


16. de Boer, G. Plasticity in food preference and diet-induced differential weighting of brain components across species, therefore, would be facilitated by a size measure that is independent of body parameters. To address this need for an internal normalization, we calculated the volume fraction (F) for each component, defining the total brain volume within each species as 1. We called the set of all the volume fractions for a species the cerebrotype. Brains therefore constitute a scalable architecture.


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**Scalable architecture in mammalian brains**

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Comparison of mammalian brain parts has often focused on differences in absolute size—, revealing only a general tendency for all parts to grow together. Attempts to find size-independent effects using body weight as a reference variable obscure size relationships owing to independent variation of body size and give phylogenies of questionable significance. Here we use the brain itself as a size reference to define the cerebrotype, a species-by-species measure of brain composition. With this measure, across many mammalian taxa the cerebellum occupies a constant fraction of the total brain volume (0.13 ± 0.02), arguing against the hypothesis that the cerebellum acts as a computational engine principally serving the neocortex. Mammalian taxa can be well separated by cerebrotype, thus allowing the use of quantitative neuroanatomical data to test evolutionary relationships. Primate cerebrotypes have progressively shifted and neocortical volume fractions have become successively larger in lemurs and lorises, New World monkeys, Old World monkeys, and hominoids, lending support to the idea that primate brain architecture has been driven by directed selection pressure. At the same time, absolute brain size can vary over 100-fold within a taxon, while maintaining a relatively uniform cerebrotype. Brains therefore constitute a scalable architecture.

Components of the brain make many more connections to one another than to any external structure. Furthermore, body size is under separate developmental and environmental control, rendering it an approximate reference measure at best. Comparison of brain components across species, therefore, would be facilitated by a size measure that is independent of body parameters. To address this need for an internal normalization, we calculated the volume fraction (F) for each component, defining the total brain volume within each species as 1. We called the set of all the volume fractions for a species the cerebrotype. A database of insectivores and primates showed significant variation of the cerebrotype (Fig. 1a, c). In insectivores, F did not vary systematically with brain size for any principal developmental brain division (Fig. 1a), therefore providing a baseline trend for comparison. By contrast, primates showed a strong trend for telencephalic growth. Fmed (volume fraction of telencephalon) was 60 ± 4% (mean ± s.d.; n = 28 species) in insectivores and 61 ± 1% in Scandentia (tree shrews; n = 3) but increased to 74 ± 5% in primates (n = 44). This increase occurred largely at the expense of medulla, mesencephalon and diencephalon: Fmed + Fmean + Fdien, respectively, was 27 ± 3% in insectivores and 26 ± 1% in tree shrews, and decreased to 14 ± 4% in primates. This trade-off was emphasized by the ratio Fmed/(Fmed + Fmean + Fdien), which was 2.3 ± 0.4 in insectivores, 2.3 ± 0.2 in tree shrews, 6.3 ± 3.2 in primates and 20.8 in *Homo sapiens*.  

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**Figure 1** Analysis of volumetric data from mammalian brains. a. Volume fractions of principal brain divisions plotted against total brain volume. The colours are as indicated in c. b. Volume fractions of telencephalon subcomponents. The colours are as indicated in c. d. Area plot of principal brain-division volume fractions sorted by increasing telencephalic volume fraction within each taxon, demonstrating relative changes in neocerebellar structures. Horizontal bars indicate the taxonomic groupings of insectivores (Insectiv.), tree shrews (Tree) and primates. d. Area plot of telencephalic subcomponent volume fractions sorted by increasing neocortical volume fraction of the telencephalon.
Studies using body-weight-referenced scaling or comparison of absolute volumes suggested that cerebellum and neocortex vary together, leading to the proposition that the cerebellum is a computational engine that serves the needs of the neocortex. By contrast, we found that $F_{cbl}$ (volume fraction of cerebellum) remained nearly constant across groups in the ref. 6 data set (13.2 ± 2.2% in insectivores, 12.7 ± 0.2% in tree shrews and 12.4 ± 2.1% in primates; Fig. 1c, red points). To explore the generality of this finding, we examined $F_{cbl}$ across 19 mammalian taxa (Fig. 2). $F_{cbl}$ values across taxa remained constant (13.5 ± 2.4%) even as $F_{neo}$ (volume fraction of neocortex) expanded from 16 ± 6% in Soricomorpha to 74 ± 5% in Hominioidea (apes and humans). Across this range of species, we found no correlation between $F_{neo}$ and $F_{cbl}$ ($r^2 = 0.013$, $n = 19$). The constancy of cerebellum in mammals suggests that its functional role must be evaluated not in terms of neocorticalization, but rather with either coordinated whole-brain function or some aspect of body plan that correlates strongly with total brain size.

