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Homo sapiens and Neanderthals share high cerebral cortex integration into adulthood

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Article

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Figure #	Figure title One sentence only	Filename This should be the name the file is saved as when it is uploaded to our system. Please include the file extension. i.e.: <i>Smith_ED_Fig1.jpg</i>	Figure Legend If you are citing a reference for the first time in these legends, please include all new references in the main text Methods References section, and carry on the numbering from the main References section of the paper. If your paper does not have a Methods section, include all new references at the end of the main Reference list.
Data Table 1	K-PLS values	Extended_Data_1able1.xisx	R-PLS of separate PLS analyses performed on the different postnatal ontogenetic stages of <i>H. sapiens</i> , <i>P. troglodytes</i> , <i>G. gorilla</i> and <i>Pongo</i> species.
Extended Data Fig. 1	Endocast local integration assessment	Extended_Data_Fig1.tif	The set of each semilandmark (a) and its 9 closest semilandmarks define the N-Core (b). The remaining semilandmarks define the C-Core (c). The N-Core and R-Core are subjected to two independent GPAs and the covariation between the two- blocks is calculated by PLS (d). With CR the GPA is computed the entire set (e). The values from PLS and CR analyses are used to create a colour map of integration (f) and modularity (g).
Extended Data Table 2	Effect sizes of separate PLS analyses	Extended_Data_Table2.xlsx	Effect sizes of separate PLS analyses performed the different postnatal ontogenetic stages of <i>H. sapiens</i> , <i>P.</i> <i>troglodytes</i> , <i>G. gorilla</i> and <i>Pongo</i> when accounting for size effect.
Extended Data Fig. 2	Evolutionary rates of CR values within the Cercopithecinae clade.	Extended_Data_Fig2.tiff	
Extended Data Fig. 3	Evolutionary rates of CR values within the Strepsirrhini.	Extended_Data_Fig3.tiff	
Extended Data Fig. 4	Evolutionary rates of CR values within the family Cebidae.	Extended_Data_Fig4.tiff	
Extended Data Table 3	CR values measured after size and phylogenetic correction.	Extended_Data_Table3.xlsx	

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Supplementary Information	Yes	Supplementary_information.pdf	Supplementary Figures 1-4, Supplementary Tables 1-3 Supplementary methods; Specimens institutional codes; Tree in Newick format.
Reporting Summary	Yes	Sansalone_RS.pdf	

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3

Homo sapiens and Neanderthals share high cerebral cortex integration into 4 adulthood 5

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Abstract: There is controversy around the mechanisms that guided the change in brain shape 31 during the evolution of modern humans. It has long been held that different cortical areas evolved 32 independently from each other to develop their unique functional specializations. However, some 33 recent studies suggest that high integration between different cortical areas could facilitate the 34 emergence of equally extreme, highly specialized brain functions. Here, we analyze the evolution of 35 brain shape in primates using 3D geometric morphometrics of endocasts. We aim to determine 36 firstly, whether modern humans present unique developmental patterns of covariation between brain 37 cortical areas and secondly, whether hominins experienced unusually high rates of evolution in 38 brain covariation as compared to other primates. Based on analyses including modern humans and 39 other extant great apes at different developmental stages, we first demonstrate that, unlike our 40 closest living relatives, Homo sapiens retains high levels of covariation between cortical areas into 41 adulthood. Among the other great apes, high levels of covariation are only found in immature 42 individuals. Secondly, at the macroevolutionary level, our analysis of 400 endocasts, representing 43 148 extant primate species and 6 fossil hominins, shows that strong covariation between different 44 areas of the brain in H. sapiens and Homo neanderthalensis evolved under distinctly higher 45 evolutionary rates than in any other primate, suggesting that natural selection favored a greatly 46 integrated brain in both species. These results hold when extinct species are excluded and allometric 47 effects are accounted for. Our findings demonstrate that high covariation in the brain may have 48 played a critical role in the evolution of unique cognitive capacities and complex behaviors in both 49 modern humans and Neanderthals. 50

51 Introduction

52

The modern human brain is remarkable in its size, unusually globular shape, and extreme left-right 53 asymmetry which are all thought to have contributed to the evolution of our exceptional cognitive 54 capacities^{1–5}. Historically, two main models have been invoked to explain the evolution of the brain: 55 i) the 'concerted' model, assuming that developmental integration affects brain evolution globally 56 and ii) the 'mosaic' model, that is the idea that functional units of the brain may co-evolve or evolve 57 independently according to the distribution of selection pressures acting on them^{6–9}. By deploying 58 mosaicism, a brain module could be fine-tuned by selection to optimize specific tasks regardless of 59 what happens in other areas of the brain¹⁰⁻¹⁴. Volumetric and morphometric analyses have 60 demonstrated that selective expansion of discrete brain areas closely reflects the establishment of 61 functional connections between them, enabling specific cognitive tasks^{14–16}. It has also been 62 proposed that mosaicism may have promoted behavioural flexibility and increased the ability to 63 respond to changes in selective regimes¹³. However, the hypothesis of the brain modular evolution 64 has been challenged by the recent observation that traits' covariation can favour the rapid evolution 65 of extreme, highly specialised morphotypes, provided that selection vectors align with major axes 66 of phenotypic variation^{17,18}. Within this 'concerted' framework, it has been argued that the multiple, 67 high-level functional specialisations of the modern human brain could originate from selection for 68 69 fine coordination between different brain units to shared functional ends, without effecting any major changes in the relative proportions of specific brain areas^{10,19,20}. Despite their apparent 70 polarisation the concerted and mosaic brain hypotheses are not mutually exclusive¹⁶. Mosaicism 71 72 does not rule out co-variation between brain units, as long as this reflects a response to shared functional demands, and a concerted brain can be the result of an adaptive process rather than the 73 product of developmental constraints²¹. 74

