A small world of weak ties provides optimal global integration of self-similar modules in functional brain networks

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The human brain is organized in functional modules. Such an organization presents a basic conundrum: Modules ought to be sufficiently independent to guarantee functional specialization and sufficiently connected to bind multiple processors for efficient information transfer. It is commonly accepted that small-world architecture of short paths and large local clustering may solve this problem. However, there is intrinsic tension between shortcuts generating small worlds and the persistence of modularity, a global property unrelated to local clustering. Here, we present a possible solution to this puzzle. We first show that a modified percolation theory can define a set of hierarchically organized modules made of strong links in functional brain networks. These modules are “large-world” self-similar structures and, therefore, are far from being small-world. However, incorporating weaker ties to the network converts it into a small world preserving an underlying backbone of well-defined modules. Remarkably, weak ties are precisely organized as predicted by theory maximizing information transfer with minimal wiring cost. This trade-off architecture is reminiscent of the “strength of weak ties” crucial concept of social networks. Such a design suggests a natural solution to the paradox of efficient information flow in the highly modular structure of the brain.

One of the main findings in neuroscience is the modular organization of the brain, which in turn implies the parallel nature of brain computations (1–3). For example, in the visual modality, more than 30 visual areas analyze simultaneously distinct features of the visual scene: motion, color, orientation, space, form, luminance, and contrast, among others (4). These features, as well as information from different sensory modalities, have to be integrated, as one of the main aspects of perception is its unitary nature (1, 5).

This leads to a basic conundrum of brain networks: Modular processors have to be sufficiently isolated to achieve independent computations, but also globally connected to be integrated in coherent functions (1, 2). A current view is that small-world networks provide a solution to this puzzle because they combine high local clustering and short path length (7–9). This view has been fueled by the systematic finding of small-world topology in a wide range of human brain networks derived from structural (10), functional (11–13), and diffusion tensor MRI (14). Small-world topology has also been identified at the cellular-network scale in functional cortical neuronal circuits in mammals (15, 16) and even in the nervous system of the nematode Caenorhabditis elegans (8). Moreover, small-world property seems to be relevant for brain function because it is affected by disease (17), normal aging, and by pharmacological blockade of dopamine neurotransmission (13).

Although brain networks show small-world properties, several experimental studies have also shown that they are hierarchical, fractal and highly modular (2, 3, 18). As there is an intrinsic tension between modular and small-world organization, the main aim of this study is to reconcile these ubiquitous and seemingly contradictory topological properties. Indeed, traditional models of small-world networks cannot fully capture the coexistence of highly modular structure with broad global integration. First, clustering is a purely local quantity that can be assessed inspecting the immediate neighborhood of a node (8). On the contrary, modularity is a global property of the network, determined by the existence of strongly connected groups of nodes that are only loosely connected to the rest of the network (2, 3, 19, 20). In fact, it is easy to construct modular and unclustered networks or, reciprocally, clustered networks without modules.

Second, the short distances of a small world may be incompatible with strong modularity, which typically presents the properties of a “large world” (21–27) characterized by long distances that effectively suppress diffusion and free flow in the system (26). Although a clustered network preserves its clustering coefficient when a small fraction of shortcuts are added (converting it into a small world) (8), the persistence of modules is not equally robust. As we show below, shrinking the network diameter may quickly destroy the modules.

Hence, the concept of small world may not be entirely sufficient to explain the modular and integration features of functional brain networks on its own. We propose that a solution to modularity and broad integration can be achieved by a network in which strong links form large-world fractal modules, which are shortcut by weaker links establishing a small-world network. A modified percolation theory (28, 29) can identify a sequence of critical values of connectivity thresholds forming a hierarchy of modules that progressively merge together. This proposal is inspired by a fundamental notion of sociology termed by Granovetter as “the strength of weak ties” (30, 31). According to this theory, strong ties (close friends) clump together forming modules. An acquaintance (weak tie) becomes a crucial bridge (a shortcut) between the two densely knit clumps (modules) of close friends (30).

Interestingly, this theme also emerges in theoretical models of large-scale cognitive architecture. Several theories suggest integration mechanisms based on dynamic binding (6, 32) or on a workspace system (1, 33). For instance, the workspace model (1, 33) proposes that a flexible routing system with dynamic and comparably weaker connections transiently connects modules with very strong connections carved by long-term learning mechanisms.

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Results

Experimental Design and Network Construction. We capitalize on a well-known dual-task paradigm, the psychological refractory period (34). A total of 16 subjects responded with the right hand to a visual stimulus and with the left hand to an auditory stimulus (see SI Text). The temporal gap between the auditory and visual stimuli varied in four stimulus onset asynchrony, SOA = 0, 300, 900, and 1,200 ms. The sequence of activated regions that unfolds during the execution of the task has been reported in a previous study (35).

The network analysis relies on the time-resolved oxygen level-dependent functional magnetic resonance imaging (BOLD-fMRI) response based on the phase signal obtained for each voxel of data (36). We first compute the phase of the BOLD signal for each voxel with methods developed previously (36). For each subject and each SOA task, we obtain the phase signal of the ith voxel of activity, \( \theta_i(t) \), over \( T = 40 \) trials performed for a particular SOA value and subject. We use these signals to construct the network topology of brain voxels based on the equal-time cross-correlation matrix, \( C_{ij} \), of the phase activity of the two voxels (see SI Text). The accuracy of the calculated \( C_{ij} \) values was estimated through a bootstrapping analysis (see SI Text and Fig. S1).

To construct the network, we link two voxels if their cross-correlation \( C_{ij} \) is larger than a predefined threshold value \( p \) (11, 12, 37). The resulting network for a given \( p \) is a representation of functional relations among voxels for a specific subject and SOA. We obtain 64 cross-correlation networks resulting from the four SOA values presented to the 16 subjects.

Percolation Analysis. Graph analyses of brain correlations relies on a threshold (11), which is problematic because small-world-like properties are sensitive to even a small proportion of variation in the connections. The following analysis may be seen as an attempt to solve this problem.

The thresholding procedure explained above can be naturally mapped to a percolation process (defined in the \( N \times N \) space of interactions \( C_{ij} \)). Percolation is a model to describe geometrical phase transitions of connected clusters in random graphs; see ref. 28, chapters 2 and 3, and refs. 29 and 38.

