Integrative Annotation of Variants from 1092 Humans: Application to Cancer Genomics

Ekta Khurana, Yao Fu, Vincenca Colonna, Xinmeng Jasmine Mu, Hyun Min Kang, Tuuli Lappalainen, Andrea Sboner, Lucas Lochofsky, Jiemei Chen, Arif Harmanli, Jishnu Das, Alexej Abyzov, Suganthi Balasubramanian, Kathryn Beal, Dimple Chakravarty, Daniel Challis, Yuan Chen, Declan Clarke, Laura Clarke, Fiona Cunningham, Uday S. Evani, Paul Flicek, Robert Fraga, Erik Garrison, Richard Gibbs, Zeynep H. Güümüş, Javier Herrero, Naoki Kitabayashi, Yong Kong, Kasper Lage, Vaja Liluashvili, Steven M. Lipkin, Daniel G. MacArthur, Gabor Marth, Donna Muzny, Tune H. Pers, Graham R. S. Ritchie, Jeffrey A. Rosenfeld, Cristina Sisu, Xiaomu Wei, Michael Wilson, Yali Xue, Fuli Yu, 1000 Genomes Project Consortium, Emmanouil T. Dermitzakis, Haiyuan Yu, Mark A. Rubin, Chris Tyler-Smith,* Mark Gerstein*

Introduction: Plummeting sequencing costs have led to a great increase in the number of personal genomes. Interpreting the large number of variants in them, particularly in noncoding regions, is a current challenge. This is especially the case for somatic variants in cancer genomes, a large proportion of which are noncoding.

Methods: We investigated patterns of selection in DNA elements from the ENCODE project using the full spectrum of variants from 1092 individuals in the 1000 Genomes Project (Phase 1), including single-nucleotide variants (SNVs), short insertions and deletions (indels), and structural variants (SVs). Although we analyzed broad functional annotations, such as all transcription-factor binding sites, we focused more on highly specific categories such as distal binding sites of factor ZNF274. The greater statistical power of the Phase 1 data set compared with earlier ones allowed us to differentiate the selective constraints on these categories. We also used connectivity information between elements from protein-protein-interaction and regulatory networks. We integrated all the information on selection to develop a workflow (FunSeq) to prioritize personal-genome variants on the basis of their deleterious impact. As a proof of principle, we experimentally validated and characterized a few candidate variants.

Results: We identified a specific subgroup of noncoding categories with almost as much selective constraint as coding genes: “ultra-sensitive” regions. We also uncovered a number of clear patterns of selection. Elements more consistently active across tissues and both maternal and paternal alleles (in terms of allele-specific activity) are under stronger selection. Variants disruptive because of mechanistic effects on transcription-factor binding (i.e., “motif-breakers”) are selected against. Higher network connectivity (i.e., for hubs) is associated with higher constraint. Additionally, many hub promoters and regulatory elements show evidence of recent positive selection. Overall, indels and SVs follow the same pattern as SNVs; however, there are notable exceptions. For instance, enhancers are enriched for SVs formed by nonallelic homologous recombination. We integrated these patterns of selection into the FunSeq prioritization workflow and applied it to cancer variants, because they present a strong contrast to inherited polymorphisms. In particular, application to ~90 cancer genomes (breast, prostate and medulloblastoma) reveals nearly a hundred candidate noncoding drivers.

Discussion: Our approach can be readily used to prioritize variants in cancer and is immediately applicable in a precision-medicine context. It can be further improved by incorporation of largescale population sequencing, better annotations, and expression data from large cohorts.