75 A key question regarding the uniqueness of the modern human brain is whether its evolution 76 branched away from the developmental programme characterising our living relatives. Studying the developmental patterns of morphological concertedness (or integration) as opposed to mosaicism 77 (or modularity) between human brain areas and comparing this pattern to those of other great apes 78 would help us determine to what extent the organization of cortical areas in Homo sapiens may 79 actually be remarkable^{1,22-25}. Another important issue is to understand whether, at the macro-80 evolutionary scale, humans display higher evolutionary rates toward either brain modularity or 81 integration. This would offer direct evidence of selection favouring the emergence of major changes 82 83 in the patterns of co-variation between cortical areas.

84

To address these questions, we have applied three dimensional geometric morphometrics to 85 measure and visualise the relative magnitudes of morphological co-variation in primate virtual brain 86 endocasts. Traditionally, investigations into patterns of covariation between different regions of the 87 brain have relied on comparative volumetric analyses (i.e., of relative sizes) of brain subunits. 88 89 However, volumetric comparisons are silent on the shape component (position and orientation) of brain form, which potentially captures aspects of brain evolution not predicted by size alone¹³. 90 91 Furthermore, in contrast to volumetric data, shape data are comparatively rare for extinct species. Hence, studying patterns of covariation directly on cranial endocasts represents the single most 92 93 informative means of gaining direct evidence on the evolutionary patterns of brain evolution across 94 hominins (Homo sapiens and its extinct close relatives). To gain this insight, we have combined a phylogenetic comparative method based on phylogenetic ridge regression to determine the presence 95 of shifts in the evolutionary rates across primate history with a novel strategy to measure and map 96 97 phenotypic covariation on brain cortical areas. As brains do not fossilise, evidence of fossil species' brain evolution can be derived from the virtual fillings of the bony braincase - or endocasts - which 98 99 can adequately approximate the outer brain morphology.

100 Our datasets comprise 127 postnatal virtual endocasts, sampled from the eruption of 101 deciduous dentition through adulthood, for *H. sapiens*, *Pan troglodytes*, *Gorilla gorilla* and two 102 species of *Pongo* for the analysis of developmental patterns and, for the macroevolutionary study

103 400 endocasts representing 154 extant and extinct species, including Australopithecus africanus,

104 Paranthropus boisei, Homo ergaster, Homo erectus, Homo heidelbergensis and H.

neanderthalensis. We explicitly tested whether i) specific patterns of modularity or integration

106 between cortical areas can be identified through human brain development and how these relate to

107 those of extant great apes; and ii) whether the hominin brain displays higher rates of evolution

108 toward either increased integration or modularity.

109

110 **Results**

111 *Question 1. Does the human brain cortical covariation differ to that of other great apes?*

112 We performed separate partial least squares (PLS) analyses on 4 successive postnatal

developmental stages of Gorilla gorilla, Pan troglodytes and Homo sapiens. We further included in

114 the analysis *Pongo abelii* and *Pongo pygmaeus* (Fig. 1). Yet, given the paucity of available

orangutan specimens we had to group them together and therefore did not explore covariation

between individual brain modules in *Pongo*. The developmental stages were defined following

117 refs.^{26,27}: Stage 2 = all deciduous dentition fully erupted; Stage 3 = deciduous dentition and at least

fully erupted M1; Stage 4 = M2 fully erupted; Adult = full permanent dentition. The PLS method allows the exploration of covariation patterns between different sets of shape variables (here brain

subunits), whereas r-PLS (measured using the r^2 derived from PLS analysis based on 999

permutations, Methods and Extended Data Tables 1-2 and Extended Data Fig. 1) is the correlation

coefficient and can be used as a measure of the magnitude of covariation.

123 Our results show that integration of the brain in *H. sapiens* and *P. troglodytes* is similar

through the pre-adult stages (Stages 2 to 4; Fig. 1a and Extended Data Table 1). Yet, in
chimpanzees (and in gorillas from stage 4 onwards), r-PLS significantly drops in adulthood,

chimpanzees (and in gorillas from stage 4 onwards), r-PLS significantly drops in adulthood,
 whereas in *H. sapiens* the brain remains significantly integrated into adulthood (Fig. 1 and Extended)

Data Table 1). The patterns of covariation between brain cortical modules are almost identical in

adult *Pan* and *Gorilla* individuals, pointing to strong covariation between the occipital and

temporal, and frontal and parietal modules, respectively (Fig. 1b). Comparable results are obtained

130 when controlling for brain size (Extended Data Table 2), and whether *Pongo* species (grouped as

131 one) are included. These results suggest that the shape covariation patterns observed during

development are largely independent from allometric effects and that humans significantly depart

133 from the brain developmental patterns shared by the other greater apes.

