Next generation tools for the annotation of human SNPs

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Abstract

Computational biology has the opportunity to play an important role in the identification of functional single nucleotide polymorphisms (SNPs) discovered in large-scale genotyping studies, ultimately yielding new drug targets and biomarkers. The medical genetics and molecular biology communities are increasingly turning to computational biology methods to prioritize interesting SNPs found in linkage and association studies. Many such methods are now available through web interfaces, but the interested user is confronted with an array of predictive results that are often in disagreement with each other. Many tools today produce results that are difficult to understand without bioinformatics expertise, are biased towards non-synonymous SNPs, and do not necessarily reflect up-to-date versions of their source bioinformatics resources, such as public SNP repositories. Here, I assess the utility of the current generation of webservers; and suggest improvements for the next generation of webservers to better deliver value to medical geneticists and molecular biologists.

Keywords: SNP; bioinformatics; prediction methods; webservers; review

INTRODUCTION

The rapid growth of genomic tools such as single nucleotide polymorphism (SNP) allele genotyping arrays and next-generation DNA sequencing has produced unprecedented amounts of information about the genotypes of individuals in many species. Yet even when association or linkage studies detect statistically significant correlations between a genomic region and a phenotype, the identity of the causative polymorphism often remains unknown. Tracking down functional SNPs is one of the key challenges of modern genetics, and a new branch of computational biology has emerged to support this effort.

The first computational methods designed to predict the biological impact of SNPs appeared almost a decade ago [1–5]. In subsequent years, a variety of methods have been introduced, reviewed in [6–9], and many now provide websites that take SNPs of interest as input and return annotations,

including classifications of biological importance [10–18]. Medical genetics and molecular biology researchers are increasingly turning to these methods and websites as an inexpensive way to prioritize SNPs of interest, prior to functional tests [19–29], and even to select *tag SNPs* for linkage and association studies [30–33]. These methods incorporate material from computer science, applied mathematics and population genetics, including machine learning, probabilistic modeling, statistics, software engineering and phylogeny. To make technical material accessible, specialized terms have been italicized and are defined in a glossary (Table 1).

The SNP function prediction community currently lacks a gold standard. Available methods have been trained and benchmarked on many different data sets (Table 2), and many methods are applicable to only a subset of all SNPs, such as non-synonymous (amino-acid changing) SNPs, or

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Table 1: Glossary of technical terms

Multiple sequence alignment

ancestor.

Affymetrix genotyping array	A microarray designed to identify known single nucleotide polymorphisms, given genomic DNA from an individual. Each SNP allele is represented by multiple short oligonucleotide probes, to which the individual's DNA binds through Watson-Crick base pairing, yielding fluorescence intensities. The intensity of these probes can be used to analyze which DNA base is present at a SNP position and whether the individual of interest carries two identical bases (homozygous) or two different bases (heterozygous) at the position.
Amino acid column distributions	A probability distribution describing the amino acid residues seen in an individual aligned column of a protein multiple sequence alignment. This distribution characterizes the probability that each of the 20 amino acids found in proteins will appear in the column.
Central or common data model	A data integration paradigm used in software engineering. The central or common data model specifies common rules about how software must access data structures. While individual databases have their own data models, called schemas, the central or common data model defines all the data relationships that exist in a particular software environment. Relationships between source data and the central data model are known as metadata.
Clade	A definition used in phylogenetic analysis. It is a taxonomic group that contains a single common ancestor and all the descendants of that ancestor.
Coefficient of determination	A statistical term, also known as R^2 . It is the proportion of variability in a data set that is accounted for by a statistical model.
Conserved	A property of entities undergoing evolution. Evolution is change over time, through descent with modification. 'Conserved' describes the entities that do not change over evolutionary time. For example, if an amino acid residue type is always seen in a particular position of the sequences from a protein family, it is said to be conserved. A 'conserved SNP' is a SNP that is at a conserved sequence position.
dbSNP reference identifiers	These are identifiers given to SNPs by the dbSNP database at NCBI. They begin with the letters 'rs' and are followed by digits.
Decision tree	A machine learning method in which attributes to be tested are organized in a tree structure and a decision is made at each branch point. Each series of decisions results in an overall prediction or classification.
Delaunay tesselation	A method used in computational geometry to reconstruct a continuous surface or volume from a discrete set of points.
Distributed data integration	Software engineering technology in which multiple databases are not integrated exclusively through a central server (a hub surrounded by spokes), but rather through multiple servers.
Domain interface	In a protein structure that has more than one protein domain, a region where amino acid residues from two or more domains are close enough to interact (\sim 6 A°) apart.
Executor kernel	Software engineering term used to describe code that controls the execution process of other code and keeps track of system state.
Gene Ontology (GO)	A controlled vocabulary, defined by an international consortium of scientists, known as the Gene Ontology Consortium, which describes gene and gene product attributes. It defines a hierarchy of biological processes and molecular functions.
Genome correlation structure	General term to describe non-random associations among DNA sequences either on the same chromosome or on different chromosomes.
Genomic range	Distance measured according to number of DNA base pairs, or starting and ending coordinates on a chromosome.
Haplotype	Regions of genomic DNA on the same chromosome, which are transmitted together from generation to generation. Also used to describe a set of SNPs on a strand of DNA that are statistically associated.
HapMap HapMap population	An international project that has identified human haplotypes in four ethnic populations. One of the four ethnic populations studied by the HapMap project (HapMap): Yoruba Africans, Han Chinese, Japanese, and the 'CEPH' families, who are Caucasians from Utah in the United States.
Hidden Markov model	A probabilistic model that was first used in speech recognition and has been useful in representation of related groups of biological sequences. The models contain a series of states, each with its own probability distribution, which estimates the probability that a particular amino acid residue (or nucleotide base) will appear at a position. There are also probabilities assigned to transitions between states. These models can be used to represent protein families.
Homolog	A biological sequence with the property that it has a common ancestor with a protein sequence of interest.
Homologous protein structure	An experimentally determined (through X ray crystallography or nuclear magnetic resonance spectroscopy) protein structure that is a homolog of a protein structure of interest.
HTTP servlet	Software engineering term to describe a Java application that runs on a web server and provides server-side processing, such as database access and HTTP requests from a web browser.
Illumina bead array	A technology for identifying SNPs using oligonucleotide probes to which an individual's DNA binds, yielding fluorescence. The probes are attached to silica beads.
Intelligent agents	Software agents that can do data mining on the internet, either by following rules or by learning and adapting as they see new data.
Ligand binding site	Region on a protein structure where amino acid residues are close enough to a ligand to interact with its atoms (~5 A°).
Linkage disequilibrium Machine learning	Non-random association between regions of DNA either on the same chromosome or not. A research area within artificial intelligence that focuses on algorithms that are able to learn. A common learning task is classifying examples from two or more categories.

