RESEARCH ARTICLE

An Integrative Computational Approach for Prioritization of Genomic Variants

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Abstract

An essential step in the discovery of molecular mechanisms contributing to disease phenotypes and efficient experimental planning is the development of weighted hypotheses that estimate the functional effects of sequence variants discovered by high-throughput genomics. With the increasing specialization of the bioinformatics resources, creating analytical workflows that seamlessly integrate data and bioinformatics tools developed by multiple groups becomes inevitable. Here we present a case study of a use of the distributed environmental integrating four complementary specialized resources, namely the Lynx platform, VISTA RViewer, the Developmental Brain Disorders Database (DBDB), and the RaptorX server, for the identification of high-confidence candidate genes contributing to pathogenesis of spina bifida. The analysis resulted in prediction and validation of deleterious mutations in the SLC19A placental transporter in mothers of the affected children that causes narrowing of the outlet channel and therefore leads to...
the reduced folate permeation rate. The described approach also enabled correct identification of several genes, previously shown to contribute to pathogenesis of spina bifida, and suggestion of additional genes for experimental validations. The study demonstrates that the seamless integration of bioinformatics resources enables fast and efficient prioritization and characterization of genomic factors and molecular networks contributing to the phenotypes of interest.

Introduction

The identification of genomic variations contributing to specific phenotypes of direct medical relevance is an ultimate goal of numerous studies in human genetics. The development of solid weighted hypotheses on the functional effects of a sequence variant is an essential step for gaining insights into a genetic architecture of a disease and for the efficient planning of experiments. However, as the volume and complexity of biological information increases, it demands sophisticated analytical workflows involving multitude of steps for extraction of actionable knowledge. In the past years much attention in the bioinformatics literature was given to data integration [1–3]. Seamless integration of complementary services and tools provided by multiple groups in workflow pipelines is essential for comprehensive data analysis. It provides the means for substantial reduction of time and effort required for analysis of translational data and significant increase in the efficiency of knowledge extraction.

A number of excellent bioinformatics platforms and tools have been developed in the recent years to support various steps of analysis of high-throughput data and prioritization of genomic variants (reviewed in [4–6]). These include, but not limited to GeneMANIA [7], STRING [8, 9], ToppGene [10], Endeavour [11] widely used by the scientific community. The eXtasy platform developed by Sifrim et al. [12] prioritizes mutations for follow-up validation studies by integrating variant-impact and haploinsufficiency predictions with phenotype-specific information. Another scientific environment, SPRING [13], has been designed to facilitate the prioritization of pathogenic non-synonymous SNVs associated with the disorders whose genetic bases are either partly known or completely unknown. It is achieved by integrating the results of analyses by multiple publicly available and developed in-house bioinformatics tools. There are more analytical platforms, such as Jannovar [14], KGGSqA [15], MToolBox [16] and FamAnn [17].

Moreover, multiple resources support the analysis of non-coding regions and their regulatory roles [18]. Most of these existing resources, understandably, address either the analysis of coding sequences or the characterization of non-coding regions.

The analytical environment described here however is different from these resources. It is based on seamless integration of data and services across multiple independently developed analytical systems and databases, namely the Lynx [19]
and the VISTA [20] systems, the Developmental Brain Disorders Database (DBDB) [21], and the RaptorX server [22, 23]. This environment, depicted in Fig. 1, allows end users to easily direct and analyze their data among all these systems. The benefits of such integration are manifold. They include the integration of the vast knowledge bases developed by each system to support the annotation of the experimental data and the subsequent analyses. Complementary analytical tools and the Web services-based collaborative interfaces provide flexible analytical pipelines seamlessly operating across the participating systems.

As an example, we have demonstrated an ability of the reported pipeline to identify polymorphisms that make plausible candidates for factors contributing to spina bifida (SB), using whole genome next generation sequence (NGS) data for
affected patients and their parents. We show advantages of an integrated approach for both hypothesis-based and discovery-based methods for identification and prioritization of genetic factors contributing to complex developmental phenotypes. The presented example also serves as a proof of concept for the integration of various computational resources for the high-throughput analysis of genomic variants.

**Materials and Methods**

1. **Integrative Analytical Approach**

   We have integrated the following analytical resources developed by four groups: (1) VISTA RViewer [24] for the annotation and comparative and evolutionary analysis of coding and non-coding regions of the genomes; (2) the Lynx platform [19] supporting enrichment analysis and networks-based gene prioritization, (3) the Developmental Brain Disorders Database (DBDB) [21], and (4) RaptorX [23] for predicting 3D structure and functional properties of identified candidate gene products. Combining knowledge bases and knowledge-extraction services into a seamlessly integrated analytical pipeline creates a one-stop solution for generating weighted hypotheses regarding the molecular mechanisms contributing to the phenotypes of interest.

2. **Data submission**

   The approach supports multiple entry points for annotation and analysis of translational data (e.g. genes, pathways, disorders), as well as batch queries via Web-based user interfaces or Web-services (see Fig. 1). The following queries can be submitted to Lynx or VISTA RViewer for annotation or downstream analysis (Fig. 1): (a) single gene queries to Lynx, RViewer or RaptorX; (b) search-based queries to Lynx or VISTA to retrieve information from these systems knowledge bases, (c) batch queries for experimental data in the form of SNPs, genomic coordinates, or gene lists. VISTA RViewer also supports analysis of the Variant Call Format (VCF) files. The results of gene expression analyses in a form of a table containing gene symbol – expression value pair may be directly uploaded into the Lynx Network-based prioritization engine. Lynx Web site provides access to detailed tutorials.

