb] and low 114 115 [Al+Fe]/[Si+K+Ti] values, confirming that these layers are, like those in the younger part of the record, associated with aridity (Figure 3, Extended Data Figure 4). Both 116 ln[Zr/Rb] and [Al+Fe]/[Si+K+Ti] values suggest that African climate was more arid 117 during the Late Miocene than the Pliocene, however estimated dust flux maxima are 118 119 lower. Given the importance of deflation-promoting high wind speeds (gustiness) in dust transport, we attribute this result to warmer Miocene high latitudes weakening 120 121 latitudinal temperature gradients (Figure 1c), resulting in lower wind speeds and less effective deflation and dust transport²²⁻²⁵. Nevertheless, our records document 122 substantial dust emissions that predate the oldest land-based evidence (~7 Myr ago) 123 for Miocene initiation of northern African hyperaridity⁶ by at least 4 Myr, and predate 124 Late Pliocene intensification of Northern Hemisphere Glaciation (iNHG)^{2, 3} by at least 125 8 Myr. 126

127

A shift towards increased [Al+Fe]/[Si+K+Ti] values followed by decreased ln[Zr/Rb] 128 from ~6.9 Ma marks the transition into stage II (which covers the latest Miocene to 129 the late Pliocene, ~5.75–3.5 Ma), indicating wetter conditions with more fine-grained 130 sedimentary inputs (Figure 1f and g). Increasing variability in [Al+Fe]/[Si+K+Ti], 131 ln[Zr/Rb] and dust fluxes is also recorded implying a heightened sensitivity of 132 northern African humidity to astronomical forcing, although cycle amplitudes may 133 134 also be more attenuated by bioturbation in Stage I where sedimentation rates are lower (Supplementary Figure 2). Dust fluxes during stage II are lower during humid 135 events associated with insolation maxima but higher during arid insolation minima 136 than seen for equivalent forcing in stage I. We attribute this result to the development 137

of extensive Pliocene river systems and lakes (including Lake Chad which has a
 Latest Miocene origin²⁶) during 'wet orbits' that, once desiccated, became dust producing hot-spots^{27,28} during succeeding 'dry orbits'.

141

The transition to Stage III, which covers the last ~2.25 Myrs (early Pleistocene to 142 Recent) of African climate history, is characterized by a shift towards more arid 143 conditions (decreasing [Al+Fe]/[Si+K+Ti] centred around 3.1 Ma) followed, ca. 400 144 kyr later, by increased mean values of ln[Zr/Rb] and dust fluxes (which also show 145 higher amplitude variability), closely contemporaneous with the intensification of 146 glacials as revealed by benthic oxygen isotope records²⁹ (Figure 1b). The highest dust 147 fluxes in our records during Stage III are consistent with the suggestion that the 148 growth of large continental ice sheets in the Northern Hemisphere promoted the 149 development of more arid and dusty conditions on Africa through steepening 150 latitudinal temperature gradients and strengthening winds^{3,30}. 151

152

153 **Consistency of source**

We investigated the source of terrigenous material accumulating at ODP Site 659 over 154 the last 11 Myr using radiogenic isotopes. Today, dust is generated widely across 155 three geochemically distinct northern African preferential source areas $(PSAs)^{31}$. 156 Strontium and neodymium isotopic signatures of the lithogenic fraction of Site 659 157 sediments are remarkably consistent compared to the range in modern PSA signatures 158 159 and lie between those reported for the Central and currently undersampled Western source regions³¹ (Figure 4). These results imply that the terrigenous source regions 160 remained broadly unchanged over the past 11 Myr and that variability in grain size is 161

not a major control on our data (Extended Data Figures 8–10 and Supplementary

163 Information). This observation, together with the high ln[Zr/Rb] and

164 [Al+Fe]/[Si+K+Ti] values recorded during Miocene insolation minima, indicates that

dust-producing regions existed in the inland Sahara region and were activated by

astronomical forcing back to at least 11 Myr ago. We conclude that while iNHG in the

167 late Pliocene (~3.3–2.5 Myr ago) drove increased dust fluxes (Figure 1h), it was not

responsible for the inception of the Saharan desert.

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170 Rainfall climate, C₄ savannas and evolutionary outcomes

Savannas dominated by C₄ grasses expanded extensively during the late Miocene^{32,33}, 171 but the cause of this major reorganization of the terrestrial biosphere is controversial. 172 Both aridification and decreasing pCO_2 are proposed drivers of this grassland 173 ecosystem shift, aided by strong positive feedbacks associated with fire, and 174 regionally modified by latitude, altitude and rainfall seasonality³²⁻³⁷. To better 175 understand the potential role of hydroclimate variability in driving the expansion of 176 C_4 grasslands, we compare our geochemical datasets to stable carbon ($\delta^{13}C$) and 177 hydrogen (δD) isotope signatures of leaf wax long-chain *n*-alkanes at Site 659 178 (Figures 1d and 2). The carbon isotope signature of plant waxes is strongly controlled 179 by the photosynthetic pathway of the parent plant: *n*-alkanes from C₄ plants are 180 isotopically heavier $(-21.4\pm2.2 \text{ }\%)$ than C₃ plants $(-34.9\pm2.7 \text{ }\%)^{38}$. 181 182

Africa and long-term hydroclimate trends (Figure 1). The plant wax records integrate material from across a large source area that was dominated by C_4 vegetation during

We find little similarity between the gradational expansion of C₄ grasses in northern

186	the Quaternary ^{39,40} , potentially contributing to the expression of only modest change
187	in <i>n</i> -alkane δ^{13} C in response to insolation forcing in the youngest part of the record.
188	Nevertheless, the hydroclimate signal in North Africa ([Al+Fe]/Si+K+Ti], ln[Zr/Rb]
189	and <i>n</i> -alkane $\delta D^{36,39,40}$) is dominated by astronomical variability, while the vegetation
190	signal (<i>n</i> -alkane $\delta^{13}C^{36,39,40}$) is dominated by the gradual long-term trend that defines
191	the C ₃ -C ₄ transition and is not replicated by the hydroclimate reconstructions (Figures
192	1 and 2, Supplementary Figure 3). Therefore, we find that a simple hydroclimate
193	driver cannot explain C ₄ grassland expansion in northwest Africa through the
194	Neogene. A similar conclusion was reached by ref. 36. However, while that study
195	ruled out a hydroclimate control because of little inferred change in regional rainfall,
196	we document major changes in hydroclimate unaccompanied by a strong response in
197	C_4 plant dominance (Figure 2). This decoupling between aridity and C_3 - C_4
198	composition does not rule out an influence of aridity on C ₄ vegetation dominance, but
199	suggests that the relationship was not simple. An improved record of Neogene pCO_2
200	variability (Figure 1a) is needed to better understand coupling between global climate
201	state, hydroclimate and the rise of C ₄ grasslands.

An inferred progressive aridification of Africa and spread of savanna ecosystems has 203 been linked to speciation and development of long-range mobility in hominins^{3, 41} and 204 extinction of many large mammals^{32,42}. Our long, detailed, continuous marine records 205 help to develop the environmental framework needed to assess these evolutionary 206 outcomes on land, including the hypothesized relationships between habitat instability 207 and development in hominins of characteristics that promote adaptability^{43,44}. We 208 highlight the longevity and primacy of the astronomical signal in controlling 209 hydroclimate in northern Africa. Our findings of three distinct phases in North 210

African hydroclimate, including relatively dry conditions during the warm Miocene, 211 suggest no simple escalation in aridity through the late Cenozoic associated with 212 213 global cooling, and argue against aridity as the primary driver of C₄ grassland expansion. Our records show evidence of changes in mean state and/or amplitude of 214 variability in hydroclimate close to the oldest suggested fossil evidence of hominin 215 bipedalism⁴⁵ and the earliest records of the genus *Homo*⁴⁶ (Figure 1). However, the 216 confidence interval on the origination of the principal taxa, especially suggested 217 bipeds, is large, implying earlier emergence⁴⁷. New fossil finds will inevitably revise 218 219 the hominin chronology, enabling a stronger understanding of the evolutionary link to climate. Meanwhile, our study reveals astronomically-paced dry and dusty conditions 220 on Africa long before both the oldest land-based evidence of a Saharan desert and the 221 intensification of Northern Hemisphere glaciation. 222

223

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247

248 Author contributions

A.J.C., P.A.W., C.P.O. and D.J.B. designed the study. A.J.C. and T.W. performed the

250 XRF measurements and developed the age model. B.D.A.N. generated the *n*-alkane

 δ^{13} C data and M.J.C. and A.J.C. generated the radiogenic isotope data. A.J.C.

252 performed the sediment endmember unmixing. A.J.C. and P.A.W. led the analysis and

interpretation of results with input from all authors. A.J.C., P.A.W., D.J.B. and C.P.O.

led the writing of the manuscript with contributions from all other authors.

255

Competing interests

257 The authors declare no competing interests.

258 Figures





264	isotope dataset, with 20 kyr smoothing ²⁹ . c Regional stacks of sea surface temperature
265	difference to modern annual means ²³ , blue: >50°N, red: tropics. d δ^{13} C signature of <i>n</i> -
266	alkanes from Site 659 (dark purple: this study, light purple: ref. 36) and offshore E.
267	Africa ⁴⁸ (grey). Error bars indicate 1σ . e Major events in hominid evolution. Solid
268	bars: taxon ranges, dashed lines: confidence interval on taxon origin ⁴⁷ . f
269	[Al+Fe]/[Si+K+Ti] of calibrated elemental abundances. Modern endmember values
270	updated from ref. 17 (see Supplementary Information). Dashed line marks mean over
271	last 1 Myr. g ln[Zr/Rb] XRF core scan ratios. Dashed line marks mean over last 1
272	Myr. h Site 659 estimated dust flux (3pt smoothed). Median value in red; 1%, 5%,
273	25%, 75%, 95% and 99% percentiles also shown in shades of orange/yellow. i Major
274	global climate events. MPT: mid-Pleistocene transition, iNHG: intensification of
275	Northern Hemisphere glaciation, mPWP: mid-Pliocene warm period, MSC:
276	Messinian Salinity Crisis. Vertical grey shading indicates transitions between climate
277	stages.



Figure 2| Relationship between lithophile element ratios and hydrogen and 281 carbon isotopic compositions of plant waxes at Site 659. a Last glacial cycle, b late 282 Pliocene, c early Pliocene. [Al+Fe]/[Si+K+Ti] of bulk sediment in orange (this study). 283 Leaf wax C_{31} *n*-alkane $\delta^{13}C$ data in green^{39,40} and δD in blue^{39,40}, error bars mark 1 284 standard deviation. High [Al+Fe]/[Si+K+Ti] and low *n*-alkane δD values both 285 indicate more humid conditions. **d-f** Violin plots indicating the distribution of leaf 286 wax C_{31} *n*-alkane $\delta^{13}C$ (green) and δD (blue) data in high-resolution snapshots in the 287 Late Quaternary (d) and Pliocene (e) (data from refs. 39 and 40), and low-resolution 288 records covering the last 11 Myr (f, data from this study and ref. 36). White dots 289 indicate median values, black rectangles indicate interquartile range. 290



298 more humid intervals (see also Extended Data Figure 4 and Supplementary

299 Information).



Figure 4 Radiogenic isotope signature of Site 659 lithic fraction compared to

304 values of preferential source regions (PSAs) reveals consistent source. a Site 659

305 bulk sediment data (black crosses) compared to source region measurements (circles)

306 coloured by PSA³¹. Data point size indicates dust source activation frequency

307 (DSAF)²⁸. Crosses indicate mean signatures of each PSA weighted by annual DSAF,

308 bars denote \pm one weighted standard deviation³¹. Samples for which only ε_{Nd} or

 87 Sr/ 86 Sr data exist are included in the calculation of mean values but are not plotted.

b Map of North African PSAs, adapted from ref. 31. Dust sources with activation

311 frequencies >5% shown in bold colours, <5% in pale colours.

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313 **References**

- Thomas N, Nigam S. Twentieth-Century Climate Change over Africa: Seasonal Hydroclimate Trends and Sahara Desert Expansion. *Journal of Climate* 2018, **31**(9): 3349-3370.
- Maley J. Les changements climatiques de la fin du Tertiaire en Afrique: leur
 conséquence sur l'apparition du Sahara et de sa végétation. *The Sahara and the Nile* 1980: 63-86.
- 322 3. deMenocal PB. Plio-Pleistocene African Climate. *Science* 1995, 270(5233):
 323 53-59.

- Trauth MH, Larrasoaña JC, Mudelsee M. Trends, rhythms and events in PlioPleistocene African climate. *Quaternary Science Reviews* 2009, 28(5–6): 399411.
- Muhs DR, Meco J, Budahn JR, Skipp GL, Betancort JF, Lomoschitz A. The
 antiquity of the Sahara Desert: New evidence from the mineralogy and
 geochemistry of Pliocene paleosols on the Canary Islands, Spain. *Palaeogeography, Palaeoclimatology, Palaeoecology* 2019, **533**: 109245.
- Schuster M, Duringer P, Ghienne J-F, Vignaud P, Mackaye HT, Likius A, et *al.* The Age of the Sahara Desert. *Science* 2006, **311**(5762): 821.

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- Zhang Z, Ramstein G, Schuster M, Li C, Contoux C, Yan Q. Aridification of
 the Sahara desert caused by Tethys Sea shrinkage during the Late Miocene. *Nature* 2014, **513**(7518): 401-404.
- 341 8. Kroepelin S, Swezey CS. Revisiting the Age of the Sahara Desert. *Science* 2006, **312**(5777): 1138-1139.
- McQuarrie N, van Hinsbergen DJJ. Retrodeforming the Arabia-Eurasia
 collision zone: Age of collision versus magnitude of continental subduction. *Geology* 2013, 41(3): 315-318.
- Allen MB, Armstrong HA. Arabia–Eurasia collision and the forcing of mid Cenozoic global cooling. *Palaeogeography, Palaeoclimatology, Palaeoecology* 2008, 265(1–2): 52-58.
- Tiedemann R, Sarnthein M, Shackleton NJ. Astronomic timescale for the
 Pliocene Atlantic δ18O and dust flux records of Ocean Drilling Program Site *659. Paleoceanography* 1994, 9(4): 619-638.
- Tjallingii R, Claussen M, Stuut J-BW, Fohlmeister J, Jahn A, Bickert T, *et al.*Coherent high- and low-latitude control of the northwest African hydrological
 balance. *Nature Geosci* 2008, 1(10): 670-675.
- Skonieczny C, Paillou P, Bory A, Bayon G, Biscara L, Crosta X, *et al.* African
 humid periods triggered the reactivation of a large river system in Western
 Sahara. *Nat Commun* 2015, 6.
- Ruddiman WF, Sarnthein M, Baldauf JG, et al. *Proc. ODP, Init. Repts., 108.*College Station, TX (Ocean Drilling Program), 1989.
- Skonieczny C, McGee D, Winckler G, Bory A, Bradtmiller LI, Kinsley CW, *et al.* Monsoon-driven Saharan dust variability over the past 240,000 years. *Science Advances* 2019, 5(1): eaav1887.
- McGee D, deMenocal PB, Winckler G, Stuut JBW, Bradtmiller LI. The
 magnitude, timing and abruptness of changes in North African dust deposition

373over the last 20,000 yr. Earth and Planetary Science Letters 2013, 371-372:374163-176.