Alignment of three or more sequences that are assumed to be related through descent from a common

Table I: Continued

Multiple sequence alignment column	A column in a multiple sequence alignment that represents part of the conserved core structure of a group of related proteins.
Neighbor-joining by sequence identity	y A method for constructing a phylogenetic tree from a multiple sequence alignment. Sequences are clustered in a bottom-up, iterative algorithm that uses percent sequence identity (fraction of identical positions) as a similarity measure.
Neural network	A machine learning method based on neural organization in the human brain.
OMIM	Online Mendelian Inheritance in Man. A database of inherited human mutations in single genes that are known to cause disease.
Phylogenetic tree	A tree that represents the evolutionary relationships among a group of nucleotide or protein sequences, assumed to have a common ancestor.
Protein domain	A region of a protein that is an independent folding unit.
Protein homologs	Proteins that have been inferred to descend from a common ancestor through phylogenetic methods or because their sequences are similar.
Protein homology model	A computational model that predicts the 3D Euclidean coordinates of all heavy atoms in a protein of interest, based primarily on an experimentally determined structure of a homolog.
Pseudocounts	Technique used in probabilistic modeling in which low probability events are given small probabilities of occurring, even when they are not observed in sample data. Also known as background counts.
Random forest	Machine learning method in which hundreds of decision trees (Decision Tree) are combined into an ensemble and a prediction or classification is arrived at by a vote of the entire ensemble.
Reconfigurable web wrapper agents	Software engineering tool to automate web browsing sessions using agents, which discover the rules and extract the structure of a web page.
Regulatory motifs	Patterns of DNA or mRNA sequence that are the signatures of binding sites for protein or RNA molecules, involved in transcriptional and translational regulation.
Sequence profile	Representation of a group of related biological sequences, which estimates the probability that a particular amino acid residue (or nucleotide base) will appear at each position.
Sequence weighting	A method used to improve the generalization ability of statistical models of biological sequences. To avoid a group of similar sequences in a data set from dominating the model, sequence 'subfamilies' that are overrepresented in the data set are downweighted and sequences that are dissimilar to the rest of the data set are upweighted.
Single-marker and two-marker correlations	Metrics of correlation (linkage disequilibrium) between pairs of SNPs (single-marker) and triples of SNPs (two-marker) that are used in selecting the most informative ('tag') SNPs for whole-genome association studies.
SNP probe libraries	A collection of short oligonucleotides that are used in genotyping microarray and bead technologies. They are designed to bind to pre-designated SNPs, culled from sources such as dbSNP and the HapMap project.
Splicing enhancer motif	A probabilistic model of a short sequence of mRNA bases which are the binding target of proteins ('splicing factors') involved in splicing. The <i>canonical</i> motif is the most frequently seen sequence of mRNA bases at the binding site of interest.
Support vector machine	A machine learning method that is based on 'decision planes' to yield maximal linear separation of different classes of data, often through projection into higher dimensions. Reviewed in [77].
Tag SNPs	The most informative SNPs for genome-wise association studies. Tag SNPs have the highest statistical power to detect association.
TaqMan	A single-tube PCR (polymerase chain reaction) assay that can be used for SNP genotyping. A SNP is represented by oligonucleotide probes with fluorophores on each end. When the probes hybridize to their targets in sample DNA, a fluorescent dye-specific signal is generated.
Web navigation description language Web wrapper agent	An XML-based language useful in automating data mining over the WWW. Software that automates a user web-browsing session. It visits a website, fills out query forms, and extracts returned data [78].
XML formatted data	Data structured with XML (extensible markup language). XML is a flexible text format with its own grammar, useful for sharing data, primarily over the internet. Like HTML, it has markup symbols, but unlike HTML, these symbols are unlimited and self-defining.

non-synonymous SNPs that can be mapped onto protein structures. Fair assessment of which methods are best is beyond the scope of this review. Instead, I present a survey of available services, discuss trends in the field, and highlight strengths and weaknesses that may be of interest to a potential user of SNP function prediction webservers.

SNP webservers: strategies and communities

Today's SNP prediction servers generally use one of three strategies (Table 2):

(i) methods servers that disseminate results of original computational method(s);

 Table 2: Computational biology SNP prediction webservers fall into three categories

Name	SIFT	PolyPhen	SNAP	PMUT	PANTHER	nsSNP Analyzer	PhD-SNP	Auto-Mute
Table 2A: Metho	ds servers							
Computational Methods	conservation among protein homologs	decision tree	Neural network	Neural network	Hidden Markov model of protein family	Random forest	Decision tree coupled to two support vector machines	Random forest, Delaunay tesse- lation [79] of protein structure.
WebsiteURL	http:// blocks.fhcrc. org/sift/ SIFT.html	http://genetics. bwh.harvard. edu/pph	http://cubic.bioc. columbia. edu/ services/SNAP/	http:// mmb2.pcb.ub. es:8080/PMut/	http:// www.pantherdb. org/tools/ csnpScoreForm.jsp	http://snpanalyzer. utmem.edu	http://gpcr2.biocomp. unibo.it/~emidio/ PhD-SNP/	http://proteins. gmu.edu/auto mute/AUTO- MUTE.nsSNPs. html
Datatypes	Protein sequences and multiple sequence alignments	Protein sequences, mul- tiple sequence alignments, protein structures	Protein sequen- ces,multiple sequence align- ments, pre- dicted protein secondary structures	Protein sequen- ces,multiple sequence align- ments, pre- dicted protein secondary structures, protein structure	Protein sequences, hidden Markov models	Protein sequences, multiple sequence alignments, homo- logous protein structures, protein secondary structure	Protein sequences, sequence profiles	Protein structures
Bench-mark or training data	Saturation mutagenesis data sets of two bacterial and one retroviral protein [80–82]	Disease variants and mutagens from SwissProt Variant Pages [83] and pre- sumed netural between- species replace- ments in multi- ple sequence alignments	The 80 000+ mutations from Protein Mutant Database [84], two bacterial and one retro- viral protein [80-82], enzymes with experimentally annotated func- tion in SwissProt [35]	Disease variants from SwissProt Variant Pages [83] and pre- sumed netural between-spe- cies replace- ments in multi- ple sequence alignments	Human Gene Mutation Database [85] for disease mutations and dbSNP [53] for presumably neutral variants	Selected mutants from SwissProt Variant Pages [83] that are mapped onto homologous protein structures	SwissProt Variant Pages [83]	The I790 disease and neutral variants from SwissProt Variant Pages [83] that can be mapped onto PDB [36] structures
Batch input of SNPs?	Yes	No	Yes	Yes	Yes	Yes	No	Up to five SNPs at one time

