Computational Neuroscience

**LinkRbrain: Multi-scale data integrator of the brain**

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**HIGHLIGHTS**

- The LinkRbrain platform cumulates information from several open databases.
- It integrates these multi-scale data into a common framework.
- It systematically links a set of brain coordinates to a set of cognitive labels.
- It systematically link a genetic expression profile to a set of cognitive labels.
- Based on the topographical overlap, the linkRbrain provides relational graphs.

**ABSTRACT**

**Background:** LinkRbrain is an open-access web platform for multi-scale data integration and visualization of human brain data. This platform integrates anatomical, functional, and genetic knowledge produced by the scientific community.

**New method:** The LinkRbrain platform has two major components: (1) a data aggregation component that integrates multiple open databases into a single platform with a unified representation; and (2) a website that provides fast multi-scale integration and visualization of these data and makes the results immediately available.

**Results:** LinkRbrain allows users to visualize functional networks or genetic expression over a standard brain template (MN152). Interrelationships between these components based on topographical overlap are displayed using relational graphs. Moreover, LinkRbrain enables comparison of new experimental results with previous published works.

**Comparison with existing methods:** Previous tools and studies illustrate the opportunities of data mining across multiple tiers of neuroscience and genetic information. However, a global systematic approach is still missing to gather cognitive, topographical, and genetic knowledge in a common framework in order to facilitate their visualization, comparison, and integration.

**Conclusions:** LinkRbrain is an efficient open-access tool that affords an integrative understanding of human brain function.

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1. Introduction

Sensorimotor and cognitive processing in humans is dependent on the anatomical and functional organization of the brain (Fox et al., 2005; Laird et al., 2011; Mesmoudi et al., 2013). Numerous studies have been conducted that focus on a single level of brain organization such as the anatomical (Catani et al., 2013), the functional (Blumensath et al., 2013) or the genetic level (Hawrylycz et al., 2012). However, to really understand the emergence of human cognition, it is necessary to investigate the relationships between these different scales and study the brain as a complex system. The main challenge is thus the integration of available multi-scale knowledge about brain function. Previous studies have tried to link the expression of certain genes with specific cognitive functions (Zeng et al., 2012; Grange et al., 2014; Hawrylycz et al., 2012; Glahn et al., 2010). In parallel, fMRI studies have related brain hemodynamic activity to cognitive and sensorimotor function. Manual and automated meta-analysis of large databases of neuroscience papers have integrated results into synthetic representations (BrainMap [Laird et al., 2005] and NeuroSynth [Yarkoni et al., 2011], respectively). These methods are based on extraction of fMRI activation peaks (i.e. coordinates) as reported in published articles. They support construction of probabilistic maps of specific cognitive functions.

To visualize and analyze the complex convolutions of the cerebral cortex, the Caret platform provides a powerful approach to quantitatively represent both the consistency and variability in the pattern of convolutions and functional activation from any given task. This tool also allows supports comparisons across species and evaluation of candidate homologies between cortical areas or functionally delineated regions (Van Essen, 2004).

Brainscanner was developed in order to analyze abstracts from scientific papers published in peer-reviewed journals. It measures lexical associations between neuroscientific concepts, then extracts relationships between brain structures, functions, and diseases (Voytek and Voytek, 2010). Other tools such as the Genetic Association Database (GAD) (Becker et al., 2004), GeneCard (Safran et al., 2010), and GeneBank (Benson et al., 2005) collect, standardize, and archive valuable genetic information. The GeneMANIA tool can be used to find other genes that are related to a set of input genes, based on a very large set of functional association data such as protein and genetic interactions, pathways, co-expression, co-localization and protein domain similarity (Warde-Farley et al., 2010).

Recently, the Allen Institute for Brain Science (ABI) released an open access database called the “Allen Human Brain Atlas” (Jones et al., 2009) that contains expression data for 21,000 different genes across 1000 brain regions.

In addition, the Neuroscience Information Framework (NIF) (Aki et al., 2011) provides a unified portal to several neuroscience databases, facilitating access to existing knowledge from neuroimaging, neuroanatomy, cognitive, and genetics databases.