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387

390

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406

411

- Mulitza S, Heslop D, Pittauerova D, Fischer HW, Meyer I, Stuut J-B, *et al.*Increase in African dust flux at the onset of commercial agriculture in the
 Sahel region. *Nature* 2010, **466**(7303): 226-228.
- 18. Drake NA, Blench RM, Armitage SJ, Bristow CS, White KH. Ancient
 watercourses and biogeography of the Sahara explain the peopling of the
 desert. *Proceedings of the National Academy of Sciences* 2011, **108**(2): 458 462.
- 19. Larrasoaña JC, Roberts AP, Rohling EJ. Dynamics of green Sahara periods
 and their role in hominin evolution. *PloS one* 2013, 8(10): e76514.
- 388 20. Tierney JE, Pausata FSR, deMenocal PB. Rainfall regimes of the Green
 389 Sahara. *Science Advances* 2017, 3(1).
- 391 21. Mori F. The earliest Saharan rock-engravings. *Antiquity* 1974, **48**(190): 87-92.
- McGee D, Broecker WS, Winckler G. Gustiness: The driver of glacial dustiness? *Quaternary Science Reviews* 2010, **29**(17–18): 2340-2350.
- Herbert TD, Lawrence KT, Tzanova A, Peterson LC, Caballero-Gill R, Kelly
 CS. Late Miocene global cooling and the rise of modern ecosystems. *Nature Geosci* 2016, 9(11): 843-847.
- 400 24. Abell JT, Winckler G, Anderson RF, Herbert TD. Poleward and weakened
 401 westerlies during Pliocene warmth. *Nature* 2021, 589(7840): 70-75.
- Burls NJ, Fedorov AV. Wetter subtropics in a warmer world: Contrasting past
 and future hydrological cycles. *Proceedings of the National Academy of Sciences* 2017, **114**(49): 12888-12893.
- 407 26. Moussa A, Novello A, Lebatard A-E, Decarreau A, Fontaine C, Barboni D, *et al.* Lake Chad sedimentation and environments during the late Miocene and
 409 Pliocene: New evidence from mineralogy and chemistry of the Bol core
 410 sediments. *Journal of African Earth Sciences* 2016, **118**: 192-204.
- 412 27. Washington R, Todd M, Middleton NJ, Goudie AS. Dust-Storm Source Areas
 413 Determined by the Total Ozone Monitoring Spectrometer and Surface
 414 Observations. *Annals of the Association of American Geographers* 2003,
 415 93(2): 297-313.
- 417 28. Schepanski K, Tegen I, Macke A. Comparison of satellite based observations
 418 of Saharan dust source areas. *Remote Sensing of Environment* 2012, 123: 90419 97.
 420

- 421 29. Westerhold T, Marwan N, Drury AJ, Liebrand D, Agnini C, Anagnostou E, *et al.* An astronomically dated record of Earth's climate and its predictability
 423 over the last 66 million years. *Science* 2020, **369**(6509): 1383-1387.
- 30. Sarnthein M, Thiede J, Pflaumann U, Erlenkeuser H, Fütterer D, Koopmann
 B, et al. Atmospheric and Oceanic Circulation Patterns off Northwest Africa
 During the Past 25 Million Years. In: von Rad U, Hinz K, Sarnthein M,
 Seibold E (eds). *Geology of the Northwest African Continental Margin*.
 Springer Berlin Heidelberg: Berlin, Heidelberg, 1982, pp 545-604.
- 431 31. Jewell AM, Drake N, Crocker AJ, Bakker NL, Kunkelova T, Bristow CS, *et al.* Three North African dust source areas and their geochemical fingerprint.
 433 *Earth and Planetary Science Letters* 2021, **554:** 116645.
- 435 32. Cerling TE, Harris JM, MacFadden BJ, Leakey MG, Quade J, Eisenmann V,
 436 *et al.* Global vegetation change through the Miocene/Pliocene boundary.
 437 *Nature* 1997, **389**(6647): 153-158.
- 439 33. Feakins SJ, Levin NE, Liddy HM, Sieracki A, Eglinton TI, Bonnefille R.
 440 Northeast African vegetation change over 12 m.y. *Geology* 2013, 41(3): 295–
 441 298.
- 443 34. Pagani M, Freeman KH, Arthur MA. Late Miocene Atmospheric CO2
 444 Concentrations and the Expansion of C4 Grasses. *Science* 1999, 285(5429):
 445 876-879.
- Beerling DJ, Osborne CP. The origin of the savanna biome. *Global Change Biology* 2006, **12**(11): 2023-2031.
- 450 36. Polissar PJ, Rose C, Uno KT, Phelps SR, deMenocal P. Synchronous rise of
 451 African C4 ecosystems 10 million years ago in the absence of aridification.
 452 Nature Geoscience 2019, 12(8): 657-660.
- 454 37. Hoetzel S, Dupont L, Schefuß E, Rommerskirchen F, Wefer G. The role of
 455 fire in Miocene to Pliocene C4 grassland and ecosystem evolution. *Nature*456 *Geosci* 2013, 6(12): 1027-1030.
- 38. Naafs BDA, Hefter J, Acton G, Haug GH, Martínez-Garcia A, Pancost R, *et al.* Strengthening of North American dust sources during the late Pliocene (2.7 Ma). *Earth and Planetary Science Letters* 2012, **317–318**(0): 8-19.
- 462 39. Kuechler RR, Dupont LM, Schefuß E. Hybrid insolation forcing of Pliocene
 463 monsoon dynamics in West Africa. *Climate of the Past* 2018, 14(1): 73-84.
- 465 40. Kuechler RR, Schefuß E, Beckmann B, Dupont L, Wefer G. NW African
 466 hydrology and vegetation during the Last Glacial cycle reflected in plant-wax467 specific hydrogen and carbon isotopes. *Quaternary Science Reviews* 2013,
 468 82(0): 56-67.

430

434

438

442

446

449

453

457

461

- 470 41. Cerling TE, Wynn JG, Andanje SA, Bird MI, Korir DK, Levin NE, *et al.*471 Woody cover and hominin environments in the past 6 million years. *Nature*472 2011, 476(7358): 51-56.
- 474 42. Faith JT, Rowan J, Du A, Koch PL. Plio-Pleistocene decline of African megaherbivores: No evidence for ancient hominin impacts. *Science* 2018, 362(6417): 938-941.
- 478 43. Potts R. Hominin evolution in settings of strong environmental variability.
 479 *Quaternary Science Reviews* 2013, **73**: 1-13.
- 481 44. Maslin MA, Brierley CM, Milner AM, Shultz S, Trauth MH, Wilson KE. East
 482 African climate pulses and early human evolution. *Quaternary Science*483 *Reviews* 2014, **101**(0): 1-17.
- 485 45. Zollikofer CPE, Ponce de León MS, Lieberman DE, Guy F, Pilbeam D, Likius
 486 A, *et al.* Virtual cranial reconstruction of Sahelanthropus tchadensis. *Nature*487 2005, **434:** 755.
- 489 46. DiMaggio EN, Campisano CJ, Rowan J, Dupont-Nivet G, Deino AL, Bibi F,
 490 *et al.* Late Pliocene fossiliferous sedimentary record and the environmental
 491 context of early *Homo* from Afar, Ethiopia. *Science* 2015, 347(6228): 1355492 1359.
- 494 47. Bobe R, Wood B. Estimating origination times from the early hominin fossil
 495 record. *Evolutionary Anthropology: Issues, News, and Reviews* 2021: 1–11.
- 497 48. Uno KT, Polissar PJ, Jackson KE, deMenocal PB. Neogene biomarker record
 498 of vegetation change in eastern Africa. *Proceedings of the National Academy*499 *of Sciences* 2016: 201521267.
- 49. Laskar J, Robutel P, Joutel F, Gastineau M, Correia ACM, Levrard B. A longterm numerical solution for the insolation quantities of the Earth. *Astronomy & Astrophysics* 2004, 428(1): 261-285.

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505 Methods

506 Site Location

507 ODP Site 659 is situated in the tropical North Atlantic Ocean on the Cape Verde Rise, offshore Mauritania (18.077°N 21.026°W, 3070 m water depth, for map see 508 Supplementary Figure 1). Today, this region receives dust delivered by the trade 509 winds during winter and spring, with additional dust transported over the site by the 510 Saharan Air Layer predominantly during boreal summer⁵⁰⁻⁵⁶. The Algeria-Morocco 511 region, Mali, Mauritania, Libya, Niger, Western Sahara and the Sahel have all been 512 identified as regions that may currently contribute dust to the Cape Verde Islands and 513 the surrounding ocean sediments, while evidence for a major contribution from the 514 Bodélé depression is weaker^{50-53,56-65}. The elevated position of Site 659 on the Cape 515 516 Verde Rise protects it from major mass transport deposits, although grain size data reveal that multiple sites along the northwestern African margin including 517 GeoB7920¹² and Site 659 received a significant fine-grained lithogenic component 518 during both the Holocene and last glacial cycle that is distinct from modern dust 519 inputs and is often attributed to a distal influence from African (palaeo)rivers¹² 520 (Extended Data Figures 9–10 and Supplementary Information). 521

522

523 X-ray fluorescence

524 Data were collected from a total of about 275 m of drill core from three holes at ODP

525 Site 659 using the XRF Core Scanner II (AVAATECH Serial No. 2) at MARUM -

526 University of Bremen. Measurements were taken every 1–5 cm down core to give a

- 527 consistent approximate temporal resolution of 1.5–2 kyr. Count times were 20
- seconds and generator settings of 30 and 10 kV with currents of 0.75 and 0.2 mA
- 529 respectively were used. The split core surface was covered with a 4 micron thin

530	SPEXCerti Prep Ultralene foil to avoid contamination of the XRF measurement unit
531	and desiccation of the sediment. The data were acquired using a Canberra X-PIPS
532	Silicon Drift Detector (SDD; Model SXD 15C-150-500) with 150eV X-ray
533	resolution, the Canberra Digital Spectrum Analyzer DAS 1000, and an Oxford
534	Instruments 50W XTF5011 X-Ray tube with rhodium (Rh) target material. Raw X-
535	ray spectra were processed using an iterative least square software (WIN AXIL)
536	package from Canberra Eurisys. Repeat runs of several core sections were used to
537	correct for any drift in counts over time.

Ca/Fe ratios provide an indicator of the relative proportions of marine (dominated by 539 biogenic calcium carbonate) and terrigenous (the main source of iron) sediment 540 (Supplementary Figure 4). We use two geochemical ratios to determine the relative 541 contribution of aeolian material to the sediment. Modern Saharan dust has much 542 lower [Al+Fe]/[Si+K+Ti] values than samples of suspended sediment from the 543 Senegal River (Figure 1f), the nearest and largest active river to Site 659, which 544 drains deeply weathered lateritic tropical soils much richer in Al and Fe than the Si-545 rich Sahelian/Saharan dust sources¹⁷. We also employ Zr/Rb as an independent grain 546 size proxy because the lithogenic dust fraction is significantly coarser than riverine-547 derived material on the northwest African margin^{12,66}, and contains a higher 548 proportion of Zr-bearing zircon grains than Rb-rich river clays⁶⁷. Additional 549 geochemical ratios are shown in Supplementary Figures 5-6. Our XRF records have 550 an average temporal resolution of approximately 2000 years. 551 552

A suite of discrete sediment samples from ODP Site 659 incorporating a wide range of lithologies were analysed by energy dispersive polarised x-ray fluorescence

spectrometry (EDP-XRF) at the University of St Andrews to convert the semi-

⁵⁵⁶ quantitative XRF scanner counts into concentrations (Supplementary Figure 7). The

downcore records plotted are calculated from ratios of these calibrated values, except

for $\ln[Zr/Rb]$ where counts of these trace elements are plotted.

559

560 Stratigraphy

We created a new spliced composite section for ODP Site 659 to a depth of 203.17 561 revised metres composite depth (rmcd) by correlating both the XRF core scan data 562 563 and core images between the three holes (A-C) drilled at Site 659, using the Code for Ocean Drilling Data macros⁶⁸. An age framework was developed from paleomagnetic, 564 nannofossil and planktonic foraminiferal datums^{14,69} with ages updated to incorporate 565 more recent improvements in astrochronology⁷⁰⁻⁷². We then correlated previously 566 published oxygen isotope values of the benthic foraminifera Cibicides wuellerstorfi at 567 Site 659¹¹ transferred onto our composite depth scale to the LR04 benthic oxygen 568 isotope stack⁷³ to improve age control for the youngest 5 Myr. The final age model 569 was then generating by tuning our XRF count data (ln[Ca/Fe]) to summer insolation 570 calculated at 65°N in the La2004 astronomical solution⁴⁹, guided by the framework 571 provided by the biostratigraphic, magnetostratigraphic and benthic oxygen isotope 572 record. The construction of this age model is illustrated in Supplementary Figure 8. In 573 Supplementary Figure 2, we compare the age-depth relationship of our 574 astronomically-tuned age model to existing bio- and magnetostratigraphic datums. 575 Continuous wavelet power spectra⁷⁴ of both astronomically-tuned and non-576 astronomically tuned time series data are shown in Extended Data Figure 5, with 577 results from REDFIT⁷⁵ spectral analysis shown in Extended Data Figure 6. 578 579

580 Geochemical end-member unmixing

Dust fluxes were estimated using an end-member unmixing approach to deconvolve 581 the relative proportions of different contributions to the bulk marine sediment, largely 582 following the method of Mulitza et al. $(2010)^{17}$. In this approach, bulk sediment is 583 assumed to consist of three components: aeolian, riverine and marine inputs. The 584 relative abundances of Al, Si, Fe, K, Ti and Ca concentrations at Site 659 (calibrated 585 by a multivariate log calibration method^{76,77} using the AvaaXelerate software⁷⁸) were 586 compared to the chemistry of endmember compositions estimated from analyses of 587 588 modern sediments. The riverine endmember was constructed from ten measurements of suspended sediment from the Senegal River⁷⁹, the present day catchment of which 589 stretches both north and south of Site 659. A total of 48 measurements were used to 590 construct a dust endmember, covering a wide geographical area and incorporating 591 data from dust traps, atmospheric sampling and bulk sediment from known dust 592 source areas with a range of grain sizes and local climates^{59,62,80-91}. A marine 593 endmember composition of 2% Si and 98% Ca was assumed¹⁷. To incorporate the 594 effect of temporal changes in active terrigenous source regions and lithic fraction 595 chemistry into our dust flux estimates, a bootstrapping approach was applied. During 596 every realization, 10 of the 48 dust compositions and 4 of the 10 river compositions 597 were selected by a bootstrap with replacement routine, with the unmixing procedure 598 carried out for a total of 500 realizations of endmember compositions. Proportions of 599 aeolian and riverine endmembers were converted to fluxes using sedimentation rates 600 calculated from our composite depth scale and age model, and dry bulk sediment 601 densities estimated from shipboard measurements of gamma-ray attenuation¹⁴ which 602 were calibrated using the discrete dry bulk density measurements of Tiedemann 603 $(1991)^{92}$. We show that there is good agreement between our dust flux estimates and 604

those based on ²³⁰Th normalization¹⁵ and lithogenic %¹¹ in Extended Data Figures 2–
3, Supplementary Figure 4 and Supplementary Information.