134 Using two-block PLS to measure the degree of association between different cortical areas we

135 confirmed the proposition that the brain of *H. sapiens* retains high levels of morphological

136 integration throughout growth, unlike other great apes (Fig. 1b,c). We applied for the first time a

137 novel approach to map the magnitude of morphological integration (Methods and Extended Data

Fig. 1) directly onto the endocast surface without defining any a priori module. This approach

involves parcelling out the brain endocasts into small independent "modulets" centred around a

single semilandmark and calculating the level of morphological integration of the modulets with the rest of the endocast. The average values of integration calculated at each semilandmark is

subsequently used to create maps of integration intensity.

143 Charting the magnitude of integration over the endocasts at different developmental stages 144 reveals clear differences between *H. sapiens* and *P. troglodytes* (Fig. 1a,c). At stage 2 the human 145 brain displays high integration over the parietal and occipital regions. At stages 3 and 4 strong 146 integration centres on the frontal and occipital lobes. In the adult stage (4) humans show the greatest 147 level of integration over the parietal, temporal and prefrontal regions. Chimpanzees follow a 148 different developmental pattern, showing poorly integrated frontal and parietal areas throughout 149 postnatal growth and relatively stronger integration at the level of temporal and prefrontal areas.

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- 151 152

- 153 *Question 2. Did hominins evolve towards high cortical integration?*
- 154 We measured the covariation between four brain subunits (corresponding to frontal, parietal,
- temporal and occipital regions, see Methods and Extended Data Table 3) at the macroevolutionary
- local level, by means of Covariance Ratio (CR; a measure of the overall covariation between modules
- 157 divided by the overall covariation within modules, see Methods).
- 158 Our results show that hominoid (apes) brains are morphologically distinct in shape (Fig. 2a) and
- display higher levels of covariation between brain cortical areas (CR = 1.01 indicating high
- 160 covariation, see Methods) than any other primate group (Fig. 2b, Methods and Extended Data Table
- 161 2). Platyrrhini and Strepsirrhini display the lowest magnitude of covariation (CR = 0.76 and 0.72
- respectively) between brain modules, whereas Cercopithecinae and Colobinae fall in between
- hominoids and all other primates (CR = 0.83 and 0.91 respectively). Accounting for allometry did
- not alter the described pattern, suggesting that size has a limited impact on the brain covariation
- 165 patterns observed at the macroevolutionary level (Extended Data Table 3).
- 166 In keeping with our ontogenetic analyses, we devised a novel approach to map the metrics for the
- 167 magnitude of covariation, the CR, over the digital endocasts. These brain maps show that hominins
- are characterized by the highest evolutionary rates in CR (Fig. 2b). Great apes display higher values
- 169 of covariation in the occipital and parieto-frontal regions and lower levels over the temporal areas.
- 170 In contrast, lesser apes show lower covariation in the pre-frontal areas closer to the olfactory bulbs
- and over the temporal region (Fig. 2b).
- 172 Among Cercopithecinae, high evolutionary rates are recorded in Papionini (Fig. 2b and Extended
- 173 Data Fig. 2). Conversely, Strepsirrhini (two-tailed p = 0.001) are characterized by a rate slowdown,
- 174 as were capuchin and squirrel monkeys (family Cebidae, two-tailed p = 0.002) among New World
- monkeys (Fig. 2b and Extended Data Figs. 3-4). Mapping CR values over the endocast surfaces
- 176 reveals different patterns in different primate clades. Cercopithecinae show higher integration in the
- 177 occipital and frontal regions than elsewhere on the brain. Colobinae, Platyrrhini and Strepsirrhini
- display similar distribution of the CR values over the endocast, with the areas corresponding to the
- frontal and pre-frontal cortical areas and the temporal regions showing moderate covariation (Fig.2b).
- 181

182 Within Hominoidea, *H. neanderthalensis* and *H. sapiens* show the highest rate of evolution of brain 183 covariation (two tailed p = 1.00, Fig. 3). Interestingly, *A. africanus* was characterized by

- 184 evolutionary rates comparable to those of *P. troglodytes* suggesting a graded trend for increased rate
- of CR evolution among hominins (Fig. 3a), leading to the highly integrated brain of *Homo*,
- 186 especially evident in the parietal area (Fig. 3b).
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- 190 191