In general, the size of the largest component of connected nodes in a percolation process remains very small for large \( p \). The crucial concept is that the largest connected component increases rapidly through a critical phase transition at \( p_c \), in which a single incipient cluster dominates and spans the system (28, 29, 38). A unique connected component is expected to appear if the links in the network are occupied at random without correlations.

However, when we apply the percolation analysis to the functional brain network, a more complex picture emerges revealing a hierarchy of clusters arising from the nontrivial correlations in brain activity.

For each participant, we calculate the size of the largest connected component as a function of \( p \). We find that the largest cluster size increases progressively with a series of sharp jumps (Fig. 1A, SOA = 900 ms, all participants, other SOA stimuli are similar). This suggests a multiplicity of percolation transitions where percolating modules subsequently merge as \( p \) decreases rather than following the typical uncorrelated percolation process with a single spanning cluster. Each of these jumps defines a percolation transition focused on groups of nodes that are highly correlated, constituting well-defined modules.

Fig. 1B shows the detailed behavior of the jumps in a typical individual (subject labeled #1 in our dataset available at http://lev.ccny.cuny.edu/~hmakse/brain.html, SOA = 900 ms). At high values of \( p \), three large clusters are formed localized to the medial occipital cortex (red), the lateral occipital cortex (orange), and the anterior cingulate (green). At a lower \( p = 0.979 \), the orange and red clusters merge as revealed by the first jump in the percolation picture. As \( p \) continues to decrease this mechanism of cluster formation and absorption repeats, defining a hierarchical process as depicted in Fig. 1B Upper. This analysis further reveals the existence of “stubborn” clusters. For instance, the anterior cingulate cluster (green), known to be involved in cognitive control (39, 40) and which hence cannot commit to a specific functional module, remains detached from the other clusters down to low \( p \) values. Even at the lower values of \( p \), when a massive region of the cortex—including motor, visual and auditory regions—has formed a single incipient cluster (red, \( p = 0.94 \)), new clusters emerge; one involving subcortical structures including the thalamus and striatum (cyan) and the other involving the left frontal cortex (purple). This mechanism reveals the iteration of a process by which modules form at a given \( p \) value and merged by comparably weaker links. This process is recursive. The weak links of a given transition become the strong links of the next transition, in a hierarchical fashion.

Below, we focus our analysis on the first jump in the size of the largest connected component, for instance, \( p_c = 0.979 \) in Fig. 1B. We consider the three largest modules at \( p_c \), with at least 1,000 voxels each. This analysis results in a total of 192 modules among all participants and stimuli, which are pooled together for the next study. An example of an identified module in the medial occipital cortex of subject #1 and SOA = 900 ms is shown in Fig. 1C in the network representation and in Fig. 1D in real space. The topography of the modules reflects coherent patterns across the subjects and stimuli as analyzed in SI Text (see Fig. S2).

Scaling Analysis and Renormalization Group. To determine the structure of the modules we investigate the scaling of the “mass” of each module (the total number of voxels in the module, \( N_c \)) as a function of three length scales defined for each module: \((I)\)
the maximum path length, \( \ell_{\text{max}} \); (ii) the average path length between two nodes, \( \langle \ell \rangle \); and (iii) the maximum Euclidean distance among any two nodes in the cluster, \( r_{\text{max}} \). The path length, \( \ell \), is the distance in network space defined as the number of links along the shortest path between two nodes. The maximum \( \ell_{\text{max}} \) is the largest shortest path in the network.

Fig. 2A indicates power-law scaling for these quantities (21, 28). For instance,

\[
N_c(\ell_{\text{max}}) \sim (\ell_{\text{max}})^{d_f}
\]

defines the Euclidean Hausdorff fractal dimension, \( d_f = 2.1 \pm 0.1 \). The scaling with \( \ell_{\text{max}} \) and \( \langle \ell \rangle \) is consistent with Eq. 1, as seen in Fig. 2A. The exponent \( d_f \) quantifies how densely the volume of the brain is covered by a specific module.

Next, we investigate the network properties of each module, applying renormalization group (RG) analysis for complex networks (21–25). This technique allows one to observe the network at different scales transforming it into successively simpler copies of itself, which can be used to detect characteristics that are difficult to identify at a specific scale of observation. We use this technique to characterize submodular structure within each brain module (2).

We consider each module identified at \( p_c \) separately. We then tile it with the minimum number of boxes or submodules, \( N_B \), of a given box diameter \( \ell_B \) (21); i.e., every pair of nodes in a box has shortest path length smaller than \( \ell_B \). Covering the network with minimal \( N_B \) submodules represents an optimization problem that is solved using standard box-covering algorithms, such as the Maximum Excluded Mass Burning algorithm, MEMB, which was introduced in refs. 21, 22, and 41 to describe the self-similarity of complex networks ranging from the World Wide Web, biological, and technical networks (see SI Text and Fig. 2B describing MEMB; the entire experimental dataset and modularization and fractal codes are available at http://lev.ceny.cuny.edu/~hmakse/brain.html). The requirement to minimize the number of boxes is important because the resulting boxes are characterized by the proximity between all their nodes and minimization of the links connecting the boxes (26). Thus, the box-covering algorithm detects boxes/submodules that also tend to maximize modularity.

The repetitive application of box-covering at different values of \( \ell_B \) is an RG transformation (21) that yields a different partition of the brain modules in submodules of varying sizes (Fig. 2B). Fig. 2C shows the scaling of \( N_B \) versus \( \ell_B \) averaged over all the modules for all individuals and stimuli. This property is quantified in the power-law relation (21):

\[
N_B(\ell_B) \sim \ell_B^{-d_{B}}
\]

where \( d_B \) is the box fractal dimension (21–25), which characterizes the self-similarity between different scales of the module where smaller-scale boxes behave in a similar way as the original network. The resulting \( d_B \) averaged over all the modules is \( d_B = 1.9 \pm 0.1 \).

**Morphology of the Brain Modules.** The RG analysis reveals that the module topology does not have many redundant links, and it represents the quantitative statement that the brain modules are large worlds. However, this analysis is not sufficient to precisely characterize the topology of the modules. For example, both, a two-dimensional complex network architecture and a simple two-dimensional lattice are compatible with the scaling analysis and the value of the exponents described in the previous section.

