

## Database of NIH grants using machine-learned categories and graphical clustering

**To the Editor:** Information on research funding is important to various groups, including investigators, policy analysts, advocacy organizations and, of course, the funding agencies themselves. But informatics resources devoted to research funding are currently limited. In particular, there is a need for information on grants from the US National Institutes of Health (NIH), the world's largest single source of biomedical research funding, because of its large number of awards (~80,000 each year) and its complex organizational structure. NIH's 25 grant-awarding Institutes and Centers have distinct but overlapping missions, and the relationship between these missions and the research they fund is multifaceted. Because there is no comprehensive scheme that characterizes NIH research, navigating the NIH funding landscape can be challenging.

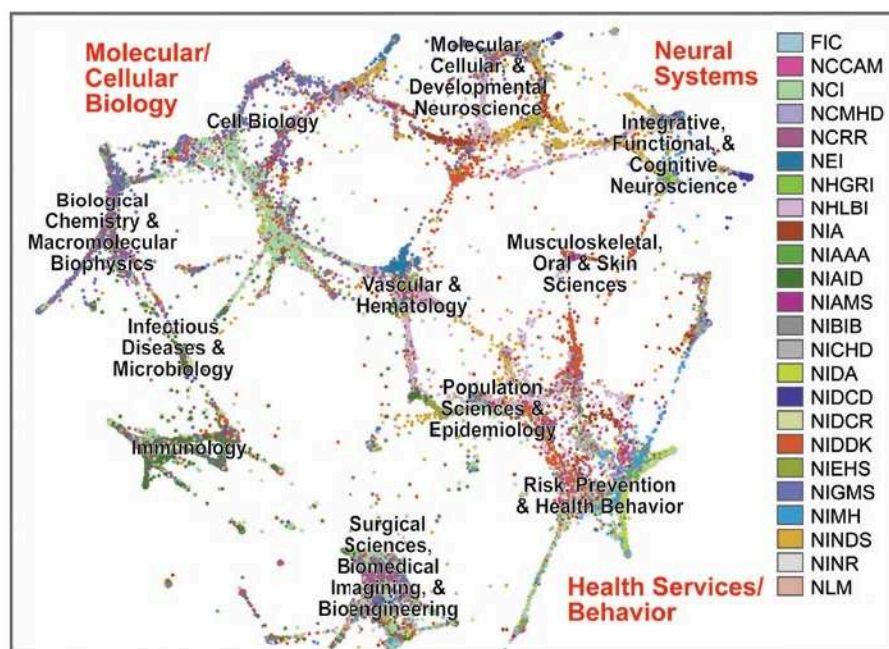
At present, NIH offers information on awarded grants via the RePORTER website (<http://projectreporter.nih.gov/>). For each award, RePORTER provides keyword tags, plus ~215 categorical designations assigned to grants via a partially automated system known as the NIH research, condition and disease categorization (RCDC) process (<http://report.nih.gov/rcdc/categories/>). But keyword searches are not optimal for various information needs and analyses, and the RCDC categories are only intended to meet specific NIH reporting requirements, rather than to comprehensively characterize the entire NIH research portfolio.

To facilitate navigation and discovery of NIH-funded research, we created a database (<https://app.nihmaps.org/>) in which we use text mining to extract latent categories and clusters from NIH grant titles and abstracts. This categorical information is discovered using two unsupervised machine-learning techniques. The first is topic modeling, a Bayesian statistical method that discerns meaningful categories from unstructured text. The second is a graph-based clustering method that produces a two-dimensional visualized output, in which grants are grouped based on their overall topic- and word-based similarity to one another. The database allows specific queries within a contextual

framework that is based on scientific research rather than NIH administrative and categorical designations.

We found that topic-based categories are not strictly associated with the missions of individual Institutes but instead cut across the NIH, albeit in varying proportions consistent with each Institute's distinct mission (**Supplementary Table 1**). The graphical map layout (**Fig. 1**) shows a global research structure that is logically coherent but only loosely related to Institute organization (**Supplementary Table 1**).

We describe four example use cases (**Supplementary Data**). First, we show a query using an algorithm-derived category relevant to angiogenesis (**Supplementary Fig. 1**). Unlike standard keyword-based searches, this type of query allows retrieval of grants that are truly focused on a particular research area. In addition, the resulting graphical clusters reveal clear patterns in the relationships between the retrieved grants and the multiple Institutes funding this research. Second, we examine a NIH peer review study section. The database categories and clusters clarify the complex relationship between the NIH Institutes and the centralized NIH peer review system, which is distinct and independent from the Institutes. Third, we show an analysis of the NIH RCDC category 'sleep research' in conjunction with the database topics, the latter



**Figure 1** | Graphically clustered NIH grants, as rendered from a screenshot of the NIHMaps user interface. NIH awards (here showing grants from 2010; ~80,000 documents) were scored for their overall topic and word similarity, and the resulting document distance calculations were used to seed a graphing algorithm. Grants are represented as dots, color-coded by NIH Institute and are clustered based on shared thematic content. For acronyms and separate views with each Institute highlighted, see the legend for **Supplementary Table 1**. Labels in black were automatically derived from review assignments of the underlying documents. Labels in red indicate a global structure that was reproducible using multiple different algorithm settings.

providing salient categorical information in greater detail than the officially reported category. Finally, we show how the database can be used for unbiased discovery of research trends, and we document the remarkable increase in funding for research on micro-RNA biology from 2007 to 2009. Changes in topics associated with this burgeoning area demonstrate a transition in the nature of the research, from basic cellular and molecular biology to investigations of complex physiological processes and disease diagnoses.