(Continued)

Name	SNP@Domain	PolyDoms	MutDB	Snap	StSNP
Table 2B: Meta servers					
Computational Methods	Integration of data from multi- ple sources	Integration of data from multi- ple sources	Integration of data from multi- ple sources	Integration of data from multi- ple sources	Integration of data from multiple sources
Website URL Datatypes	http://snpnavigator.net Protein multiple sequence alignments, protein struc- tures, predicted functional effects, disease annotations	http://polydoms.cchmc.org Protein multiple sequence alignments, protein struc- tures, GO categories [86], dis- ease annotations, pathways, interacting protein net- works, mammalian pheno- types, predicted functional effects. Includes synon- ymous SNPs	http://mutdb.org Genomic DNA, mRNA transcripts, protein sequence, protein multiple sequence alignments, protein structures, pathways, disease annotations. Includes intronic, untranslated region, and and synonymous SNPs	http://snap.humgen.au.dk Genomic DNA, mRNA tran- scripts, protein sequence, phylogenetic trees, interact- ing protein networks, dis- eases, post translational modifications, splice sites	http://glinka.bio.neu.edu/StSNP Genomic. DNA, protein sequence, pathways, protein structures, pro tein homology models
Benchmark or training data	N/A	A total of I338 SNPs from 6II candidate genes with known disease mutations (ftp://ftp.ncbi.nih.gov/snp/Entrez/snp.omimvar.txt)	N/A	N/A	N/A
Batch input of SNPs?	Yes	No	Yes	Yes, if in the same gene	Yes
Name	PupaSuite	SNP function portal	SNPselect	F-SNP	
Computational Methods	Integration of data from multiple sources	Integration of data from multi- ple sources	Integration of data from multi- ple sources	Integration of data from multi- ple sources	
Website URL	http://pupasuite.bioinfo.cipf.es	http://brainarray.mbni.med. umich.edu/Brainarray/ Database/SearchSNP/ snpfunc.aspx	http://snpselector.duhs. duke.edu/hqsnp36.html	http://compbio.cs.queensu.ca/ F-SNP/	
Data types	Genomic DNA, mRNA transcripts, protein sequence, haplotypes. Regulatory SNPs, synonymous SNPs, intronic SNPs, untranslated region SNPs, intergenic SNPs, nonsense and frameshift mutations, protein structure, cellular processes, functional sites, evolutionary selection strength dN/dS, and epigenetic effects (triplex DNA regions). Human, mouse and rat included	Genomic DNA, mRNA transcripts, protein sequences, protein structures and homology models, pathways, diseases, and haplotypes, (gene expression is under construction)	Applied Biosystems and Illumina SNP data, genomic DNA and haplotypes	Protein sequences, protein structures protein homology models, mRNA transcripts, predicted functional impact on protein structure, splicing regulation, post translational modifications and evolutionary conservation	
Benchmark or training data	N/A	N/A	A total of 700 SNPs from I40 genes associated with cardio-vascular disease in [87]	N/A	
Batch input of SNPs?	Yes	Yes	Yes	Yes, if in same gene or genomic region	

Table 2: Continued

Name	LS-SNP	SNPeffect	SNPs3D	FastSNP	MutaGeneSys
Table 2C: Hybrids					
Computational Methods	Support vector machine	Integration of data from mul- tiple sources	Support vector machine	Decision tree	Identifies indirect correla- tions between SNPs and mutations from OMIM [13]
Website URL	http://karchinlab.org/LS-SNP	http://snpeffect.vib.be/	http://www.snps3d.org/	http://fastsnp.ibms. sinica.edu.tw	http://magnet.c2b2.columbia. edu/mutagenesys
Datatypes	Protein sequences, multi- ple sequence alignments, protein homology models, predicted domain interfaces, ligand binding sites, hidden Markov models, genes, path- ways, genomic DNA	Predicted changes in protein stability and folding, aggregation and amyloidosis, catalytic sites and binding sites, phosphorylation and glycosylation sites, cellular localization and protein turnover.	Protein sequences, multiple sequence alignments, profiles, protein structures, genes, gene networks, disease candidate genes, GO categories [86], mouse knockout data	Genes, genomic DNA, mRNA transcripts, protein sequences, protein domains	Genomic sequence, haplo- types, linkage disequilibrium data
Benchmark or training data	A total of I457 disease-associated variants from SwissProt [35] which could be mapped to the OMIM database [59] and and 2504 putatively neutral nsSNPs from dbSNP [53]	N/A	A total of 10263 deleterious mutants in 731 proteins from Human Gene Mutation Database [85] and 16682 control substitutions in 348 proteins from aligned positions of close orthologs	A total of I569 SNPs from the SNP500 Cancer database [88]	N/A
Batch input of SNPs?	Yes	Yes	Yes	Yes	Yes

⁽A) Methods servers primarily disseminate an original computational method for SNP function prediction. (B) Meta-servers pull information from many severs, including general purpose protein and genomic annotation bioinformatics servers and servers from category. (C) Hybrids both disseminate original method(s) and pull information from other servers. Technical terms have been italicized and can be looked up in the Glossary (Table I).

- (ii) metaservers that pull information from many servers, including general purpose protein and genomic annotation bioinformatics servers; and
- (iii) hybrids that both disseminate original method(s) and pull information from other servers.