$F_{cbl}$ may be constrained by the high rate of energy consumption by brain tissue, which would exert selection pressure for structures not to exceed a minimum essential size. Conversely, any taxon-specific increases in $F_{cbl}$ might reflect necessary adaptations related to the addition of new functions. The cerebellum has been suggested to transform sensory information as a means of guiding motor activity. We found significantly higher values of $F_{cbl}$ in mammals with sophisticated echolocating abilities: Cetacea (dolphins and whales, 19 ± 3%), 7 species; different from $F_{cbl}$ in other taxa by two-tailed significance test, $P < 0.02$) and Microchiroptera (microbats, 22 ± 5%, 225 species; different from other taxa, $P < 0.01$). In contrast to Microchiroptera, the Megachiroptera (megabats), which lack the ability to echolocate, did not have unusually large cerebella ($F_{cbl} = 14 ± 1%$, 47 species; $P = 0.97$), suggesting that adaptations to bat body plan alone were not responsible for the cerebellar enlargement. The observed size differences may be region specific: Cetacea and Microchiroptera are hypertrophied in specific homologous parafloccular and medial regions, which in bats are responsive to acoustic target position and velocity (reviewed in ref. 14). Similarly, among teleost fishes, the largest cerebella are found in mormyrids, which electrolocate to detect object size and distance. The common factor of echolocation and electrolocation is the interpretation of subtle timing differences in echoes returned from transmitted pulses. Thus, regardless of its overall functional role, the cerebellum seems in these species to provide additional processing power for sophisticated sensory adaptations. Our results also suggest a more general speculation: that the cerebellum provides guidance to the entire brain on the basis of fine features of sensory input.

Our observations in the whole brain led us to search for size trade-offs and constancy within the telencephalon. Normalizing to a total telencephalic volume of 1, telencephalic volume fractions $T$ also showed high variability among orders (Fig. 1b, d). $T_{neo}$ increased from 28 ± 10% ($n = 28$) in insectivores to 55 ± 1% ($n = 3$) in tree shrews, 81 ± 8% ($n = 44$) in primates and 95% in Homo (Fig. 1b, blue points). This expansion in neocortex occurred at the expense of hippocampus, septum, schizocortex, piriform cortex and olfactory bulbs: the sum of these fractions decreased from 64 ± 10% in insectivores to 37 ± 1% in tree shrews, 12 ± 7% in primates and 3% in Homo.

In the face of this neocortical expansion, the telencephalic structure that showed the least change was the striatum (Fig. 1d, red points). The striatum maintained a volume fraction of $T_{stri}$ = 8 ± 2% in insectivores, 8 ± 1% in tree shrews, 6 ± 1% in primates and 3% in Homo. The striatum forms extensive interconnections with the hippocampus, amygdala and nearly all of the cerebral cortex. These connections form part of an extensive loop corticothalamic network that, when damaged, leads to extrapyramidal motor disorders such as Parkinson’s and Huntington’s disease. The relative preservation of $T_{stri}$ is consistent with a function in motor activity that depends on the amount of telencephalic input and requires some minimum necessary amount of striatal processing.