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Integrative Annotation of Variants from 1092 Humans: Application to Cancer Genomics

Ekta Khurana,1,2* Yao Fu,1,3 Vincenza Colonna,3,4 Xinmeng Jasmine Mu,1,4 Hyun Min Kang,5 Tuuli Lappalainen,7,7,9 Andrea Shoner,7,10 Lucas Lockovsky,1,9 Jiemin Chen,1,11 Arif Harmanci,1,2 Jishnu Das,12,13 Alexej Abzyov,1,2 Suguantli Balasubramanian,1,2,14 Kathryn Beal,14 Dimple Chakravarty,9 Daniel Challis,15 Yuan Chen,9 Declan Clarke,16 Laura Clarke,14 Fiona Cunningham,14 Uday S. Evan,15 Paul Flicek,14 Robert Fraga,13,17 Erik Garrison,18 Richard Gibbs,17 Zeynep H. Güümü,10,19 Javier Herrero,20 Naoki Kitabayashi,7 Yong Kong,20 Kasper Lage,21,22,24,25 Vaja Liluashvili,10,19 Steven M. Lipkin,26 Daniel G. MacArthur,22,27 Richard Gibbs,15 Zeynep H. Gümüş,10,19 Javier Herrero,20 Naoki Kitabayashi,7 Yong Kong,20 Kasper Lage,21,22,24,25 Vaja Liluashvili,10,19 Steven M. Lipkin,26 Daniel G. MacArthur,22,27 Gabor Marth,18 Donna Muzny,15 Tune H. Pers,28,29 Graham R. S. Ritchie,14 Jeffrey A. Rosenfeld,30,31,32 Cristina Siu,1,2 Xiaomui Wei,13,26 Michael Wilson,1,33 Yali Xue,1 Fuji Yu,15 1000 Genomes Project Consortium,1,16 Emmanouil T. Dermitzakis,6,7,8 Haiyuan Yu,12,13 Mark A. Rubin,9 Chris Tyler-Smith,3,‡ Mark Gerstein1,2,34 ‡

Interpreting variants, especially noncoding ones, in the increasing number of personal genomes is challenging. We used patterns of polymorphisms in functionally annotated regions in 1092 humans to identify deleterious variants; then we experimentally validated candidates. We analyzed both coding and noncoding regions, with the former corroborating the latter. We found regions particularly sensitive to mutations ("ultrasensitive") and variants that are disruptive because of mechanistic effects on transcription-factor binding (that is, "motif-breakers"). We also found variants in regions with higher network centrality tend to be deletions. Insertions and deletions followed a similar pattern to single-nucleotide variants, with some notable exceptions (e.g., certain deletions and enhancers). On the basis of these patterns, we developed a computational tool (FunSeq), whose application to ∼90 cancer genomes reveals nearly a hundred candidate noncoding drivers.

Whole-genome sequencing has revealed millions of variants per individual. However, the functional implications of the vast majority of these variants remain poorly understood (1). It is well established that variants in protein-coding genes play a crucial role in human disease. Although it is known that noncoding regions are under negative selection and that variants in them have been linked to disease, their role is generally less well understood (2–9).

In particular, whereas some studies have demonstrated a link between common variants from genome-wide association studies (GWASs) and regulatory regions (2, 3), the deleterious effects of rare inherited variants and somatic cancer mutations in noncoding regions have not been explored in a genome-wide fashion. Recently, three studies reported noncoding driver mutations in the TERT promoter in multiple tumor types, including melanomas and gliomas (10–12). In light of these studies and the growing availability of whole-genome cancer sequencing (13–20), an integrated framework facilitating functional interpretation of noncoding variants would be useful.

One may think to identify noncoding regions under strong selection purely through mammalian sequence conservation, and ultraconserved elements have been found in this fashion (21). However, signatures of purifying selection identified by using population-variation data could provide better insights into the importance of a genomic region in humans than evolutionary conservation. This is because many regions of the genome show human-specific purifying selection, whereas other regions conserved across mammals show a lack of functional activity and selection in humans (7). Thus, identifying the specific elements under particularly strong purifying selection among humans could provide novel insights.

Besides single-nucleotide polymorphisms (SNPs), the human genome also contains other variants, including small insertions and deletions (indels) and larger structural variants (SVs) (22). They account for more nucleotide differences among humans than SNPs; hence, an understanding of their relationship with functional elements is crucial (23).

We used the full range of sequence polymorphisms (ranging from SNPs to SVs) from 1092 humans to study patterns of selection in various functional categories, especially noncoding regulatory regions (24). We identified specific genomic regions where variants are more likely to have strong phenotypic impact. The list of these regions includes groups of coding genes and specific sites within them and, importantly, particular noncoding elements. By further comparing patterns of polymorphisms with somatic mutations, we show how this list can aid in the identification of cancer drivers. We used multiple experimental methods for validation, including yeast two-hybrid experiments, Sanger sequencing of independent cancer samples, and relevant gene-expression measurements. Furthermore, we provide a software tool that allows researchers to prioritize noncoding variants in disease studies.