These tools illustrate the opportunities for data mining across multiple sources of neuroscience and genetic information. However, the field is currently lacking a global systematic approach for gathering cognitive, topographical, and genetic knowledge in a common framework that can facilitate visualization, comparison, and integration.

In order to take on the challenge of integrating cognitive, genetic, and anatomical knowledge about brain function, we developed the linkRbrain platform (www.linkrbrain.org). This platform (1) accumulates information from several databases, and (2) integrates these multi-scale data into a common framework so that every point in the brain is characterized by a cognitive profile, a gene expression profile, and a neuroanatomical label. Thus, linkRbrain systematically links: (1) a set of activation peaks over the brain to a set of cognitive labels; (2) a genetic expression profile to a set of cognitive labels; and (3) a set of cognitive labels or genetic expression profile to neuroanatomical labels.

The linkRbrain platform provides brain mapping and relational graphs comprising current available information on brain activity, genetic expression and cognitive functions. This integrative platform is available to the whole community, through an open collaborative website.

2. Data sources

Functional MRI data. LinkRbrain relies on the database of activation peaks generated by the Neurosynth framework (Yarkoni et al., 2010). The version of the database used in linkRbrain contains 194,387 activation peaks automatically extracted from over 5000 published neuroimaging papers, with roughly 140,974 coordinates correctly labeled in Talairach or MNI (Talairach and Tournoux, 1993; Evan et al., 1993).

Article abstracts. In order to extract the terms used by authors to describe cognitive tasks in a bottom-up manner, we used the abstracts and titles of over 5000 neuroimaging articles contained in the Neurosynth database.

Gene expression data. The Allen Human Brain Atlas (ABA) (Jones et al., 2009) produced by the Allen Brain Institute (ABI) provides microarray expression profiles of almost every gene in the human genome at hundreds of locations in the brain. Two complete postmortem brains (H0351.2001 and H0351.2002) are available. Genetic profiles of the two brains are highly compatible (Hawrylycz et al., 2012). LinkRbrain used the H0351.2001 ABA, which reports the genetic profiles for a set of 947 samples, representing the structures within the human brain in approximate proportion to the volumetric representation of each cortical, subcortical, cerebellar, and brain stem structure. This first version of linkRbrain supports visualization of about 21,000 genes expression profiles (as studied in (Hawrylycz et al., 2012)) across the complete set of 947 brain regions.

The H0351.2001 ABA dataset contains about 451 cortical and 496 subcortical regions. To avoid driving results that could be induced by the over-sampling of the subcortical structure, the first version of linkRbrain focuses exclusively on human cortical organization. Hence, the graphs used to quantify gene/cognition overlap and gene/gene overlap take into account only the cortical samples (451 regions).

Neuroanatomical data. We used the 3D Talairach Atlas (Lancaster et al., 2000) to label neuroanatomical structures. This atlas was created from the reference images using resampling following the x and y axis and nearest neighbor interpolation in the z direction. The 3D Talairach Atlas is available as a NIfTI image “Talairach.nii”, where every voxel refers to one of the area labels.

3. Methods

3.1. Text mining

Two types of relations link the sensorimotor/cognitive tasks. First, lexical relations can be directly extracted from the text of articles, thereby providing an overview of the relations between cognitive domains as conceptualized by the scientific community as a whole. Second, topographical relations can be estimated by quantifying the spatial overlap between task-related activations for different functions. In this work, we focus on topographical
relations between cognitive networks, genetic expression profiles, and neuroanatomical structures.

Cognitive task labels. We used the lexical extraction capabilities of the CorText platform to identify a set of pertinent terms. Text processing involves both grammatical\(^2\) and statistical (Kageura, 1996; Frantzi and Ananiadou, 2000) phases to automatically extract candidate n-grams (also called multi-tems).\(^3\) We analyzed the textual content found in titles and abstracts of the 5000 papers within Neurosynth (corpus). The most pertinent noun phrases in our corpus were then curated by a neuroscientist who manually selected the 300 most frequent sensorimotor/cognitive tasks (e.g. spatial working memory, emotional facial expressions).