2. **Participating Analytical Resources**

   The sections below will describe the components of the integrated distributed analytical pipeline in more details.

**VISTA Region Viewer (RViewer)**

VISTA RViewer [24], one of the VISTA comparative genomics programs [20] widely adopted by the biomedical community [25–27], employs a new concept of comparative analysis for automating the prioritization of functional variants based on comparative genomics. VISTA RViewer allows for comparison and
prioritization of the entire functional content of several genomic regions in parallel. Examples of such uses could be sets of CNVs regions from experiments, genomic neighborhoods of SNPs from GWAS or other studies, genes of expression studies, etc. RViewer has several functions not found in other currently available tools [28–30], namely it enables the simultaneous comparison of functional features in multiple genomic intervals, i.e. provides capabilities for quicker analysis, prioritization and visual inspection. RViewer takes as input genetic variation data from different biomedical studies (e.g.: GWAS, exomes), and determines a number of functional parameters for both coding and noncoding sequences in each region. Each gene in the region is characterized using the following contexts: (a) biological function; (b) features of protein products of these genes; (c) tissue expression; (d) binding partners (e) developmental stage; (f) pathways and networks information from pathways databases and literature (g) known disease associations; and (h) known genetic variations. For noncoding regions, RViewer provides: homotypic clusters of transcription factor binding sites, a key component of promoters and enhancers [31]; experimentally verified enhancers from the VISTA Enhancer browser [32]; heart- and hindbrain- specific enhancers derived computationally [33, 34]; and conserved TFBS in the promoters of all human genes. In addition, a wide range of comparative genomics data based on pairwise and multiple alignments [35] is accessible.

In the LYNX - VISTA integrative system, RViewer plays a dual role, both calculating a number of functional parameters used by LYNX in further analysis, and visualizing a set of genomic regions with prioritized variant provided to a user as an output of the system. In particular, it finds a genomic position of a submitted by a user variant (coordinate and a specific position in intron, exon, UTRs, intergenic), associates it with UCSC isoforms, calculates deleteriousness of the non-synonymous coding SNPs using Polyphen2 [36], and defines occurrence of SNPs in clusters of TFBS [31], enhancers [32], and highly conserved intervals in inter-species pairwise and multiple alignments. Prioritized genes and other functional features in resulting genomic intervals are displayed in RViewer with all relevant functional content for further interactive exploration by a user.

**Lynx annotation and knowledge extraction engine**

Lynx (http://lynx.ci.uchicago.edu) is an integrated bioinformatics platform and a knowledge extraction engine for annotation and analysis of high-throughput biomedical data [19]. Lynx receives user data as genomic variants whose coding and non-coding signals have been characterized by RViewer (Fig. 1). The platform supports both hypothesis-based and discovery-based approaches to prediction of genetic factors and networks associated with phenotypes of interest. It provides a knowledge extraction engine and a supporting knowledge base (LynxKB) combining various classes of information from over thirty-five public databases and private collections. Lynx knowledge retrieval engine offers advanced search capabilities and a variety of algorithms for gene enrichment analysis and network-based gene prioritization. Lynx’s XML schema-driven annotation service
supports extraction of annotations for an individual object (e.g. a gene) or batch queries (e.g. list of genes) from LynxKB. Annotations include inter alia associated pathways, diseases, phenotypes, molecular interactions, Gene Ontology categories, toxicogenomic information displayed according to the user preferences. All information related to the objects is easily accessible via user interface and available for download in tab-delimited, XML or JSON formats (Web Services).

Lynx gene enrichment analysis supports Bayes factor and p-value estimates for identification of functional categories over-represented in the query data sets (see B. Xie et al. for more details [37]). Lynx enrichment analysis is based on a large variety of features (e.g. Gene Ontology terms, toxicogenomic information, tissues), as well as unique for the system customized brain connectivity ontology, symptoms-level phenotypes and associated non-coding signals from VISTA (e.g. enhancers and clusters of transcription factors binding sites). Lynx also supports context-sensitive enrichment analysis (e.g. against genes expressed on a particular developmental stage or in a particular tissue) that may substantially increase the accuracy of the results.

Additionally, Lynx integrates five network propagation algorithms (simple random walk, heat kernel diffusion, PageRank with priors, HITS with priors and K-step Markov) as initially developed in the gene prioritization tool PINTA [38]. These algorithms were modified for Lynx to replace continuous gene expression data with binary data from seed genes. This modification accommodates the use of a variety of weighted data types for gene prioritization including ranked gene to phenotype associations, weighted canonical pathways, gene expression, results of sequencing analyses and others. STRING v 9.0 [8] is used as the underlying protein interaction network. Networks-based gene prioritization facilitates prioritization of promising candidate genes from large gene sets or even from the entire genome to provide a preliminary step for network reconstruction. Lynx Service Oriented Architecture provides public access to LynxKB and its analytical tools via user-friendly web services and interfaces.

**Developmental Brain Disorders Database (DBDB).** In a lot of cases the analysis of translational data requires integration of the domain- and project specific data (e.g. phenotypic and clinical data, tissue-specific information). Significant amount of this information is already integrated into the Lynx knowledge base. However, the described analytical pipeline can be customized to integrate additional information resources (e.g. public databases and private collections) to meet the user requirements. Here we present the integration of the information from the Developmental Brain Disorders Database (DBDB) [39] as an example of satisfying the needs of neurodevelopmental research.