**Figure 2** Constancy of cerebellar volume fraction across mammalian taxa. Cerebellar volume fractions are plotted at order level with the exception of the primates, which were divided into Strepsirhini (lemurs and lorises), Platyrrhini (New World monkeys), Cercopithecoidae (Old World monkeys) and Hominioidea (apes and human). The cerebellar volume fractions are indicated by solid symbols. Data are sorted by neocortical volume fraction (open symbols). The error bars indicate s.d.; the shaded bar indicates mean ± 1 s.d. for the pooled data. Statistically significant differences between an individual taxon and the entire group are identified in bold (Student’s t test: asterisk, $P < 0.02$; double asterisk, $P < 0.01$).

**Figure 3** Clustering of cerebrotypes by taxonomy. a, Neocortical fraction of total brain volume plotted against total brain volume for insectivores (filled squares), tree shrews (open squares), lemurs, lorises and tarsier (filled circles), and monkeys and apes (open circles). b, Variation in the 11-component cerebrotypes for the species plotted in a displayed in a plane using multidimensional scaling. Compared with true Euclidean distance between cerebrotypes, the root-mean-squared (r.m.s.) fractional error in displayed interspecies distance is 8.4%.
The cerebrotype variation we observe suggests that each taxon might be associated with its own distinct brain architecture. This can be seen at a coarse level by plotting a single quantity, $F_{rneo}$ against brain volume\(^1\). Although absolute brain volume varied several hundredfold within each order, there was little variation of $F_{rneo}$ (Fig. 3a), indicating that this fraction takes on specific values for major taxonomic divisions\(^3\).

To facilitate the identification of finer taxonomic distinctions, we used all total-brain and telencephalic volume fractions to generate a relational map of mammalian brain architectures. We did this using multidimensional scaling, an algorithm that shows the variation of all 11 volume fractions in a plane with maximum fidelity to the true distances between cerebrotypes. This method allows species to be compared without making \textit{a priori} assumptions about allometric scaling relationships between components. Figure 3b shows clear separation between insectivores and primates. Scandentia, once categorized as primates\(^6\), were nearer to insectivores than to primates in cerebrotype, but there was no overlap between either group. This assessment agrees with recent morphological and molecular evidence, and supports the identification of Scandentia as a separate mammalian lineage.

The eight ‘insectivore’\(^15\) orders represented in this data set also appeared as distinct groups of points (Fig. 4a), with the exception of the overlapping cerebrotype distributions of Soricidae (shrews) and Tenrecidae (tenrecs). This overlap arose entirely from the cerebrotypes of three tenrecs that live and/or hunt in the water (Fig. 4a). This difference from other tenrecs suggests that these three species have brain specializations that reflect an aquatic lifestyle, such as reductions in the size of the olfactory bulbs and piriform cortex (see Supplementary Information). Thus, cerebrotype analysis provides a means of taxonomically grouping mammals by their brain structure. The existence of these characteristic within-taxon cerebrotypes across large variations in absolute brain size suggests that brain components are scalable; that is, a functionally optimized brain has the same proportions independent of absolute volume. Furthermore, the fact that 11-component cerebrotypes can be mapped into a two-dimensional plane but still accurately represent distance information suggests that the range of possible brain architectures is strongly constrained.