192 **Discussion**

- *Homo sapiens* and the other great apes share high covariation between different cortical areas of the brain throughout most postnatal development. However, only *Homo sapiens* retains such strong
- 195 morphological integration into adulthood. This finding is consistent with other reports indicating
- that the cortical areas of the human brain are tightly integrated throughout the adult life^{12,28}.
- 197 Connectome analysis suggests an evolutionary shift in the human brain to enhance global network
- integration over that of the chimpanzee²⁹, indicating that humans evolved strong covariation even
- among spatially distant brain regions³⁰ (which is consistent with our Fig. 1c). This evolutionary
- 200 pattern seems to have deep evolutionary roots. Hominins show a trend for an increased magnitude
- 201 of covariation between different brain regions, escalating through Middle to Late Pleistocene
- 202 human species (*H. sapiens* and *H. neanderthalensis*). This finding contradicts the common
- 203 perception that functional specialisation in the modern human brain arises from a modular

architecture (e.g., semi-independent evolution of different cortical areas)¹³, but is in agreement with
studies of encephalised non-mammalian vertebrates suggesting that high integration may drive
functional specialisation in the brain, even among distantly related taxa and under very different
selective scenarios³¹. Our findings similarly suggest that coordinated changes in brain shape may
have played a major role in maintaining the functional association between brain subunits,
ultimately leading to the derived cognitive specialisation observed in *Homo*.

Charting morphological integration over the endocasts shows that the great apes are clearly 210 distinct from the lesser apes, suggesting that a shift in the spatial patterns of covariation (and not 211 just in the magnitude of integration or relative brain size) occurred at the time of divergence 212 between the two groups. Hominins show a high degree of covariation in the parietal and frontal 213 214 regions, which are thought to have played a fundamental role in the evolution of cognitive capacities unique to humans^{32,33}. Modifications in the parietal regions are thought to represent a 215 derived condition apparent only within the most recent *Homo sapiens* populations^{23,34}. The parietal 216 cortex is involved in different association tasks such as dexterity, self-awareness and visual 217 imaging³⁵. These functions confer the capacity to translate cognition into novel behavioural 218 attributes, allowing the incorporation of tools and technology into behavioural patterns^{33,36}. 219

Australopithecus africanus, H. ergaster and H. erectus display evolutionary rates like, or 220 slightly higher than those showed by P. troglodytes and P. paniscus (Fig. 3a). In general, larger-221 222 bodied species, mostly occurring among hominoids and papionins, are marked by higher rates of covariation among brain areas³⁷. Yet, even after correcting for brain size, the *Homo* clade still 223 shows the highest levels and rates of brain cortex covariation (Extended Data Table 3). This 224 suggests that the major shift in the pattern of brain shape covariation emerged independently from 225 size and, likely, occurred within these species only. This increased level of interconnection between 226 different cortical areas of the brain may have facilitated the emergence of derived cognitive 227 228 capacities in Neanderthals as suggested by the palaeoanthropological record^{38–41}. However, modern 229 humans and Neanderthals have distinctly different brain morphologies, suggesting that high levels of covariation might have been inherited from their last common ancestor and that brain shape 230 evolution then followed divergent trajectories in *H. neanderthalensis* and *H. sapiens*⁴². This 231 evidence brings into question the role of globularity in the emergence of high cognitive abilities in 232 233 Homo sapiens. Neanderthals, and the other great apes, did not go through a "globularisation phase" during the earliest postnatal growth stages, retaining the plesiomorphic, antero-posteriorly elongated 234 adult brain common to archaic *Homo* species^{43,44}. The development of a globular brain is exclusive 235 to modern humans⁴⁵ and its role in maintaining high levels of integration into adulthood deserves 236 further investigation. 237

Our findings do not favour either the mosaic or the concerted model of brain evolution, 238 suggesting that the debate between these two hypotheses of brain evolution should be reframed 239 within in a more inclusive proposition. We evidenced that a shared or conserved pattern of 240 covariation could have an adaptive value or be instrumental to the emergence of derived modern 241 242 humans functional capacities, rather than being considered a mere developmental or phylogenetic constraint²¹. In contrast, this study suggests that departure from an established pattern does not 243 244 necessarily involve the presence of a modular behaviour and that high covariation may favour the emergence of functional specialisation, as predicted by the mosaic model. 245

In conclusion, we propose that the persistence of high levels of morphological covariation into adulthood in modern humans and Neanderthals is linked to the evolution of derived cognitive abilities. In addition, modern humans show high levels of integration between cortical areas throughout development. Unfortunately, the scarcity of immature Neanderthals with well-preserved skulls prohibits us from conclusively determining whether *H. neanderthalensis* brain followed the same developmental path as ours^{43,44}. Yet, the strong covariation in adult brains shared by Neanderthals and *H. sapiens* only, suggests this is arguably the case.

Neural plasticity and innovative-explorative behaviours are typically associated with juvenile life stages, as well as the extension of childhood learning^{45,46} and are at central to Mithen's

- theory of cognitive fluidity^{47,48}, which postulates that only modern humans are capable of fully 255
- integrating diverse dominions of knowledge. Our evidence supports the argument that juvenilisation 256
- of the human (and possibly to some extent Neanderthal's as well) brain was driven by prolonged 257
- brain growth, mediated by the retention of unusually high degree of covariation between the 258
- different brain units into adulthood. 259
- 260
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264 Methods

Endocast segmentation 265

Virtual endocasts of primate crania were generated from CT image stacks using a combination of 266 Mimics 21.0 (Materialise, Ann Arbor, MI, USA) and Geomagic Studio 2014 (Research Triangle 267 Park, NC). For each specimen, cranial bone was first segmented in Mimics with the gray-value 268