DBDB ([https://www.dbdb.urmc.rochester.edu/home](https://www.dbdb.urmc.rochester.edu/home)) is a publicly available, curated on-line repository of genes, phenotypes, and syndromes associated with human neurodevelopmental disorders [21]. The discovery of genetic mechanisms contributing to pathogenesis of diseases relies on pre-existing knowledge regarding complex phenotype-genotype relationships accumulated in the libraries of disease candidate genes (e.g. OMIM [40], AutDB [41]), databases describing genetic variations (e.g. GAD [42], SLEP [43]) or journal publications. The
accuracy of these data varies significantly and affects the quality of the developed models. DBDB addresses this issue by employing a novel system for estimating levels of evidence for over 850 gene-phenotype associations curated by domain knowledge experts. While a useful tool for clinical diagnostics, DBDB is also increasingly valuable for annotation of variants identified from next-generation sequencing experiments and the development of predictive computational models for the disorders of interest. Here, DBDB was used for assigning of the levels of confidence to the gene-phenotype associations used for the reconstruction of molecular pathways potentially contributing to pathogenesis of Spina bifida.

RaptorX

RaptorX ([http://raptorx.uchicago.edu](http://raptorx.uchicago.edu)) [23] is a protein structure and function prediction web server, excelling at predicting 3D structures for protein sequences without close homologs in the Protein Data Bank (PDB). Given a query sequence, RaptorX predicts its secondary and tertiary structures as well as solvent accessibility and disordered regions. RaptorX also predicts the binding sites of a protein sequence, based upon the predicted 3D model.

RaptorX applies a template-based approach to protein structure prediction. To deal with cases where no close template exists, RaptorX employs a couple of novel modeling strategies. First, RaptorX integrates a variety of context-specific biological signals in a non-linear probabilistic scoring function by using a powerful machine learning model Conditional Neural Fields (CNF) [44]. Second, RaptorX uses a multiple-template threading (MTT) procedure to significantly improve both alignment and modeling accuracy for some targets with multiple similar templates [45]. The predictions and tertiary structure models produced by RaptorX can serve as starting points for further analysis in a number of diverse application areas. For example, the predicted 3D models used for binding site prediction can also be used for epitope prediction as well as in protein-protein interaction studies. Given a protein sequence, RaptorX predicts its binding sites by aligning its predicted 3D model to a database of ligand-binding protein structures using a structure alignment tool DeepAlign [46].

3. Integrated Environment

The systems described above provide complementary services for identification and characterization of causative factors and molecular mechanisms associated with phenotypes of interest. While they all act as individual entry points for the end users to start the analysis, the implemented integrative tools provide an environment for seamless data transfers between the systems. Following are a few examples of the interfaces built towards integrating these systems:

(i) **Lynx and RViewer.** After entering a list of genes, genetic variants (SNPs) or regions (CNVs) in the Lynx system, the users are provided with the interface to submit these datasets to RViewer for the annotations of the genomic regions including variants, transcription factor binding sites, experimental and computational enhancers and many others. The annotations data for
transcription factor binding sites and enhancers is periodically fetched from the RVViewer and integrated into the Lynx knowledge base and is used for the enrichment analysis of the genes of interest.

(ii) **Lynx and DBDB.** The richly curated data in DBDB make possible providing weights to the genes of interest for particular phenotypes. These data can be used in Lynx’s networks based prioritization interface. DBDB also provides direct links to the Lynx’s single gene and multi-gene annotation pages.

(iii) **Lynx and RaptorX.** Lynx provides a novel interface for submission of protein sequences directly to RaptorX, thus creating capabilities for RaptorX to predict the structure and binding site for high-confidence genes flawlessly.

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**Results and Discussion**

Use Case: Identification of genetic factors contributing to the pathogenesis of spina bifida from whole genome sequencing data using coding and non-coding variant analysis

Spina bifida is a common congenital birth defect with an average worldwide prevalence of one case per 2000 births. It manifests itself as failed closure of the embryonic neural tube \(^{47,48}\) resulting in a breach of several vertebrae leaving spinal cord and/or spinal nerves exposed or covered by only a thin membrane. To date the genetic mechanisms underlying this disorder still remain elusive, but many studies indicate the convergence of gene-gene and gene-environment interactions to produce the defect \(^{49,50}\).

In our study the Laboratory of Neurogenetics and Development (Weil Cornell Medical College) performed the whole genome sequencing for four patients with spina bifida, and four parents with normal phenotype from the same consanguineous family. The resulting SNPs were analyzed using both discovery-based, and hypothesis-based approaches for identification of the genetic factors contributing to the birth defect phenotype. The analysis included the following steps (schematically shown in Fig. 2):

**Step 1. Discovery-based approach: annotation of genomic variants using VISTA RVViewer.** SNPs were filtered based on quality and annotated using VISTA RVViewer \(^{24}\) with the following information: SNP to gene association (UCSC isoforms), deleteriousness of the non-synonymous SNPs using Polyphen2 \(^{36}\), occurrence of SNPs in clusters of TFBS \(^{31}\) and enhancers \(^{32}\), genomic features associated with the region of the variant (intron, exon, UTRs). The Minor Allele Frequency (MAF) was estimated using ANNOVAR \(^{51}\). The resulting annotated SNPs were then filtered based on homozygosity, Minor Allele Frequency (MAF), and deleteriousness of mutations in exonic regions. All of the variants that were homozygous in probands but heterozygous in parents were extracted and used for further analysis to identify de novo mutations in proband but not in the parents. A total of 743 variants in proband genomes were identified, out of which 6 variants were in the exonic region, 10
variants fell into UTRs, 337 variants were in the introns and the rest of the variants were in the intergenic un-annotated regions of the genome.