To identify the network architecture of the modules we follow established analysis (18, 42) based on the study of the degree distribution of the modules, \( P(k) \), and the degree-degree correlation \( P(k_1,k_2) \) (22, 43). The form of \( P(k) \) distinguishes between a Euclidean lattice (delta function), an Erdos–Renyi network (Poisson) (29), or a scale-free network (power law) (42). We find that the degree distribution of the brain modules is a power law (11, 42) \( P(k) \sim k^\gamma \) over an interval of \( k \) values. In the SI Text and Fig. S3 we describe a statistical analysis based on maximum likelihood methods and KS analysis, which yield the value of the degree exponent \( \gamma = 2.1 \pm 0.1 \) and the interval and error probability of the hypothesis that the data follow a power law (Fig. S4). The analysis rules out an exponential distribution (see SI Text).

How can fractal modularity emerge in light of the scale-free property, which is usually associated with small worlds (18)? In a previous study (22), we introduced a model to account for the simultaneous emergence of scale-free, fractality, and modularity in real networks by a multiplicative process in the growth of the number of links, nodes, and distances in the network. The dynamic follows the inverse of the RG transformation (22) where the hubs acquire new connections by linking preferentially with less connected nodes rather than other hubs. This kind of “repulsion between hubs” (23) creates a disassortative structure—with hubs spreading uniformly in the network and not crumpling in the core as in scale-free models (42). Hubs are buried deep into the modules, while low degree nodes are the intermodule connectors (23).

A signature of such mechanism can be found by following hubs’ degree during the renormalization process. At scale \( \ell_B \), the degree of a hub \( k \) changes to the degree of its box \( k' \), through the relation \( k' = s(\ell_B)k \). The dependence of the scaling factor \( s(\ell_B) \) on \( \ell_B \) defines the renormalized degree exponent \( d_k \) by \( s(\ell_B) \sim \ell_B^{-d_k} \) (21). Scaling theory defines precise relations be-
between the exponents for fractal networks (21), through $\gamma = 1 + dB/d_d$. For the case of brain modules analyzed here (Fig. S4A), we find $d_d \approx 1.5 \pm 0.1$. Using the values of $dB$ and $d_d$ for the brain clusters, the prediction is $\gamma = 2.26 \pm 0.11$, which is within error bars of the calculated value $\gamma = 2.1 \pm 0.1$ from Fig. S4B.

The previous analysis reveals the mechanism of formation of a scale-free network, but it does not assure a fractal topology, yet. Fractality can be determined from the study of the degree-degree correlation through the distribution, $P(k_1,k_2)$ to find a link between nodes with $(k_1,k_2)$ degree (22, 43). This correlation characterizes the hub-hub repulsion through scaling exponents $d_c$ and $e$ (see SI Text and Fig. S6) (22, 43). In a fractal, they satisfy $e = 2 + d_c/d_d$. A direct measurement of these exponents yields $d_c = 0.51 \pm 0.08$ and $e = 2.1 \pm 0.1$ (Fig. S6). Using the measured values of $d_c$ and $d_d$, we predict $e = 2.3 \pm 0.1$, which is close to the observed exponent. Taken together, these results indicate a scale-free fractal morphology of brain modules. Such structure is in agreement with previous results of the anatomical connectivity of the brain (2, 3) and functional brain networks (11).

**Quantifying Submodular Structure of Brain Modules.** Standard modularity decomposition methods (19, 20) based on maximization of the modularity factor $Q$ as defined in refs. 2, 19, 20, 26, and 27 can uncover the submodular structure. For example, the Girvan–Newman method (19) yields a value of $Q \sim 0.82$ for the brain clusters, indicating a strong modular substructure. Additionally, the box-covering algorithm benefits from detecting submodules (the boxes) at different scales. Then, we can study the hierarchical character of modularity (2, 26, 27) and detect whether modularity decomposition methods (19, 20) based on maximization of the largest number of intramodule links and the fewest intermodular links out $\sum_{i=1}^{N_B} L_i^{in}$.

$$Q(\epsilon_B) \equiv \frac{1}{N_B} \sum_{i=1}^{N_B} L_i^{in} - L_i^{out}.$$  

which is a variation of the modularity factor, $Q$, defined in refs. 19 and 20. Here, $L_i^{in}$ and $L_i^{out}$ represent the intra- and intermodular links in a submodule $i$, respectively. Large values of $Q$ (i.e., $L_i^{out} = 0$) correspond to high homogeneity (26). We make the whole modularization method available at http://lev.ccny.cuny.edu/~hmakse/brain.html.

Fig. 2D shows the scaling of $Q(\epsilon_B)$ averaged over all modules at percolation revealing a monotonic increase with a lack of a characteristic value of $\epsilon_B$. Indeed, the data can be fitted with a power-law form (26):

$$Q(\epsilon_B) \sim \epsilon_B^{d_M},$$  

which is detected through the modularity exponent, $d_M$. We study the networks for all the subjects and stimuli and find $d_M = 1.9 \pm 0.1$ (Fig. 2D). The lack of a characteristic length scale expressed in Eq. 4 implies that submodules are organized within larger modules such that the interconnections between those submodules repeat the basic modular character of the entire brain network.

The value of $d_M$ reveals a considerable modularity in the system as it is visually apparent in the sample of Fig. 3A Left, where different colors identify the submodules of size $\epsilon_B = 15$ in a typical fractal module. For comparison, a randomly rewired network (Fig. 3A Right and Center) shows no modularity and has $d_M \approx 0$. Scaling analysis indicates that $d_M$ is related to $L\text{rew} \sim \epsilon_B^{d_M}$, which defines the outbox exponent $d_c$, characterizing the number of intermodular links for a submodule (26) [$d_c$ is related to the Rent exponent in integrated circuits (3)]. From Eq. 4, we find:

$$d_M = dB - d_c,$$  

which indicates that the strongest possible modular structure has $d_M = dB$ ($d_c = 0$) (26). Such a high modularity induces very slow diffusive processes (subdiffusion) for a random walk in the network (26). Comparing Eq. 4 with Eq. 2, we find $d_c = 0$, which quantifies the large modularity in the brain modules.