In each case, the machine-learned topics are robustly correlated with funding by specific NIH Institutes, highlighting the importance of the underlying categories to the NIH. The patterns elucidated in this framework are consistent with Institute policies, but obtaining similar information in the absence of the current database would require extensive exploration of Institute websites, followed by time-consuming research on appropriate keywords for queries of specific categories. Our database offers an alternative approach that enables rapid and reproducible retrieval of meaningful categorical information.

To ensure transparent and accurate representations of the algorithm-derived topics, we provide extensive contextual information derived from the documents associated with each topic, in a format conducive to spot checks and to detailed examination for cases requiring precise categorical distinctions. Additionally, we implemented a new technique for automatically assessing topic quality using statistics of topic word co-occurrence (Supplementary Methods), which we used for curating the database to identify poor quality topics.

Our use of this graphing algorithm is somewhat different from previous gene expression analyses and scientometric studies based on journal citation linkages (see Supplementary Methods for references). We assessed the information-retrieval capabilities of the graphs and found that they performed well relative to the document similarity measures that served as inputs. Notably, rather than forming isolated clusters, in this case the algorithm produced a lattice-like structure, in which clusters are linked by strings of aligned documents whose topical content is jointly relevant to the clusters at either end of each string (Supplementary Fig. 1). In addition to providing extra 'subcluster' resolution of content that falls between clusters, this lattice-like framework formed a logical organizational structure, merging the local, intermediate and global levels of the graph.

The categories and clusters represented in this database are comprehensive and thus provide reference points from which various information requirements can be addressed by users with divergent interests and needs. Perhaps more importantly, they provide a basis for discovery of interrelationships among concepts and documents that otherwise would be obscure.

Note: Supplementary information is available on the Nature Methods website.

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The authors declare competing financial interests: details accompany the full-text HTML version of the paper at <http://www.nature.com/naturemethods/>.

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## Predicting protein associations with long noncoding RNAs

**To the Editor:** Only a small fraction of the human transcriptome (~1%) encodes proteins<sup>1</sup>, but a large portion of transcripts is long noncoding RNAs (lncRNAs) and is an unexplored component of mammalian genomes<sup>2</sup>. Here we introduce a method to perform large-scale predictions of protein-RNA associations. Our algorithm, 'fast predictions of RNA and protein interactions and domains at the Center for Genomic Regulation, Barcelona, Catalonia' (catRAPID), evaluates the interaction propensities of polypeptide and nucleotide chains using their physicochemical properties. The algorithm is freely available at [http://big.crg.cat/gene\\_function\\_and\\_evolution/services/catrapid](http://big.crg.cat/gene_function_and_evolution/services/catrapid).

We trained catRAPID on 592 protein-RNA pairs available in the Protein Data Bank to discriminate interacting and non-interacting molecules using only information contained in their sequences (Supplementary Table 1). Secondary structure propensities accounted for 72% of catRAPID ability to predict protein-RNA associations, followed by hydrogen bonding (58%) and van der Waals (26%) contributions. Occurrence of hairpin loops in nucleotide sequences and presence of helical elements in polypeptide sequences positively correlated with interaction propensities. Protein and RNA binding sites had higher interaction propensities than other regions in complexes (Fig. 1a, Supplementary Methods and Supplementary Tables 2 and 3).

We validated our algorithm on a large collection of protein associations with lncRNAs<sup>3</sup>, the NPInter dataset (Supplementary Methods and Supplementary Table 4). Using catRAPID we correctly predicted 89% of experimentally supported interactions linked to physical evidence of binding (Fig. 1b). We observed less significant performance ( $P \sim 0.1$ ) for interactions inferred from indirect evidence (Supplementary Methods). To test catRAPID's ability to identify non-interacting molecules, we generated random lists of RNA associations with proteins involved in DNA and protein-binding (DNA BP and protein BP datasets, respectively; Fig. 1c and Supplementary Table 5). We predicted interactions only for <40% of cases, which suggests that these associations are unlikely to take place (RNA BP dataset; Fig. 1c and Supplementary Table 5). With regard to random associations with RNA-binding proteins, we observed slightly higher interaction propensities (~52%), which indicates occurrence of spurious binding.

To validate the ability of catRAPID to identify binding regions, we analyzed the human ribonuclease mitochondrial RNA processing (MRP) complex<sup>4</sup> (Supplementary Methods). The MRP assembly comprises ten protein subunits: hPop1, hPop5, Rpp14, Rpp20, Rpp21, Rpp25, Rpp29, Rpp30, Rpp38 and Rpp40. We predicted