**Step 2.** Hypothesis driven approach to SNPs prioritization: Defects in folate metabolism and transport were suggested to be contributing factors in pathogenesis of this birth defect [52, 53] however the exact genes associated with this disorder are not known. Here, thirty-nine genes involved in folate metabolism and transport were extracted from the literature [54, 55] and used as seed genes for network-based gene prioritization as implemented in Lynx [38]. STRING 9.0 [8] was used as a global probabilistic network. Network predicted genes with significant p-values <0.005 were used for subsequent analysis.

**Evaluation of the results.** Direct overlap with the known folate metabolism genes as well as the results of the network-based gene prioritization yielded a number of genes with rare variants (MAF=0) in the exonic and UTR regions of both the patients and unaffected parents genes (Table 1 and Table S1 in S1 Materials).
In the recent years the genetic variations in genes involved in the folate-homocysteine metabolism were assumed to be possible risk factors for spina bifida [56–58]. These studies, however, are problematic due to the complexity of epistatic relationships between the affected genes and potential involvement of both the maternal and the offspring genotype in determining the pathogenic effect of these mutations.

As seen in Table 1, a number of patient genes involved in folate biosynthesis contained deleterious mutations. These included folate hydrolase 1 (FOLH1), mesagenin (MSGN1) and solute carrier family 19 folate transporter, member 1 (SLC19A1). The mesagenin 1 (MSGN1 gene) is known to be involved in specification of the paraxial mesoderm and regulation of the expression of T-box transcription factors required for mesoderm formation and differentiation. The mouse mutant of this gene is implicated in spina bifida phenotype. It was also suggested [59] that MSGN1 serves as one of the Wnt target genes in Wnt/β-catenin signaling that plays a well-established role in the regulation of embryonic and adult stem cell homeostasis. Folate hydrolase 1 (FOLH1), also found to contain deleterious mutations in two patients is known to act as a glutamate carboxypeptidase on different substrates, including the nutrient folate and the neuropeptide N-acetyl-l-aspartyl-l-glutamate [60].

### Table 1. Identified genetic variants in folate metabolism genes in spina bifida patients and unaffected parents.

| Gene     | NW P-value* | Identified Variations | Variation Type          | Variation occurrence in patients** | Variation occurrence in parents | Reference of association with SB |
|----------|-------------|-----------------------|-------------------------|------------------------------------|---------------------------------|---------------------------------|
| FOLH1    | 0           | rs61886492            | Missense, possibly damaging | 6C1, 6C2                           |                                 | Morin, Devlin et al. 2003 [58]  |
| DMGDH    | 0           | rs532964              | Missense, possibly damaging | 2C1, 2C2, 6C1, 6C2                 | 2M, 2F, 6M, 6F                  | Marini, Hoffmann et al. 2011 [53] |
| MTR      | 0           | rs1805087             | Missense, possibly damaging | 2C1, 2C2, 6C1                     | 2M, 2F, 6M, 6F                  | Doolin, Barbaux et al. 2002 [59] |
| MSGN1    | 0.002       | rs35858730            | Missense, possibly damaging | 2C1, 2C2, 6C1, 6C2                |                                 | Chalamalasetty, Dunty et al. 2011 [66] |
| CUBN     | 0.002       | rs1801228             | Benign                   | 2M, 6M                             |                                 | Kozyraki, Fye et al. 1999, Wahlstedt-Froberg, Pettersson et al. 2003, Whitehead, 2006 [64, 68, 69] |
| SLC19A   | 0.0028      | rs1051266             | Missense, possibly damaging | 2M, 2F, 6M, 6F                    |                                 | Shaw, Lammer et al. 2002, Morin, Devlin et al. 2003 [58, 67] |

*The P-values in Table 1 are generated by 10 000 random permutations of the input data scored according to the strength of association with the phenotype using DBDB recommendations (random reassignment of the scores to network nodes and computation of the corresponding randomized scores for all candidate genes) [38].

**Family 1: affected children 2C1, 2C2; mother 2M, father 2F. Family 2: affected children 6C1, 6C2; mother 6M, father 6F.

![image](image_url)
Our analysis also identified a number of parental genes containing deleterious mutations. With respect to offspring, the effects of maternal genetic mutations may be considered to be environmental risk factors. Identified parental genes potentially contributing to spina bifida in affected children included CUBN1, MTR, SLC19A and DMGDH genes. Some of these genes were previously found to be associated with spina bifida. Doolin et al. [61], have demonstrated that methionine synthase (MTR) variants influence the risk of spina bifida via the maternal rather than the embryonic genotype. Moreover, in our study both mothers also showed an exonic variant (rs1051266) in the SLC19A1 gene encoding placental solute carrier family 19 folate transporter, member 1 (RFC) as it was demonstrated by RViewer. Polymorphisms in vitamin B receptor (CUBN) and in SLC19A1 (RFC) in mothers identified by networks-based gene prioritization have been previously shown to be associated with spina bifida or other neural tube defects in offspring [62–64] supporting the above inferences obtained in the course of our analysis.