**The Conundrum of Brain Networks: Small-World Efficiency or Large-World Fractal Modularity.** An important consequence of Eqs. 1 and 2 is that the network determined by the strong links above the first $p_c$ jump lacks the logarithmic scaling characteristic of small worlds and random networks (8):

$$\langle \ell \rangle \sim \log N_c.$$  

A fractal network poses much larger distances than those appearing in small worlds (21): A distance $\ell_{\text{max}} \approx 100$ observed in Fig. 24 (red curve) would require an enormous small-world network $N_c \sim 10^{1000}$, rather than $N_c \sim 10^4$, as observed for fractal networks in Fig. 24. The structural differences between a modular fractal network and a small-world (and a random network) are starkly revealed in Fig. 3A. We rewire the fractal module in Fig. 3A Left by randomly reconnecting a fraction $p_{\text{rew}}$ of the links while keeping the degree of each node intact (8). Fig. 3B quantifies the transition from fractal ($p_{\text{rew}} = 0$) to small world ($p_{\text{rew}} \approx 0.01$–0.1) and eventually to random networks ($p_{\text{rew}} = 1$), illustrated in Fig. 3A: We plot $\ell_{\text{max}}(p_{\text{rew}})/\ell_{\text{max}}(0)$, the clustering coefficient $C(p_{\text{rew}})/C(0)$, and $\ell(1) = \ell(0)$ for a typical $\epsilon_B = 15$ as we rewire $p_{\text{rew}}$ links in the network. As we create a tiny fraction $p_{\text{rew}} = 0.01$ of shortcuts, the topology turns
into a collapsed network with no trace of modularity left, while the clustering coefficient at $p_{rew} = 0.01$ still remains quite high (Fig. 3B). The rewired networks present the exponential behavior of small worlds (8) and also random networks as $p_{rew}$ increases, obtained from Eq. 5:

$$N_c \sim \exp((\ell)/\ell_0),$$  

[6]

where $N_c$ is averaged over all the modules (Fig. 3C). The characteristic size is very small and progressively shrinks to $\ell_0 = 1/7$ when $p_{rew} = 1$. The hallmark of small worlds and random networks, exponential scaling (Eq. 6), is incompatible with the hallmark of fractal large-worlds, power-law scaling (Eq. 2). More importantly, although we find a broad domain where short network distances coexist with high clustering forming a small-world behavior, modularity does not show such a robust behavior to the addition of shortcuts (Fig. 3B).

**Shortcut Wiring is Optimal for Efficient Flow.** Fig. 3B suggests that modularity and small world cannot coexist at the same level of connectivity strength. Next, we set out to investigate how the small world emerges.

When we extend the percolation analysis lowering further the threshold $p$ below $p_c$, weaker ties are incorporated to the network connecting the self-similar modules through shortcuts. A typical scenario is depicted in Fig. 4A, showing the three largest percolation modules identified just before the first percolation jump in the subject #1 shown in Fig. 1B at $p = 0.98$. For this connectivity strength, the modules are separated and show submodular fractal structure indicated in the colored boxes obtained with box-covering. When we lower the threshold to $p = 0.975$ (Fig. 4B) the modules are now connected and a global incipient component starts to appears. A second global percolation-like transition appears in the system when the mass of the largest component occupies half of the activated area (see, e.g., Fig. 1). For different individuals, global percolation occurs in the interval $p = [0.945, 0.996]$ as indicated in Fig. 1A Inset.

Our goal is to investigate whether the weak links shortcut the network in an optimal manner. When the cumulative probability distribution to find a Euclidean distance between two connected nodes, $r_{ij}$, larger than $r$ follows a power law,

$$P(r_{ij} > r) \sim r^{-\alpha+1},$$  

[7]

statistical physics makes precise predictions about optimization schemes for global function as a function of the shortcut exponent $\alpha$ and $d_f$ (25, 44, 45). Specifically, there are three critical values for $\alpha$, as shown schematically in Fig. 4C. If $\alpha$ is too large then shortcuts will not be sufficiently long and the network will behave as fractal, equal to the underlying structure. Below a critical value determined by $\alpha < 2d_f$ (25), shortcuts are sufficient to convert the network in a small world. Within this regime there are two significant optimization values:

i. **Wiring cost minimization with full routing information.** This considers a network of dimension $d_f$, over which shortcuts are added to optimize communication, with a wiring cost constraint proportional to the total shortcut length. It is also assumed that coordinates of the network are known (i.e., it is the shortest path that it being minimized). Under these circumstances, the optimal distribution of shortcuts is $\alpha = d_f + 1$ (45). This precise scaling is found in the US airport network (46), where a cost limitation applies to maximize profits.

ii. **Decentralized greedy searches with only local information.** This corresponds to the classic Milgram’s “small-world experiment” of decentralized search in social networks (44), where a person has knowledge of local links and of the final destination but not of the intermediate routes. Under these circumstances, which also apply to routing packets in the Internet, the problem corresponds to a greedy search, rather than to optimization of the minimal path. The optimal relation for greedy routing is $\alpha = d_f$ (25, 44).

Hence, the analysis of $P(r_{ij} > r)$ provides information both on the topology of the resulting network and on which transport procedure is optimized. This distribution reveals power-law behavior Eq. 7 with $\alpha = 3.1 \pm 0.1$ when averaged over the modules below $p_c$ (Fig. 4D). Given the value obtained in Eq. 1, $d_f = 2.1$, this implies that the network composed of strong and weak links is small-world ($\alpha < 2d_f$) (25) and optimizes wiring cost with full knowledge of routing information ($\alpha = d_f + 1$) (45).

**Discussion**

The existence of modular organization that becomes small world when shortcut by weaker ties is reminiscent of the structure found to bind dissimilar communities in social networks. Granovetter’s work in social sciences (30, 31) proposes the existence of weak ties to bind dissimilar communities in social networks. Granovetter’s work in social sciences (30, 31) proposes the existence of weak ties to bind dissimilar communities in social networks. Granovetter’s work in social sciences (30, 31) proposes the existence of weak ties to bind dissimilar communities in social networks. The observation of such an organization in brain networks suggests that it may be a ubiquitous natural solution to the puzzle of information flow in highly modular structures.
Over the last decades, wire length minimization arguments have been used successfully to explain the architectural organization of brain circuitry (47–51). Our results are in agreement with this observation, suggesting that simultaneous optimization of network properties and wiring cost might be a relevant principle of brain architecture (see SI Text). In simple words, this topology does not minimize the total wire per se, simply to connect all the nodes; instead, it minimizes the amount of wire required to achieve the goal of shrinking the network to a small world. A second intriguing aspect of our results, which is not usually high-lighted, is that this minimization assumes that broadcasting and routing information are known to each node. How this may be achieved—what aspects of the neural code convey its own routing information—remains an open question in neuroscience.