\(^2\) The POS-tagging step assigns grammatical tags to each word, which are then exploited to find noun phrases (in the “chunking” step). The NLTK python library (Bird et al., 2009) was used to perform these tasks.

\(^3\) Further information about textual processing, is available in the CorText documentation http://docs.cortext.net/lexical-extraction.
Linking activation peaks and task labels. Neurosynth data and software⁴ were used to generate a meta-analytic reverse inference map (Yarkoni et al., 2011; Poldrack, 2006) for each of the previously extracted n-grams. This map quantifies the degree to which each brain region is preferentially activated in studies tagged with a particular sensorimotor/cognitive task label.

3.2. From probes to expression of genes

Genetic expression profiles of the H0351.2001 Atlas were published by ABI as a matrix of 58000 probes × 947 regions of the brain, and includes a list of correspondences between probes and genes (Jones et al., 2009). The final expression profile of every gene was computed by averaging the corresponding probes.

3.3. Topographical distances

The topographical overlap among cognitive activations, between cognitive activations and gene expression regions, or between cognitive activations/gene expression regions and anatomical structure of brain, is expressed as a distance based on a correlation metric. Each network (corresponding to a cognitive task, gene expression region, or anatomical structures of brain) is a set of points or nodes. Let two nodes (sets of weighted points) A and B:

\[ A = \{ (M_i, \mu_i) \mid i \in [1, m] \} \]
\[ B = \{ (N_j, \nu_j) \mid j \in [1, n] \}, \]

where \( \mu_i \) and \( \nu_j \) are the weights of \( M_i \) and \( N_j \) respectively.

The weights express statistical or genetic expressions values related to the activation peaks or cortical regions respectively.

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⁴ Available at (http://github.com/neurosynth).
The correlation score \( \text{cor}(A, B) \) between the nodes A and B was defined by the formula:

\[
\text{cor}(A, B) = \sum_{d(M_i, N_j) \leq r} \mu_i \cdot v_j \frac{d(M_i, N_j)}{r},
\]

(1)

where \( r \) is the reference radius (10 mm in this case, consistent with the meta-analysis), and \( d(M_i, N_j) \) is the distance between two points \( M_i \) and \( N_j \).

The overlap between the two nodes A and B, was obtained by normalizing the correlation score with their autocorrelation. This overlap was based on the RV coefficient (Robert and Escoufier, 1976):

\[
s(A, B) = \frac{\text{cor}(A, B)}{\sqrt{\text{cor}(A, A) \cdot \text{cor}(B, B)}}
\]

(2)

3.4. Visualization

All the activation peaks extracted from the literature (Yarkoni et al., 2010, 2011), the gene expressions regions (Jones et al., 2009), or anatomical structures of brain (Lancaster et al., 2000) are plotted on a 3D and 2D reconstructed brain from the T1-weighted MNI152 (Evan et al., 1993). 3D visualization allows a global localization of studied networks or regions of brain, while 2D visualization allows more precise location of various anatomical structures of the brain.

Visualization of activation peaks. We use the MAP inference procedure to generate sensorimotor/cognitive networks from the activation peaks (Yarkoni et al., 2010). To better illustrate the spatial distribution and overlapping, only one color was assigned to each network. Every task can be represented in two different ways: as spheres, or as a continuous surface.

Fig. 1 illustrates the spatial distribution of the networks involved in a sensorimotor transformation relevant to speech: the relation between a sound and the motor production of this sound by speech. The observation of this graph suggests a relation between the networks specialized in syllable production, sensory tone processing, and motor control of the lips.

Visualization of expression genes regions. For each gene, we identified 451 regions of cortical and 496 subcortical regions of gene expression. In this version of LinkRbrain, we focused on the cortical region to evaluate the overlap between task-based networks and genetic expression profiles. Plane visualization is anyway available for all the 947 regions that cover the cortical and subcortical brain regions. The expression of each of the 21,000 genes is represented by spheres located at each of the 947 brain samples; the diameter of each sphere is proportional to the intensity of expression of the given gene at this position in the brain (see Figs. 2 and 3).