All of these servers are built on top of an infrastructure of general bioinformatics resources that curate SNPs, genomic and protein sequences, protein structures, interactions, pathways and regulatory elements (such as sites important for transcription factor binding and accurate splicing). The relationships among SNP webservers and other bioinformatics resources can be represented as a directed graph (Figure 1). Partitioning the graph with an algorithm based on local modularity [34]

yields three main communities, which can be loosely defined as: the protein community (ellipse), connected to the large, core bioinformatics databases UniProt [35], Protein Data Bank (PDB) [36], Structural Classification of Proteins (SCOP) [37], Biomolecular Interaction Network Database (BIND) [38], Molecular Interactions Database (MINT) [39], Gene Ontology (GO) [40], Kyoto Encyclopedia of Genes and Genomes (KEGG) [41] and BioCarta (http://www.biocarta.com); the regulatory community (trapezoid) connected to webservers that predict post-translational modifications, splicing enhancers and repressors and transcription factor binding sites (TFBSs); and a regulatory plus linkage disequilibrium community (rectangle), which is connected to the HapMapwebserver (http://www.hapmap.org). Core resources such as National Center for

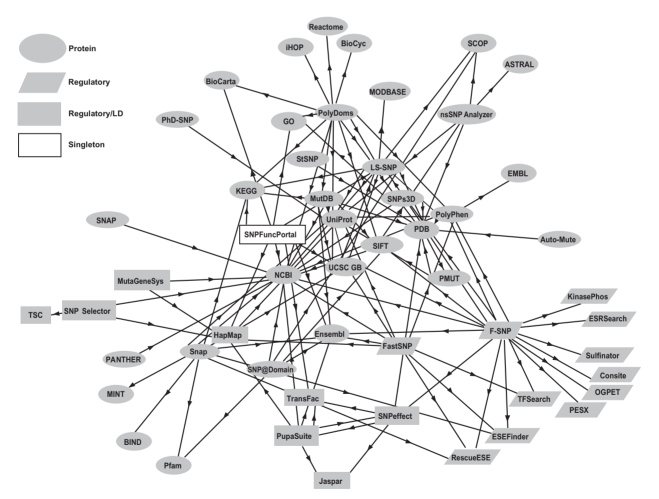


Figure I: Directed graph of relationships among SNP prediction webservers and their bioinformatics sources. A heuristic partition of the graph identifies three communities. They are loosely defined as (I) focus on protein properties (ellipse); (2) focus on regulation (trapezoid); and (3) connect to HapMap and consider linkage disequilibrium among SNPs (rectangle). SNPFunctionPortal is a singleton, perhaps an emerging community (white rectangle) that is equally connected to the protein and LD communities. LD = linkage disequilibrium.

Biotechnology Information (NCBI) databases (http://www.ncbi.nlm.nih.gov), UCSC Genome Browser [42] (http://genome.ucsc.edu) Ensembl [43] (http://www.ensembl.org) are used by all three communities. A singleton server (white rectangle), the SNP Function Portal [14], is connected to both protein and linkage disequilibrium communities, and perhaps represents an emergent fourth community. Protein community webservers primarily predict the biological importance of non-synonymous SNPs, using properties such as evolutionary conservation of amino acid sequence, protein structure and protein binding interactions. These properties are often combined in 'black box' machine learning algorithms—neural networks, support vector machines and random forests—vielding predictions that are difficult to understand from a biological point of view. The regulatory community primarily harvests predictions from external servers that specialize in identification of regulatory motifs. Although these methods were not designed specifically for SNPs, they can be used, at least in theory, to predict the effect of the SNP on normal patterns of regulation. The third community contains websites connected to resources that provide information about genomic linkage disequilibrium structure.

The 'protein community' is the largest and the oldest. But the general landscape is shifting towards inclusion of regulatory SNPs and consideration of inter-SNP associations through linkage disequilibrium (Figure 2a). The landscape may also be shifting away from methods servers towards meta-servers and hybrids (Figure 2b).

The webserver graph (Figure 1) shows that there is not much feedback to the servers from their sources, although this may change with time. There is currently one exception—a feedback loop connecting two SNP servers in the regulatory/linkage disequilibrium community—SNPeffect [13] and PupaSuite [12]. These servers are synchronized and describe their relationship as a joint effort to cover both protein and regulatory related SNPs. Such relationships may become more common in the next generation.

FIELD TESTING OF CURRENT WEB SERVERS

To assess their usability and scientific utility, I evaluated 22 severs by submitting to each a set of SNPs that were reported to be associated with disease in recently published medical literature. All

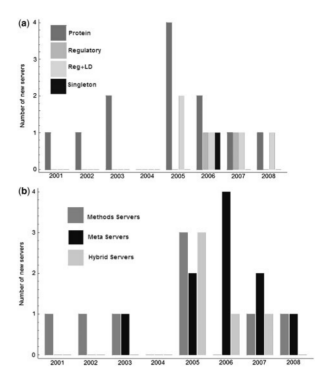


Figure 2: Trends in scope of SNP webservers. (a) Prior to 2006, protein-based servers that only handle non-synonymous SNPs were predominant. Newer servers include regulatory SNPs and annotate associations among SNPs through linkage disequilibrium estimates. (b) The earliest servers implemented original computational methods, but the current trend is towards meta-servers and hybrids. Dates associated with each server are based on date of first journal publication, unless an alternate date is documented on its website.

submissions were done using Firefox 2.0.0.13 on Windows XP Professional Edition. The field tests were done during the week of 28 April 2008. One server returned no results and inquiry emails went unanswered. It was eliminated from the assessment (Pmut [44]). Detailed descriptions of all field tests are provided (Supplementary Tables S2, S3 and S4) with the main results summarized in this section. Since all of these websites were designed by bioinformaticians, it is not surprising that all of them require some bioinformatics expertise on the part of the user. For each server tested, I provide an assessment of the expected user skill set. General definitions of basic bioinformatics skills and expert bioinformatics skills are also provided (Table 3).

ALS/FTLD study: novel SNPs discovered in sequencing

Novel SNPs are often discovered through DNA sequencing studies that compare individuals with a

Table 3: Basic and expert bioinformatics skill levels

Basic skill level Basic 'data types'—sequences and structures of DNA, RNA and protein and their representations in bioinformatics

Retrieving information from bioinformatics web databases.

Pairwise and multiple sequence alignments.

Higher level organization of and relationships between the basic data types—biochemical pathways, biochemical functions, gene and protein families, protein folds, motifs and regulation.

Expert skill level Algorithms used for biological data search and analysis.

Relationship of protein sequence, structure and function.

Protein structure modeling and its limitations.

Sequence alignment and motif-matching algorithms, how to interpret their output scores and also their limitations. Probability and statistics—extreme value distributions, hidden Markov models, Bayesian networks, profiles, Fundamentals of molecular evolution.

Fundamentals of machine learning methods used in bioinformatics. Neural networks, support vector machines, random forests, decision trees. Strengths and weaknesses of these methods.