The present results provide a unique view by placing modularity under the three pillars of critical phenomena: scaling theory, universality, and renormalization groups (52). In this framework, brain modules are characterized by a set of unique scaling exponents, the setuplet \( (d_1, d_2, d_3, d_4, d_5, d_6, d_7, d_8) \approx (2.1, 1.9, 1.5, 0.5, 1.9, 2.1, 1.3, 1.1) \), and the scaling relations \( d_5 = d_8 - d_7 \), relating fractality with modularity: \( d_7 = 1 + 1 \), relating global integration with modularity; \( \gamma = 1 + d_6/d_7 \), relating scale-free with fractality; and \( \epsilon = 2 + d_6/d_7 \), relating degree correlations with fractality.

One advantage of this formalism is that the different brain topologies can be classified into universality classes under RG (52) according to the setuplet \( (d_1, d_2, d_3, d_4, d_5, d_6, d_7, d_8) \). Universality applies to the critical exponents but not to quantities like \( \langle p_1, C_\alpha \rangle \), which are sensitive to the microscopic details of the different experimental situations. In this framework, noncritical small worlds are obtained in the limit \( (d_1, d_2, d_3, d_4, d_5, d_6, d_7, d_8) \to (\infty, \infty, \infty, 0.0, \infty) \). A path for future research will be to test the universality of the setuplet of exponents under different activities covering other areas of the brain (e.g., the resting-state correlation structure (53)).

In conclusion, we propose a formal solution to the problem of information transfer in the highly modular structure of the brain. The answer is inspired by a classic finding in sociology: the strength of weak ties (30). The present work provides a general insight into the physical mechanisms of network information processing at large. It builds up on an example of considerable relevance to natural science, the organization of the brain, to establish a concrete solution to a broad problem in network science. The results can be readily applied to other systems—where the coexistence of modular specialization and global integration is crucial—ranging from metabolic, protein, and genetic networks to social networks and the Internet.

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Supporting Information

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SI Text

fMRI methods and network construction. A total of 16 participants (7 women and 9 men, mean age 23, ranging from 20 to 28) were asked to perform two tasks with the instruction that they had to respond accurately and fast to each of them. The first task was a visual task of comparing a given number (target T1) to a fixed reference, and the second task was an auditory task of judging the pitch of an auditory tone (target T2) (1). The two stimuli are presented with a stimulus onset asynchrony (SOA); i.e., the delay in the onset of T1 and T2, varying from: SOA = 0, 300, 900, and 1,200 ms. In the number-comparison task, a number varying randomly among four values (28, 37, 53, 62) was flashed on a computer screen and subjects had to respond, with a key press using the right hand, whether the number was larger or smaller than 45. In the auditory task, subjects had to respond whether the tone was high or low frequency with a key press using the left hand. Full details and preliminary statistical analysis of this experiment have been reported in ref. 1. The study is part of a larger neuroimaging research program headed by Denis Le Bihan and approved by the Comité Consultatif pour la Protection des Personnes dans la Recherche Biomédicale, Hôpital de Bicêtre (Le Kremlin-Bicêtre, France).

Subjects performed a total of 160 trials (40 for each SOA value) with a 12 s intertrial interval (2). The 160 trials were performed in five blocks of 384 s with a resting time of approximately 5 min between blocks. For each trial, we recorded whole-brain functional magnetic resonance imaging (fMRI) images at a sampling time, TR = 1.5 s producing eight fMRI images between two consecutive trials. From these images we computed the phase and amplitude of the hemodynamic response of each trial as a sequence, inversion time $T_1 = 5$ min between blocks. For each trial, we recorded whole-brain functional images sensitive to blood oxygenation level-dependent (BOLD) contrast were obtained with a fMRI system (Bruker). Functional images were acquired in 24 slices with a slice thickness of 5 mm. High-resolution images (three-dimensional gradient echo inversion-recovery sequence [repetition time (TR) = 1.5 s; echo time = 40 ms; angle = 90°; field of view (FOV) = 192 × 256 mm; matrix = 64 × 64]). The whole brain was acquired in 24 slices with a slice thickness of 5 mm. High-resolution images were also acquired.

To estimate the periodicity and phase of the event-related BOLD response, the data from each subject were submitted to a nonparametric bootstrap analysis. The correlation calculations, we performed the bootstrap analysis for each possible pair of voxels. The correlation between two voxels for each of those trials serves as our original sample of 40 correlation values. We then draw 10,000 resamples from this sample with substitution. The arithmetic mean is calculated for each resample. Calculating the average value of all these means gives the bootstrap estimate for the mean correlation. The 95% confidence interval for the mean correlation is calculated by the distribution of the 10,000 mean values at the 0.05 and 0.95 points of the distribution, respectively.

The above process yields the confidence interval for the correlation value between two voxels. A different pair of voxels may have very different value of correlation, so in Fig. SI we present the 95% bootstrap confidence interval as a function of the average value of correlation. The interval becomes smaller (i.e., the accuracy of the calculation increases) for larger $p$ values. Considering the networks of Fig. 4A and B, for example, the 95% confidence intervals for $p = 0.975$ and $p = 0.98$ correspond to (see Inset) [0.9744, 0.9760] and [0.9795, 0.981], respectively. The 95% confidence interval becomes more narrow for higher $C_{ij}$ values The corresponding standard deviation is of the order of 0.003. Thus, we typically distinguish between values that differ by 0.005.

By construction, the elements satisfy $-1 \leq C_{ij} \leq 1$, where $C_{ij} = 1$ corresponds to perfect correlations, $C_{ij} = -1$ corresponds to perfect anticorrelations, and $C_{ij} = 0$ describes a pair of uncorrelated voxels. The entire experimental dataset is available at http://lev.ccnx.cuny.edu/~hmakse/brain.html.

For our analysis, we create a mask where we keep voxels which were activated in more than 75% of the cases, i.e., in at least 48 instances out of the 64 total cases considered. The obtained number of activated voxels is $N \approx 60,000$, varying slightly for different individuals and stimuli. The “activated or functional map” exhibits phases consistently falling within the expected response latency for a task-induced activation (1). As expected for an experiment involving visual and auditory stimuli and bimanual responses, the responsive regions included bilateral visual occipito-temporal cortices; bilateral auditory cortices; motor, premotor, and cerebellar cortices; and a large-scale bilateral parieto-frontal structure, see section Spatial Projection of the Modules below. In the present analysis, we do not explore the differences in networks between different conditions. Rather, we consider them as independent experiments, generating a total of 64 different networks, one for each condition of temporal gap and subject.