- 269 range set conservatively to avoid extensive manual corrections later in the process. The endocranial
- cavity was then closed off at the foramen magnum using a flat plane spanning basion to opisthion. 270
- Next, a 3D object was generated, and all gaps below 1 mm in diameter were closed using the 271
- "Wrap" function before closing off all remaining openings (e.g., foramen ovale, optic canal) near 272
- the endocranial surface. This created a sealed cavity that was filled using the "Cavity Fill" tool. 273
- Endocasts were then imported as stereolithography (STL)-formatted surface files into Geomagic 274
- where excess material protruding through cranial foramina was removed and the polygon meshes 275
- were lightly smoothed using the "QuickSmooth" function. Endocast volumes were then measured in 276 cubic centimeters (cm3) using the "Compute Volume" function. 277
- 278
- 279 Automatic landmarking procedure
- 280 The points on the template (*Piliocolobus badius*) were projected on all the other specimens using
- the function *placePatch()* from the R package 'Morpho^{'49}. In order to remove any incorrect 281
- projection the semi-landmarks on the curves were set bold-distanced using the function 282
- *pointsOnBezier()* from the 'bezier' R package ⁵⁰ then the curves present on the sides of the endocast 283 284
- geometry were mirrored using the function *symmetrize()* from the R package 'Morpho'. After this process was complete the semi-landmarks were slid along the curves by minimising the bending 285
- energy of a thin plate spine deformation (semi-landmarks relaxation) using the *slider3d()* function 286
- from the R package 'Morpho'. This approach follows the algorithm described by Gunz et al.⁵¹ and 287
- has been shown to be the most appropriate method to slide semi-landmarks on curves and surfaces 288 according to Bookstein 52.
- 289
- 290

Shape analysis 291

On each endocast (Supplementary Fig. 1), we manually digitised 21 anatomical and homologous 292 293 landmarks, then performed a principal component analysis (PCA) to identify the individual closest to the consensus shape (Piliocolobus badius USNM 481795). We manually digitised 76 semi-294 295 landmarks placed equidistantly along curves and surfaces on the consensus specimen endocast and used it as the template individual (Supplementary Table 1). All landmarks were placed by using 296 297 IDAV Landmark software. Once all the semi-landmarks were automatically placed, we imported 298 the landmark coordinates into R version 4.0.1 for further analyses. We performed generalised Procrustes analysis (GPA) on all landmarks, implemented in the function procSym from the R 299 package 'Morpho' to rotate, translate and scale landmark configurations to unit centroid size (CS), 300 that is the square root of squared differences between landmark coordinates and centroid 301 coordinates ⁵³. To visualise the multivariate ordination of the aligned Procrustes coordinates, we 302

- used a phylomorphospace using the first two regular non-phylogenetic PCA scores. We classified 303
- 304 the species using similar taxonomic groups to those defined in Sansalone et al.⁴ and Neaux et al.⁵⁴:
- Hominoidea, Cercopithecinae, Colobinae, Platyrrhini, Strepsirrhini. Shape data have been 305

306 controlled for size (Extended Data Tables 2-3), sexual dimorphism effects and for measurement

- 307 error.
- 308
- 309 Phylogeny

310 The phylogenetic tree used in our analyses is a time-calibrated tree based on a Bayesian estimate

obtained from the 10kTrees Project $v3^{55}$ for the 146 extant species in our dataset. A maximum clade

- 312 credibility tree of the extant species in the analysis was constructed from a set of 1000 molecular
- trees using the function MaxCredTree() from the R package 'phangorn'⁵⁶. Finally, the eight fossil
- species included in our dataset were manually added to the tree (available in Newick format in
- Supplementary Information) following the topological arrangement in refs.^{2,57,58} using the RRphylo
- function *tree.merger*⁵⁹. The full list of the accessed specimens is indicated in Supplementary Table 2.
- 317 318
- 319 Measurement error

320 The measurement error associated with the digitisation of landmarks was measured on three

321 replicates of 60 specimens representative of the total dataset variation. For each specimen we

- 322 digitized only the homologous landmarks, subsequently we automatically applied the semi-
- 323 landmarks following the procedure previously described. We calculated the mean Procrustes
- 324 distances for each triplet of the same specimen occurring in the three replicas. We then computed
- the averages of all the mean values of the minimum and maximum values of each triplet. The
- amount of digitisation error, with respect to the total variation in the shape, can be expressed as a

327 percentage. We calculated the ratio of the mean value for total digitisation and the mean of the total

- dataset. We found the digitization error in the endocast dataset was as low as 0.36% of the total
- variation, respectively. Because the measurement error was smaller than 5% in both datasets it
 could be safely assumed its effect on the results was negligible.
- 331
- 332 Sexual dimorphism
- In order to account for the potential effect of sexual dimorphism on the shape data, we performed a
- 334 Procrustes ANOVA to test for the presence of significant shape and size differences between males
- and females. The analysis returned a non-significant result ($r^2 = 0.01$, p = 0.28), suggesting that, at
- 336 macroevolutionary scale, sexual dimorphism is not impacting the brain shape variation in Primates.