Visualization of user results. LinkRbrain users can visualize and correlate their neuroimaging results with the sensorimotor-cognitive networks, neuroanatomical regions, or genetic expression regions by loading a NIFTI file or MNI coordinates (using NIFTI files, input coordinate, or text files option). Moreover, by using the input coordinate or text file options, the user can (1) quickly and
easily compare the expression among several genes by mapping only the cortex regions where they are the most expressed, and (2) upload her genetic results or a specific probe from the ABA dataset to correlate it with sensorimotor/cognitive networks.

**Graph-based visualization of correlations.** The topographical overlaps (i.e. correlations) between selected task(s), gene(s), or neuroanatomical regions (current node) and the other cognitive tasks, genes, or neuroanatomical regions (nodes) can be represented as a graph. The graph is computed using the force-directed layout algorithm (Fruchterman and Reingold, 1991). Fig. 4 shows two types of links: (1) colored links connect the current node to another node and reveal that an overlap exists between these two nodes; (2) grey links express the overlap between the other nodes without involving the current node. The thickness of the link is proportional to the magnitude of the correlation (overlap).

**Visualization of correlations with identified neuroanatomical structures.** To identify the regions involved in cognitive activation or genetic expression regions, linkRbrain quantified the overlap (i.e. correlations) between this input and the structures of brain identified in the Talairach atlas. These results can be represented as colored regions in the brain, or a graph with cognitive activation or genetic expression regions (see Fig. 5).

4. **LinkRbrain as a multi-scale, integrative explorer**

4.1. From cognitive functions to functional-anatomical architecture

To separately assess the performance of linkRbrain as an exploratory tool at the cognitive and the topographical scales, and as an integrated multiscale tool, we used the example of the model called “the dual intertwined ring architecture” (Mesmoudi et al., 2013). This model originated from a systematic analysis of anatomical, resting-state (RSN), and task based networks (TBN). According to this model, the human cerebral cortex comprises two large ensembles shaped like two rings. The first ring, called the Visual–Sensorimotor–Auditory ring (VSA ring),

![Brain mapping in 2D 3D](image)

**Fig. 7.** Mapping of the regions where the genes coding for oxytocin receptor (OXTR) and dopamine receptor D5 (DRD5) (in blue and red, respectively) are the most expressed. The regions in purple represent the overlap between the two gene expressions. The corresponding graph shows the sensorimotor/cognitive task-based networks that are localized in the same regions where the genes OXTR and DRD5 are most strongly expressed. Red links connect the DRD5 expressions with the topographically nearest tasks, whereas the blue links connect the OXTR expressions with the topographically nearest tasks. The gray links connect tasks with each other on the basis of their topographical overlaps. (For interpretation of the references to color in this text, the reader is referred to the web version of the article.)
comprises visual, auditory, somatosensory and motor cortices, including intermediate bimodal regions. The second ring, called the Parieto–Temporo–Frontal ring (PTF ring), comprises parietal, temporal, and frontal regions. The VSA ring is continuous and forms a circle around the parietal areas BA 39 and 40, whereas the PTF ring, which is not fully continuous over the cortical mantle, is closed by the long-range association fiber tracts (longitudinal parieto-frontal, arcuate, uncinate, and cingulum) that complete the intertwining. The two rings share a set of common regions mostly localized along the precentral, intraparietal, and superior temporal sulci.

Based on the overlap between the resting state networks and different areas and cortices, we can infer that the first ring (VSA ring) links various sources of auditory, visual and somatomotor information together and to control actual behavior. These interactions are important, not only within each modality, but also for all of their bimodal interactions: between visual and motor (e.g., grasping, reaching, imitation), between auditory and somatomotor information (e.g., recognizing and producing phonemes) and between auditory and visual information (important for communication). To illustrate these functional interactions, we used linkRbrain to build the VSA ring by plotting on the brain the task based networks (TBN) corresponding to these modalities (visual, auditory, and motor) and bimodal interactions (motor–visual, visual–auditory, etc.). As shown in Fig. 6, we plot in blue the activations corresponding to: motor function, pictures, hand gestures, limb, somatomotor stimulation, auditory, etc. These networks are activated when we directly stimulated the visual, motor and auditory cortices. In addition, to obtain the bimodal interactions, we plot activations corresponding to speech, oculomotor, etc.