The use of fMRI neighboring voxels can be expected to carry some shared signal due to spatial autocorrelations (vascular, subject motion, or scanner noise), which could give rise to spurious correlations over short distance. To test for this effect, we double the lattice spacing, increasing the voxels volume by a factor of 8, and repeat the calculations. The results are consistent with the percolation picture of Fig. 1, albeit with a lower $p_c$, while the main results on long-range links are insensitive to this type of artifacts.

Bootstrap Analysis to Determine the Accuracy of $C_{ij}$. In order to estimate the accuracy of the correlation calculations, we performed a nonparametric bootstrap analysis. We consider the set of the 40 trials per subject and SOA value. We perform the bootstrap analysis for each possible pair of voxels. The correlation between two voxels for each of those trials serves as our original sample of 40 correlation values. We then draw 10,000 resamples from this sample with substitution. The arithmetic mean is calculated for each resample. Calculating the average value of all these means gives the bootstrap estimate for the mean correlation. The 95% bootstrap confidence interval is calculated by the distribution of the 10,000 mean values at the 0.05 and 0.95 points of the distribution, respectively.

We calculate cross-correlations between different brain areas based on these phases (3–5). We determine the equal-time cross-correlation matrix $C$ with elements $C_{ij}$ measuring the cross-correlation between the phase activity $\phi_i(t)$ of the $i$th and $j$th voxel over $T = 40$ trials for each subject and SOA condition:
Spatial Projection of the Modules. The complex network representation reveals functional links between brain areas, but cannot directly reveal spatial correlations. Because voxels are embedded in real space, we also study the topological features of modules in three dimensions, where now voxels assume their known positions in the brain and links between them are transferred from the corresponding network (i.e., they are assigned according to the degree of correlation between any two voxels, Eq. S1), which is independent of the voxels proximity in real space. The above procedure yields a different spatial projection of the modules for each subject; an example for subject #1 and SOA = 900 ms in the medial-occipital cortex is shown in Fig. 1D. We study each of these percolation modules separately and find that they all carry statistically similar patterns. The topography of the identified modules reflects coherent patterns across different subjects, as shown next.

Fig. S2A shows a medial sagittal view of the largest four percolation modules for all the participants under stimulus SOA = 0. In virtually all subjects, we observe a module covering the anterior cingulate (AC) region, a module covering the medial part of the posterior parietal cortex (PPC), and a module covering the medial part of posterior occipital cortex (area V1/V2), along the calcarine fissure.

We measure the likelihood that a voxel appears in the largest percolation module among all the participants in Fig. S2A by counting, for each voxel, the number of individuals for which it was included in one of the four percolation modules. The spatial distribution of the first percolation modules averaged over all the subjects depicted in Fig. S2 B and C shows that modules in the three main modes, V1/V2, AC, and PPC, are ubiquitously present in percolation modules and, to a lesser extent, voxels in the motor cortex (along the central sulcus) are slightly more predominant on the left hemisphere. The correlation networks obtained from each subject yield modules with consistent topographic projections.

Box-Covering Algorithm MEMB for Fractal Dimension in Network Space. For a given percolation module, the detection of submodules or boxes follows from the application of the box-covering algorithm for self-similar networks (6, 7). The algorithm can be downloaded at http://lev.ccny.cuny.edu/~hmakse/brain.html. In box-covering, we assign every node to a box or submodule, by finding the minimum possible number of boxes, \( N_B(\ell_B) \), that cover the network and whose diameter (defined as the maximum distance between any two nodes in this box) is smaller than \( \ell_B \).

We implement the maximum excluded mass burning (MEMB) algorithm from ref. 7 for box-covering. The algorithm uses the basic idea of box optimization, where we require that each box should cover the maximum possible number of nodes, and works as follows: We first locate the optimal “central” nodes, which will act as the origins for the boxes. This is done by first calculating the number of nodes (called the mass) within a distance \( r_B \) from each node. We use \( \ell_B = 2r_B + 1 \). The node that yields the largest mass is marked as a center. Then, we mark all the nodes in the box of this center node as “tagged.” We repeat the process of calculating the mass of the boxes starting from all noncenter nodes, and we identify a second center according to the largest remaining mass, while nodes in the corresponding box are tagged, and so on. When all nodes are either centers or tagged, we have identified the minimum number of centers that can cover the network at the given \( r_B \) value. Starting from these centers as box origins, we then simultaneously burn the boxes from each origin until the entire network is covered; i.e., each node is assigned to one box we call this process burning because it is similar to burning algorithms developed to investigate clustering statistics in percolation theory (8, 9)]. In Fig. 2A, we show how box-covering works for a simple network at different \( \ell_B \) values. RG is then the iterative application of this covering at different \( \ell_B \).

Statistical Analysis for the Exponents Calculation Based on Maximum Likelihood Methods and Bootstrap Analysis. In Fig. 2D of the main text, we show an aggregate average of the degree distributions for all clusters. This curve exhibits the general trends of the \( P(k) \) distribution, demonstrating, for example, the heavy tail.

Here, we present a rigorous statistical test based on maximum likelihood methods and bootstrap analysis. We follow ref. 10 for maximum likelihood estimator for discrete variables. We study the properties of 192 network clusters, as described in the main text. The calculation of the scaling exponents is done separately for each network. The resulting set of 192 values is then analyzed through nonparametric bootstrap analysis, in order to get the average value of the exponent and the corresponding confidence intervals.

Some of the distributions are shown in Fig. S3 for nine different clusters, together with the best fittings, in logarithmic and semilogarithmic plots. We fit these distributions assuming a power law within a given interval. For this, we use a generalized power-law form

\[
P(k; k_{\min}, k_{\max}) = \frac{k^{-\gamma}}{\xi(\gamma, k_{\min}) - \xi(\gamma, k_{\max})},
\]

where \( k_{\min} \) and \( k_{\max} \) are the boundaries of the fitting interval, and the Hurwitz \( \xi \) function is given by \( \xi(\gamma, x) = \sum_{i=0}^{x} i^{-\gamma} \).