Analysis and validation of the functional impact of the SLC19A1 mutation
The known average distribution of the exonic variant (rs1051266; Arg27His, 80G>A) in SLC19A1 placental folate transporter in general population is 30:25:45 [65]. Further investigation of a potential impact of this variation on function was done using RaptorX. Fig. 3 presents the results of the predictions of SLC19A1 3D structure and the locations of the binding sites by RaptorX.

The reconstruction has shown that SLC19A1 protein has a typical MFS membrane transporter architecture with N- and C-terminus domains containing six trans membrane helices each [66]. It was previously demonstrated [67] that the RFC1 80A-allele is associated with reduced plasma folate. This phenomena could be explained by the reconstructed model in the following way: the 80G>A substitution leads to a change in position 27 on TM1 from histidine (80A allele) that has a relatively medium volume and neutral charge with Arg27 (80G allele) that has a larger volume and positive charge. A variation is located on the transmembrane helix (TM1) close to the intracellular outlet site. Such substitution would likely lead to the narrowing of the outlet channel due to the repulsion of Arg151 on TM5 due to electrostatic repulsion force and therefore to the reduced folate permeation rate (see Fig. 3).

Indeed, an in-silico simulation and energy minimization study of the mutation (see Text S1 in S1 Materials for more details) shows that the area surrounding Arg151 becomes more compact and its contacting residues change after the mutation (see Fig. 4, and Figures S1 and S2 in S1 Materials). This results in 33% decrease in the volume of the cleft (from 807 Å³ to 541 Å³) and 28% decrease in the surface area of the cleft (from 442 Å² to 328 Å²) (see Table 2 for details).

Furthermore, a Pro-kink at Pro146 forces the side-chain of Phe141 to turn, narrowing the channel in front of the cleft (Figures S1 and S2 in S1 Materials). Further docking studies confirm that this narrowing results in a different conformation of binding for the folate (see Fig. 5 and Figures S3 and S4 in S1 Materials).
Annotation of SLC19A1 using Lynx showed that the expression and functionality of this gene is negatively affected by a number of pharmacological compounds, such as indomethacin, phenobarbital, nicotine and vitamin E. Analysis of potential use of these medications by mothers of affected children during pregnancy may provide additional clues regarding high occurrence of spina bifida in the consanguineous family under study.

The described approach has let us to correctly identify CUBN and SLC19A1 genes previously shown to contribute to pathogenesis of spina bifida [62–64] and to suggest additional genes for the next round of experimental validations.

![Ribbon diagram of SLC19A1 protein model generated by RaptorX.](image)

Rainbow coloring from blue to red indicates the N- to C-terminal positions of the residues in the model. The docking location of Folic Acid (FOL), shown in a spacefill form, was predicted by RaptorX-Binding. Numbers in black correspond to key residues, shown in spacefill form, which related to the functional impact of an exonic variant (rs1051266; Arg27His, 80G>A). The diagram was generated using PyMOL.

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Fig. 3. The changes to the binding site caused by the mutation (left: native, right: H27R mutant). The mutation results in changing contact landscape inside the cleft, especially for the Arg151 residue.

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Conclusions

The presented approach is an example of multilevel integration of the bioinformatics resources that offers seamless access to the knowledge bases, analytical tools and user interfaces independently developed by participating groups. Such integration is critically important for the progress of the translational studies since it significantly reduces the time and effort required to efficiently extract knowledge from the exponentially growing data sets produced by numerous genomics projects.

The spina bifida example demonstrates one of the possible analytical scenarios supported by the described computational framework. The presented here integrative approach however, can be generalized to support prioritization of the high-throughput experimental results and prediction of novel candidate genetic factors for any disorder of interest to the user. The power of the approach lies in massive integration of various classes of data from the Lynx (e.g. functional, phenotypic, pathways information), VISTA (e.g. genomic, evolutionary information), and RaptorX (proteomic and structural information). In the spina bifida example, the DBDB knowledge base was used as a source of domain-specific

| Cleft Volume     | Native    | H27R Mutant |
|------------------|-----------|-------------|
| Volume           | 807 Å³    | 541 Å³      |
| Surface area     | 442 Å²    | 328 Å²      |
| Sphericity       | 0.95      | 0.98        |
| Effective radius | 5.47 Å    | 4.94 Å      |

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![Fig. 5. Docked folate conformations (blue: native, red: H27R mutant-bound folate molecules) showing the distinct change in the optimal conformation between native and the mutant.](doi:10.1371/journal.pone.0114903.g005)
neurodevelopmental data. The resulting combined knowledge base may be used for extensive annotation of data and the results of analyses by these multiple resources. Another advantage offered by the described platform is the seamless integration of tools that supports the workflows spanning across the contributing resources (see Fig. 6). Datasets provided by the user in the form of SNPs, gene lists, genomic coordinates, VCF files, results of gene expression experiments, or obtained via queries to the participating knowledge bases may be analyzed by a variety of tools used in combinations suitable for the goals of a particular experiment.

The following types of analysis are currently integrated: RViewer provides an extensive annotation of genomic intervals of interest (e.g. known variations, TFBS); Lynx enrichment analysis allows the user to identify biological functions and phenotypes over-represented in the user datasets; Lynx network-based gene prioritization predicts high-confidence genes contributing to disease phenotypes; RaptorX server gives predictions of the 3D protein structure for protein sequences without close homologs in PDB, solvent accessibility, and disordered regions thus facilitating understanding of protein-ligand interactions.