607

Lithogenic grain size measurements and geochemical data strongly support the 608 concept of distal riverine inputs to Site 659. The proportion of the terrigenous fraction 609 that is attributed to aeolian inputs by this geochemical unmixing approach varies 610 strongly down core, with estimates of 0-87% dust origin. These estimates are broadly 611 in line with those of Tjallingii et al. $(2008)^{12}$ who find that dust accounts for 5–95% of 612 613 the lithic fraction through the last glacial cycle at nearby site GeoB7920 using a unmixing approach based upon grain size distributions. We cannot, however, rule out 614 contributions to the fine-grained, high Al+Fe end-member from very fine dust or 615 resuspended shelf material. Thus, our dust fluxes are minimum (conservative) 616 estimates (see Supplementary Information). 617

618

619 Radiogenic isotopes

The Sr and Nd isotope composition of terrigenous sediments from ODP Site 659 were
 measured at the University of Southampton National Oceanography Centre.

622 Approximately 1g of crushed and homogenised dried bulk sediment was decarbonated

using 10 % acetic acid solution for 24 hours on a shaker table and then rinsed in

deionized water. Samples were placed in a 5 % hydrogen peroxide solution for 48

hours at 65 °C to remove organic matter, then rinsed three times in deionized water.

1M MgCl₂ was added to the samples to remove any Sr adsorbed from seawater, which

- 627 were shaken overnight then rinsed three times. A buffered hydroxylamine HCl, Na-
- EDTA and acetic acid solution was added to remove authigenic coatings, with
- samples mixed for 24 hours and then rinsed. A magnesium ion solution was added to

aid flocculation if a colloid phase formed at any stage during processing. The

⁶³¹ 'cleaned' sediments were then totally digested in concentrated nitric acid (HNO₃) and
⁶³² hydrofluoric acid (HF), followed by 6M HCl.

633

Digested sediment solutions for Sr isotope analysis were purified on columns filled with Sr-spec Resin in 3M HNO₃. Samples were loaded onto tantalum (Ta) filaments with a 1M HCl solution after a tantalum chloride (TaCl) activator solution was loaded. Isotopic measurements were made by thermal ionization mass spectrometry (Thermo-Fisher TRITON Plus). Repeated measurements of the Sr isotope standard NBS987 gave 87 Sr/ 86 Sr = 0.710243 ± 0.000021 (2 σ).

Neodymium was purified from the digested sediments for isotope analysis using a 641 642 standard column chemistry procedure, based upon the methods of Cohen et al. $(1988)^{93}$. Cation exchange resin was used to strip iron and titanium from the samples, 643 then the remaining material was then run through using Ln-SpecTM resin columns⁹⁴ to 644 purify and concentrate neodymium. Neodymium isotope ratios were measured by 645 multi-collector inductively coupled plasma mass spectrometry (Thermo NEPTUNE). 646 Instrumental mass bias was corrected using the procedure of Vance and Thirlwall 647 (2002)⁹⁵, adjusting to a ¹⁴⁶Nd/¹⁴⁴Nd ratio of 0.7219 and using cerium-doped standards 648 to correct for interference of ¹⁴²Ce on ¹⁴²Nd. All ratios were normalised to the JNdi-1 649 standard $(^{143}Nd/^{144}Nd = 0.512115 \pm 0.00007)^{96}$, with independent measurements of 650 JNdi-1 = 0.512115 ± 0.000006 (2 σ). Isotopic signatures are expressed in epsilon 651 notation, relative to the chondritic uniform reservoir value of 0.512638⁹⁷. 652 653

654 Leaf wax δ^{13} C

655	Compound specific stable carbon isotopes (δ^{13} C) of the C ₃₁ and C ₃₃ <i>n</i> -alkane were
656	obtained from 30 samples from ODP Site 659 at the Organic Geochemistry Unit,
657	University of Bristol. For this purpose, between 5 and 10 gram of freeze-dried
658	sediment was extracted using a Microwave extraction system (Milestone Ethos Ex)
659	and a mixture of dichloromethane (DCM) and methanol (MeOH) (9:1, v/v). Fine
660	grained sediment was removed by eluting the total lipid extract (TLE) over a short (4
661	cm) inert NaSO ₄ column using 4 ml of DCM:MeOH (9:1). The TLE was dried under
662	a gentle flow of N_2 . The TLE was subsequently derivatized using pyridine and
663	BSFTA (1 hr at 70 °C) and within 24 hrs analysed on an Isoprime 100 gas
664	chromatograph combustion isotope ratio mass spectrometer (GC-C-IRMS). Samples
665	were measured in triplicate and $\delta^{13}C$ values converted to VPDB by bracketing with an
666	in-house gas (CO ₂) of known δ^{13} C value. Instrument stability was monitored by
667	regular analysis of an in-house fatty acid methyl ester standard mixture and indicates
668	that long-term instrument stability was better than \pm 0.3 ‰. Injection volume was 1
669	or 2 μl onto to a Zebron-I nonpolar column (50 m x 0.32 mm i.d., 0.10 μm film
670	thickness). The GC oven was programmed as: 70 °C (1 min hold) to 130 °C at 20 °C
671	min ⁻¹ , then to 300 °C at 4 °C min ⁻¹ , and a final hold at 300 °C for 25 min. Samples
672	were automatically integrated using the Ion Vantage software.
673	

- For some samples, the concentration of the C_{33} *n*-alkane was not enough to obtain a reliable $\delta^{13}C$ value. The $\delta^{13}C$ of the C_{31} *n*-alkane was consistently lighter than that of the C_{33} *n*-alkane from the same sample by on average 2 ‰.
- 677

678 Lithogenic grain size distributions at ODP Site 659

679	The distribution of grain sizes in lithogenic material can be used to identify different
680	terrigenous components in marine sediment. 55 samples from ODP Site 659 were
681	selected to generate grain size distribution data, incorporating a range of sediment
682	ages and lithologies. Calcium carbonate, authigenic coatings and organic matter were
683	removed following the procedure used for radiogenic isotope analysis. In addition, a
684	biogenic silica removal step was applied by sonicating samples in 1.5M NaOH.
685	Calgon was then added to the samples to prevent flocculation before analysis of the
686	lithogenic grain sizes using a Laser Coulter Sizer at the University of Southampton
687	Waterfront Campus, NOCS. The results of these grain size analyses are shown in
688	Extended Data Figure 9d.
689	
690	The signatures of different terrigenous sources were separated from the bulk sediment
691	signature following the end-member modeling analysis approach (EMMA) of Dietze
692	et al. (2011) ⁹⁸ . A two lithogenic end-member solution explains 88% of the variance in
693	the Site 659 data set, with the end-members shown in Extended Figure 9. The coarser
694	of these two endmembers shows a strong match with measurements of African dust
695	(Extended Data Figure 9a-b)
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697 Data availability

⁶⁹⁸ The data presented in this study are available in the Zenodo repository (DOI:

699 10.5281/zenodo.6594643).

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701 Methods references

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Kumar A, Abouchami W, Galer SJG, Singh SP, Fomba KW, Prospero JM, *et al.* Seasonal radiogenic isotopic variability of the African dust outflow to the

tropical Atlantic Ocean and across to the Caribbean. *Earth and Planetary Science Letters* 2018, **487**: 94-105.

707

712

717

723

728

732

737

741

746

- 51. Gama C, Tchepel O, Baldasano JM, Basart S, Ferreira J, Pio C, *et al.* Seasonal patterns of Saharan dust over Cape Verde a combined approach using observations and modelling. *Tellus B: Chemical and Physical Meteorology* 2015, 67(1): 24410.
- 52. Patey MD, Achterberg EP, Rijkenberg MJ, Pearce R. Aerosol time-series
 measurements over the tropical Northeast Atlantic Ocean: Dust sources,
 elemental composition and mineralogy. *Marine Chemistry* 2015, **174**: 103119.
- 53. Skonieczny C, Bory A, Bout-Roumazeilles V, Abouchami W, Galer SJG,
 Crosta X, *et al.* A three-year time series of mineral dust deposits on the West
 African margin: Sedimentological and geochemical signatures and
 implications for interpretation of marine paleo-dust records. *Earth and Planetary Science Letters* 2013, **364**: 145-156.
- 724 54. Ratmeyer V, Fischer G, Wefer G. Lithogenic particle fluxes and grain size
 725 distributions in the deep ocean off northwest Africa: Implications for seasonal
 726 changes of aeolian dust input and downward transport. *Deep Sea Research*727 *Part I: Oceanographic Research Papers* 1999, **46**(8): 1289-1337.
- 55. Bory A, Dulac F, Moulin C, Chiapello I, Newton PP, Guelle W, *et al.*Atmospheric and oceanic dust fluxes in the northeastern tropical Atlantic
 Ocean: how close a coupling? *Ann Geophys* 2002, **20**(12): 2067-2076.
- 56. Chiapello I, Bergametti G, Chatenet B, Bousquet P, Dulac F, Soares ES.
 Origins of African dust transported over the northeastern tropical Atlantic. *Journal of Geophysical Research: Atmospheres* 1997, 102(D12): 1370113709.
- 57. Schepanski K, Tegen I, Macke A. Saharan dust transport and deposition towards the tropical northern Atlantic. *Atmos Chem Phys* 2009, 9(4): 1173-1189.
- 742 58. Caquineau S, Gaudichet A, Gomes L, Legrand M. Mineralogy of Saharan dust
 743 transported over northwestern tropical Atlantic Ocean in relation to source
 744 regions. *Journal of Geophysical Research: Atmospheres* 2002, **107**(D15):
 745 AAC 4-1-AAC 4-12.
- Formenti P, Rajot JL, Desboeufs K, Caquineau S, Chevaillier S, Nava S, *et al.*Regional variability of the composition of mineral dust from western Africa:
 Results from the AMMA SOP0/DABEX and DODO field campaigns. *Journal of Geophysical Research: Atmospheres* 2008, **113**(D23): D00C13.
- Friese CA, van Hateren JA, Vogt C, Fischer G, Stuut J-BW. Seasonal
 provenance changes in present-day Saharan dust collected in and off
 Mauritania. *Atmospheric Chemistry and Physics* 2017, **17**(16): 10163.

755 61. McConnell CL, Highwood EJ, Coe H, Formenti P, Anderson B, Osborne S, et 756 al. Seasonal variations of the physical and optical characteristics of Saharan 757 dust: Results from the Dust Outflow and Deposition to the Ocean (DODO) 758 759 experiment. Journal of Geophysical Research: Atmospheres 2008, 113(D14). 760 62. Salvador P, Almeida SM, Cardoso J, Almeida-Silva M, Nunes T, Cerqueira 761 762 M, et al. Composition and origin of PM10 in Cape Verde: Characterization of long-range transport episodes. Atmospheric Environment 2016, 127: 326-339. 763 764 Skonieczny C, Bory A, Bout-Roumazeilles V, Abouchami W, Galer SJG, 765 63. Crosta X, et al. The 7-13 March 2006 major Saharan outbreak: Multiproxy 766 characterization of mineral dust deposited on the West African margin. 767 768 Journal of Geophysical Research: Atmospheres 2011, 116(D18). 769 64. Stuut J-B, Zabel M, Ratmeyer V, Helmke P, Schefuß E, Lavik G, et al. 770 Provenance of present-day eolian dust collected off NW Africa. Journal of 771 Geophysical Research: Atmospheres 2005, 110(D4): D04202. 772 773 65. Zhao W, Balsam W, Williams E, Long X, Ji J. Sr-Nd-Hf isotopic 774 fingerprinting of transatlantic dust derived from North Africa. Earth and 775 Planetary Science Letters 2018, 486: 23-31. 776 777 66. Holz C, Stuut J-BW, Henrich R. Terrigenous sedimentation processes along 778 the continental margin off NW Africa: implications from grain-size analysis of 779 seabed sediments. Sedimentology 2004, 51(5): 1145-1154. 780 781 67. Matthewson AP, Shimmield GB, Kroon D, Fallick AE. A 300 kyr high-782 resolution aridity record of the North African continent. *Paleoceanography* 783 784 1995, 10(3): 677-692. 785 68. Wilkens RH, Westerhold T, Drury AJ, Lyle M, Gorgas T, Tian J. Revisiting 786 the Ceara Rise, equatorial Atlantic Ocean: isotope stratigraphy of ODP Leg 787 154 from 0 to 5 Ma. Clim Past 2017, 13(7): 779-793. 788 789 790 69. Manivit H. Calcareous nannofossil biostratigraphy of Leg 108 sediments. In: Ruddiman W, Sarnthein, M., et al., (ed). Proceedings of the Ocean Drilling 791 792 Program, Scientific Results, Vol. 108, 1989, pp 35-69. 793 70. 794 Raffi I, Backman J, Fornaciari E, Pälike H, Rio D, Lourens L, et al. A review 795 of calcareous nannofossil astrobiochronology encompassing the past 25 million years. Quaternary Science Reviews 2006, 25(23-24): 3113-3137. 796 797 Ogg JG. Chapter 5 - Geomagnetic Polarity Time Scale. In: Gradstein FM, Ogg 71. 798 JG, Schmitz MD, Ogg GM (eds). The Geologic Time Scale. Elsevier: Boston, 799 800 2012, pp 85-113. 801 72. Wade BS, Pearson PN, Berggren WA, Pälike H. Review and revision of 802 Cenozoic tropical planktonic foraminiferal biostratigraphy and calibration to 803

the geomagnetic polarity and astronomical time scale. Earth-Science Reviews 804 2011, **104**(1): 111-142. 805 806 73. Lisiecki LE, Raymo ME. A Pliocene-Pleistocene stack of 57 globally 807 808 distributed benthic δ 180 records. *Paleoceanography* 2005, **20**(1). 809 74. Grinsted A, Moore JC, Jevrejeva S. Application of the cross wavelet transform 810 and wavelet coherence to geophysical time series. Nonlinear processes in 811 geophysics 2004, 11(5/6): 561-566. 812 813 814 75. Schulz M, Mudelsee M. REDFIT: estimating red-noise spectra directly from unevenly spaced paleoclimatic time series. Computers & Geosciences 2002, 815 **28**(3): 421-426. 816 817 Weltje GJ, Tjallingii R. Calibration of XRF core scanners for quantitative 76. 818 geochemical logging of sediment cores: Theory and application. Earth and 819 820 Planetary Science Letters 2008, 274(3-4): 423-438. 821 77. Weltje GJ, Bloemsma M, Tjallingii R, Heslop D, Röhl U, Croudace IW. 822 Prediction of geochemical composition from XRF core scanner data: a new 823 multivariate approach including automatic selection of calibration samples and 824 quantification of uncertainties. In: Croudace; IW, Rothwell RG (eds). Micro-825 XRF Studies of Sediment Cores. Springer, 2015, pp 507-534. 826 827 78. Bloemsma MR. Development of a Modelling Framework for Core Data 828 Integration using XRF Scanning. TU Delft, Delft University of Technology, 829 830 2015. 831 79. Gac J-Y, Kane A. Le fleuve Sénégal: I. Bilan hydrologique et flux 832 continentaux de matières particulaires à l'embouchure. Sciences Géologiques, 833 bulletins et mémoires 1986: 99-130. 834 835 80. Scheuvens D, Schütz L, Kandler K, Ebert M, Weinbruch S. Bulk composition 836 of northern African dust and its source sediments - A compilation. Earth-837 Science Reviews 2013, 116(0): 170-194. 838 839 81. Orange D, Gac J-Y. Bilan géochimique des apports atmosphériques en 840 domaines sahélien et soudano-guinéen d'Afrique de l'Ouest (bassins supérieurs 841 du Sénégal et de la Gambie). Géodynamique 1990, 5(1): 51-65. 842 843 82. Orange D, Gac J-Y, Diallo MI. Geochemical assessment of atmospheric 844 deposition including Harmattan dust in continental West Africa. Tracers in 845 Hydrology; 1993: IAHS; 1993. 846 847 83. Guieu C, Thomas AJ. Saharan Aerosols: From the Soil to the Ocean. In: 848 Guerzoni S, Chester R (eds). The Impact of Desert Dust Across the 849 Mediterranean. Springer Netherlands: Dordrecht, 1996, pp 207-216. 850 851

852 84. Criado C, Dorta P. An unusual 'blood rain' over the Canary Islands (Spain).
853 The storm of January 1999. *Journal of Arid Environments* 2003, 55(4): 765854 783.