Genomic Elements Under Strong Purifying Selection: Ultrasensitive Regions

Enrichment of rare variants can be used to estimate the strength of purifying selection in different functional categories (24). As expected, we found that having variants from 1092 individuals allowed us to detect specific functional categories under strong purifying selection with greater confidence than possible with smaller datasets.
power than previously possible (2, 7, 9). In particular, the increased number of samples provided a better estimate of allele frequencies, making possible the measurement of differential selective constraints between specific categories [e.g., between motifs of transcription-factor (TF) families HMG and MADS box] (Figs. S4 and S5).

Estimates of purifying selection obtained by using enrichment of rare nonsynonymous SNPs (derived allele frequency or DAF < 0.5%) showed that different gene categories exhibit differential selection consistent with their known phenotypic consequences (data S1). Genes tolerant of loss-of-function (LoF) mutations are under the weakest selection, whereas cancer-causal genes are under the strongest (Fig. 1A and table S1). GWAS genes associated with complex disorders lie in between these extremes, consistent with the presence of common genetic variants in them.

We then analyzed selective constraints in non-coding regions, trying to find elements under very strong selection (i.e., with a fraction of rare variants similar to that of coding genes, ~67%). We first estimated the strength of negative selection in broad categories [e.g., in all TF binding sites (TFBSs), deoxyribonuclease I (DNaseI)-hypersensitive sites (DHSs), noncoding RNAs (ncRNAs), and enhancers] (Fig. 2A). As observed previously, most of these categories show slight but statistically significant enrichment of rare SNPs compared with the genomic average; in contrast, pseudogenes demonstrate a depletion (Fig. 2A and data S2) (2).

We further divided the broad categories into 677 high-resolution ones. These span various genomic features likely to influence the extent of selection acting on the element. For example, TFBSs of different TF families are divided into proximal versus distant and cell-line-specific versus nonspecific (fig. S7). We find heterogeneous degrees of negative selection for specific categories (Fig. 2B and data S2). For instance, core motifs in the binding sites of TF families HMG and Forkhead are under particularly strong selection, whereas those in the CBF/NFY family do not exhibit selective constraints (relative to the genomic average) (Fig. 2B). Among all the pseudogenes, polymorphic ones have the highest fraction of rare alleles, consistent with their functional coding roles in some individuals (25). Overall, we found that 102 of the 677 categories show statistically significant selective constraints (data S2) (Figs. S8 to S10).

Among these 102 categories, we defined the top ones covering ~0.02% and ~0.4% of the genome as ultrasensitive and sensitive, respectively (fig. S11) (data S3). Thus, these regions were defined such that they possess a high fraction of rare variants comparable to that for coding sequences (67.2% for coding and 65.7% for ultrasensitive) (Fig. 2C). We validated the rare variants in them by comparison with Complete Genomics data. Sensitive regions include binding sites of some chromatin and general TFs (e.g., BRF1 and FAM48A) and core motifs of some important TF families (e.g., JUN, HMG, Forkhead, and GATA). For some TFs, there is a strong difference between proximal and distant binding sites—for example, for ZNF274, proximal binding sites are under strong selection and belong to the ultrasensitive category, whereas distal sites are not under negative selection.

In order to validate the functional importance of sensitive and ultrasensitive regions, we examined the presence of inherited disease-causing mutations from HGMD (Human Gene Mutation Database) in them (26). We found ~40- and ~400-fold enrichment of disease-causing mutations in sensitive and ultrasensitive regions, respectively (compared with the entire noncoding sequence, P < 2.2 × 10^-16) (Fig. 2E). Thus, these documented disease-causing variants provide independent validation for the functional importance of sensitive regions. As a specific example, the disease congenital erythropoietic porphyria is caused by disruption of a binding site classified as sensitive (the GATA1 motif upstream of uroporphyrinogen-III synthase) (27). Similarly, the well-known disease-causing ncRNA RMRP is in the binding site of BRF2, classified as ultrasensitive (28).