Cerebrotype-based measures might also be useful in finding directional changes in brain architecture. According to the social intelligence hypothesis of primate brain evolution\(^4\), a larger neocortex may confer selection advantages owing to increased cognitive capacity for use in social dynamics. This would lead to progressive, relative enlargement of the neocortex over time. Fossil and molecular evidence and morphological evidence from non-brain characters shows that successively more derived primate taxa have arisen over time: lemurs/lorises, followed by tarsiers, New World monkeys, Old World monkeys, and then hominoids\(^8\). The arrangement of the corresponding cerebrotypes (Fig. 4b) is collinear with this order of primate taxa, with the exception of some overlap between New World and Old World monkeys. This overlap arises from the four New World monkeys in this database—\textit{Ateles geoffroyi}, \textit{Lagothrichia lagothricha}, \textit{Cebus} sp. and \textit{Saimiri sciureus}—with larger group sizes and complex social structures resembling those of Old World monkeys\(^2\). The cerebrotypes of New World monkeys with large group sizes differ from other New World monkeys principally in $F_{rneo}$ (large group size 69.3 ± 0.5%, 4 species; other 61.8 ± 1.7%, 8 species; $P < 0.001$) but are similar to Old World monkeys (70.4 ± 2.4%, 11 species; $P = 0.1$; see Supplementary Information). Taken together, these orderings suggest that increased derivation was indeed accompanied by progressive changes in the cerebrotype, perhaps relating to social intelligence.

Cerebrotype shifts in primates are accompanied by significant expansion of the neocortical volume fraction\(^1\) (Fig. 3a). To identify other, independent changes in architecture, we eliminated neocortex from the data set and renormalized the remaining ten regions. Multidimensional scaling still showed clear separation of primate taxa (Fig. 4c). We obtained similar results when we also removed the cerebellum from the data set (not shown). Thus, the rise of new primate taxa has been accompanied by substantial shifts in brain architecture, marked both by increases in relative neocortical size and by distinct changes in the distribution of brain volume among other regions.

Because cerebrotype shifts are progressive, cerebrotype differences should be largest between groups with the oldest last common ancestor. We emphasized changes in forebrain structures by considering the telencephalic cerebrotype. At principal points of divergence for which dates have been estimated multiple times from DNA and fossil evidence\(^16\) (Fig. 5a), the Euclidean cerebrotype distance was calculated between groups of species on either side of each node (see Methods) (Fig. 5b). The most recent divergence points were consistently associated with shorter distances between cerebrotypes (Spearman’s rank correlation coefficient $r_s = 1.00$, $P < 0.01$). By contrast, a previous metric of body-size-scaled indices\(^5\) has suggested homology between the brains of humans and spider monkeys, a result irreconcilable with known phylogeny. Therefore, among quantitative brain parameters examined to date, only the cerebrotype provides a measure of architecture that correlates with date of divergence of advanced primates.

To test if changes in brain architecture tracked primate evolution within each major group, the method of independent contrasts (see Methods)\(^18\) was used to make comparisons among groups of species (Fig. 5c). Overall, longer cerebrotype distances were associated...
with older divergence dates (Spearman’s rank correlation coefficient $r = 0.45$, $P < 0.01$, $n = 28$ independent comparisons). This strong correlation led us to use distance between individual species to reconstructure the taxonomy of four hominoids (Fig. 5d). The resulting tree was identical to that obtained from DNA sequence (Fig. 5e, left), (((Homo, Pan), Gorilla), Pongo), but not to a tree obtained from bone and tooth morphology (Fig. 5e, right), (Homo, (Pan, (Pongo, Gorilla))). This lends support to the suggestion that soft tissues represent a more accurate route than hard tissues to phylogenetic reconstruction.

Our observations suggest a model for primate brain evolution in which adaptive radiation within a taxon has generated changes in the absolute brain (and body size) while preserving an approximately fixed cerebrotype. Shifts in cerebrotype occurred at infrequent (about 10 Myr) intervals coinciding with the rise of new taxa. These shifts were marked primarily by relative expansion of the neocortex, especially the neocortex. This expansion has reached an extreme in Homo, which possesses the largest $F_{\text{neu}}$ value known (0.80).