855

859

864

869

873

877

881

886

890

894

- 856
 85. Viana M, Querol X, Alastuey A, Cuevas E, Rodríguez S. Influence of African
 857 dust on the levels of atmospheric particulates in the Canary Islands air quality
 858 network. *Atmospheric Environment* 2002, **36**(38): 5861-5875.
- 860 86. Formenti P, Elbert W, Maenhaut W, Haywood J, Andreae MO. Chemical composition of mineral dust aerosol during the Saharan Dust Experiment (SHADE) airborne campaign in the Cape Verde region, September 2000.
 863 Journal of Geophysical Research: Atmospheres 2003, 108(D18): 8576.
- 865 87. Linke C, Möhler O, Veres A, Mohácsi Á, Bozóki Z, Szabó G, *et al.* Optical
 866 properties and mineralogical composition of different Saharan mineral dust
 867 samples: a laboratory study. *Atmospheric Chemistry and Physics* 2006, 6(11):
 868 3315-3323.
- 870 88. Khiri F, Ezaidi A, Kabbachi K. Dust deposits in Souss–Massa basin, South871 West of Morocco: granulometrical, mineralogical and geochemical
 872 characterisation. *Journal of African Earth Sciences* 2004, **39**(3): 459-464.
- 874 89. Moreno T, Querol X, Castillo S, Alastuey A, Cuevas E, Herrmann L, *et al.*875 Geochemical variations in aeolian mineral particles from the Sahara–Sahel
 876 Dust Corridor. *Chemosphere* 2006, **65**(2): 261-270.
- Mounkaila M. Spectral and Mineralogical Properties of Potential Dust Sources
 on a Transect from the Bodélé Depresseion (Central Sahara) to the Lake Chad
 in the Sahel. *Hohenheimer Bodenkundliche Hefte* 2006, **78:** 1-311.
- Herrmann L, Jahn R, Maurer T. Mineral dust around the Sahara—from source
 to sink. A review with emphasis on contributions of the German soil science
 community in the last twenty years. *Journal of Plant Nutrition and Soil Science* 2010, **173**(6): 811-821.
- 887 92. Tiedemann R. Acht Millionen Jahre Klimageschichte von Nordwest Afrika
 888 und Paläo-Ozeanographie des angrenzenden Atlantiks: Hochauflösende
 889 Zeitreihen von ODP-Sites 658-661. Christian-Albrechts-Universität, 1991.
- 891 93. Cohen AS, O'Nions RK, Siegenthaler R, Griffin WL. Chronology of the
 892 pressure-temperature history recorded by a granulite terrain. *Contributions to*893 *Mineralogy and Petrology* 1988, **98**(3): 303-311.
- Pin C, Zalduegui JS. Sequential separation of light rare-earth elements, thorium and uranium by miniaturized extraction chromatography: Application to isotopic analyses of silicate rocks. *Analytica Chimica Acta* 1997, **339**(1–2): 79-89.
- 900 95. Vance D, Archer C. Isotopic constraints on the origin of Heinrich event precursors. *Goldschmidt Conference Abstract*; 2002.

902		
903	96.	Tanaka T, Togashi S, Kamioka H, Amakawa H, Kagami H, Hamamoto T, et
904		al. JNdi-1: a neodymium isotopic reference in consistency with LaJolla
905		neodymium. Chemical Geology 2000, 168(3-4): 279-281.
906		
907	97.	Jacobsen SB, Wasserburg GJ. Sm-Nd isotopic evolution of chondrites. <i>Earth</i>
908		and Planetary Science Letters 1980, 30 (1): 139-135.
909	08	Dietze E at al An end member algorithm for decinhering modern detrital
910 011	90.	processes from lake sediments of Lake Donggi Cona, NE Tibetan Plateau
911 912		China Sedimentary Geology 243-244 169-180 (2011)
913		China. Seamentary Geology 243-244 , 109-100 (2011).
914	99	Wood S N Generalized additive models: an introduction with R (CRC
915	<i>))</i> .	press 2017)
916		press, 2017).
917	100.	Grinsted, A., Moore, J. C. & Jevrejeva, S. Application of the cross wavelet
918		transform and wavelet coherence to geophysical time series. <i>Nonlinear</i>
919		processes in geophysics 11, 561-566 (2004).
920		
921	101.	Hammer, Ø., Harper, D. A. T. & Ryan, P. D. PAST: Paleontological Statistics
922		Software Package for Education and Data Analysis. Palaeontologia
923		<i>Electronica</i> 4 , 9 (2001).
924		
925	102.	Castillo, S. et al. Trace element variation in size-fractionated African desert
926		dusts. Journal of Arid Environments 72, 1034-1045 (2008).
927		
928		



Extended Data Figure 1| Comparison between hydroclimate proxies measured at Site 659. a, b, c Bulk sediment [Al+Fe]/[Si+K+Ti] (orange), δD signatures of C₃₁ *n*-alkanes (pale blue, 1 σ error bars)^{39,40}, ln[Zr/Rb] (green, pale colours indicate low element counts). d, e, f Generalised additive model (GAM) fit of Site 659 [Al+Fe]/[Si+K+Ti] data shown by black line with grey shaded confidence band (2 standard error). Black crosses indicate original data points with resampled data points used in Kendall's tau-b correlation tests indicated by red circles (see Supplementary Information). a,d late Pleistocene, b,e late Pliocene, c,f early Pliocene.



Extended Data Figure 2| Coherency spectra comparing our data from Site 659 to published dust and hydroclimate records from the same and nearby sites. Green dashed line marks 90% Monte Carlo false-alarm level. **a** & **b** Coherency between the dust % estimates from Site 659 of Tiedemann et al. $(1994)^2$ and our ln[Ca/Fe] (**a**) and dust flux (**b**) estimates over the last 8 Myr. **c** Coherency between our estimated dust fluxes and those of Skonieczny et al. $(2019)^{15}$ from nearby site MD03-2705 over the last 240 kyr. **d**, **e**, **f** Coherency between our [Al+Fe]/[Si+K+Ti] values and C₃₁ *n*-alkane δ D values from Site 659 from Kuechler et al. $(2013, 2018)^{39,40}$ for three time slices in the Quaternary (**d**) and Pliocene (**e**, **f**).



Extended Data Figure 3| Comparison between methods for calculating dust fluxes to marine sediments over the last 250 kyr. a Red: modal dust flux estimates from Site 659 based on a geochemical end-member unmixing approach, with orange lines marking ± 1 standard deviation of 500 realizations. Blue: Dust flux estimates from site MD03-2705 (directly adjacent to Site 659) calculated by ²³⁰Th normalization¹³ with error bars indicating ± 1 standard deviation. **b** Generalized Additive Model⁹⁹ fit of Site 659 median dust fluxes shown by black line with grey shaded confidence band (2 standard error). Black crosses indicate original data points with resampled data points used in Kendall's tau-b correlation tests indicated by red circles (see Supplementary Information).



Extended Data Figure 4| Box and whisker plot illustrating co-variation in sediment colour and geochemistry at Site 659. Top: [Al+Fe]/[Si+K+Ti], bottom: ln[Zr/Rb]. Data are plotted from "light" and "dark" sediment layers (as defined by the method described in the Supplementary Information) for each of the three time stages (Stage I: 11.15 - 6.7 Ma, Stage II: 5.75 - 3.5 Ma, Stage III: 2.25 - 0 Ma). Box indicates interquartile range (IQR) with line marking median value, outliers (>1.5 IQR from median) marked with circles and whiskers drawn to the maximum/minimum values excluding outliers. Mann-Whitney-Wilcoxon tests were used to test the null hypothesis that samples from light and dark layers have identical continuous distributions with equal medians for each time interval. All resulting *p*-values were <0.001, giving >99.9% confidence that the differences between light and dark layers are significant.



Extended Data Figure 5 Wavelet analysis of Site 659 geochemical records. Comparison between continuous wavelet power spectra of calibrated $\ln[Ca/Fe]$ (top) and [Al+Fe]/[Si+K+Ti] (bottom) data on the astronomically-tuned age model (left) and an untuned age model based solely on biostratigraphic and magnetostratigraphic datums^{14,69} (right). Thick black contours designate the 5% significance level against red noise and the cone of influence is shown as a lighter shade, where edge effects may cause distortion. Data were detrended and smoothed (5-pt moving average) prior to the wavelet analyses. Separate spectra were also generated for the older and younger sections of the full record to reduce the impact of temporal changes in cycle amplitude on the detected frequencies. Analyses were performed and figures generated using the Matlab code of Grinsted et al. (2004)¹⁰⁰. See Supplementary Information for further discussion.



Extended Data Figure 6 REDFIT spectral analysis⁷⁵ **of Site 659 geochemical records.** Top: $\ln[Ca/Fe]$ ratios, middle: calibrated [Al+Fe]/[Si+K+Ti] ratios, bottom: median dust flux values. Data are divided into the three time stages discussed in the text. Left: Stage III (2.25–0 Ma), centre: Stage II (5.75–3.5 Ma), right: Stage I (11.15–6.9 Ma). Green curves mark the false-alarm level at the 95% confidence level, red curves indicate AR(1) red noise models. Orange lines and numbers indicate the frequencies equivalent to periods (in kyr) of major astronomical cycles (precession, obliquity and eccentricity). Analysis performed and figures created using PAST software¹⁰¹. See Supplementary Information for further discussion.



Extended Data Figure 7| Running statistical analysis of Site 659 geochemical data, comparing 1 Myr data bins. a Mann-Whitney-Wilcoxon test log(p) values to detect shifts in central tendency (see Supplementary Information). Low values indicate extremely low probabilities that the two data bins have the same central tendency. **b** Estimated difference in location between the two data bins divided by the interquartile range of the complete data set, with 95% confidence interval plotted. Note that [Al+Fe] is plotted on an inverted axis. **c** Ansari-Bradley test log(p) values to detect shifts in dispersion. Low values indicate extremely low probabilities that the two data bins have the same dispersion. d Ratio of scales between the two data bins, with 95% confidence interval plotted. Orange: calibrated [Al+Fe]/[Si+K+Ti], green: ln[Zr/Rb] (with XRF counts <300 removed), red: median dust flux (g cm⁻² kyr⁻¹). Grey shading indicates intervals of greatest change in the geochemical time series revealed by statistical analyses.



Extended Data Figure 8 Cross-plots of strontium and neodymium isotopic signature of lithic fraction of Site 659 sediments. Data coloured by: **a** the proportion of the lithic fraction attributed to dust by end-member unmixing ([dust]/[dust+riverine]), **b** co-registered ln[Zr/Rb] values. Red marks samples dominated by dust/coarse grains, blue marks samples dominated by riverine inputs/fine grains. **c** Data coloured by age, where stage I (pink) is the oldest (>6.9 Ma) and stage III (blue) is the youngest (<2.25 Ma) and grey indicates samples from the transition between stages II and III (3.5–2.25 Ma). Individual samples are marked by crosses and mean values for each age range shown by circles, with error bars indicating 1 standard deviation.



Extended Data Figure 9| Grain size distributions of the end-members calculated from the lithic fraction of ODP Site 659. Grain size derived end-member 1 (EM1) in blue and grain size derived end-member 2 (EM2) in red compared to: **a** Modern dust samples recorded offshore NW Africa from Stuut et al. (2005) in grey⁶⁴. **b** As (a) but with just the most proximal measurements to Site 659 (M41/1 D4, centered at 19.73°N, 17.91°W) plotted. **c** The grain size end members of Tjallingii et al. (2008) from site GeoB7920-2 (20.75°N, $18.58^{\circ}W)^{12}$. Yellow and orange dashed lines indicate the end members attributed to fine and coarse dust respectively, with the riverine end member shown in green. **d** All Site 659 lithogenic grain size distributions (black).



Extended Data Figure 10| Comparisons between grain size and geochemical proxies at Site 659. a & b [Al+Fe]/[Si+K+Ti] ratios of the coarse and fine fractions of (a) 12 sediment samples from Site 659 sieved at 10 μ m and analysed by discrete XRF analysis with sample ages are listed along the top and (b) desert surface soil and aeolian dusts from four locations in the Sahara-Sahel dust corridor, analysed by inductively coupled plasma mass spectrometry and grouped into <12 μ m and >12 μ m size fractions, from Castillo et al. (2008)¹⁰². CB: Chad Basin, HM: Hoggar Massif, WS: Western Sahara, HAR: Harmattan. c and d The proportion of lithogenic grain size derived end-member 1 (grain size EM1, attributed to fine riverine inputs) plotted against the sediment geochemical ratios c [Al+Fe]/[Si+K+Ti] and d ln[Zr/Rb]. Data points are coloured by sample age, where the youngest samples are in blue and the oldest in orange.