**Purifying Selection and Other Aspects of Regulatory Regions**

We analyzed sites at which SNPs break or conserve core-binding motifs. As expected, we found that disruptive motif-breaking SNPs are significantly enriched for rare alleles compared with motif-conserving ones (P = 2.2 × 10^-16; Fig. 2D; a motif-breaking SNP is defined as a change that decreases the matching score in the motif position weight matrix). This result is over all TF families; moreover, we find the difference between constraints on motif-breaking versus -conserving SNPs varies considerably for different TF families, possibly reflecting differences in the topology of their DNA binding domains (data S4).

We also found that expression quantitative trait loci (eQTLs) are enriched in the binding sites of many TF families (Fig. 2B); the association of TF binding and gene expression at these loci provides a plausible explanation for their phenotypic effects.

An analysis of SNPs from a personal genome (NA12878) exhibiting allele-specific TF binding in chromatin immunoprecipitation sequencing (ChIP-Seq) data or allele-specific expression in RNA-seq data (with the allele-specific “activity” tagging a difference between maternal and paternal chromosomes at the genomic region in question) showed that these sites are depleted for rare variants (relative to a matched control) (Fig. 2F). This suggests that regions where differential allelic activity is not observed may be under stronger purifying selection (29).

In a similar fashion, we found that core-motif regions bound in a “ubiquitous manner” (i.e., where differential cell-type-specific binding is not observed) are under stronger selection than those bound by TFs in a single cell line (data S2), consistent with the greater functional importance of ubiquitously bound regions. In relation to this, we further examined how selective constraints vary among coding genes and DHSs with tissue-specific activity (Fig. 1B). We found there are pronounced differences between tissues: For example, genes with ovary- and brain-specific ex-
pression are under significantly stronger selection than the average across all tissues (Fig. 1B and table S4). Similarly, some DHSs are under significantly stronger selection, whereas others are under relaxed constraints relative to the average (brain- and kidney-specific versus urothelium- and breast-specific, respectively: Fig. 1B and table S4). Last, matched expression and DHS data for six tissues indicate that purifying selection in tissue-specific genes and their corresponding regulatory regions is likely correlated (fig S15). Thus, our results suggest that the deleteriousness of both coding and regulatory variants depends on the tissues they affect.

**Purifying Selection in the Interactome and Regulome**

We found a significant positive correlation between the fraction of rare SNPs and the degree centrality of genes in networks: physical protein-protein interaction (PPI) (rho = 0.15; P < 2.2 × 10^{-16}) and regulatory (rho = 0.07; P = 6.8 × 10^{-09}). Thus, consistent with previous studies, we found that hub genes tend to be under stronger negative selection (29–31). Indeed, centralities of different gene categories in the PPI network follow the same trend as differential selective constraints on them: Cancer-causal genes show the highest connectivity, and LoF-tolerant genes, the least, with GWAS genes in the middle (Figs. 1A and 3A). These results indicate that the interactions of a gene likely influence the selection acting on it.

Hub proteins tend to have more interaction interfaces in the PPI network (31). A corollary of this is that interaction interfaces are themselves under strong selection, in turn leading to stronger constraints on hub proteins. Indeed, we found that SNPs disrupting interaction interfaces are enriched for rare alleles (P < 2.2 × 10^{-16}) (Fig. 3B).

To further corroborate this, we tested a specific case, the Wiskott-Aldrich syndrome protein (WASP), using yeast two-hybrid (Y2H) experiments (32). All of the three tested single-nucleotide variants (SNVs) at WASP interaction interfaces disrupted its interactions with other proteins (Fig. 3C). We observed similar behavior for two other proteins: Mutations at their interfaces disrupted specific protein interactions (fig. S16).

**Relationship of Functional Elements with Indels and Larger SVs**

We analyzed the association of functional annotations with small indels (<50 base pairs (bp)) and large SVs (deletions). Similar to the results for nonsynonymous SNPs, we found that genes linked with diseases show stronger selection against indels whereas LoF-tolerant genes show weaker constraints (relative to all genes), with a consistent trend for indels overall and frameshift indels, in particular (Fig. 4A, fig. S17, and table S1).