The PTF ring (in red) interfaces systems dedicated to higher cognitive functions with systems dedicated to emotions, biological needs and rhythms. In Fig. 6 we showed that the PTF ring (in red) was reconstructed functionally by plotting the activations peaks corresponding to functions such as recollection, autobiographical memory, semantic memory, working memory, episodic memory, etc.

4.2. From genetic expression to cognitive functions

As explained in Section 3, the linkRbrain platform computes the topographical overlap on the brain between different cognitive activations, gene expression regions, or between cognitive activations and gene expression regions. This topographical overlap is based on a correlation metric, and is represented by linkRbrain as a graph of topographical correlations. To illustrate these correlations, we compare the gene expression profiles of the Oxytocin receptor (OXTR) (Grillon et al., 2013; Hurlemann et al., 2010), and the Dopamine D5 Receptor (DRD5) (Lak et al., 2014; Sunahara et al., 1991) (see Fig. 7). Results show large overlaps, with an important difference: DRD5 is more highly expressed in the anterior cingulate cortex, and OXTR is more highly expressed in the medial anterior temporal pole. To compare the differential spatial correlations (overlap) with cognitive tasks, linkRbrain provides the graphs of topographical similarities. This graph is obtained from the correlation between regions of expression of the OXTR and DRD5 genes and all the activations corresponding to the 300 cognitives tasks extracted from the 5000 neuro–cognitive papers. We see that both regions of gene expression are related to Reward, but OXTR expression is more strongly associated with overlapping emotional networks, and in particular facial expression (important for social interaction), while DRD5 is more related to autobiographical memory, self, and interoceptive awareness. These results obtained from the overlap between genetic expressions and cognitive networks, are confirmed by the literature. In fact, several studies suggest a modulatory role of oxytocin on amygdala responses to facial expressions (Domanski et al., 2007; Lischke et al., 2012) (see Fig. 4). Conversely, other studies implicate dopaminergic pathways in regulation of neuronal systems associated with reward sensitivity, self-control, and interoceptive awareness (Volkow et al., 2013).

5. LinkRbrain as comparator of brain networks of cognitive functions

The cognitive functions “speech” and “sentences” are considered as two distinct functions. The “speech” function is related to
ICN7 involve visuospatial processing and reasoning, with a strong weighting for tasks such as the mental rotation, and counting or calculation.

Based on correlations between the ICN7 image and each cognitive network in the linkRbrain database (see Section 3), we produced a proximity graph. According to the graph of topographical proximities (see Fig. 9), the volume analyzed corresponds to cognitive functions such as oculomotor, visuospatial working memory, saccades, anti-saccades, arithmetic operation and mental calculation. Theses functions extracted by linkRbrain are quite similar with those described in Laird et al. (2011).

7. Conclusion

The number of shared fMRI databases and genetic studies is continuously increasing thanks to the collective efforts of the scientific community. In this work we exploited several new open access databases, extending their value by integrating them in a common framework: a multi-scale data integrator. In fact, with the linkRbrain platform we can: (1) integrate anatomical, functional, and genetic knowledge, (2) visualize and compare functional networks and/or genetic expression, and (3) make sense of new experimental results produced by the community by comparing them with previously published work. In the future, we will extend the linkRbrain platform to include: (1) the new NeuroSynth database with 9721 studies, (2) fiber tracts data, to extend the present integration of multiscale data, and (3) new data coming from the community of users to extend and improve the existing database of cognitive task networks.

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6. LinkRbrain as a comparator of user results with the literature

Laird et al. reconstructed 20 intrinsic connectivity networks (ICN) from the BrainMap database which archived the peak coordinates and metadata associated with 8637 functional brain imaging experiments (Laird et al., 2011). These experiments were extracted from 1840 publications that reported 69,481 activation locations across 31,724 subjects. The masks of these 20 ICN3. Based on (Laird et al., 2011), we used the ICN7 as example which include dorso-lateral prefrontal (BA 46) and posterior parietal cortices (BA 7).


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