Supplementary Information

For: Astronomically controlled aridity in the Sahara since at least 11 million years ago

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Supplementary Discussion

Comparison to published dust reconstructions

Reconstructions of the flux of dust from the African continent deposited in ODP Site 659 sediments over the past 5 Myr were originally published by Tiedemann et al. $(1989)^{1}$ and Tiedemann et al. $(1994)^2$. Their records are based upon the assumption that the proportion of dust in the sediment can be approximated from the non-carbonate fraction (=100-%CaCO₃). We find an excellent correlation between the discrete measurements of these earlier studies (which include a range of analytical techniques)¹ and ln[Ca/Fe] values measured by XRF core scanning (Supplementary Figure 4). A Kendall's tau-b correlation test between the dust % estimates of Tiedemann et al. $(1994)^2$ and those presented here reveals a positive, extremely significant correlation ($\tau = 0.669$, *p*-value < 0.001), with a comparison between published dust % to $\ln[Ca/Fe]$ revealing an even stronger (but negative) correlation ($\tau = -0.711$, p-value <0.001). These results show that our ln[Ca/Fe] data provide reliable down core records of terrigenous sediment content at ODP Site 659. We note, that these datasets, in common with many palaeoclimate records, likely show autocorrelation (where there is a degree of similarity between a time series and a lagged version of itself). This acts to elevate the calculated correlation coefficients between data series. We therefore also show coherency spectra, calculated using the REDFIT-X program of Ólafsdóttir et al. (2016)³. This approach uses a Lomb-Scargle Fourier transform to perform cross-spectral analyses between unevenly spaced data sets. Monte Carlo simulations that mimic key properties of the datasets (e.g. autocorrelation and time spacing) are used to estimate the false-alarm level for the calculated coherency. These spectra are shown in Extended Data Figure 2. Significant coherency between the dust % estimates of Tiedemann et al. (1994) and both our ln[Ca/Fe] and dust % estimates across almost all frequencies supports the strong agreement among the different techniques (Extended Data Figure 2a,b).

Recent work has shown that, in addition to dust inputs, fine-grained riverine sediment is a likely contributor to Site 659 and the nearby continental margin^{4,5}. We therefore partitioned the bulk terrigenous sediment component into two separate components (classified as riverine and aeolian) by applying an unmixing approach (Methods). Our absolute dust fluxes are therefore lower than those of Tiedemann et al. $(1994)^2$ (Supplementary Figure 4), which were calculated by assuming that all terrigenous material at Site 659 was wind-blown in origin. Nevertheless, for the last 5 Myrs, where there are detailed records from Tiedemann et al. $(1994)^2$, the down core trends in the data yielded by the two different approaches are generally very consistent over both secular and astronomical timescales.

For sediments younger than about 200 to 300 kyr, thorium-normalization of sediment accumulation rates has been used to provide age control independent of sediment focusing and dissolution along the NW African margin⁶⁻¹⁰. To assess whether these processes likely exerted a major control over our dust flux reconstructions we compared our calculated fluxes to those from site MD03-2705⁹ (18°05′N, 21°09′W). Because dust flux estimates were calculated at

different age points between the two sites, we calculated generalized additive models $(GAMs)^{11}$ using R^{12} to create a smoothed fit to our datasets and resampled this at ages corresponding to the ²³⁰Th-normalised data points (Extended Data Figure 3). This approach provides a good representation of the original dataset, except for the most recent 20 kyr, where GAMs struggle to capture the high frequency, high magnitude variability seen at both Site 659 and MD03-2705. The degree of correlation between the two dust flux reconstructions was then assessed using a Kendall's tau-b correlation test. Despite uncertainty introduced by the GAM approximations and chronological uncertainties between the two sites, we find a positive, strongly significant correlation ($\tau = 0.298$, *p*-value < 0.001) between our dust fluxes at Site 659 and those calculated by ²³⁰Th-normalisation at proximal site MD03-2705⁹ (Extended Data Figure 3). We also performed cross-spectral analysis between the two datasets using REDFIT- X^{3} (where no resampling is required). The results of these analyses indicate that there is largely significant coherency between data sets across the frequency range 0.025-0.23 cycles/kyr (Extended Data Figure 2c), equivalent to periodicities of 40–4 kyr. These results suggest that our dust flux reconstructions are representative of aeolian transport from the continent and not major sediment redistribution in the ocean or rapid pulses of dissolution, at least over the past 240 kyr where ²³⁰Th-normalised dust flux estimates exist.

Comparison to established XRF ratios and other hydroclimate proxies

Our discussion focuses on two geochemical ratios ([Al+Fe]/[Si+K+Ti] and ln[Zr/Rb]) provided by our XRF work but our interpretations gain further support from three other elemental ratios that have also been shown to closely track climatic gradients along the African coastline today. Ti/Al is proposed as a proxy for aeolian versus fluvial input and Fe/K and Al/Si are suggested as tracers of terrestrial climatic zones¹³, although Al/Si can also be influenced by high siliceous microfossil concentrations at times of intense upwelling¹⁴. All three of these ratios agree very closely with one another and with [Al+Fe]/[Si+K+Ti] over astronomical timescales throughout our records (Supplementary Figures 5 and 6). Dark sediment layers consistently record high ln[Ti/Al] values and low ln[Fe/K] and ln[Al/Si] values, indicating a dominance of aeolian over riverine sediment and a decrease in the intensity of chemical weathering, a trend documented with increasing aridity associated with increasing distance from the equator today¹³. These findings strongly support our interpretation that the dark sediment bands represent dust pulses associated with increased aridity throughout the last 11 Myr.

The different elemental ratios also show clear similarity over multi-million year timescales. The onset of more extreme wet events between 7 and 6 Myr can be seen in all ratios, as can a shift towards more arid conditions over the past \sim 3 Myr. A decrease in ln[Al/Si] through the Pliocene and Pleistocene is more pronounced than the other ratios, and, given the very low concentrations of siliceous microfossils recorded in the Site 659 sediments¹⁵, may indicate an increase in the proportion of silica-rich terrigenous material reaching the site, such as pale quartz grains from coastal sources¹⁶.

We also compare our [Al+Fe]/[Si+K+Ti] and ln[Zr/Rb] data to an independent hydroclimate proxy: the hydrogen isotopic (δ D) signature of *n*-alkanes (Extended Data Figure 1). These compounds are found as leaf waxes in terrestrial plants, with δ D values at Site 659 suggested to largely depend upon the amount of rainfall, although differences in plant physiology (e.g. photosynthetic pathway, rooting depth and leaf anatomy) and climate (e.g. moisture source region, evaporation) can also influence plant wax δ D signatures¹⁷. Extended Data Figure 1 illustrates the strong agreement between C₃₁ *n*-alkane δ D values and our XRF-based hydroclimate proxies during several snapshots of the Pliocene and Pleistocene (where δ D data are available)^{18,19}. We also test the similarity between the datasets statistically. Our data and the reported δD values are from the same site but different sample sets were used so we apply GAMs to create a smoothed fit to our [Al+Fe]/[Si+K+Ti] record and resample it at the matching depths to the δD data (Extended Data Figure 1d-f). Separate fits were made for 0.14–0 Ma, 3.27–3 Ma, 3.63–3.34 Ma, 4.83–4.61 Ma and 4.99–4.83 Ma to ensure that the GAMs captured astronomical-scale variability. We combined the data from all of the time windows to perform a Kendall's tau-b correlation test with the full δD dataset, which revealed a negative, strongly significant correlation with [Al+Fe]/[Si+K+Ti] ($\tau = -0.218$, *p*-value <0.001). Note that a negative correlation indicates agreement between the proxies, with higher [Al+Fe]/[Si+K+Ti] values but more negative δD values indicating a wetter climate. The strength of the relationship between our XRF data and the δD results (Extended Data Figure 1a-c) support the use of [Al+Fe]/[Si+K+Ti] to reconstruct past changes in continental hydroclimate in these sediments.

We also performed cross-spectral analysis to assess how the coherency of [Al+Fe]/[Si+K+Ti] and δD varies with frequency (Extended Data Figure 2d-f). These analyses were carried out for three separate time slices due to the discontinuous nature of available high-resolution δD data. We find the strongest coherency occurs at frequencies of ~0.03–0.05 cycles/kyr, implying that these records are coherent over precessional timescales, both responding to insolation-forced variability in hydroclimate. Outside this range, the coherency is lower. This result may indicate the influence of secondary processes acting on one or both proxies over different timescales.

Geochemical differences between light and dark layers

Colour data were generated from the central portion of ODP core photograph images¹⁵ along our splice using Code for Ocean Drilling Database macros²⁰, with an average sample spacing of \sim 2mm. Lightness values of >80 or more than 3 standard deviations above or below a 4 cm rolling mean value were excluded to filter out artefacts such as cracks and core gaps from the dataset. Lightness data were then smoothed by calculating the mean value over a 4 cm rolling window.

To identify "light" and "dark" sediment layers, we filtered out longer-term variability in the lightness data by calculating residuals from a 5 m rolling mean. The residual dataset was then resampled to match the depths corresponding to XRF data using simple linear interpolation. "Light" layers were defined as those with residual lightness values greater than the 60% percentile of the rolling 5 m window, while dark layers were defined as those with lightness values below the 40% percentile.

Extended Data Figure 4 illustrates the geochemical differences between sedimentary layers classified as "light" and "dark" in the Site 659 XRF datasets. Mann-Whitney-Wilcoxon tests were used to identify whether there were significant geochemical differences between the light and dark layers for each of our three time intervals. Extremely low *p*-values (<0.001) are documented for both [Al+Fe]/[Si+K+Ti] and ln[Zr/Rb] in each of the three time windows, which shows that there are clear, highly significant geochemical differences between light and dark sediment layers at Site 659.

Identification of change points in time series

To identify the intervals of greatest change in our geochemical time series, we applied running Mann-Whitney-Wilcoxon (also known as Wilcoxon rank sum) and Ansari-Bradley statistical tests, following the approach of Trauth et al. $(2009)^{21}$. We test whether samples taken from 1 Myr ranges either side of a selected age come from distributions with the same properties, or

whether there were significant differences between them. We used 1 Myr bins to smooth out the influence of shorter frequency astronomical variability and highlight secular changes in the datasets.

[Al+Fe]/[Si+K+Ti], ln[Zr/Rb] and median dust flux data sets were each resampled using simple linear interpolation to give an average data resolution of 2 kyr. A Mann-Whitney-Wilcoxon test was applied to test the null hypothesis that samples from the million years preceding and postdating the chosen date were selected from identical continuous distributions with equal medians.

The Ansari-Bradley test determines whether two samples come from continuous distributions with the same median and shape but different dispersions. As the results of the Mann-Whitney-Wilcoxon test indicate that there are significant shifts in the median values of all three of our proxy data sets between different time windows, we calculate the difference of each data point from the whole-core mean and then adjust these residuals by the difference in location between the two time windows (the median of the difference between a sample from each of the two distributions). Due to the size of the datasets, a normal approximation was used when calculating the p-values.

The results of the running Mann-Whitney-Wilcoxon and Ansari-Bradley tests on our three proxy datasets reveal two major intervals of change in our time series; 6.9-5.75 Ma and 3.5-2.25 Ma (Extended Data Figure 7). The first interval is marked by a major increase in the variability of all three proxy records from 6.9-6.3 Ma, accompanied by an increase in the central tendency of [Al+Fe]/[Si+K+Ti] values. This is followed by a decrease in ln[Zr/Rb] values (6.2-5.75 Ma).

During the second key interval of change, a decrease in the central tendency of [Al+Fe]/[Si+K+Ti] (centred around 3.2–3.1 Ma) again leads change in the ln[Zr/Rb] data (which shows an increase centred around 2.8–2.25 Ma), although this time there is also an increase in the mean dust flux values approximately synchronous with the change in ln[Zr/Rb]. There is a shift to greater dispersion in the dust fluxes centred around 2.7 Ma, but no concurrent change in the amplitude of variability of [Al+Fe]/[Si+K+Ti] or ln[Zr/Rb].

Attribution of sedimentological end-members

Our two lithogenic grain size end-members identified at Site 659 have clearly distinct grain size distributions. End-member EM1 is by far the finer of the two, with modal grain sizes of 0.7–4 μ m. End-member EM2 has a sharp, unimodal distribution dominated by grains of 10–40 μ m diameter.

There is remarkable agreement between the grain size distributions of Site 659 lithogenic endmember 2 (EM2) and dust samples taken offshore NW Africa²², particularly those sampled closest to Site 659 (Extended Data Figure 9). EM2 is also similar to the fine dust end-member identified at NW African site GeoB7920-2 by Tjallingii et al. (2008)⁴ (Extended Data Figure 9). We therefore attribute EM2 to windblown dust exported from the African continent.

The origin of EM1 is less immediately obvious. Distal dusts transported long distances from their sources are suggested to have volume median grain sizes of ca. $2-7 \ \mu m^{23-26}$, therefore, it is possible that EM1 at Site 659 also represents a dust component, albeit much finer than EM2. However, Site 659 end-member 1 (EM1) contains a much higher proportion of grains <2 μm than any of the atmospheric dust samples recorded offshore northwestern Africa by Stuut et al. $(2005)^{22}$ (Extended Data Figure 9a), and is finer than either samples dominated by either wet or dry deposition captured by sediment traps offshore Cape Blanc²⁷. EM1 is also finer than

modern African dust captured across the Atlantic throughout the year and more distal Africansourced dust in Puerto Rico^{23,28,29}. These observations strongly suggest that EM1 is unlikely to represent a pure dust component, at least under wind conditions similar to modern.

We note that EM1 has a very strong similarity to the grain size distribution of the fine lithogenic end-members identified by both Holz et al. $(2004)^{30}$ and Tjallingii et al $(2008)^4$, which are attributed to distal rain-out of fine particulates from riverine sources. Networks of river channels, active during past intervals of humidity have been identified across North Africa, many of which drain into the Atlantic^{5,31}, with submarine channels transporting this material into the deep ocean^{32,33}. A distal riverine interpretation for fine-grained sediments at Site 659 is supported by high proportions of the fine end-member in core top sediments offshore Morocco which receive sediment from the Souss River³⁰ and the fact that highest proportions of this fine end-member occur at times where enhanced rainfall is well evidenced in North Africa, such as African Humid Period 1^{4,34}, although increased fluxes of fine sediments are not consistently observed at all sites on the northwest African margin at this time⁸. The elevated location of Site 659 on the Cape Verde Plateau means that hypopycnal transport would likely be required for material of riverine origin to reach the site. EM1 at Site 659 is slightly finer than the riverine end-member identified at site GeoB7920- 2^4 (Extended Data Figure 9c), which is consistent with the greater distance from the continent of Site 659 compared to GeoB7920-2 and the requirement for sediment to be suspended within the water column to reach Site 659 due to its elevated position on the Cape Verde Rise (Supplementary Figure 1).