The wide range of SV sizes (~50 bp to ~1 Mb) leads to their diverse modes of intersection with functional elements; for example, a single SV breakpoint can split an element, a smaller SV can cut out a portion of a single element, and a large SV can engulf an entire element. To analyze the diverse effects of SVs, we computed the enrichment or depletion of SVs overlapping each

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**Fig. 2. Fraction of rare SNPs in noncoding categories.** Red dotted lines represent genomic average. Error bars denote 95% binomial confidence intervals. Total numbers of SNPs in each category shown. (A) Broad categories. Ultratransient and sensitive regions are those under very strong negative selection. TFSS, sequence-specific TFs. Categories tested for enrichment of HGM D sites (Fig. 5A) marked by using hollow triangles on the left. (B) Example of high-resolution categories. TFBS motifs separated into 15 families, e_superscripts in red denote enrichment of eQTLs in TFBSs of specific families. (C) Examples of TFBSs included in ultraconservative category. (D) SNPs breaking TF motifs show an excess of rare alleles compared with those conserving them. Representative motifs for two families are shown. (E) Enrichment of HGM D regulatory disease-causing mutations in ultratransient, sensitive, and annotated regions compared with all noncoding regions. (F) SNPs not exhibiting allele-specific behavior (−) are enriched in rare alleles compared with SNPs exhibiting allele-specific behavior (+).
DAF (HighD sites) (candidates for positive selection: sites where con- 
standing variation) likely also played a major 
selection via other modes (such as selection on 
the classic selective-sweep model (22)). However, when we broke down the 
mode of SV intersection with genes into partial versus whole (an SV breakpoint splitting a gene versus an SV engulfing a whole gene), we unex-
pectedly found that SVs are enriched for whole-
but depleted for partial-genome overlap. This suggests that 
partial-genome overlap is under stronger selec-
tion than whole-genome overlap, possibly because 
whole-genome deletions may be compensated by 
duplications. Furthermore, another category of gene-
related elements, pseudogenes, are enriched for 
SVs, consistent with their formation mechanism 
involving either duplication or retrotransposition.

In relation to nongenic elements, we found that 
SVs tend to be depleted in regulatory elements 
such as binding-site motifs and enhancers (Fig. 
4B), consistent with our expectations from SNPs. 
However, enhancer elements are enriched for SVs 
formed by nonallelic homologous recombination 
(NAHR). This observation is further supported by 
the high signal of activating histone marks asso-
ciated with enhancers (e.g., H3K4me1) around 
NAHR breakpoints (Fig. 4C and fig. S18). The 
association of enhancers and NAHR deletions 
may be explained by the three-dimensional struc-
ture of chromatin bringing enhancer elements 
into close proximity with the gene transcription 
start site (via DNA “looping”). If these two “non-
allelic” loci contain homologous sequences, it 
would be favorable for NAHR to occur.

Functional Implications of Positive Selection 
Among Human Populations
Negative selection is widespread in the genome; 
nevertheless, some positions within negatively 
selected regions also experience positive selec-
tion (33–36). We have previously identified and 
validated one category of variants that are strong 
candidates for positive selection: sites where con-
tenational populations show extreme differences in 
DAF (HighD sites) (24). By analyzing these HighD 
sites, we are focusing on positive selection under 
the classic selective-sweep model (37). Positive 
selection via other modes (such as selection on 
standing variation) likely also played a major 
role in recent human evolution (38). Nonethe-
less, functional annotation of HighD sites can 
provide important insights about recent adapta-
tions (39).

We examined positive selection in the same 
fashion as we have done for negative selection: 
in coding genes, noncoding regulatory elements, 
and networks of gene interactions. The func-
tional analysis of positive selection using highly 
differentiated sites is limited to SNPs, because of 
the low numbers of such indels and SVs in 
fundamental elements.