Similar results were obtained when we tested for size differences between males and females ($r^2 = 0.01, p = 0.24$).

- 339
- 340 Size and phylogenetic correction

341 The relationship between size (measured as CS; independent variable) and shape (measured as

342 aligned Procrustes coordinates; dependent variable) was tested by means of multivariate

- regressions. We repeated all the following analyses by using residuals of the multivariate regressionof shape vs size.
- 345 Specifically, to account for size effects on the ontogenetic series, we used shape residuals computed
- 346 from separate, per developmental stage, multivariate regression. The shape residuals were used to
- 347 perform size-free PLS analyses, and the results are summarised in Extended Data Table 2. Overall,
- we did not observe any difference from the pattern described by the standard version of the PLS.
- However, it must be noted the r-PLS were lower for each group. This is in agreement with previous
- 350 findings reporting allometry and development as integrating factors, therefore the removal of the
- 351 size component may reduce the observed levels of covariation⁶⁰.
- 352 The same holds for the macroevolutionary analyses, which we repeated using residuals of
- 353 multivariate regressions of shape vs size performed within a phylogenetic context using PGLS
- 354 (Phylogenetic Generalised Least Squares) regression. Specifically, shape residuals have been
- 355 computed using the function *PGLS_fossil()* from the R package RRphylo. It must be noted that the
- 356 PGLS analysis using shape as the respondent and size as the predictor variables and accounting for

- 357 phylogenetic variance covariance matrix, returned marginally significant results (p-value = 0.042; r²
- = 0.101) suggesting that size is explaining a relatively small fraction of the total shape variation,
- this result is in line with previous investigations evidencing a limited effect of size on primates' $\frac{4}{22}$
- 360 brain shape 4,22 .
- 361 We computed the Covariance Ratio (CR, see below for more details) values using shape residuals
- 362 (results are summarized in Extended Data Table 3) for the different primate clades while accounting
- 363 for phylogeny using the function *phylo.modularity()* from the R package geomorph. Furthermore,
- 364 we used shape residuals to compute per-species CR values to then compute size-free evolutionary
- rates of covariation. Again, we did not notice any alteration in the pattern produced by the standard
- RRphylo analyses of evolutionary rates, with the major shifts identified on the same nodes.
- 367
- 368 Assessing brain covariation
- 369 We measured the magnitude of covariation between the different ontogenetic stages by employing
- the standard PLS analysis. PLS differs from linear regression by treating the two variables
- 371 symmetrically rather than using one set of variables (independent) to predict the other. Instead, PLS
- 372 constructs new variables that are linear combinations of the variables within each of the sets,
- accounting for as much as possible of the covariation between the two original sets of variables.
- 374 The magnitude of morphological covariation in the brain at the macroevolutionary context has been
- assessed using the CR coefficient measured accounting for shared ancestry applying the function
- 376 *phylo.modularity* from the R package 'geomorph'⁶¹. The CR coefficient is a measure of the overall
- 377 covariation between modules divided by the overall covariation within modules. The CR coefficient
- ranges from 0 to positive values, where lower values indicate low covariation and high values
- indicate higher covariation, here departure from the null hypothesis of random association between
- modules is assessed via permutation. Furthermore, measuring the CR coefficient is insensitive to
- variation in sample size and number of variables as the variance of each module is not included.
- These analyses were repeated after accounting for the effect of size measured as logarithm of centroid size.
- Finally, it has been recently noted⁶² that sliding semi-landmarks using the minimum bending energy
- 385 (BEN) approach may result in increased covariation between modules. Because we used semi-
- 386 landmarks in our dataset, we repeated all the following integration analyses using shape coordinates
- derived using both the minimum BEN and minimum Procrustes distances (PRD) approaches to
- evaluate any potential discrepancy in the results. We found no significant discrepancies when using
- either sliding methods, hence we present only the results obtained from the analyses performed on
- the shape coordinates derived after using the minimum BEN approach.
- 391

392 Assessing endocast modular partitioning

- 393 Brain covariation was measured by dividing the brain into 6 distinct subunits following previously
- ³⁹⁴ published protocols and on the recognition of traits on the cortical surface areas identified from the ³⁹⁵ 3D reconstruction^{2,4,23,35,63-65} (see Supplementary Fig. 2).
- 595 5D reconstruction², 25,555 or (see Supplementary Fig. 2).
- 1-2) The frontal and pre-frontal regions extend from the frontal pole anteriorly to the central sulcus
- 397 posteriorly. The central sulcus is a longitudinal unfolding beginning on the medial surface of the
- brain. The frontal region borders with the postcentral gyrus of parietal lobe, and it is separated from
 the temporal lobe by the lateral sulcus⁶⁶.
- 400 3) The anterior border of the parietal region is demarcated by the central sulcus and the inferior
- 401 border is demarcated by the Sylvian fissure. It extends posteriorly where it meets the occipital areas.
- 402 4) The parietal lobe can be further subdivided into major subareas which can be identified from the
- 403 endocranial surface (supramarginal gyrus, angular gyrus, intraparietal sulcus, superior parietal
 404 lobule) ⁶⁵.
- 405 5) The temporal lobe is separated from the other cortical area by the Sylvian fissure, a feature
- 406 unique to primates⁶⁷.