A third possible contributor to EM1 is resuspension of sediments from the continental shelf. Although shelf sediments are dominated by grains much coarser than the aeolian inputs, potentially representing relict dune fields³⁵, fine grained components are preferentially transported in nepheloid layers³⁶. Nepheloid layer activity is currently very low around Cape Verde^{37,38}, however, a contribution of these processes to Site 659 cannot be ruled out.

To further explore the origin of EM1, we test for geochemical differences between the fine and coarse components of the sediments. 12 samples with a range of ages and lithologies were wet sieved at 10 μ m (close to the boundary between EM1 and EM2 dominance) to separate out the coarse and finer fractions. Samples were then oven dried and homogenized, then mixed with di-lithium tetraborate and fused to create beads. These were analysed by X-ray fluorescence using a Phillip MagiX Pro WD-XRF at the University of Southampton Waterfront Campus, NOCS, calibrated using international rock standards prepared in the same way as our samples.

All 12 of our samples show a clear offset in [Al+Fe]/[Si+K+Ti] between the >10 µm and <10 µm size fractions (Extended Data Figure 10a). The <10 µm [Al+Fe]/[Si+K+Ti] values are 0.11–0.46 greater than the <10 µm fraction, which means that the fine grained components plot much closer to the value of suspended sediments from the Senegal River than the coarse fractions do. The largest size fraction differences are generally recorded in the Miocene aged samples from depths with high terrigenous contents (indicated by high ln[Fe/Ca] values and a dark colour). In addition, there is a strong correlation between the relative proportions of grain-size end-members and our geochemical ratios for past dust inputs (Extended Data Figure 10c,d). High proportions of EM1 are associated with both high [Al+Fe]/[Si+K+Ti] values (similar to modern riverine inputs) and low ln[Zr/Rb] values which are commonly interpreted as indicating a dominance of fine-grained clays over coarse dust^{4,39}. These results are strongly supportive of a distal riverine contribution to EM1.

We note that differences in major element compositions might also be expected between coarse and fine fractions of a purely aeolian sample due to the susceptibility of different minerals with different chemical compositions to be weathered into finer particles. However, samples of dusts and dust source sediments from across the Sahara-Sahel dust corridor show negligible offsets in [Al+Fe]/[Si+K+Ti] between size fractions (Extended Data Figure 10b)⁴⁰. Therefore, we suggest that the differences we record in major element signatures between the <10 μ m and >10 μ m size fractions reflect inputs from multiple sources rather than size-dependent fractionation from a single source.

The higher [Al+Fe]/[Si+K+Ti] values of the <10 μ m sediment component (Extended Data Figure 10) mean that our geochemical unmixing approach acts to remove this very fine component. Therefore, even if EM1 is dominantly derived from resuspended shelf material rather than distal riverine material, the geochemical unmixing approach will still act to remove its contribution from the total lithogenic component to produce our dust flux estimates. If EM1 contains a component of fine-grained dust, then our dust flux estimates are likely to be underestimates.

In conclusion, a dominantly distal riverine origin for EM1 is indicated by: (i) grain size distribution analyses showing that EM1 is very similar to the fine end-member previously identified at higher concentrations during humid time intervals^{4,34} and at latitudes to the north of the Sahara where precipitation is higher³⁰ (ii) EM1 is much finer than either proximal or distal North African dust (including samples from as far as the Caribbean)^{22,23,29,41}, (iii) large geochemical differences between the coarse and fine fractions of Site 659 sediments are too great to be explained by size-dependent fractionation from a single source, and (iv) the current absence of major nepheloid layer activity around Cape Verde^{37,38}.

Effects of grain size on radiogenic isotope signatures

A grain size effect on Sr isotope signatures of sedimentary lithics has previously been documented including in African dust, with fine-grained samples having higher ⁸⁷Sr/⁸⁶Sr values, although a similar effect in Nd isotopes is not commonly observed⁴²⁻⁴⁴. We find no strong relationship between the radiogenic isotopic signature and either the proportion of dust or the grain size of the sediment as approximated by ln[Zr/Rb] (Extended Data Figure 8a,b), although we note a tendency for samples with the most radiogenic Sr and unradiogenic Nd isotope signatures to have low ln[Zr/Rb] values and be dominated by riverine inputs rather than dust (according to the results of our geochemical end-member unmixing), with the inverse also true, particularly in ln[Zr/Rb]. Radiogenic Sr values may be attributed to enhanced chemical weathering under humid climates resulting in a higher proportion of fine clay minerals and micas reaching Site 659^{42} . The covariation with unradiogenic ε_{Nd} values suggest there is also a shift in provenance, potentially attributable to transport of an increased proportion of proximal material sourced from the West African Craton transported by rivers compared to aeolian inputs, which may contain a greater contribution from the interior of the continent (or potentially Morocco, although the reported Sr isotope signatures⁴⁵ suggest this scenario is less likely).

Temporal variability in radiogenic isotope signatures

The radiogenic isotope signatures of the lithic fraction of the Site 659 sediments hint at subtle differences with sediment age, but we record no clear temporal shifts in the provenance of lithogenic material through the last 11 Myr to suggest major changes in the source regions of terrigenous sediment. The oldest samples (stage I) generally show more non-radiogenic isotope values in ⁸⁷Sr/⁸⁶Sr (Extended Data Figure 8c), which means that these Miocene-aged samples plot slightly closer to the Central source area than to the Western region, as defined by the currently available data⁴⁶ (Figure 4). Therefore, we infer that the strong signal of dust

inputs during stage I cannot be attributed to input from coastal sand dunes (which have previously been documented back to the Early Miocene and potentially much earlier^{47,48}) and instead indicate the presence of strong dust generating regions in the continental interior back to at least the Late Miocene.

Fidelity of an astronomically-tuned age model at Site 659

Pronounced lithological and geochemical variability is well documented at Site 659, particularly over the younger intervals, and has been linked to astronomical forcing^{2,19,49-51} (e.g. Figure 3, Supplementary Figure 6). These observations therefore provide an opportunity to capitalize on the potential to gain a high-fidelity age model for Site 659 by tuning the clear lithological cycles to an astronomical signal, as with many records of Mediterranean climate and low-latitude continental margins where a similar tuning approach has been used for chronology⁵²⁻⁵⁶. However, this approach requires care to ensure that age is assigned to our environmental records as independently as possible to guard against circularity. In our final age model, we used the core images and the ratio ln[Ca/Fe] to tune to insolation and [Al+Fe]/[Si+K+Ti] and ln[Zr/Rb] as our main records of environmental change.

To assess the extent to which astronomical power is present throughout the Site 659 records, we first analysed the frequencies present in our data using wavelet analysis⁵⁷ with an age model was based solely on bio- and magnetostratigraphic datums^{15,58} and no tuning applied to the records (Extended Data Figure 5). The resulting continuous wavelet power spectrum show extremely similar patterns of variability on astronomical periodicities to our final, astronomically tuned record in both ln[Ca/Fe] and [Al+Fe]/[Si+K+Ti] across the full 11 Myr datasets (Extended Data Figure 5). This result indicates that the astronomical periodicities identified in the lithology and geochemistry of Site 659 sediments are genuine and have not been artificially tuned into the data. We also note that significant power is often detected concurrently at two or more frequencies, with the ratios between these frequencies matching those of major astronomical cycles (Extended Data Figures 5 and 6). This result would be very difficult to tune into our records artificially, providing further evidence for the lithological and geochemical variability at Site 659 being astronomically-paced.

Changing astronomical pacing of hydroclimate variability in North African over the past 11 Myr

To assess how the dominant frequencies of variability change throughout our >11 Myr records, we performed spectral analysis on our $\ln[Ca/Fe]$, [Al+Fe]/[Si+K+Ti] and dust flux data using REDFIT⁵⁹ to identify significant periodicities in our unevenly spaced data series (Extended Data Figure 6). Analyses were carried out using the PAST software⁶⁰

REDFIT analysis reveals a strong signature of precession throughout the last 11 Myr in ln[Ca/Fe], [Al+Fe]/[Si+K+Ti] and dust flux data from Site 659 (Extended Data Figure 6), with obliquity also present in all three time intervals examined (stages I to III). Significant eccentricity is only recorded during stage I (>6.9 Ma), rather than the youngest 800 kyr when strong 100 kyr glacial cycles are recorded in benthic oxygen isotope records⁶¹. The signatures of obliquity and precession are seen in all three stages of our record, but are generally weakest in the oldest part of the record (stage I), likely, at least in part, because of the stronger smoothing effects of bioturbation where sedimentation rates are lower (Supplementary Figure 2).

The relative importance of high and low latitude forcing in driving African hydroclimate, particularly during the Quaternary Period, is debated^{2,9,62-67}. The strong presence of precessional frequencies (23 and 19 kyr) in our data (Extended Data Figure 6) supports a major role for insolation forcing of the West African monsoon. Obliquity (41 kyr) is also

strongly present, which may indicate that the ice volume at high latitudes also exerts a major control on North African hydroclimate, perhaps by influencing wind strength. However, obliquity can also influence low latitude hydroclimate directly through modulation of the cross-equatorial insolation gradient⁶⁸. The absence of an obvious strengthening of the obliquity signature associated with increasing amplitude of glacial-interglacial cycles in the Late Pliocene/early Pleistocene⁶¹ in our hydroclimate proxy [Al+Fe]/[Si+K+Ti] (e.g. Extended Data Figure 5) supports this interpretation. The signal of obliquity in the latest Pliocene/Pleistocene appears slightly stronger in the ln[Ca/Fe] and dust flux records than [Al+Fe]/[Si+K+Ti] which may indicate the influence of glacial-interglacial shifts in overturning circulation bringing more corrosive bottom waters to the site driving carbonate dissolution⁹ and/or higher latitude forcing exerting a stronger control over wind strength (and hence dust transport) than continental hydroclimate.

Interpreting plant wax history at Site 659

Long-chain *n*-alkanes originating from the epicuticular leaf waxes of terrestrial plants can be removed from the leaf surface by the action of wind or rain and transported to the ocean via both dust and rivers⁶⁹⁻⁷¹. A strong correlation between concentrations of *n*-alkanes and dust percentages at Site 659 suggests that transport by wind is the dominant process at this location^{2,19}. The source regions of plant waxes are likely very similar to those of the terrigenous material reaching Site 659, however, given the very low net primary production across much of North Africa, the plant wax signature may be biased to areas with higher biomass densities, particularly during arid time intervals.

Previous studies suggest that Site 659 records plant waxes dominantly sourced from the northern Sahel, with biomarker and pollen records suggesting a minor additional input of terrestrial material transported by the northeast trade winds from more northerly latitudes, evidenced during times of aridity and increased trade wind strength such as the Last Glacial Period, but not during the warmer and wetter Pliocene^{18,19,72,73}. The Sahel is currently dominated by C₄ grasslands, while the vegetation in northernmost Africa is instead dominated by C₃ plants⁷⁴. Therefore, the northwest African margin currently receives a mixture of plant waxes from these two sources. Several studies^{72,75} estimate approximately equal proportions of inputs from C₄ and C₃ plants to recent and Late Quaternary aged marine sediments around the latitude of Site 659, however other, more recent studies point to a much higher contribution from C₄ plants^{18,76}.

Combined evidence from plant wax carbon isotopes and pollen indicates C₄ savanna grasslands directly replaced savannas dominated by C₃ grasses in both eastern and southern Africa as C₄ vegetation expanded through the Neogene^{77,78}. Evidence of Poaceae (grasses) in the Niger delta as far back as the early Miocene⁷⁹ suggests a similar sequence of events unfolded in the Sahel, with a gradational increase in plant wax δ^{13} C of ca. 10 ‰ at Site 659 indicating an expansion of C₄ vegetation starting approximately 10 Myr ago⁸⁰. Superimposed on this shift (at least during the Pliocene and Late Quaternary where high-resolution data exist) is variability in δ^{13} C driven by precession, obliquity and eccentricity¹⁹. These cycles have a typical magnitude of 0.5–2.5 ‰, much smaller than the multi-million year trend in plant wax δ^{13} C (Figure 2, Supplementary Figure 3).

We note that the relatively muted signature of astronomically-forced variability recorded at Site 659 is not observed at all locations across North Africa. Some localities record variability in plant wax δ^{13} C of >8 ‰ over astronomical timescales⁸¹⁻⁸⁶, most commonly records from lakes in East Africa where the topography can lead to diverse ecosystems in closer proximity to one another than further west. Lacustrine sediments also typically record larger variability in plant wax δ^{13} C than distal marine sediment cores⁸⁷ because lake archives document more

proximal vegetation sources, can be influenced by local tectonics and hydrography particularly in tectonically active regions such as East Africa, and may also be biased by microbiallyderived *n*-alkanes⁸⁷. Marine records integrate source material from across a large region and are less sensitive to subtle vegetation shifts than lake records although they can be influenced by transport processes^{19,87}.

A larger range of variability in *n*-alkane δ^{13} C in the Late Quaternary than Site 659 is also recorded at northwest African margin sites GeoB9508-5 (15.5°N, -21.2 to -26.5 ‰, ~65-100 % C₄)⁷⁶ and GeoB9528-3 (9.2°N, -21.8 to -28 ‰, ~55–100 % C₄)⁸⁸ (21.0°N, -23 to -25.7 ‰, ~70-90% C₄). Both GeoB9508-5 and GeoB9528-3 lie further south and closer to the continent than Site 659. Therefore, a scarcity of sources of waxes carrying very low carbon isotopic signatures in the Sahara due to a regional dominance of C₄ vegetation could be responsible for low amplitude variability in δ^{13} C at Site 659. That said, we note that the amplitude of astronomically-forced variability in *n*-alkane δ^{13} C at both GeoB9508-5 and GeoB9528-3 is only around half the magnitude of the multi-million year shift associated with C₄ grassland expansion. We also observe that the magnitude of astronomically-forced variability in δ^{13} C at Site 659 appears slightly smaller in the Pliocene than in the Last Glacial Cycle, despite lower δ^{13} C values indicating a greater proportion of C₃ plants in the older time period (Figure 2, Supplementary Figure 3). The *n*-alkane δ^{13} C data from the Late Miocene are not of sufficient resolution to assess the magnitude of astronomically-forced variability, however we do not observe an obvious relationship between δ^{13} C and our geochemical ratios to suggest that there was strong aridity-driven cyclicity at a time when isotopic signatures indicate significant inputs from both C₃ and C₄ plants.

In summary, we conclude that it is unlikely that the muted astronomically-forced variability in δ^{13} C recorded in both the Pliocene and Quaternary at Site 659 (compared to the high amplitude astronomically-forced changes in hydroclimate) can simply be explained by the absence of a C₃ plants in the regions of Northern Africa sampled by our records.