We are working on the expansion of the array of available analytical services integrated in the described environment, specifically to provide access to additional domain-specific resources (e.g. cancer and cardiovascular studies). As the volume and complexity of biological information continues to increase, the seamless integration of bioinformatics platforms will offer a practical solution for the needs of biomedical studies.
Supporting Information

Materials S1. Supporting text, figures and table. Text S1. Methodology. Figure S1. The overall view of the energy-minimized structures of SLC19A1, native (a) and H27R mutant (b). Figure S2. The changes to the binding site caused by the mutation (left: native, right: H27R mutant). (a and b) The mutation results in changing contact landscape inside the cleft, especially for the Arg151 residue. (c) Shift of Pro146 causes a kink in the loop, causing the helix to kink and the sidechain of Phe141 to turn, reducing the size of the cleft entrance. Figure S3. Visualization of the cleft volume and shape in both the native (left) and the mutant (right) structures. The H27R mutation reduces the volume of the cleft by 33%. Table S1. Volume and surface area of the cleft in the native and the mutant structures as calculated by 3V (ref).

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Author Contributions

Conceived and designed the experiments: ID MER CEM NM TCG. Performed the experiments: CM LA. Analyzed the data: SB SW AP JM. Contributed reagents/materials/analysis tools: DS DB BX AT ARP GMM PD GA JX. Wrote the paper: ID NM TCG SB SW MER CEM CM.

References

1. Boucher B, Jenna S (2013) Genetic interaction networks: better understand to better predict. Front Genet 4: 290.
2. Pastrello C, Pasini E, Kotlyar M, Otasek D, Wong S, et al. (2014) Integration, visualization and analysis of human interactome. Biochem Biophys Res Commun 445: 757–773.
3. Seoane JA, Lopez-Campos G, Dorado J, Martin-Sanchez F (2013) New approaches in data integration for systems chemical biology. Curr Top Med Chem 13: 591–601.
4. Wang S, Xing J (2013) A primer for disease gene prioritization using next-generation sequencing data. Genomics Inform 11: 191–199.
5. Cordero F, Beccuti M, Donatelli S, Calogero RA (2012) Large disclosing the nature of computational tools for the analysis of next generation sequencing data. Curr Top Med Chem 12: 1320–1330.
6. Hong H, Zhang W, Shen J, Su Z, Ning B, et al. (2013) Critical role of bioinformatics in translating huge amounts of next-generation sequencing data into personalized medicine. Sci China Life Sci 56: 110–118.
7. Warde-Farley D, Donaldson SL, Comes O, Zuberi K, Badrawi R, et al. (2010) The GeneMANIA prediction server: biological network integration for gene prioritization and predicting gene function. Nucleic Acids Res 38: W214–220.
8. Franceschini A, Szklarczyk D, Frankild S, Kuhn M, Simonovic M, et al. (2013) STRING v9.1: protein-protein interaction networks, with increased coverage and integration. Nucleic Acids Res 41: D808–815.
9. Szklarczyk D, Franceschini A, Kuhn M, Simonovic M, Roth A, et al. (2011) The STRING database in 2011: functional interaction networks of proteins, globally integrated and scored. Nucleic Acids Res 39: D561–568.
10. Chen J, Bardes EE, Aronow BJ, Jegga AG (2009) ToppGene Suite for gene list enrichment analysis and candidate gene prioritization. Nucleic Acids Res 37: W305–311.

11. Tranchevent LC, Barriot R, Yu S, Van Vooren S, Van Loo P, et al. (2008) ENDEAVOUR update: a web resource for gene prioritization in multiple species. Nucleic Acids Res 36: W377–384.

12. Sifrim A, Popovic D, Tranchevent LC, Ardeshirdavani A, Sakai R, et al. (2013) eXtasy: variant prioritization by genomic data fusion. Nat Methods 10: 1083–1084.

13. Wu J, Li Y, Jiang R (2014) Integrating multiple genomic data to predict disease-causing nonsynonymous single nucleotide variants in exome sequencing studies. PLoS Genet 10: e1004237.

14. Jager M, Wang K, Bauer S, Smedley D, Krawitz P, et al. (2014) Jannovar: a java library for exome annotation. Hum Mutat 35: 548–555.

15. Li MX, Gui HS, Kwan JS, Bao SY, Sham PC (2012) A comprehensive framework for prioritizing variants in exome sequencing studies of Mendelian diseases. Nucleic Acids Res 40: e53.

16. Calabrese C, Simone D, Diroma MA, Santorsola M, Gutta C, et al. (2014) MToolBox: a highly automated pipeline for heteroplasmy annotation and prioritization analysis of human mitochondrial variants in high-throughput sequencing. Bioinformatics.

17. Yao J, Zhang KX, Kramer M, Pellegrini M, McCombie WR (2014) FamAnn: an automated variant annotation pipeline to facilitate target discovery for family-based sequencing studies. Bioinformatics [Epub ahead of print].

18. Li X, Montgomery SB (2013) Detection and impact of rare regulatory variants in human disease. Front Genet 4: 67.

19. Sulakhe D, Balasubramanian S, Xie B, Feng B, Taylor A, et al. (2014) Lynx: a database and knowledge extraction engine for integrative medicine. Nucleic Acids Res 42: D1007–1012.