We observed enrichment of HighD sites in 
UTRs and missense SNPs in coding regions 
(Fig. 5A). Next, we observed that some disease 
gene groups (Online Mendelian Inheritance in 
Man, HGMD, and GWAS) are enriched for HighD 
SNPs (fig. S20). Mutations in disease genes are 
likely to have strong phenotypic impact; thus, it 
is possible that some of these mutations confer 
advantage for local adaptation. For example, 
whereas LoF mutations in ABCA12 lead to the 
severe skin disorder harlequin ichthyosis (40), 
we found that a SNP within the second intron of 
this gene is a HighD site (DAF > 90% in Europe 
and East Asia; 13% in Africa), possibly reflecting 
adaptations of the skin to levels of sunlight 
outside of Africa.

Similar to our analysis of negative selection, 
we analyzed the enrichment of HighD sites in 
broad and specific noncoding categories, finding 
significant enrichment in many noncoding cate-
gories (Fig. 5A). These enriched categories in-
clude DDISS (particularly distal ones) and binding 
sites of sequence-specific TFs (specifically those 
in ZNF and NR families). Out of the seven en-
riched categories, five are also under significant 
negative selection (Figs. 2A and 5A and data 
S2). Thus, even though an entire category might 
be under negative selection, some particular sites 
within it can be targets of positive selection. In 
this respect, our results are consistent with pre-
vious studies for missense SNPs: Overall they 
are under strong negative selection, but a small 
group of them have targets of positive selec-
tion (36).

We found that, as expected, coding genes with 
HighD SNPs tend to have lower degree centrality 
in both PPI and regulatory networks (although 
the small number of these cases does not produce 
statistical significance) (Fig. 5B and fig. S21) (41). 
In an opposite trend to genes (where positive se-
lection occurs on the network periphery), HighD 
sites in TFBSs tend to occur in hub promoters ($P = 
0.02$ with 23 promoters and $P = 3.2 \times 10^{-3}$ with

![Fig. 3. SNPs in protein-protein interaction (PPI) network. (A) Degree centrality of coding-gene categories in PPI network. (B) Fraction of rare missense SNPs at protein-interaction interfaces is higher than all rare missense SNPs (error bars show 95% binomial confidence intervals; total number of SNPs also shown). (C) Effects of SNVs at interaction interfaces on interactions of WASP with other proteins tested by Y2H experiments. Wild-type (WT) WASP interacts with all proteins shown, whereas each missense SNV disrupts its interaction with at least one protein.](image-url)
proximal TFBSs) (Fig. 5B). It was previously proposed that mutations in cis elements in regulatory networks may play an important role in development (42, 43); our study supports this by suggesting that some hub promoters may have undergone recent adaptive evolution.

Contrasting Patterns of Somatic Mutations with Inherited Variants

After analyzing inherited polymorphisms in functional elements, we examined somatic variants. Because somatic variants from diverse tumors exhibit different sets of properties, we analyzed variants from a wide range of cancer types: prostate, breast, and medulloblastoma (17, 19, 20). We found that ~99% of somatic SNVs occur in noncoding regions, including TFBSs, ncRNAs, and pseudogenes (fig. S22).

Analysis of matched tumor and normal tissues from the same individuals showed that somatic variants tend to be enriched for missense (~5×), LoF (~14×), sensitive (~1.2×), and ultrasensitive (~2×) variants (Fig. 6A, fig. S24, and table S6). Consistent with this trend, we found higher TF-motif-breaking/conserving ratios for somatic variants compared with germline ones across many different samples and cancer types (~3 for somatic versus ~1.4 for germline) (Fig. 6B and table S7). Thus, somatic-cancer variants are generally enriched for functionally deleterious mutations.

This enrichment of functionally deleterious mutations among somatic variants is understandable because they are not under organism-level natural selection (unlike inherited-disease mutations, including GWAS variants). Indeed, among all somatic mutations, those most deviating from patterns of natural polymorphisms are the most likely to be cancer drivers. Consistent with this, our analysis has shown that, among all disease mutations, those causing cancer occur in genes under strongest negative selection (and with highest network connectivity) (Figs. 1A and 3A). Thus, we argue that somatic variants in the noncoding elements under strongest selection are the most likely to be cancer drivers.
Another feature of somatic mutations associated with their potential role as drivers is their recurrence in the same genomic element across multiple cancer samples. We found that some non-coding elements from our functional categories show recurrent mutations (fig. S23). For example, the pseudogene RP5-857K21.6 is mutated in three out of seven prostate cancer samples, and the promoter of RP1 is mutated in two.  