6) The occipital lobe is the most posterior region of the brain and borders the parieto-occipital fissure which separate it from the parietal areas⁶⁸.

409 However, describing different modules on the endocasts can be challenging and to better define the

- 410 different regions we accounted for the uncertainties of assessing clear boundaries between the
- 411 different modules we applied two different strategies.
- 412 1) We defined four different modular configurations and evaluate between them by using the
- 413 standardised test statistics based on the comparison of the Covariance Ratio (CR) measurement.
- This assesses the covariances within and among hypothesised modules and compares this ratio with
- a null hypothesis of random assignment of shape variables to partitions^{69,70}. We found that the most
- supported configuration was the one formed by four distinct modules (Supplementary Fig. 2 and
- 417 Supplementary Table 3).
- 418 2) We devised a novel strategy to measure the intensity of local modularity and integration without
- defining modules a priori. In geometric morphometrics applications, a module is defined as a
- 420 discrete region characterised by greater integration internally than externally. To locate brain areas
- 421 matching this condition, for each semilandmark we selected its 9 closest semilandmarks, forming a
- 422 candidate modulet (N-Core) of 10 semilandmarks. All the other semilandmarks of the entire set
- 423 define a second module (R-Core) (Extended Data Fig. 1). We calculated the Covariance Ratio (CR)
- 424 between N-Core and R-Core, repeated the operation over all semilandmarks for the entire set and
- 425 mapped CR values on a reference mesh. The CR between each N-Core and its corresponding R-
- 426 Core indicated how much N-Core is likely to form a discrete module (see Supplementary Fig. 3-4).
- 427 A similar procedure was used to calculate the local integration by computing the correlation of the
- first PLS axis between N-Cores and R-Cores. At each iteration, the Procrustes Generalised Analysis (GPA) is performed separately on each of the two blocks (N and R-Cores). This way, by using PLS
- (GPA) is performed separately on each of the two blocks (N and R-Cores). This way, by using P.
 the level of integration was calculated iteratively over all semilandmarks of the entire sample.
- 431
- 432 *RRphylo and overfitRR*
- 433 We derived rates of brain shape evolution by the *RRphylo* method ⁷¹, available within the R package
- 434 'RRphylo' (v.2.5.0). Under *RRphylo*, consequent phenotypic changes occurring along a phyletic
- 435 line, from the root to a species are given by the equation $\Delta P = \beta_1 l_1 + \beta_2 l_2 + ... + \beta_n l_n$ where β_{ith} and
- 436 *lith* represent the regression coefficient and branch length, respectively, for each *i*th branch along the
- 437 phyletic line. Being regression slopes, the β coefficients represent the magnitude of phenotypic
- 438 change occurring along each branch, that is the actual rate of phenotypic evolution. The matrix
- 439 solution to find the vector of β coefficients for all the branches is given by the equation $\hat{\beta} =$
- 440 $(L^T L + \lambda I)^{-1} L^T y$; where L is the matrix of species to root time distances of the tree (the branch
- lengths), having tips as rows, \hat{y} is the vector of species phenotypes, and $\hat{\beta}$ is the vector of rates. λ is
- 442 a penalisation factor which prevents overfitting by penalising extremely large rates. Lambda (λ) is
- derived by means of maximum likelihood estimation by minimising rate variance within clades ascompared to variance between clades.
- 445 To locate clade-wise shifts in evolutionary rates, we used the function *search.shift* from the package
- 446 'RRphylo' ⁷¹. *search.shift* is specifically meant to automatically scan the phylogeny to identify
- shifts in absolute phenotypic evolutionary rates. Given rates as produced by *RRphylo*, *search.shift*
- starts by selecting all the subclades within the tree ranging from one tenth to one half of the total
- tree size. For each clade, it computes the difference between the mean absolute rate pertaining the
- 450 branches within the clade and the same figure for all other branches within the tree. Each difference
- is compared to a random distribution of 1,000 differences derived by randomly swapping ratevalues among the branches.
- 453 To account for sampling, phylogenetic uncertainty in tree topology and branch lengths, we used the
- 454 RRphylo function *overfitRR*. Over 100 consecutive iterations, the function randomly removes a
- 455 number of tips corresponding to 25% of the tree size and swaps species phylogenetic position of the
- 456 10% of the remaining species. For instance, a topology of the kind ((A, B), C) might change to ((C,
- B), A) or ((A, C), B). In addition, the age of 10% of the tree nodes is changed 'moving' the node in

between the age of its direct ancestor and the age of its oldest daughter node. At each iteration,

459 *overfitRR* performs *search.shift* on pruned tree and data testing whether the pattern found with the

460 original data is robust to sampling and phylogenetic uncertainty issues. The results of the analysis of

- rates of CR evolution were confirmed after accounting for phylogenetic uncertainty, by randomly
- swapping tree branches and node ages, suggesting they are not a consequence of the tree topology
- 463 we used (Hominoidea: p = 0.99; Strepsirrhini: p = 0.01; Cebidae: p = 0.01).
- 464

465 **Data availability:** All data required to replicate this study are available at (definitive Figshare link)

- 466 **Code availability:** The code required to replicate this study is available at (definitive Figshare link)
- 467

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471

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473 AP, JL, MM and KA processed the endocasts. SL and GS digitized the landmarks. GS, PR, AP, CS,

474 SC, MM and AM analyzed the data. GS, PR, AP and SW wrote the manuscript with significant

- contribution from all the other authors.
- 476

477 **Competing Interests Statement:** The authors declare no competing interests.