Metrics for comparing the performance of classification methods—receiver operating characteristic curves, specificity versus sensitivity.

Skill using a molecular visualization package, such as Chimera [89], PyMol [90] or RasMol [91].

Webservers tested in this study implicitly assume that users have familiarity with the concepts listed under the appropriate skill level.

condition of interest to a control population. In a recent study of familial amyotrophic lateral sclerosis (ALS) with frontotemporal lobal degeneration (FTLD), researchers investigated sequence variation in the gene TARDBP [45]. All coding exons, most of the 5'-untranslated region, and approximately 100 intronic bases upstream and downstream of each exon were sequenced for 259 ALS/FTLD patients and 1127 controls. The TARDBP (NM_007375) variants 869G->C (amino acid change G290A) and 892G->A (amino acid change G298S) were found to be statistically associated with disease, and putatively linked both with loss and/or gain of protein function.

All of the 'Methods Servers' are capable of handling novel non-synonymous SNPs, because they offer the ability to submit a protein sequence along with a residue position and amino acid substitution. None of the 'Hybrid Severs' or 'Meta-servers' allows submission of protein sequences, but one of the 'Meta-servers' (FAST-SNP [15]) handles novel SNPs of all kinds, by allowing the user to submit a DNA sequence plus base position and nucleotide substitution. The TARDBP SNPs were submitted to the SIFT [4], PolyPhen [10], SNAP [17], PMUT, PANTHER [18], nsSNPAnalyzer [46], PhD-SNP [47], Auto-mute [48] and FAST-SNP servers. In cases where servers offered a choice of parameter settings, defaults were used. Generally, the servers reported results that were understandable, if accepted on face value. Most predicted that both SNPs are

neutral, and the predictions that disagreed with neutrality were low confidence (Table 4). The servers varied widely in terms of communicating prediction reliability. Some have no confidence measures and some have a simple binary (yes/no) confidence measure. The SNAP server provides the most detailed confidence information, including both a reliability index and an estimated accuracy rate for each prediction. In general, the results are qualitative, rather than quantitative, reflecting the current state-of-the-art of webserver-based SNP function prediction.

Required user skills

- (i) Basic bioinformatics skills (Table 3) such as ability to find and handle data, accession numbers and reference identifiers in web databases such as UniProt, PDB and NCBI.
- (ii) Bioinformatics expertise (Table 3) is required to understand server errors. Two servers returned error messages that assumed users know about hidden Markov models and protein structural homology.
- (iii) Bioinformatics expertise is required to think critically about how to interpret server results and their significance.

Interpreting server results SIFT

In addition to predicting SNP functional impact, SIFT builds a protein *multiple sequence alignment* of the

Table 4: Field test of novel SNPs discovered in sequencing

	Classification		Score		Prediction confidence			
	G290A	G298S	G290A	G298S	G290A	G298S		
SIFT	Affects protein function	Tolerated	0.03	0.41	Low	Good		
PolyPhen	Benign	Benign	1.016	0.038	_	_		
SNAP	Neutral	Neutral	_	_	53%	89%		
anther	_	_	_	_	_	_		
sSNP analyzer	_	_	_	_	_	_		
hD-SNP	Neutral	Disease	_	_	4	0		
Auto-mute	_	_	_	_	_	_		
AST-SNP	_	_	_	_	_	_		

The TARDBP SNPs 869G->C (amino acid change G290A) and 892G->A (amino acid change G298S) were found to be statistically associated with disease in a case/control study of familial ALS with FTLD and putatively linked to loss and/or gain of protein function. Both SNPs were submitted to eight "Methods Servers" (Table 2A) for SNP function prediction to evaluate required user skills and agreement with associations from the case/control study. Three of the webservers were unable to classify these SNPs for technical reasons (detail in Supplementary Tables 2). FAST-SNP does not classify novel SNPs with respect to overall impact on disease risk, but it predicted that a TFBS might be affected by both of these SNPs. '-' = not provided.

protein of interest and emails it to the user, allowing alignment analysis with bioinformatics software. I used the SIFT TARDBP alignment to build a phylogenetic tree, using neighbor-joining by sequence identity in JALVIEW [49]. Human TARDBP is located in a distinct clade on this tree. The G290A and G298S SNPs are in a glycine-rich domain that is present only in this clade and appears to be an evolutionary late comer in the TARDBP protein family. Sequence annotations, available through JALVIEW links to European Bioinformatics Institute (EBI) resources, indicate that human proteins in the clade are expressed in brain tissues, rendering plausible the hypothesis that at least one of these SNPs, or a SNP in this protein domain that is in linkage disequilibrium with these SNPs, could contribute to ALS/FTLD, a brain disorder.

FAST-SNP

According to FAST-SNP, the sequence surrounding the SNPs is a significant match to a predicted TFBS, but TFBS predictions are generally not reliable unless the prediction is in a known promoter region. Given that this region is within a coding exon, one should be suspicious of this prediction. FAST-SNP submitted the sequence to three splicing regulatory analysis servers: ESEFinder [50], Rescue-ESE [51] and FAS-ESS [52]. Only one of the three predicted anything. That prediction is that 869G->C (G290A) introduces a significant match (CTAATAG) to the canonical *splicing enhancer motif* CAGAGGG, which is bound by SF2/ASF proteins. Altogether, this raises

the interesting possibility of impact on the regulatory level rather than the protein level.

A user of these SNP methods servers who sees their outputs only on a surface level would conclude that the two ALS/FTD SNPs are neutral. However, a user with bioinformatics expertise (Table 3) might use the server results to suggest testable hypotheses about how these SNPs could affect biological function.

Schizophrenia study: common intronic SNPs

When case–control studies are done with microarray or *TaqMan* technologies that use *SNP probe libraries*, researchers may find SNPs in which the frequency differences between cases and controls are statistically significant. These SNPs are not novel, and are already indexed in large databases such as dbSNP [53]. A recent study compared two large schizophrenia populations to ethnically matched controls [54]. Seven SNPs in the introns of PDE4B, which encode a large phosphodiesterase involved in cAMP signaling regulation, were found to be significantly associated with schizophrenia (*dbSNP reference identifiers*: rs4320761, rs910694, rs1354064, rs1321177, rs2144719, rs1040716 and rs78038).