**FunSeq: Tool for Identification of Candidate Drivers in Tumor Genomes**  
On the basis of the integrative analysis above, we developed a tool to filter somatic variants from tumor genomes and obtain a short list of candidate driver mutations (funseq.gersteinlab.org). FunSeq first filters mutations overlapping 1000 Genomes variants and then prioritizes those in regions under strong selection (sensitive and ultraselective), breaking TF motifs, and those associated with hubs. It can score the deleterious

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**Fig. 5. Functional implications of positive selection.** (A) (Left) Frequency of HighD SNPs versus matched sites for broad categories (marked by hollow triangles in Fig. 2A). (Right) Specific categories, e.g., specific TF families. Asterisk denotes significant enrichment after multiple-hypothesis correction. e superscripts in red denote the enrichment of eQTLs. (B) (Left) The in-degree of genes with HighD missense SNPs is lower than that of all genes. (Center) The in-degree of genes with HighD SNPs in their promoters is higher than all genes. (Right) The human regulatory network with edges in gray. Red nodes represent genes with HighD SNPs in their promoters, and blue nodes represent genes with HighD missense SNPs. Size of nodes scaled based on their degree centrality. Nodes with higher centrality are bigger and tend to be in the center, whereas those with lower centrality are smaller and tend to be on the periphery.
Fig. 6. Functional interpretation of disease variants. (A) Enrichment of functionally deleterious mutations among somatic compared with germline SNVs. Mean values from seven prostate cancer samples shown (variation shown in fig. S16). (B) Ratios for the number of SNVs that conserve versus break TF-binding motifs depicted for NA12878, the average of 1000 Genomes Phase I samples, and the average of somatic and germline samples from different cancers. Error bars represent 1 SD. MB, medulloblastoma. (C) Filtering of somatic variants from a breast (PD4006, left) and a prostate (PR-2832, right) cancer sample leading to identification of candidate drivers. (D) A part of the FAM48A binding site sequenced by Sanger sequencing in an independent cohort of 19 prostate cancer samples shown in green (with the coordinates of mutations observed in one sample). (E) Application of variants filtering scheme to Venter personal genome. Number of SNVs in various categories shown.
potential of variants in single or multiple genomes and output the results in easy-to-use formats (i.e., “decorated” variant call format files, fig. S29 and data S6). The scores for each noncoding variant vary from 0 to 6, with 6 corresponding to maximum deleterious effect. When multiple tumor genomes are given as input, FunSeq also identifies recurrent mutations in the same element. Although our emphasis is on noncoding variants, it also outputs scores for coding variants.

We demonstrate the application of FunSeq as a workflow on representative breast and prostate cancer genomes (Fig. 6C). In the breast cancer sample, the workflow yielded one noncoding SNV likely to have strong phenotypic consequences: This SNV (i) occurs in an ultrasegment region (BRF2 binding site); (ii) interacts with a PAX-5 TF binding motif; (iii) is associated with a network hub (44); and (iv) is recurrent—that is, the regulatory module contains somatic mutations in multiple breast-cancer samples. In a similar fashion, the prostate-cancer sample revealed two noncoding SNVs predicted to have strong functional consequences (Fig. 6C). One of these is in an ultrasegment region (FAM48A binding site) and lies in the promoter of HNR74 gene (a hub in the PPI network with degree centrality = 56). We further tested the presence of mutations in this binding site by polymerase chain reaction followed by Sanger sequencing in an independent cohort of 19 prostate-cancer samples (45). We found that one sample in the cohort also harbors mutations in this region (Fig. 6D and fig. S25). Furthermore, we also observed increased expression of WDR74 in the tumor relative to benign samples (fig. S26). These experimental results provide support for a likely functional role of this candidate driver.

A large-scale application of our tool to three medulloblastoma, 21 breast, and 64 prostate cancer genomes provided a total of 98 noncoding candidate drivers (table S8 and data S6) (17–20). Among these candidates, 68 occur in sensitive regions, 55 break TF motifs, and 90 target network hubs.