20. Frazer KA, Pachter L, Poliakov A, Rubin EM, Dubchak I (2004) VISTA: computational tools for comparative genomics. Nucleic Acids Res 32: W273–279.

21. Mirzaa GM, Millen KJ, Barkovich AJ, Dobyns WB, Paciorkowski AR (2014) The Developmental Brain Disorders Database (DBDB): A curated neurogenetics knowledge base with clinical and research applications. Am J Med Genet A.

22. Kallberg M, Wang H, Wang S, Peng J, Wang Z, et al. (2012) Template-based protein structure modeling using the RaptorX web server. Nat Protoc 7: 1511–1522.

23. Peng J, Xu J (2011) RaptorX: exploiting structure information for protein alignment by statistical inference. Proteins 79 Suppl 10: 161–171.

24. Lukashin I, Novichkov P, Boffelli D, Paciorkowski AR, Minovitsky S, et al. (2011) VISTA Region Viewer (RViewer)–a computational system for prioritizing genomic intervals for biomedical studies. Bioinformatics 27: 2595–2597.

25. Hamilton NA, Tammen I, Raadsma HW (2013) Multi-species comparative analysis of the equine ACE gene identifies a highly conserved potential transcription factor binding site in intron 16. PLoS One 8: e55434.

26. Infante CR, Park S, Mihala AG, Kingsley DM, Menke DB (2013) Pitx1 broadly associates with limb enhancers and is enriched on hindlimb cis-regulatory elements. Dev Biol 374: 234–244.

27. Ravi V, Bhatia S, Gautier P, Loosli F, Tay BH, et al. (2013) Sequencing of Pax6 loci from the elephant shark reveals a family of Pax6 genes in vertebrate genomes, forged by ancient duplications and divergences. PLoS Genet 9: e1003177.

28. Flicek P, Amode MR, Barrett D, Beal K, Brent S, et al. (2012) Ensembl 2012. Nucleic Acids Res 40: D84–90.

29. Kent WJ, Sugnet CW, Furey TS, Roskin KM, Pringle TH, et al. (2002) The human genome browser at UCSC. Genome Res 12: 996–1006.

30. Thorvaldsdottir H, Robinson JT, Mesirov JP (2013) Integrative Genomics Viewer (IGV): high-performance genomics data visualization and exploration. Brief Bioinform 14: 178–192.

31. Gotea V, Visel A, Westlund JM, Nobrega MA, Pennacchio LA, et al. (2010) Homotypic clusters of transcription factor binding sites are a key component of human promoters and enhancers. Genome Res 20: 565–577.
32. Visel A, Minovitsky S, Dubchak I, Penacchio LA (2007) VISTA Enhancer Browser—a database of tissue-specific human enhancers. Nucleic Acids Res 35: D88–92.
33. Burzynski GM, Reed X, Taher L, Stine ZE, Matsui T, et al. (2012) Systematic elucidation and in vivo validation of sequences enriched in hindbrain transcriptional control. Genome Res 22: 2278–2289.
34. Narlikar L, Sakabe NJ, Blanski AA, Arimura FE, Westlund JM, et al. (2010) Genome-wide discovery of human heart enhancers. Genome Res 20: 381–392.
35. Brudno M, Poliakov A, Minovitsky S, Ratnere I, Dubchak I (2007) Multiple whole genome alignments and novel biomedical applications at the VISTA portal. Nucleic Acids Res 35: W669–674.
36. Adzhubei IA, Schmidt S, Peshkin L, Ramensky VE, Gerasimova A, et al. (2010) A method and server for predicting damaging missense mutations. Nat Methods 7: 248–249.
37. Xie B, Agam G, Sulakhe D, Malteev N, Chitturi B, et al. Prediction of Candidate Genes for Neuropsychiatric Disorders Using Feature-based Enrichment; 2012; New York, NY, USA.
38. Nitsch D, Tranchevent LC, Goncalves JP, Vogt JK, Madeira SC, et al. (2011) PINTA: a web server for network-based gene prioritization from expression data. Nucleic Acids Res 39: W334–338.
39. Mirzaa GM, Millen KJ, Barkovich AJ, Dobyns WB, Paciorkowski AR (2014) The Developmental Brain Disorders Database (DBDB): a curated neurogenetics knowledge base with clinical and research applications. Am J Med Genet A 164A: 1503–1511.
40. Amberger J, Bocchini CA, Scott AF, Hamosh A (2009) McKusick’s Online Mendelian Inheritance in Man (OMIM). Nucleic Acids Res 37: D793–796.
41. Basu SN, Kollu R, Banerjee-Basu S (2009) AutDB: a gene reference resource for autism research. Nucleic Acids Res 37: D832–836.
42. Becker KG, Barnes KC, Bright TJ, Wang SA (2004) The genetic association database. Nat Genet 36: 431–432.
43. Konneker T, Barnes T, Furberg H, Losh M, Bulik CM, et al. (2008) A searchable database of genetic evidence for psychiatric disorders. Am J Med Genet B Neuropsychiatr Genet 147B: 671–675.
44. Ma J, Peng J, Wang S, Xu J (2012) A conditional neural fields model for protein threading. Bioinformatics 28: i59–66.
45. Peng J, Xu J (2011) A multiple-template approach to protein threading. Proteins 79: 1930–1939.
46. Wang S, Ma J, Peng J, Xu J (2013) Protein structure alignment beyond spatial proximity. Sci Rep 3: 1448.
47. Mitchell LE, Adzick NS, Melchionne J, Pasquariello PS, Sutton LN, et al. (2004) Spina bifida. Lancet 364: 1885–1895.
48. Padmanabhan R (2006) Etiology, pathogenesis and prevention of neural tube defects. Congenit Anom (Kyoto) 46: 55–67.
49. Ross ME (2010) Gene-environment interactions, folate metabolism and the embryonic nervous system. Wiley Interdiscip Rev Syst Biol Med 2: 471–480.
50. Wallingford JB, Niswander LA, Shaw GM, Finnell RH (2013) The continuing challenge of understanding, preventing, and treating neural tube defects. Science 339: 1222002.
51. Wang K, Li M, Hakonarson H (2010) ANNOVAR: functional annotation of genetic variants from high-throughput sequencing data. Nucleic Acids Res 38: e164.
52. Copp AJ, Stanier P, Greene ND (2013) Neural tube defects: recent advances, unsolved questions, and controversies. Lancet Neurol 12: 799–810.
53. Copp AJ, Greene ND (2013) Neural tube defects—disorders of neurulation and related embryonic processes. Wiley Interdiscip Rev Dev Biol 2: 213–227.
54. Boyles AL, Billups AV, Deak KL, Siegel DG, Mehltretter L, et al. (2006) Neural tube defects and folate pathway genes: family-based association tests of gene-gene and gene-environment interactions. Environ Health Perspect 114: 1547–1552.
55. Marini NJ, Hoffmann TJ, Lammer EJ, Hardin J, Lazaruk K, et al. (2011) A genetic signature of spina bifida risk from pathway-informed comprehensive gene-variant analysis. PLoS One 6: e28408.
56. Chandler AL, Hobbs CA, Mosley BS, Berry RJ, Canfield MA, et al. (2012) Neural tube defects and maternal intake of micronutrients related to one-carbon metabolism or antioxidant activity. Birth Defects Res A Clin Mol Teratol 94: 864–874.