The 'schizophrenia SNPs' were submitted to five servers that handle intronic SNPs: SNPselector [55], PupaSuite, FASTSNP, F-SNP [56] and MutaGeneSys [57] (Table 5). None of the SNPs were predicted to have functional impact by SNPselector, FASTSNP and MutaGeneSys. PupaSuite reported that rs910694 is in a DNA

Table 5: Field test of common intronic SNPs

	Classification				Score					Prediction Confidence											
	rs 4320761	rs 910694	rs1321177	rs2144719	rs1354064	rs1040716	rs783038	rs 4320761	rs 910694	rs1321177	rs2144719	rs1354064	rs1040716	rs783038	rs 4320761	rs 910694	rs1321177	rs2144719	rs1354064	rs1040716	rs783038
SNPSelector	_	_	_	_	_	_	_	0.6, 0.0	0.6, 0.0	0.6, 0.0	0.6, 0.0	0.6, 0.0	0.6, 0.0	0.6, 0.0	_	_	_	_	_	_	_
PupaSuite	_	Т	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_
F-SNP	R	_	_	_	R	R	_	_	_	_	_	_	_	_	_	_	_	_	_	_	_
FAST - SNP	LR	LR	LR	LR	_	_	_	_	_	-	_	-	-	_	_	_	_	_	_	_	_
MutaGeneSys	-	-	-	-	-	-	_	-	-	-	-	-	-	-	-	-	-	-	-	-	-

The PDE4B intronic SNPs rs432076l, rs910694, rs1354064, rs432076l, rs1321177, rs21447l9, rs10407l6 and rs78038 were found to be statistically associated with schizophrenia in a case/control study. The SNPSelector score '0.6' means that the SNP is in an intron, but not at a exon–intron junction. The SNPSelector score '0.0' means that the SNP is predicted to not be important for regulation of transcription. R = biologically important because of impact on regulation of transcription. LR = low risk (general assessment with respect to increases in disease susceptibility). T = low triplex region '-'= not provided.

triplex region, a region of DNA with three strands. These regions play a role in repression of transcription, reviewed in [58], thus this SNP could putatively disrupt normal regulation of PDE4B. F-SNP identified three of the SNPs (rs1354064, rs4320761 and rs1040716) as being involved in transcriptional regulation and rs1040716 as being at a position that is *conserved* among species.

Required user skills

- Submitting queries to these servers requires no special skills, just dbSNP reference identifiers for a SNP of interest.
- (ii) Bioinformatics skills are required to understand outputs of SNPselector, PupaSuite and F-SNP, even at a surface level.
- (iii) Unix skills are required to access SNPselector's results, which are sent by email as a compressed tarball.
- (iv) F-SNP requires expertise specifically with UCSC Genome Browser tools and terms.
- (v) The FASTSNP server outputs are integrated into a decision tree algorithm, which is clearly laid out and understandable to a general user. This feature is not available in FASTSNP's 'novel SNP' service.
- (vi) MutaGeneSys requires some knowledge of statistical genetics, as the user must select a minimum *coefficient of determination* and has the option of selecting a *HapMap population*. It reports when a SNP is correlated by *linkage disequilibrium* with an externally annotated

disease-associated SNP, based on *OMIM* [59]. Both *single-marker* and *two-marker correlations* are considered.

MutaGeneSys is a tool aimed at the medical genetics community, where the importance of linkage disequilibrium is well understood. By enabling identification of SNPs that are indirectly associated with disease, it can help users narrow down the number of SNPs likely to have a direct functional effect. The PupaSuite result for rs910694 suggests a testable hypothesis that might explain schizophrenia association.

Esophageal cancer study: mix of common exonic and intronic SNP

Esophageal and esophago-gastric junction adenocarcinomas (EAC and EGJAC) have been linked to acid reflux, obesity and smoking. Risk is also related to exposure to nitrites (found in compounds such as tobacco smoke) that alkylate DNA at the O⁶ position of guanine [60]. A recent population casecontrol study in Australia looked at SNPs in DNArepair genes MGMT, XPD, XRCC1 and ERCC1 to identify possible genetic predispositions to EAC and EGJAC. MGMT (O⁶-methylguanine-DNA methyltransferase) specifically repairs O⁶-guanine alkylation damage. Results point to MGMT SNPs rs12268840 (intronic) and rs2308321 (non-synonymous) as being statistically significant in frequency between EAC patients (n=263) and controls (n = 1337) [60].

Table 6: Field test of non-synonymous and intronic SNPs

	Classification		Score			confidence
	rs230832I	rsl2268840	rs230832I	rsI2268840	rs230832I	rsl2268840
SIFT	Tolerated	_	0.7	_	Good	_
PolyPhen	Benign	_	0.378	_	_	_
SNAP	Neutral	_		_	60%	_
Panther	_	_	0.24 ^a	_	_	_
nsSNP-analyzer	Neutral	_	_	_	_	_
PhD-SNP	Neutral	-	_	_	4 ^b	_
Auto-mute	_	_	_	_	_	_
SNP@Domain	_	_	_	_	_	_
Pmut	_	_	_	_	_	_
PolyDoms	_	_	_	_	_	_
MutDB	_	-	_	_	_	_
Snap	_	-	_	_	_	_
StSNP	_	_	_	_	_	_
PupaSuite	ESE site	_	3.13 ^c	_	_	_
Snp Function Por	tal TFBS	_	_	_	_	_
SNPselector	_	-	I ^d , 0.0 ^d	0.6 ^d , 0.0 ^d	_	_
F-SNP	ESE site ESR site	_	_	_	_	_
LS-SNP	Neutral	-	_	_	High	_
SNPEffect	No effect	-	_	_	_	_
SNPs3D	Neutral	-	_	_	_	_
FAST-SNP	Low-medium risk	Very low-low risk	_	_	_	_
MutaGeneSys	No LD to known disease mutation	ns No LD to known disease mutations	_	_	_	_

The MGMT SNPs rs230832l (non-synonymous) and rsl2268840 (intronic), found to be significantly associated with esophageal cancers, were submitted to all 22 servers in this study (those that do not handle intronic SNPs were only queried about rs230832l). ^aPanther score is probability of SNP being deleterious. ^bMeaning of the PhD-SNP prediction confidence score is not explained on their website. ^cMeaning of the ESE site prediction score is not explained on their website. ^dSNPselector score 0.6 describes an intronic SNP that is not at an exon/intron junction. 1.0 describes a non-synonymous (amino-acid changing) SNP. The '0.0' means the SNP is predicted to not be important for regulation of transcription. '—' = not provided. ESE = exonic splicing enhancer; ESR = exonic spicing repressor.