We use the maximum likelihood method, following the rigorous analysis of Clauset et al. (10). The fit was done in an interval where the lower boundary was \( k_{\min} \). For a given \( k_{\min} \) value we fix the upper boundary to \( k_{\max} = wk_{\min} \), where \( w \) is a parameter. We calculate the slopes in successive intervals by continuously increasing \( k_{\min} \) and varying the value of \( w \) from 4 to 10. In this way, we sample a large number of possible intervals. For each one of them, we calculate the maximum likelihood estimator through the numerical solution of

\[
\gamma = \arg\max \left( -\gamma \sum_{i=1}^{N} \ln k_i - N \ln \xi(\gamma, k_{\min}) - \xi(\gamma, k_{\max}) \right).
\]

where \( k_i \) are all the degrees that fall within the fitting interval, and \( N \) is the total number of nodes with degrees in this interval. The optimum interval was determined through the Kolmogorov–Smirnov (KS) test.

For the goodness-of-fit test, we use the Monte Carlo method described in ref. 10. For each possible fitting interval, we generate 10,000 synthetic random distributions following the best-fit power law. We then calculate the value of the KS test for each one of them and measure the fraction \( p_{\text{KS}} \) of realizations where the real data KS value was smaller than the synthetic KS value. We accept the power-law hypothesis when this ratio was larger than \( p_{\text{KS}} > 0.2 \). The average ratio over all clusters that were retained is \( p_{\text{KS}} = 0.65 \). The final exponent is the average of the individual exponents. Standard bootstrap analysis on the resulting set of the individual cluster values yielded the exponent \( \gamma = 2.1 \pm 0.1 \), with a 95% confidence interval \[2.039, 2.178\].

The same analysis is performed to test for a possible exponential distribution to describe the data. We scan the same intervals as for the case of power law and we use the maximum likelihood method to determine the optimum exponential fitting to the form:

\[
P(k; k_{\min}, k_{\max}) = \frac{1 - e^{-k/k_{\min}}}{e^{-k/k_{\min}} - e^{-k/k_{\max}}} e^{-\lambda k}.
\]

We use KS statistics to determine the optimum fitting intervals and also the goodness-of-fit. In all the cases where the power law was accepted, the exponential fitting gave an average ratio
of \( p_{0i} = 0.017 \), which rules out the possibility of an exponential distribution.

**Correlation Function Dependence on the Distance Between Voxels.**

The embedding of scale-free networks in a finite-dimension real space constitutes a problem that has attracted recent attention (11–13). Scale-free networks may arise from a two-dimensional lattice with added dense connectivity locally, where the weights and connectivity are inversely proportional to the Euclidean distance on the lattice. To investigate this possibility, we study the correlation function of the phases of the voxels as a function of Euclidean distance in real space: \( C(r) = \langle \cos(\theta_i - \theta_j) \rangle \) versus \( r = r_i - r_j \). This function can be interpreted as the correlation between two spins with orientation determined by the phase \( \theta_i \) of the voxel at location \( r_i \) (average is over all pairs at distance \( r \)). We find (Fig. S5) that \( C(r) \) decays algebraically with distance. Thus, our results indicate that modules are scale-free networks that can be embedded in a lattice with an added long-range connectivity.

**Scaling Analysis.**

The structure of a fractal network can be characterized by a set of scaling exponents. They define the scaling of many important system properties. Some of these properties and the corresponding exponents are as follows:

- i. The degree distribution: \( P(k) \sim k^{-\gamma} \), where \( \gamma \) is the degree exponent (14).
- ii. The scaling of the mass with size: \( N_B \sim \epsilon_B^{d_B} \), which defines the fractal exponent \( d_B \) (6).
- iii. The degree-degree distribution \( P(k_1, k_2) \sim k_1^{\epsilon_1} k_2^{\epsilon_2} \), where \( \epsilon_1 \) is the degree-degree exponent, and can be measured through \( E_B(k) \sim k^{-\gamma} \), which is the integration of \( P(k_1, k_2) \) over \( k_2 \) (15).
- iv. The probability that modules are connected through their hubs, \( \Phi \sim \epsilon_B^{d_B} \), defines the hub-hub exponent \( d_B \) (6).
- v. The scaling of the degree of the modules with the size of the modules: \( s \sim \epsilon_B^{d_B} \), which defines the \( d_E \) exponent (6).
- vi. The scaling of the modular factor as defined in Eq. S3: \( \Phi(\epsilon_B) \sim \epsilon_B^{-\epsilon_B} \), through the modularity exponent \( d_M \) (16, 17).

Scaling theory then defines precise relations between the exponents valid for fractal scale-free networks:

- vii. \( r = 1 + d_B/d_E \) (6).
- viii. \( \epsilon = 2 + d_E/d_E \) (15), and
- ix. \( d_M = d_B - d_E \) (16, 17).

We have measured directly all the exponents (see Fig. 2 and Figs. S4 and S6) for the brain modules and find: \( \gamma = 2.1 \pm 0.1 \), \( d_B = 0.51 \pm 0.08 \), \( d_E = 1.9 \pm 0.1 \), \( d_E = 1.5 \pm 0.1 \), \( \epsilon = 2.1 \pm 0.1 \), \( d_M = 1.9 \pm 0.1 \). Using these values in the scaling relations above, we predict \( \gamma = 2.25 \pm 0.11 \) and \( \epsilon = 2.34 \pm 0.06 \), which are within error bars of the calculated exponents \( \gamma = 2.1 \) and \( \epsilon = 2.1 \) from the direct measures. This set of results gives support to a scale-free fractal morphology of the brain modules. Notice that a Euclidean 2D lattice would be obtained in the limit \( \gamma \to \infty \), \( d_B = 0 \), \( \epsilon \to \infty \).

**Modularity Analysis.**

In the main text of the paper, we have described our modularity analysis of the brain clusters according to the Memb technique. The modular properties of the same clusters can be also analyzed through techniques that partition a network according to maximization of modularity. We employed the Girvan–Newman method (18), which locates the point where the modular character of the clusters. The use of the Memb, though, provides us with the extra advantage of modifying the scale at which we observe the modules to determine whether the modular structure is scale-invariant; i.e., if it is composed of modules inside modules at all scales.