478

479 Figure Captions

Figure 1. Patterns of postnatal integration in modern humans and chimpanzees. a. Postnatal growth
stages for *Homo sapiens*, *Pan troglodytes*, *Gorilla gorilla* (only stages 4 and adult) and r-PLS
values per ontogenetic stage. The meshes in the lower left corner refer to the average for each stage

values per ontogenetic stage. The meshes in the lower left corner refer to the average for each stage
 and are coloured according to the magnitude of integration. **b**. Pairwise r-PLS values between brain

and are coloured according to the magnitude of integration. **b**. Pairwise r-PLS values between bra
 modules in adult *H. sapiens*, *P. troglodytes* and *G. gorilla*. **c**. Comparison of r-PLS values per

484 includes in adult *TL suprens*, *T. troglodytes* and *O. gortua*. C. Comparison of 1-1 LS values per
 485 ontogenetic stage between *H. sapiens* and *P. troglodytes* calculated using the NR-PLS approach,

that does not require the *a priori* definition of brain modules. Warm (cold) colours refer to low

487 (high) magnitude of integration. Animal silhouettes were available under Public Domain license at

488 phylopic (http://phylopic.org/). Specifically, Homo sapiens (http://phylopic.org/image/c089caae-

489 43ef-4e4e-bf26-973dd4cb65c5/) - No Copyright - Public Domain Dedication 1.0; *Pan troglodytes*

490 (http://phylopic.org/image/2f7da8c8-897a-445e-b003-b3955ad08850/) - credit to T. Michael

491 Keesey (vectorization) and Tony Hisgett (photography) - Creative Commons Attribution 3.0

492 Unported license.

493

494 **Figure 2**. Macroevolution of primate brain morphology and covariation. **a**. PC1/PC2

⁴⁹⁵ phylomorphospace of primate brain shape variation. **b**. Distribution of CR rate shifts on the tree.

496 Magenta shades indicate a slowdown in CR rate of evolution, whereas the cyan shade indicates

497 acceleration. Brain meshes represent the average shape for each clade and are coloured according to

the magnitude of CR. Warmer colours refer to low CR values, cooler colours refer to high CR.

499

500 **Figure 3**. **a**. Distribution of CR evolutionary rates within hominoidea. The black vertical line

represents the average rate of CR evolution calculated over the entire Primate tree, orange dots

502 indicate internal nodes in the phylogeny. **b.** Evolutionary patterns of morphological integration

503 within Homo. Magenta shades indicate a slowdown in the CR rate of evolution, the cyan shade

- 504 indicates acceleration. Brain meshes represent the average shape for *H. sapiens* plus *H.*
- 505 *neanderthalensis* and all *Homo* species, respectively. The CR values are mapped over the endocast
- mesh. Warmer colours refer to low, cooler colours to high CR values.

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667





Symphalangus syndactylus

Australopithecus africanus Hoolock hoolock Pongo pygmaeus Pongo abelii Nomascus concolor Hylobates pileatus Hylobates klossii Pan troglodytes troglodytes Hylobates agilis Hvlobates lar Homo ergaster Hylobates muelleri Homo erectus Nomascus gabriellae Gorilla beringei Gorilla gorilla gorilla Nomascus leucogenys Homo heidelbergensis Homo neanderthalensis \cap 2 6 8 0 4 Rates



b









Taxon	Stage 2	Stage 3	Stage 4	Adult
Pan troglodytes	0.93	0.94	0.97	0.57
Homo sapiens	0.91	0.94	0.96	0.93
			Stages 3-4	Adult
Gorilla gorilla			0.97	0.71
Pongo sp.			0.94	0.63

Taxon	Stage 2	Stage 3	Stage 4	Adult
Pan troglodytes	0.46	1.33	1.18	0.11
Homo sapiens	1.44	3.77	3.12	2.73
			Stage 3-4	Adult
Gorilla gorilla			0.73	0.21
Pongo			1.02	0.17

Clade	CR		
Hominoidea	0.963		
Cercopithecinae	0.825		
Colobinae	0.918		
Platyrrhini	0.767		
Strepsirrhini	0.726		

Pairwise test								
	Cercopithecinae	Colobinae	Hominoidea	Platyrrhini	Strepsirrhini			
Cercopithecinae		0.468	<0.001	<0.001	<0.001			
Colobinae			<0.001	<0.001	<0.001			
Hominoidea				<0.001	<0.001			
Platyrrhini					<0.001			
Strepsirrhini								