The MGMT SNPs were submitted to all 22 servers (those that do not handle intronic SNPs were only queried about rs2308321) (Table 6). None of the servers predicted that rs2308321 has an impact on protein function. Several servers reported that this SNP was found at splicing regulation sites, but only F-SNP predicted that it would impact splicing regulation, because it changes both an exonic splicing enhancer and an exonic splicing repressor. None of the servers predicted functional impact for rs12268840.

Required user skills

- (i) The basic skills required to input queries and interpret outputs are the same as described for the TARDBP and 'schizophrenia SNPs'.
- (ii) Bioinformatics skills and knowledge of human genome structure allow users to submit advanced input queries. *Genomic range* is accepted by MutDB [61], Snap [62], PupaSuite, SNP Function Portal [14], F-SNP and LS-SNP [11]. Linkage disequilibrium can be factored into

- inputs using PupaSuite, SNP Function Portal, SNPselector and MutaGeneSys. In total, 18 distinct input data types are available on the servers tested (Table 5).
- (iii) Results output of the meta-servers (Table 2B) is generally large, heterogeneous and difficult to integrate without bioinformatics skills. One exception is the FastSNP server, which integrates its harvested data in a decision tree algorithm that is transparent and clearly explained to users.

The only testable hypothesis yielded from these server results was the possibility that splicing regulation of MGMT might be affected by rs2308321. In general, there is poor agreement among servers that harvest predictions of SNP impact on splicing, and the predictions are not associated with clear reliability measures.

Stale data

Most of the tested servers use NCBI's dbSNP [53] database as a primary source of SNP data, but are not

up-to-date, increasing the chances that annotations for SNPs of interest will not be available to users. Between 2003 and 2008, dbSNP has been updated, on average 2–3 times per year. Fourteen of the tested servers accept dbSNP rsIDs, and the current dbSNP build is version 129, May 2008. Only one server, FastSNP is using version 128. Seven servers are using version 126; three are using version 125; two are using version 124 and one is still using version 123 (from October 2004) (Supplementary Table 1).

These three field studies suggest a set of desirable features for a SNP webserver:

- (i) Options for submission input that require minimal bioinformatics expertise. Even when advanced submission options are available, offering an easy way to input SNPs ensures that a wider community will have access to the server.
- (ii) Error messages that do not require bioinformatics expertise to understand. Such messages can be confusing and frustrating to users and alienate non-bioinformaticians.
- (iii) For those with bioinformatics expertise (Table 3), the option to download server outputs such as alignments and protein structure models. The ability to use external bioinformatics software to analyze server output will help bioinformaticians develop testable hypotheses about SNP biological impact.
- (iv) Quantitative, calibrated measures of prediction reliability. If server output contains many predictions, such as impact on protein structure, impact on exonic splicing, etc., a reliability measure should be provided for each prediction. Without such information, users will have difficulty assessing which prediction is the most likely to be correct.
- (v) A method to integrate diverse outputs of heterogeneous data types and to put them in perspective. Without algorithms to integrate and prioritize information available about a SNP, many users will come away with nothing of value.
- (vi) Ability to handle all kinds of SNPs, possibly through linkouts to other servers. Users will often not know in advance whether SNPs of interest impact protein function or regulation. It is annoying to submit SNPs and find that there is no information about them because you have chosen an inappropriate server.

- (vii) Ability to report other SNPs in linkage disequilibrium with submitted SNPs. If submitted SNPs are indirectly linked to disease, users will benefit by discovering which other SNPs might be responsible, so that their biological impact can be investigated.
- (viii)Up-to-date data. The dbSNP database is updated several times a year. The number of new human SNP reference IDs ranges widely (e.g. 44 000 in Build 127, over 6 000 000 in the current Build 129). When SNP webservers do not keep up with these updates, users miss out on coverage of thousands to millions of SNPs.

HOW DIFFERENT ARE THE VARIOUS SNP ANNOTATION METHODS?

A review of current literature reveals that medical geneticists are grappling with issues surrounding the meaning of agreement and disagreement among available SNP annotation methods.

- In a meta-analysis study that included computational biology nsSNP methods, predictive scores (for 54 nsSNPs in 37 genes) were compared to lung cancer risk odds ratios from 51 published case-control studies, using a non-parametric correlation test (Spearman rank) [19]. The authors designed a summary statistic which combined scores from SIFT, PolyPhen, SNPs3D [16] and PMut and reported that the summary was more highly correlated with the lung cancer risk odds ratios (r = 0.51) than any of the individual scores. The correlation increase was modest with respect to SIFT, the most highly correlated individual score (r = -0.36). The rationale for combining scores produced by different methods was that each method uses a 'fundamentally different algorithm', and that when the algorithms agree, predictions are more trustworthy.
- (ii) In a case—control study of nsSNPs in nucleotide excision repair genes, putatively linked with prostate cancer [63], SIFT and PolyPhen were used to explore the possible biological impact of seven nsSNPs with significant association to prostate cancer and minor allele frequency > 0.05. The methods disagreed on four nsSNPs and for two out of three on which they agreed, a functional nucleotide excision repair capacity (NERC) assay disagreed with both. The authors tried to explain these disparities by

suggesting that PolyPhen uses protein structure information, while SIFT uses evolutionary sequence conservation, but this is not generally true, as described below.

Although users may perceive SNP prediction services as a set of fundamentally different methods, there are major similarities 'underneath the lid'. For example, SIFT, PolyPhen's PSIC (Position Specific Independent Counts) score and 'SNPs3D Profile SVM' (support vector machine) all base their predictions on a multiple sequence alignment of the protein of interest and related proteins. Although PolyPhen does use protein structural information when it is available, for the majority of queries, its predictions are based on amino acid residue properties and PSIC sequence alignment scores [64]. Like SIFT, the PSIC score measures the probability that a substituted amino acid will be tolerated, based on the distribution of amino acids in a multiple sequence alignment column. The measures differ mainly in technical details, such as how pseudocounts and sequence weighting are applied. When SIFT and PolyPhen outputs are substantially different, it is probably because different multiple sequence alignments were used to calculate scores, rather than these details. Inferences based on amino acid column distributions are also used in PANTHER, and as input features to machine learners LS-SNP, SNPs3D, SNAP and PMut. While the decision algorithms used by these different methods are not the same, the correlation among their outputs is the result of similarity among their inputs, and is not necessarily ground for increased confidence.