**SI Discussion**

Minimizing wire length is in fact of paramount importance, because about 60% of the cortical volume is taken up by wire (axons and dendrites) (19). This turns out to optimize conduction rate, placing a strict packing limitation of the amount of wire in cortical circuits (20). Our finding of a distribution of weak links that minimizes wiring cost is hence in line with a previous literature, consistently showing that neural circuit design is under pressure to minimize wiring length. However, some important nuances of the specific optimization procedure ought to be considered. First, we specifically showed that at the mesoscopic scale, shortcut distribution optimizes wiring cost while maintaining network proximity. This is consistent with the organization of large-scale neural networks in which total wiring can in fact be decreased by about 32% (in 95 primate cortical areas) and up to 48% in the global neuronal network of the nematode C. elegans (20). This extra wiring cost comes from long-range connections that achieve network benefits of shortening the distance between pairs of nodes (20).

**BOLD-fMRI** is an indirect measure of brain activity that relies on multiple vascular and biophysical factors that couple the neural response to the haemodynamic signal (21). Even if in fMRI research it is always assumed that haemodynamic signals reflect metabolic demand generated by local neuronal activity, recent studies have shown reliable haemodynamic signals that entrain to task structure independently of standard neural predictors of haemodynamics (22). Hence, our results, as any other fMRI analysis, may partly reflect the underlying structure of vascular motives. Specifically, the human cortical vascular system has a large number of arterial anastomoses that show a seemingly looking fractal structure in the millimeter to centimeter range (23). Precise measurements of fractality have been reported at the micrometer scales in volumes of the order of a few cubic millimeters (24, 25), which corresponds to approximately a voxel volume, where branching structure of microcapillaries then generates fractals. Hence, it is possible that the fractal organization of brain modules is inherited from the vascular system itself.

Although we cannot readily test the influence of the vascular system at a large scale, it is still possible to address this concern at a microscopic scale, by discarding neighboring correlations. Neighboring voxels are expected to carry some shared signal due to spatial autocorrelations from the microvascular network. To assure that our results do not rely on neighboring correlations that might be particularly spurious, we coarse-grained the original fMRI signal by doubling the lattice spacing, reducing the number of voxels by a factor of 8, and repeating the calculations. The results are consistent with the percolation picture of fractal modules, albeit with an expected lower \( p_c \). Such a renormalized \( p_c \) is expected from renormalization theory to change under coarse-graining, while the main results on long-range links, such as the value of the exponents, are insensitive to this type of coarse-graining.
We also investigate whether the map of fractal dimension $d_F$ reflects a meaningful organization based on known facts of functional properties of the cortex and the specific task that subjects are performing. We found a topographical organization of fractality in the human brain (Fig. S8). The right portion of the anterior cingulate, supplementary motor area (SMA), and the right posterior parietal cortex (PPC) regions involved in routing of information and cognitive control, which are expected to have a more complex functional organization, are the clusters with higher fractal dimension. The left-right asymmetry is interesting because, in this specific task, the left-hand response is queued for a few hundred milliseconds and has to be temporally connected to working memory and inhibitory circuits. Although not fully conclusive, this analysis suggests a functional role of the network architectures described here.

Another similar concern is that the recovered brain modules may be a manifestation of the fractal structure of the underlying three-dimensional vortex grid or of the cortex. However, because the dimensions of the grid ($d = 3$) and of the cortex ($d = 2.7$) (26) are both sufficiently different from 1.9 and the connectivity distribution of the modules is much broader than the typical Euclidean fractal cortex (which should be narrow around $k \sim 6$) or a 3D lattice ($k = 6$), we may safely assume that these objects have their own structure. Moreover, we also observed modules with similar fractal dimension in subcortical structures, suggesting that these results do not simply reflect anatomical properties of the cortical mantle.

Fig. S1. Bootstrap analysis. The interval between the two curves corresponds to the 95% confidence interval for the calculation of the mean fraction of links ($\rho$) as a function of ($p$). The Inset zooms in the regime around the values used in Fig. 4A.

**Fig. S2.** The emerging modules have consistent spatial projections. (A) Spatial distribution of the four largest percolation modules (yellow, orange, red, brown) appearing at the first percolation jump, $p_c$, for each subject under stimulus SOA $= 0$. Most modules are localized in the same regions: anterior cingulate, posterior medial-occipital, posterior parietal, and thalamus. (B and C) These panels show the number of times that the largest percolation cluster for each of the 16 subjects appears in a given voxel. White bleached regions correspond to voxels that are active in the 16 subjects, while the red regions correspond to voxels shared by half of the subjects. The anterior cingulate, a fundamental node in cognitive control, is the only region shared by all subjects.
Fig. S3. Degree distribution for network clusters. A number of degree distribution functions $P(k)$ are shown for different clusters. The red lines correspond to the best power-law fitting, and the blue ones to an exponential fitting. (A) Degree distribution $P(k)$ in logarithmic axes. The power-law slopes correspond to the exponent $\gamma$, and are shown on the plots. (B) The same distributions and fittings in semilogarithmic axes.

Fig. S4. (A) Degree distribution averaged over all the brain modules. The individual degree distributions for each module are shown in Fig. S3. (B) Dependence of the scaling factor $s(\ell_B)$, defined through $k' = s(\ell_B)k$ for the renormalized degree $k'$, on $\ell_B$. The exponent $d_k = 1.5$ characterizes how the node degree changes during the renormalization process.
Fig. S5. Spatial correlation function. This function measures the correlation $C(r)$ between the phase of two voxels that are at a Euclidean distance $r$ apart, as a function of $r$. As shown in the Inset, it decays as a power law.

Fig. S6. Calculation of the scaling exponents. (A) Hub-hub exponent $d_\alpha$ through the scaling of $\mathcal{E}(\ell_B)$. (B) Degree-degree exponent $\epsilon$ through the dependence of $E_b(k)$ on the degree $k$ (15).

Fig. S7. Modular properties of the brain clusters. Comparison between the partition provided by the MEMB method (at $\ell_B = 15$) with the corresponding partition using the Garvin–Newman method (11). The modularity index from the Newman definition $Q$ is around 0.82, as found by the latter method. Both methods yield similar submodules.

Fig. S8. Topographical map of module fractality. For each voxel, we calculate the average fractal dimension of the clusters to which it belongs, considering only voxels that form part of a cluster for at least eight subjects, to assure that mean values are not heavily determined by individual contributions. While the average over all clusters is $d_B = 1.9 \pm 0.1$, the dimension of each cluster exhibits small variations around this value which allows us to identify consistent differences among them. The clusters in the auditory cortex present the smaller fractal dimension $d_B$, while parietal and motor clusters show intermediate values of $d_B$. The right portion of the SMA and the right PPC were the clusters with the higher fractal dimension.