57. Fisk Green R, Byrne J, Crider KS, Gallagher M, Koontz D, et al. (2013) Folate-related gene variants in Irish families affected by neural tube defects. Front Genet 4: 223.

58. Liu J, Qi J, Yu X, Zhu J, Zhang L, et al. (2014) Investigations of single nucleotide polymorphisms in folate pathway genes in Chinese families with neural tube defects. J Neurol Sci 337: 61–66.

59. Chalamalasetty RB, Dunty WC Jr, Biris KK, Ajima R, Iacovino M, et al. (2011) The Wnt3a/beta-catenin target gene Mesogenin1 controls the segmentation clock by activating a Notch signalling program. Nat Commun 2: 390.

60. Morin I, Devlin AM, Leclerc D, Sabbagian N, Halsted CH, et al. (2003) Evaluation of genetic variants in the reduced folate carrier and in glutamate carboxypeptidase II for spina bifida risk. Mol Genet Metab 79: 197–200.

61. Doolin MT, Barbaux S, McDonnell M, Hoess K, Whitehead AS, et al. (2002) Maternal genetic effects, exerted by genes involved in homocysteine remethylation, influence the risk of spina bifida. Am J Hum Genet 71: 1222–1226.

62. Aminoff M, Carter JE, Chadwick RB, Johnson C, Grasbeck R, et al. (1999) Mutations in CUBN, encoding the intrinsic factor-vitamin B12 receptor, cubilin, cause hereditary megaloblastic anaemia 1. Nat Genet 21: 309–313.

63. Franke B, Vermeulen SH, Steegers-Theunissen RP, Coenen MJ, Schijvenaars MM, et al. (2009) An association study of 45 folate-related genes in spina bifida: Involvement of cubilin (CUBN) and tRNA aspartic acid methyltransferase 1 (TRDMT1). Birth Defects Res A Clin Mol Teratol 85: 216–226.

64. Whitehead VM (2006) Acquired and inherited disorders of cobalamin and folate in children. Br J Haematol 134: 125–136.

65. Rady PL, Szucs S, Matalon RK, Grady J, Hudnall SD, et al. (2001) Genetic polymorphism (G80A) of reduced folate carrier gene in ethnic populations. Mol Genet Metab 73: 285–286.

66. Matherly LH, Hou Z (2008) Structure and function of the reduced folate carrier a paradigm of a major facilitator superfamily mammalian nutrient transporter. Vitam Horm 79: 145–184.

67. Stanislawska-Sachadyn A, Mitchell LE, Woodside JV, Buckley PT, Kealey C, et al. (2009) The reduced folate carrier (SLC19A1) c.80G>A polymorphism is associated with red cell folate concentrations among women. Ann Hum Genet 73: 484–491.

68. Kozyraki R, Fyfe J, Kristiansen M, Gerdes C, Jacobsen C, et al. (1999) The intrinsic factor-vitamin B12 receptor, cubilin, is a high-affinity apolipoprotein A-I receptor facilitating endocytosis of high-density lipoprotein. Nat Med 5: 656–661.

69. Wahlstedt-Froberg V, Pettersson T, Aminoff M, Dugue B, Grasbeck R (2003) Proteinuria in cubilin-deficient patients with selective vitamin B12 malabsorption. Pediatr Nephrol 18: 417–421.