Our observations suggest a reconciliation of developmentalist and adaptationist models of how brain architecture is determined. Within each taxon, brain regions are scalable, tending to maintain fixed proportionality of size to one another independent of absolute total brain volume. This suggests that, within a taxon, the development of multiple brain regions is governed by a common set of mechanisms. However, because the cerebrotype varies from taxon to taxon, these developmental mechanisms must also be variable. This variation arises from genetic modification that leads to the appearance of new cerebrotypes. For instance, in primates, the enlargement of the neocortex may be caused by the action of additional rounds of cortical neurogenesis. In addition to these major shifts, variations in developmental rates in other brain regions can lead to limited adaptive radiation of the cerebrotype within each taxon. These variations may be driven by ecological and/or social requirements. Finally, the microscopic composition of neural tissue (neurons, glia, axons and dendrites) varies across mammals. An ultimate explanation of these changes in brain structure will, therefore, require a deeper understanding of both the causes for variation at a cellular level and the selective advantages provided by a changing brain architecture.

Methods
For cerebrotype analysis, data were taken from ref. 6, consisting of 28 insectivores, 3 Scandentia (tree shrews), 18 strepsirhine (lemur and loris) primates and 27 haplorhine (tarsier, New World and Old World monkey, and hominoid) primates. Definitions of primate species conformed to ref. 16 and definitions of brain regions conformed to ref. 6. Brain volumes ranged over four orders of magnitude. The volume fraction was defined for the five principal brain components and for the seven telencephalic components. Whole-brain volume fractions were calculated for medulla oblongata, cerebellum, mesencephalon, diencephalon and telencephalon by dividing by total brain volume. Volume fractions were calculated for olfactory bulb, piriform cortex, septum, striatum, hippocampus and neocortex by dividing by either the whole-brain volume (fractions $F$) or by the telencephalic volume (fractions $T$). We excluded the accessory olfactory bulb, which is not present in many primates, from the analysis. We also excluded one haplorhine (Calliethrix jacchus) from the analysis because the components failed to sum to the whole in the original data. Additional data used to compare neocortical and cerebellar volume fractions across taxa (Fig. 2) were obtained from the literature for Artiodactyla, Carnivora, Cetacea, Edentata, Lagomorpha, Marsupialia, Microchiroptera and Megachiroptera, Rodentia, Sirenia and orangutan. All values are given as mean ± s.d. unless otherwise noted.

Independent contrasts
We arranged species in a tree according to a published primate phylogeny. At each node, two mean cerebrotypes were calculated, one for each branch emanating from the node. The Euclidean distance between the two means was used to make a comparison for that node, the method of independent contrasts.

Phylogenetic reconstruction
The telencephalic cerebrotype was defined as $T$, the vector of telencephalic volume fractions. For hominoids, a phylogenetic tree was constructed from cerebrotype distances using the Fitch–Margoliash algorithm, which sequentially adds species to a test tree while minimizing the quantity $\sum (n_i - e_j)^2 / \sigma_j^2$, where $e_j$ is the Euclidean cerebrotype distance between species $i$ and $j$, and $n_i$ is the total length of the path joining the two species in the test tree. Colobine (leaf-eating monkeys) were used as the outgroup to root the tree. Optimization and display were done using the PHYLIP software package running on a PC-DOE platform.
Multidimensional scaling was done using MATLAB (The Mathworks, Natick, MA) to allow simultaneous display of variation in all 11 F components (principal brain divisions and telencephalic components) of the cerebrotype. The quantity \( \Sigma (d_{ij} - d_{ij0})^2 \) was minimized where \( d_{ij} \) is the cerebrotype distance and \( d_{ij0} \) is the displayed distance in the plane. Minimizations were done using a bootstrapping procedure. The plot was initially seeded using three points chosen by a trial run to lie near the outskirts of the final plot. We added subsequent points at a time in a random order. After each point was added, a round of error minimization was performed in which only the new point was allowed to vary, followed by a round of minimization in which all points were varied. Each minimization was performed at least ten times and the solution with the least error was accepted. In the 76-point minimization of all species (Fig. 3b), new points were added four at a time. This overall procedure resembles the Fitch–Margoliash algorithm for phylogeny reconstruction but yields a mapping in a plane rather than a connected tree.

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