References for *p***CO**² **reconstructions**

Badger et al. $(2013)^{89}$ Badger et al. $(2019)^{90}$ Bartoli et al. (2011)⁹¹ Bolton et al. $(2016)^{92}$ Chalk et al. $(2017)^{93}$ de la Vega et al. (2020)⁹⁴ Dvez et al. (2018)⁹⁵ Hönisch et al. (2009)⁹⁶ Martinez-Boti et al. (2015)⁹⁷ Pagani et al. $(2005)^9$ Pagani et al. (2010)⁹⁹ Seki et al. (2010)¹⁰⁰ Sosdian et al. (2018)¹⁰¹ Stap et al. $(2016)^{102}$ Super et al. (2018)¹⁰³ Tanner et al. $(2020)^{104}$ Zhang et al. (2013)¹⁰⁵ Zhang et al. $(2017)^{106}$ Zhang et al. (2019)¹⁰⁷

Supplementary Figures



Supplementary Figure 1 Map indicating the location of ODP Site 659 and nearby site GeoB7920, offshore NW Africa. Site locations marked by black circles, adapted from Ruddiman et al. (1989)¹⁵.



Supplementary Figure 2 Age depth plot for ODP Site 659. Age model created by benthic oxygen isotope correlation and orbital tuning used in this study (black line) compared to the original datums^{58,108} (J. Backman pers. comm.), with depths converted onto the new splice and out-of-splice sections stretched to correlate to the splice where necessary using XRF data, bulk properties and core images. Biostratigraphic and magnetostratigraphic ages updated following Raffi et al. $(2006)^{109}$, Wade et al. $(2011)^{110}$ & Ogg $(2012)^{111}$. Red crosses: planktonic foraminiferal datums, blue: nannofossil datums, purple: paleomagnetic reversals.



Supplementary Figure 3| Comparison of astronomically-forced variability and multimillion year trends in plant wax isotopes at Site 659. Top: δ^{13} C of C₃₁ *n*-alkanes, data from Kuechler et al. (2013, 2018, line only)^{18,19}, Polissar et al (2019, crosses)⁸⁰ and this study (plusses). Bottom: δ D of C₃₁ *n*-alkanes, data from Kuechler et al. (2013, 2018, line only)^{18,19} and Polissar et al (2019, crosses)⁸⁰. Error bars indicate 1 s.e.m in the data of Polissar et al. (2019) and 1 sd in the data from both Kuechler et al. (2013, 2018) and this study.



Supplementary Figure 4| Comparison of techniques for dust estimation at Site 659. Cross-plots of dust content as a percentage of the total sediment from Tiedemann et al. $(1994)^2$ plotted against: **a** % dust estimated by end-member unmixing (this study) and **b** ln[Ca/Fe] measured by XRF core scanning (this study). Measurements included in these plots were taken within 1 cm of each other. **c** Down core records of dust as a % of the total sediment by Tiedemann et al. 1994^2 (blue) and this study (red), from 0–5 Ma. All data have been converted onto the splice and age model developed in this study.



Supplementary Figure 5| Comparison of calibrated XRF ratios of Site 659 sediments spanning 0–11.2 Ma. **a** ln[Al/Si], **b** ln[Fe/K], **c** ln[Ti/Al], **d** [Al+Fe]/[Si+K+Ti], **e** ln[Ca/Fe]. Background image is a composited core photograph.



Supplementary Figure 6 Comparison of snapshots of calibrated XRF ratios of Site 659 sediment, with an example from each of zone I–III. Top: 0–1 Ma, middle 5–6 Ma, bottom 8–9 Ma. a ln[Al/Si], b ln[Fe/K], c ln[Ti/Al], d [Al+Fe]/[Si+K+Ti], e ln[Ca/Fe]. Background images are composited core photographs.



Supplementary Figure 7 | Comparison of measurements of major element abundances in the calibration sample set between Avaatech XRF core scanner counts and oxide concentrations measured by discrete sample XRF analysis. **a** Al, **b** Si, **c** K, **d** Ca, **e** Ti and **f** Fe.



Supplementary Figure 8| Stratigraphic tie-points used in the construction of the Site 659 age model. a,e LR04 benthic foraminiferal oxygen isotope stack⁶¹. b,f Site 659 *C. wullerstorfi* oxygen isotope values² converted onto on our new composite depth scale and age model. c,g,k,o Calibrated ln[Ca/Fe] values of Site 659 sediments. d,h,l,p Mean boreal summer insolation (June-August) at 65°N given by the La2004 orbital solution¹¹². i,m Calibrated

[Al+Fe]/[Si+K+Ti] ratios of Site 659 bulk sediment. **j**,**n** ln[Zr/Rb] values of Site 659 sediments (pale colour indicates low counts). Grey vertical lines mark the position of tie points used to generate the age model. The position of magnetostratigraphic boundaries as identified in the Leg 108 Scientific Results¹⁰⁸ are marked by stars, with listed ages from the Geologic Time Scale 2012 (GTS2012)¹¹¹, and biostratigraphic events (as identified in the Leg 108 Scientific Results⁵⁸ and J. Backman pers. comm.) are marked by bars, with ages updated following Raffi et al. (2006)¹⁰⁹. Magnetostratigraphic and biostratigraphic dates were used to guide tuning.

References for Supplementary Information

- 1 Tiedemann, R., Sarnthein, M. & Stein, R. in *Proceedings of the Ocean Drilling Program, Scientific Results.* Vol. 108, 241-277 (ODP, 1989).
- 2 Tiedemann, R., Sarnthein, M. & Shackleton, N. J. Astronomic timescale for the Pliocene Atlantic δ18O and dust flux records of Ocean Drilling Program Site 659. *Paleoceanography* **9**, 619-638 (1994).
- 3 Ólafsdóttir, K. B., Schulz, M. & Mudelsee, M. REDFIT-X: Cross-spectral analysis of unevenly spaced paleoclimate time series. *Computers & Geosciences* **91**, 11-18 (2016).
- 4 Tjallingii, R. *et al.* Coherent high- and low-latitude control of the northwest African hydrological balance. *Nature Geosci* **1**, 670-675 (2008).
- 5 Skonieczny, C. *et al.* African humid periods triggered the reactivation of a large river system in Western Sahara. *Nat Commun* **6**, 8751 (2015).
- 6 Francois, R., Frank, M., Rutgers van der Loeff, M. M. & Bacon, M. P. 230Th normalization: An essential tool for interpreting sedimentary fluxes during the late Quaternary. *Paleoceanography* **19** (2004).
- 7 Adkins, J., deMenocal, P. & Eshel, G. The "African humid period" and the record of marine upwelling from excess 230Th in Ocean Drilling Program Hole 658C. *Paleoceanography* **21**, PA4203 (2006).
- 8 McGee, D., deMenocal, P. B., Winckler, G., Stuut, J. B. W. & Bradtmiller, L. I. The magnitude, timing and abruptness of changes in North African dust deposition over the last 20,000 yr. *Earth and Planetary Science Letters* **371-372**, 163-176 (2013).
- 9 Skonieczny, C. *et al.* Monsoon-driven Saharan dust variability over the past 240,000 years. *Science Advances* **5**, eaav1887 (2019).
- 10 Kinsley, C. W. *et al.* Orbital- and Millennial-Scale Variability in Northwest African Dust Emissions Over the Past 67,000 years. *Paleoceanography and Paleoclimatology* **36**, e2020PA004137 (2022).
- 11 Wood, S. N. Generalized additive models: an introduction with R. (CRC press, 2017).
- 12 R: A language and environment for statistical computing. https://www.R-project.org/ (Vienna, Austria, 2020).
- 13 Govin, A. *et al.* Distribution of major elements in Atlantic surface sediments (36°N–49°S): Imprint of terrigenous input and continental weathering. *Geochemistry, Geophysics, Geosystems* **13**, Q01013 (2012).
- 14 Meckler, A. N. *et al.* Deglacial pulses of deep-ocean silicate into the subtropical North Atlantic Ocean. *Nature* **495**, 495-498 (2013).
- 15 Ruddiman, W. F., Sarnthein, M., Baldauf, J. G. & et al. Proc. ODP, Init. Repts., 108. (ODP, 1989).
- 16 Tjallingii, R. Application and quality of X-ray fluorescence core scanning in reconstructing late *Pleistocene NW African continental margin sedimentation patterns and paleoclimate variations* PhD thesis, University of Bremen, (2006).
- 17 Sachse, D. *et al.* Molecular Paleohydrology: Interpreting the Hydrogen-Isotopic Composition of Lipid Biomarkers from Photosynthesizing Organisms. *Annual Review of Earth and Planetary Sciences* **40**, 221-249 (2012).
- 18 Kuechler, R. R., Schefuß, E., Beckmann, B., Dupont, L. & Wefer, G. NW African hydrology and vegetation during the Last Glacial cycle reflected in plant-wax-specific hydrogen and carbon isotopes. *Quaternary Science Reviews* **82**, 56-67 (2013).
- 19 Kuechler, R. R., Dupont, L. M. & Schefuß, E. Hybrid insolation forcing of Pliocene monsoon dynamics in West Africa. *Climate of the Past* 14, 73-84 (2018).
- 20 Wilkens, R. H. *et al.* Revisiting the Ceara Rise, equatorial Atlantic Ocean: isotope stratigraphy of ODP Leg 154 from 0 to 5 Ma. *Clim. Past* **13**, 779-793 (2017).
- 21 Trauth, M. H., Larrasoaña, J. C. & Mudelsee, M. Trends, rhythms and events in Plio-Pleistocene African climate. *Quaternary Science Reviews* **28**, 399-411 (2009).
- 22 Stuut, J.-B. *et al.* Provenance of present-day eolian dust collected off NW Africa. *Journal of Geophysical Research: Atmospheres* **110**, D04202 (2005).
- 23 Reid, J. S. *et al.* Comparison of size and morphological measurements of coarse mode dust particles from Africa. *Journal of Geophysical Research: Atmospheres* **108(D19)**, 8593 (2003).
- Arimoto, R., Ray, B. J., Lewis, N. F., Tomza, U. & Duce, R. A. Mass-particle size distributions of atmospheric dust and the dry deposition of dust to the remote ocean. *Journal of Geophysical Research: Atmospheres* **102**, 15867-15874 (1997).
- 25 Maher, B. A. *et al.* Global connections between aeolian dust, climate and ocean biogeochemistry at the present day and at the last glacial maximum. *Earth-Science Reviews* **99**, 61-97 (2010).
- 26 Rea, D. K. The paleoclimatic record provided by eolian deposition in the deep sea: The geologic history of wind. *Reviews of Geophysics* **32**, 159-195 (1994).
- 27 Friese, C. A. *et al.* Environmental factors controlling the seasonal variability in particle size distribution of modern Saharan dust deposited off Cape Blanc. *Aeolian Research* **22**, 165-179 (2016).

- 28 van der Does, M., Pourmand, A., Sharifi, A. & Stuut, J.-B. W. North African mineral dust across the tropical Atlantic Ocean: Insights from dust particle size, radiogenic Sr-Nd-Hf isotopes and rare earth elements (REE). *Aeolian Research* **33**, 106-116 (2018).
- 29 van der Does, M., Korte, L. F., Munday, C. I., Brummer, G.-J. A. & Stuut, J.-B. W. Particle size traces modern Saharan dust transport and deposition across the equatorial North Atlantic. *Atmospheric Chemistry and Physics* **16**, 13697 (2016).
- 30 Holz, C., Stuut, J.-B. W. & Henrich, R. Terrigenous sedimentation processes along the continental margin off NW Africa: implications from grain-size analysis of seabed sediments. *Sedimentology* **51**, 1145-1154 (2004).
- 31 Drake, N. A., Blench, R. M., Armitage, S. J., Bristow, C. S. & White, K. H. Ancient watercourses and biogeography of the Sahara explain the peopling of the desert. *Proceedings of the National Academy of Sciences* **108**, 458-462 (2011).
- 32 Krastel, S. *et al.* CapTimiris Canyon: A newly discovered channel system offshore of Mauritania. *Eos, Transactions American Geophysical Union* **85**, 417-423 (2004).
- 33 Zühlsdorff, C., Wien, K., Stuut, J. B. W. & Henrich, R. Late Quaternary sedimentation within a submarine channel-levee system offshore Cap Timiris, Mauritania. *Marine Geology* **240**, 217-234 (2007).
- 34 Holz, C., Stuut, J.-B. W., Henrich, R. d. & Meggers, H. Variability in terrigenous sedimentation processes off northwest Africa and its relation to climate changes: Inferences from grain-size distributions of a Holocene marine sediment record. *Sedimentary Geology* **202**, 499-508 (2007).
- 35 Grousset, F. E. *et al.* Saharan wind regimes traced by the Sr–Nd isotopic composition of subtropical composition of subtropical Atlantic sediments: Last Glacial Maximum vs today. *Quaternary Science Reviews* **17**, 395-409 (1998).
- 36 McCave, I. N. Particulate size spectra, behavior, and origin of nepheloid layers over the Nova Scotian Continental Rise. *Journal of Geophysical Research: Oceans* **88**, 7647-7666 (1983).
- 37 Gardner, W. D., Richardson, M. J. & Mishonov, A. V. Global assessment of benthic nepheloid layers and linkage with upper ocean dynamics. *Earth and Planetary Science Letters* **482**, 126-134 (2018).
- 38 Biscaye, P. E. & Eittreim, S. L. Suspended particulate loads and transports in the nepheloid layer of the abyssal Atlantic Ocean. *Marine Geology* **23**, 155-172 (1977).
- 39 Mulitza, S. *et al.* Increase in African dust flux at the onset of commercial agriculture in the Sahel region. *Nature* **466**, 226-228 (2010).
- 40 Castillo, S. *et al.* Trace element variation in size-fractionated African desert dusts. *Journal of Arid Environments* **72**, 1034-1045 (2008).
- 41 Skonieczny, C. *et al.* A three-year time series of mineral dust deposits on the West African margin: Sedimentological and geochemical signatures and implications for interpretation of marine paleo-dust records. *Earth and Planetary Science Letters* **364**, 145-156 (2013).
- 42 Meyer, I., Davies, G. R. & Stuut, J.-B. Grain size control on Sr-Nd isotope provenance studies and impact on paleoclimate reconstructions: An example from deep-sea sediments offshore NW Africa. *Geochemistry, Geophysics, Geosystems* **12**, Q03005 (2011).
- 43 Dasch, E. J. Strontium isotopes in weathering profiles, deep-sea sediments, and sedimentary rocks. *Geochimica et Cosmochimica Acta* **33**, 1521-1552 (1969).
- 44 Aarons, S. M., Aciego, S. M. & Gleason, J. D. Variable HfSrNd radiogenic isotopic compositions in a Saharan dust storm over the Atlantic: Implications for dust flux to oceans, ice sheets and the terrestrial biosphere. *Chemical Geology* **349-350**, 18-26 (2013).
- 45 Zhao, W., Balsam, W., Williams, E., Long, X. & Ji, J. Sr–Nd–Hf isotopic fingerprinting of transatlantic dust derived from North Africa. *Earth and Planetary Science Letters* **486**, 23-31 (2018).
- 46 Jewell, A. M. *et al.* Three North African dust source areas and their geochemical fingerprint. *Earth and Planetary Science Letters* **554**, 116645 (2021).
- 47 Sarnthein, M. *et al.* in *Geology of the Northwest African Continental Margin* (eds Ulrich von Rad, Karl Hinz, Michael Sarnthein, & Eugen Seibold) 545-604 (Springer Berlin Heidelberg, 1982).
- 48 Lever, A. & McCave, I. N. Eolian components in Cretaceous and Tertiary North Atlantic sediments. *Journal of Sedimentary Research* **53**, 811-832 (1983).
- 49 Colin, C. *et al.* Late Miocene to early Pliocene climate variability off NW Africa (ODP 659). *Palaeogeography, Palaeoclimatology, Palaeoecology* **401**, 81-95 (2014).
- 50 Vallé, F., Dupont, L. M., Leroy, S. A. G., Schefuß, E. & Wefer, G. Pliocene environmental change in West Africa and the onset of strong NE trade winds (ODP Sites 659 and 658). *Palaeogeography*, *Palaeoclimatology*, *Palaeoecology* **414**, 403-414 (2014).
- 51 Ruddiman, W. F. et al. in Proceedings of the Ocean Drilling Program: Scientific Results. Vol. 108 (ODP, 1989).
- 52 Zeeden, C. *et al.* Revised Miocene splice, astronomical tuning and calcareous plankton biochronology of ODP Site 926 between 5 and 14.4 Ma. *Palaeogeography, Palaeoclimatology, Palaeoecology* **369**, 430-451 (2013).