The authors of the lung cancer meta-analysis assumed that the two 'SNPs3D SVMs' ('SVM Profile' and 'SVM structure') could be grouped together because they are more similar to each other than either one is to SIFT. Emphasis on the SVM algorithm caused them to miss the fundamental similarity between 'SVM Profile' and SIFT. A better choice for the summary statistic would be 'SVM structure', because it is based on protein structure, and provides an orthogonal prediction to methods based on sequence alignment.

As scientists outside of the bioinformatics community attempt to optimize their use of SNP prediction methods, those within the community must make an effort to better communicate the inner workings of these methods and to clarify both their similarities and differences.

SNP WEBSERVERS: CHALLENGES FOR THE NEXT GENERATION

SNP webservers of the first generation were created by bioinformaticians for bioinformaticians. A major challenge for next generation tools is how to deliver utility to medical geneticists and molecular biologists.

Flexible input tools that can handle high throughput data

Users should have the option of entering from one to thousands of SNPs, including novel SNPs. FASTSNP already allows entry of genes of interest and returns a list of all known SNPs, which can then be selected for annotation. But as the number of candidate SNPs of interest increases, manual selections will not be feasible. Users should be able to enter SNP lists in the form that they receive them from sequencing centers (DNA base change, chromosome position and transcript identifier) or to directly submit the output files from *Illumina bead arrays* or *Affymetrix genotyping arrays*.

Leverage of genome correlation structure

Users should be able to find out if their SNPs are in linkage disequilibrium with other SNPs having known or predicted functional effects. MutaGeneSys already allows users to select a preferred correlation threshold and find SNPs listed in *OMIM* that are in linkage disequilibrium with input SNPs. Such capabilities can be expanded to SNPs having predicted functional impact on regulation or protein function.

Other types of genetic variation

The causative mutation sought in association studies may turn out to be a copy number variant, an inversion, deletion, insertion or frameshift. As other kinds of genetic variation are catalogued, it will be useful both to annotate them and to provide information about linkage disequilibrium between SNPs and these variants.

Cis-regulation

Associations between phenotype and intronic, UTR, and/or promoter region SNPs are prominent in case/control and family studies published over the last several years [65-76]. Yet computational methods to predict the effects of these SNPs lag behind

those developed for their impact on proteins. We do not know yet how to accurately detect the sequence signals that identify sites important for transcription, splicing, or miRNA binding, or how to score the impact of a SNP on these sites. Advances in basic science and computational analysis of these elements will play an important role in advancing the utility of SNP webservers.

Inference framework

In addition to serving hypotheses about molecular mechanisms, servers should offer the option of integrating multiple hypotheses and molecular features into a decision algorithm. Without such a framework, and given the growing number of known regulatory mechanisms, users have difficulty for making sense of available information, particularly when harvested by meta-servers. FASTSNP already offers a decision tree framework to integrate information into a risk level (1–5).

Dynamic visualization and analysis tools

The outputs of many servers include protein sequence alignments, structural models, model viewers and structural features. But protein and SNP representations using ribbons, balls and sticks, and multiple sequence alignments cannot provide biological insight to anyone but a protein expert, even if the graphics are interactive. We can maximize the utility of these tools by designing them to help users gain intuition about SNP effects, such as the impact of amino acid substitution. Interactive protein structure graphics could be pre-annotated by 'painting' according to biologically important attributes, such as electrostatic surface potential, and the tools could allow users to see how these attributes change with amino acid substitution. Interactive multiple sequence alignment graphics could dynamically display relevant statistics, such as probability that a given amino acid substitution is tolerated in an alignment column. New tools could allow users to experiment selecting different amino acids and to view how the tolerance probability changes.

Dynamic data updates

Most current SNP webservers are based in academic labs and are not supported by full-time staff. Furthermore, these servers were designed to store data locally, requiring regular downloads from their

primary sources (such as NCBI, UCSC Genome Browser, UniProt, etc.) and subsequent rerunning of annotation pipelines. It is not surprising that most servers are 2 years or more out-of-date. FAST-SNP and SNPit (http://students.washington.edu/hyshen/ research.html which is not yet publicly available) have already made progress on this problem. FAST-SNP uses reconfigurable web wrapper agents to fetch HTML pages, extract relevant data, deliver to a Web Navigation Description Language (WNDL) executor kernel and then to its machine learning algorithm, which renders a decision about SNP risk level. SNPit's wrappers are HTTP servlets that accept queries as URLs and return XML formatted data. It uses a BioMediator 'source knowledge base', composed of a central data model and rules to translate the source data models into the common data model. These distributed data integration technologies help ensure that data delivered to the user is up-to-date, although there is nothing they can do about stale data at their sources.

SUMMARY AND CONCLUSIONS

As a whole, the 22 SNP annotation webservers assessed in this study yielded interesting hypotheses to explain why several SNPs might be statistically associated to either ALS, schizophrenia or esophageal cancer in recent medical genetics studies. However, these hypotheses were not immediately apparent and required bioinformatics expertise to sift out from a wide array of 'black box' classifications, technical details and predictive scores spanning evolutionary conservation, protein structure, splicing regulators, transcriptional regulators, etc.

The next generation of SNP annotation webservers can take advantage of the growing amount of data in core bioinformatics resources and use intelligent agents to fetch data from different sources as needed. From a user's point of view, it is more efficient to submit a set of SNPs and receive results in a single step, which makes meta-servers the most attractive choice. However, if meta-servers deliver heterogeneous data covering sequence, structure, regulation, pathways, etc., they must also provide frameworks for integrating data into a decision algorithm(s), and quantitative confidence measures so users can assess which data are relevant and which are not. Without progress along these lines, all of this data will only be useful to bioinformatics experts.

Key Points

- Computational biology methods for SNP annotations can maximize their contributions to medical genetics research by designing services that are easy for researchers who are not bioinformatics experts to use and understand.
- Medical geneticists and molecular biologists who are interested in using available SNP annotation web servers can select from: (i) servers that disseminate original methods to predict biologically important SNPs; (ii) metaservers, which yield large amounts of heterogeneous bioinformatics information from external servers; and (iii) hybrids which combine (i) and (ii).
- There is more similarity among bioinformatics SNP annotation methods than many users realize.
- Developing new algorithms for integrating heterogeneous datatypes is now essential to take advantage of the available information, which can potentially be used to infer the biological impact of SNPs.

SUPPLEMENTARY DATA

Supplementary data are available online at http://bib.oxfordjournals.org/.

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