**Generalized Identification of Deleterious Variants in Personal Genomes**

Although we envision the most effective use of our tool for tumor genomes, it can also be applied to germline sequences to identify potentially deleterious variants. We applied it to four personal genomes: Snyder, Venter, NA12878, and NA19240 (46–48). Out of ~3 million SNVs, we were able to identify ~15 (range from 6 to 26) noncoding SNVs per individual with high scores from FunSeq (>4), indicating their potential deleterious effects (Fig. 6E, tables S9 and S10, and data S6 and S7). Thus, our approach can be used to prioritize noncoding variants in personal genomes as well.

**Discussion**

We identified the sensitive and ultrasonic noncoding elements, which exhibit delemion of common polymorphisms and strong enrichment of known, inherited disease-causing mutations. Because they cover a small fraction of the entire genome (comparable to the exome), these regions can be probed alongside exome sequences in clinical studies. We found that functionally disruptive noncoding mutations tend to be under strong selection: In an analogous manner to LoF variants in coding genes, variants that break motifs in TF binding sites are selected against. There is a close relation between connectivity in biological networks and selective constraints: Higher connectivity is generally associated with higher constraint. Furthermore, selection against indels and large SVs acts in a similar fashion as against SNPs overall; however, the large size of SVs sometimes leads to a complex relation with functional elements. On the basis of these patterns of negative selection in functional elements, we developed a workflow and a corresponding software tool to prioritize noncoding variants in disease studies.

The prioritization scheme presented in our paper can be readily extended by incorporation of genomic polymorphisms from larger populations and higher-resolution functional annotations. Moreover, with the availability of RNA-seq data from large cohorts, additional genomic features such as eQTLs can be folded in. Our approach can be immediately applied in precision medicine studies to prioritize noncoding variants for follow-up characterization, particularly candidate driver mutations in cancer, and it can be further extended in the future.

**Materials and Methods**

Details of all data sets and methods are provided in the supplementary materials. A brief summary of major data sets and methods is provided here. Patterns of somatic mutations were obtained with multiple hypothesis correction while identifying regions, 55 break TF motifs, and 90 target network hubs.

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Data S1 to S7

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Integrative Annotation of Variants from 1092 Humans: Application to Cancer Genomics

Ekta Khurana, Yao Fu, Vincenza Colonna, Xinneng Jasmine Mu, Hyun Min Kang, Tuuli Lappalainen, Andrea Sboner, Lucas Lochovsky, Jieming Chen, Arif Harmanci, Jishnu Das, Alexej Abyzov, Suganthi Balasubramanian, Kathryn Beal, Dimple Chakravarty, Daniel Challis, Yuan Chen, Declan Clarke, Laura Clarke, Fiona Cunningham, Uday S. Evani, Paul Flicek, Robert Fragoza, Erik Garrison, Richard Gibbs, Zeynep H. Gm, Javier Herrero, Naoki Kitabayashi, Yong Kong, Kasper Lage, Vaja Liluashvili, Steven M. Lipkin, Daniel G. MacArthur, Gabor Marth, Donna Muzny, Tune H. Pers, Graham R. S. Ritchie, Jeffrey A. Rosenfeld, Cristina Sisu, Xiaomu Wei, Michael Wilson, Yali Xue, Fuli Yu, 1000 Genomes Project Consortium, Emmanouil T. Dermitzakis, Haiyuan Yu, Mark A. Rubin, Chris Tyler-Smith, and Mark Gerstein

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Identifying Important Identifiers
Each of us has millions of sequence variations in our genomes. Signatures of purifying or negative selection should help identify which of those variations is functionally important. Khurana et al. (1235587) used sequence polymorphisms from 1092 humans across 14 populations to identify patterns of selection, especially in noncoding regulatory regions. Noncoding regions under very strong negative selection included binding sites of some chromatin and general transcription factors (TFs) and core motifs of some important TF families. Positive selection in TF binding sites tended to occur in network hub promoters. Many recurrent somatic cancer variants occurred in noncoding regulatory regions and thus might indicate mutations that drive cancer.

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