- 53 Krijgsman, W., Hilgen, F. J., Raffi, I., Sierro, F. J. & Wilson, D. S. Chronology, causes and progression of the Messinian salinity crisis. *Nature* **400**, 652-655 (1999).
- 54 Mourik, A. A. *et al.* Astronomical tuning of the La Vedova High Cliff section (Ancona, Italy) Implications of the Middle Miocene Climate Transition for Mediterranean sapropel formation. *Earth and Planetary Science Letters* **297**, 249-261 (2010).
- 55 Krijgsman, W., Fortuin, A. R., Hilgen, F. J. & Sierro, F. J. Astrochronology for the Messinian Sorbas basin (SE Spain) and orbital (precessional) forcing for evaporite cyclicity. *Sedimentary Geology* **140**, 43-60 (2001).
- 56 van der Laan, E. *et al.* Astronomical forcing of Northwest African climate and glacial history during the late Messinian (6.5–5.5 Ma). *Palaeogeography, Palaeoclimatology, Palaeoecology* **313–314**, 107-126 (2012).
- 57 Grinsted, A., Moore, J. C. & Jevrejeva, S. Application of the cross wavelet transform and wavelet coherence to geophysical time series. *Nonlinear processes in geophysics* **11**, 561-566 (2004).
- 58 Manivit, H. in *Proceedings of the Ocean Drilling Program, Scientific Results, Vol.108* (ed W. Ruddiman, Sarnthein, M., et al.,) 35-69 (ODP, 1989).
- 59 Schulz, M. & Mudelsee, M. REDFIT: estimating red-noise spectra directly from unevenly spaced paleoclimatic time series. *Computers & Geosciences* **28**, 421-426 (2002).
- 60 Hammer, Ø., Harper, D. A. T. & Ryan, P. D. PAST: Paleontological Statistics Software Package for Education and Data Analysis. *Palaeontologia Electronica* **4**, 9 (2001).
- 61 Lisiecki, L. E. & Raymo, M. E. A Pliocene-Pleistocene stack of 57 globally distributed benthic δ180 records. *Paleoceanography* **20**, PA1003 (2005).
- 62 Tierney, J. E., deMenocal, P. B. & Zander, P. D. A climatic context for the out-of-Africa migration. *Geology* **45**, 1023-1026 (2017).
- 63 deMenocal, P. B., Ruddiman, W. F. & Pokras, E. M. Influences of High- and Low-Latitude Processes on African Terrestrial Climate: Pleistocene Eolian Records from Equatorial Atlantic Ocean Drilling Program Site 663. *Paleoceanography* **8**, 209-242 (1993).
- 64 Pokras, E. M. & Mix, A. C. Earth's precession cycle and Quaternary climatic change in tropical Africa. *Nature* **326**, 486-487 (1987).
- 65 Larrasoaña, J. C., Roberts, A. P., Rohling, E. J., Winklhofer, M. & Wehausen, R. Three million years of monsoon variability over the northern Sahara. *Climate Dynamics* **21**, 689-698 (2003).
- de Boer, B., Peters, M. & Lourens, L. J. The transient impact of the African monsoon on Plio-Pleistocene Mediterranean sediments. *Clim. Past* **17**, 331-344 (2021).
- 67 Kutzbach, J. E. *et al.* African climate response to orbital and glacial forcing in 140,000-y simulation with implications for early modern human environments. *Proceedings of the National Academy of Sciences*, **117** (5), 2255-2264 (2020).
- 68 Bosmans, J. H. C., Hilgen, F. J., Tuenter, E. & Lourens, L. J. Obliquity forcing of low-latitude climate. *Clim. Past* **11**, 1335-1346 (2015).
- 69 Schefuß, E., Ratmeyer, V., Stuut, J.-B. W., Jansen, J. H. F. & Sinninghe Damsté, J. S. Carbon isotope analyses of n-alkanes in dust from the lower atmosphere over the central eastern Atlantic. *Geochimica et Cosmochimica Acta* 67, 1757-1767 (2003).
- 70 Eglinton, T. I. & Eglinton, G. Molecular proxies for paleoclimatology. *Earth and Planetary Science Letters* **275**, 1-16 (2008).
- 71 Rommerskirchen, F. *et al.* A north to south transect of Holocene southeast Atlantic continental margin sediments: Relationship between aerosol transport and compound-specific δ 13C land plant biomarker and pollen records. *Geochemistry, Geophysics, Geosystems* **4** (2003).
- 72 Zhao, M., Dupont, L., Eglinton, G. & Teece, M. n-Alkane and pollen reconstruction of terrestrial climate and vegetation for N.W. Africa over the last 160 kyr. *Organic Geochemistry* **34**, 131-143 (2003).
- 73 Hooghiemstra, H., Bechler, A. & Beug, H.-J. Isopollen maps for 18,000 years B.P. of the Atlantic offshore of northwest Africa: Evidence for paleowind circulation. *Paleoceanography* **2**, 561-582 (1987).
- 74 Ehleringer, J. R. & Cerling, T. E. in *Global Biogeochemical Cycles in the Climate System* (eds Ernst-Detlef Schulze *et al.*) 267-277 (Academic Press, 2001).
- 75 Huang, Y., Dupont, L., Sarnthein, M., Hayes, J. M. & Eglinton, G. Mapping of C4 plant input from North West Africa into North East Atlantic sediments. *Geochimica et Cosmochimica Acta* 64, 3505-3513 (2000).
- 76 Niedermeyer, E. M. *et al.* Orbital- and millennial-scale changes in the hydrologic cycle and vegetation in the western African Sahel: insights from individual plant wax δD and $\delta 13C$. *Quaternary Science Reviews* 29, 2996-3005 (2010).
- 77 Hoetzel, S., Dupont, L., Schefusz, E., Rommerskirchen, F. & Wefer, G. The role of fire in Miocene to Pliocene C4 grassland and ecosystem evolution. *Nature Geosci* **6**, 1027-1030 (2013).
- Feakins, S. J. *et al.* Northeast African vegetation change over 12 m.y. *Geology* **41** (3), 295–298 (2013).
- 79 Morley, R. J. & Richards, K. Gramineae cuticle: a key indicator of Late Cenozoic climatic change in the Niger Delta. *Review of Palaeobotany and Palynology* 77, 119-127 (1993).

- 80 Polissar, P. J., Rose, C., Uno, K. T., Phelps, S. R. & deMenocal, P. Synchronous rise of African C4 ecosystems 10 million years ago in the absence of aridification. *Nature Geoscience* **12**, 657-660 (2019).
- 81 Tierney, J. E., Russell, J. M. & Huang, Y. A molecular perspective on Late Quaternary climate and vegetation change in the Lake Tanganyika basin, East Africa. *Quaternary Science Reviews* **29**, 787-800 (2010).
- 82 Beuning, K. R. M., Talbot, M. R., Livingstone, D. A. & Schmukler, G. Sensitivity of carbon isotopic proxies to paleoclimatic forcing: A case study from Lake Bosumtwi, Ghana, over the last 32,000 years. *Global Biogeochemical Cycles* **17**, 1121 (2003).
- 83 Sinninghe Damsté, J. S. *et al.* A 25,000-year record of climate-induced changes in lowland vegetation of eastern equatorial Africa revealed by the stable carbon-isotopic composition of fossil plant leaf waxes. *Earth and Planetary Science Letters* **302**, 236-246 (2011).
- 84 Lupien, R. L. *et al.* Vegetation change in the Baringo Basin, East Africa across the onset of Northern Hemisphere glaciation 3.3–2.6 Ma. *Palaeogeography, Palaeoclimatology, Palaeoecology* **570**, 109426 (2021).
- 85 Lupien, R. L. *et al.* A leaf wax biomarker record of early Pleistocene hydroclimate from West Turkana, Kenya. *Quaternary Science Reviews* **186**, 225-235 (2018).
- 86 Tierney, J. E. & deMenocal, P. B. Abrupt Shifts in Horn of Africa Hydroclimate Since the Last Glacial Maximum. *Science* **342**, 843-846 (2013).
- 87 Feakins, S. J., Eglinton, T. I. & deMenocal, P. B. A comparison of biomarker records of northeast African vegetation from lacustrine and marine sediments (ca. 3.40 Ma). *Organic Geochemistry* **38**, 1607-1624 (2007).
- 88 Castañeda, I. S. *et al.* Wet phases in the Sahara/Sahel region and human migration patterns in North Africa. *Proceedings of the National Academy of Sciences* **106**, 20159-20163 (2009).
- 89 Badger, M. P. S. *et al.* CO2 drawdown following the middle Miocene expansion of the Antarctic Ice Sheet. *Paleoceanography* **28**, 42-53 (2013).
- 90 Badger, M. P. S. *et al.* Insensitivity of alkenone carbon isotopes to atmospheric CO2 at low to moderate CO2 levels. *Clim. Past* **15**, 539-554 (2019).
- 91 Bartoli, G., Hönisch, B. & Zeebe, R. E. Atmospheric CO2 decline during the Pliocene intensification of Northern Hemisphere glaciations. *Paleoceanography* **26**, PA4213 (2011).
- 92 Bolton, C. T. *et al.* Decrease in coccolithophore calcification and CO2 since the middle Miocene. *Nat Commun* 7, 10284 (2016).
- 93 Chalk, T. B. *et al.* Causes of ice age intensification across the Mid-Pleistocene Transition. *Proceedings of the National Academy of Sciences* **114**, 13114 (2017).
- de la Vega, E., Chalk, T. B., Wilson, P. A., Bysani, R. P. & Foster, G. L. Atmospheric CO2 during the Mid-Piacenzian Warm Period and the M2 glaciation. *Scientific Reports* **10**, 11002 (2020).
- 95 Dyez, K. A., Hönisch, B. & Schmidt, G. A. Early Pleistocene Obliquity-Scale pCO2 Variability at ~1.5 Million Years Ago. *Paleoceanography and Paleoclimatology* **33**, 1270-1291 (2018).
- 96 Hönisch, B., Hemming, N. G., Archer, D., Siddall, M. & McManus, J. F. Atmospheric Carbon Dioxide Concentration Across the Mid-Pleistocene Transition. *Science* **324**, 1551-1554 (2009).
- 97 Martinez-Boti, M. A. *et al.* Plio-Pleistocene climate sensitivity evaluated using high-resolution CO2 records. *Nature* **518**, 49-54 (2015).
- 98 Pagani, M., Zachos, J. C., Freeman, K. H., Tipple, B. & Bohaty, S. Marked Decline in Atmospheric Carbon Dioxide Concentrations During the Paleogene. *Science* **309**, 600-603 (2005).
- 99 Pagani, M., Liu, Z., LaRiviere, J. & Ravelo, A. C. High Earth-system climate sensitivity determined from Pliocene carbon dioxide concentrations. *Nature Geoscience* **3**, 27 (2010).
- 100 Seki, O. *et al.* Alkenone and boron-based Pliocene pCO2 records. *Earth and Planetary Science Letters* **292**, 201-211 (2010).
- 101 Sosdian, S. M. *et al.* Constraining the evolution of Neogene ocean carbonate chemistry using the boron isotope pH proxy. *Earth and Planetary Science Letters* **498**, 362-376 (2018).
- 102 Stap, L. B. *et al.* CO2 over the past 5 million years: Continuous simulation and new δ 11B-based proxy data. *Earth and Planetary Science Letters* **439**, 1-10 (2016).
- 103 Super, J. R. *et al.* North Atlantic temperature and pCO2 coupling in the early-middle Miocene. *Geology* **46**, 519-522 (2018).
- 104 Tanner, T., Hernández-Almeida, I., Drury, A. J., Guitián, J. & Stoll, H. Decreasing Atmospheric CO2 During the Late Miocene Cooling. *Paleoceanography and Paleoclimatology* **35**, e2020PA003925 (2020).
- 105 Zhang, Y. G., Pagani, M., Liu, Z., Bohaty, S. M. & DeConto, R. A 40-million-year history of atmospheric CO2. *Philosophical Transactions of the Royal Society of London A: Mathematical, Physical and Engineering Sciences* **371**, 20130096 (2013).
- 106 Zhang, Y. G., Pagani, M., Henderiks, J. & Ren, H. A long history of equatorial deep-water upwelling in the Pacific Ocean. *Earth and Planetary Science Letters* **467**, 1-9 (2017).
- 107 Zhang, Y. G. *et al.* Refining the alkenone-pCO2 method I: Lessons from the Quaternary glacial cycles. *Geochimica et Cosmochimica Acta* **260**, 177-191 (2019).

- 108 Tauxe, L., Valet, J.-P. & Bloemendal, J. in *Proc. ODP, Sci. Results, 108* (ed W. Ruddiman, Sarnthein, M., et al.,) 429–439 (ODP, 1989).
- 109 Raffi, I. *et al.* A review of calcareous nannofossil astrobiochronology encompassing the past 25 million years. *Quaternary Science Reviews* **25**, 3113-3137 (2006).
- 110 Wade, B. S., Pearson, P. N., Berggren, W. A. & Pälike, H. Review and revision of Cenozoic tropical planktonic foraminiferal biostratigraphy and calibration to the geomagnetic polarity and astronomical time scale. *Earth-Science Reviews* **104**, 111-142 (2011).
- 111 Ogg, J. G. in *The Geologic Time Scale* (eds Felix M. Gradstein, James G. Ogg, Mark D. Schmitz, & Gabi M. Ogg) 85-113 (Elsevier, 2012).
- 112 Laskar, J. *et al.* A long-term numerical solution for the insolation quantities of the Earth. *Astronomy & Astrophysics* **428**, 261-285 (2004).