

The IGVF catalog – from genetic variation to function

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Abstract

Genomic variation between individuals is essential for understanding how differences in the genome sequence affect molecular and cellular processes. The Impact of Genomic Variation on Function (IGVF) Consortium aims to uncover the relationships among genomic variation, genome function, and phenotypes by combining experimental techniques, such as single-cell mapping and genomic perturbation assays, with computational approaches such as machine learning-based predictive modeling. The IGVF Data and Administrative Coordinating Centers collect, analyze, and disseminate data and results from across the consortium through an open-source platform called the IGVF Catalog. This resource includes: 1) a centralized database of data and results concerning variants, protein abundance and retention, and functional assays; 2) a graph database comprising over 50 types of data collections with nearly 3 billion nodes and over 7.5 billion edges; 3) The Catalog offers public API endpoints (<https://api.catalog.igvf.org/>) and a user-friendly interface for exploring, querying, and visualizing the data at <https://catalog.igvf.org>. We expect that this open-access platform will support the broader scientific community to advance our understanding of how genomic variation influences biology and disease.

Introduction

Genomic variation between individuals provides the foundation for understanding how differences in the genome sequence influence molecular and cellular processes, ultimately shaping human phenotypes and disease susceptibility. While large-scale genomic sequencing projects have cataloged millions of variants, the functional interpretation of these variants, particularly in noncoding regions, remains a major challenge in biological and biomedical sciences [1, 2].

Several widely used resources provide invaluable observational data. For example, gnomAD characterizes population allele frequencies [1], GWAS Catalog aggregates genotype–phenotype associations [3], GTEx links genetic variation to gene expression across tissues [2], ClinVar curates clinically observed variant interpretations [4], and Open Targets integrates multi-omic evidence for drug discovery [5]. However, these databases are primarily descriptive or observational in nature: they document associations but rarely establish mechanistic causality.

The Impact of Genomic Variation on Function (IGVF) Consortium, established by the NHGRI in 2021, was created to fill this gap by systematically uncovering causal relationships between genomic variation, genome function, and phenotypes [6]. IGVF employs a dual strategy: (i) experimental approaches, including single-cell mapping technologies and genomic perturbation assays such as Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) gene editing, massively parallel reporter assays (MPRAs), and saturation mutagenesis; and (ii) computational modeling, leveraging machine learning and artificial intelligence to predict the functional consequences of genomic variants.

To coordinate and disseminate these efforts, the Data and Administrative Coordinating Centers (DACCs) collect, standardize, and integrate data generated across the consortium. These outputs are made available through the IGVF Catalog, an open-source, graph-based resource designed to serve both researchers and computational biologists. The Catalog contains diverse datasets, including the effects of coding variants on protein abundance and function, noncoding variants on enhancer activity (measured experimentally with MPRAs or inferred computationally), and variant-to-trait associations derived from integrative analyses.

All data are organized within a large-scale graph database currently encompassing over 50 types of data collections with nearly 3 billion nodes and over 7.5 billion edges, enabling users to explore interconnected layers of genomic information. To maximize accessibility, the Catalog provides both a public API (<https://api.catalogkg.igvf.org>) and an interactive user-friendly interface (<https://catalog.igvf.org>) for searching, querying, and visualizing data in real time. By moving beyond association-based resources toward a framework built on causal experimental evidence and predictive modeling, IGVF aims to accelerate the discovery of mechanisms linking genomic variation to function and to empower the broader scientific community in advancing our understanding of human biology and disease.

Implementation

Backend database design

The knowledge graph

The IGVF Catalog database is designed as a graph in which the nodes represent biological entities or concepts, whereas

the edges represent relationships (usually measurements or predictions) between these nodes (Table 1). Many nodes are represented by genomic coordinates (in either GRCh38 for human or GRCm39 for mouse), e.g. variants, genes, transcripts, and genomic elements. Other biochemical entities, such as proteins, protein (or other coding) variants, protein complexes, and drugs, are also represented as nodes. Finally, biological concepts such as tissues, cells, cell lines, phenotypes, diseases, biochemical functions, etc., are represented as nodes derived from common public ontologies (Supplementary Table 1). Informational entities such as GWAS studies, pathways, or protein–DNA binding motifs are also represented as nodes. Edges relate various nodes together; for example, genes are linked to transcripts, which are linked to proteins. Edges between ontology terms are loaded directly from the ontologies themselves, representing parent–child relationships between concepts. Experimental and predicted data (from both the public domain and IGVF) are also represented by edges between nodes; protein–protein interactions from BioGrid [7] and IntAct [8] are edges between two proteins; expression quantitative trait loci (eQTLs) are edges between variants and genes; enhancers predicted to affect gene expression are edges between genomic elements and genes, while chromatin accessibility QTLs (caQTLs) are edges between variants and genomic elements. GWAS results and functional effect predictions are edges between variants and phenotypes (ontology terms). Supplementary Table 2 provides a list of IGVF-generated datasets currently integrated into the knowledgebase.

Graph database implementation

Using a knowledge graph representation has a few advantages over an object or relational database. While the simple API functions that return nodes and edges over 1–2 “hops” could be served from nearly any database, the graph database formulation allows many more deep hop queries and can also be used as input tokens with a large language model (LLM) for retrieval-augmented generation (RAG). We have chosen to implement this graph database in a package called ArangoDB (<https://arangodb.com/>), which has several advantages over other systems. ArangoDB is not a pure graph database; it can act as an object store or relational database as well, which gives the system significant flexibility. Foremost is the inclusion of an “MDI” multidimensional index, which allows us to efficiently query based on chromosomal coordinate ranges. This is used to find, for example, which genomic elements overlap the position of a given nucleotide variant, or which genomic elements are within 10 kb of a given gene. Without this feature, we would have to precalculate linkages in genomic coordinate space for all nodes with a genomic coordinate. The system runs on a cluster of four memory-optimized cloud computing nodes. We employ node types that are suitable for high-performance databases using the NVMe protocol for highly parallel data transfers with local storage. Data ingestion is done by parsing flat text files (usually TSV, or tab-separated values) using Python scripts into JSONL (<https://jsonlines.org/>) from all the various data sources, which are then imported into the database.

Table 1. Summary statistics of the data currently loaded into the backend graph database (version 1.0)

Entity	Type	Arango collection	Source	v1.0
Variants	node	variants	FAVOR (dbSNP 155) [9, 10]+ dbNSFP [11]	1 870 783 419
Non-synonymous variants	node	coding_variants	dbNSFP	942 417 704
Genes	node	genes	GENCODE (v43) [12]	68 881
Transcripts	node	transcripts	GENCODE (v43)	272 993
Proteins	node	proteins	UniProtKB/TrEMBL [13, 14]	188 013
Protein–protein interactions	edge	proteins_proteins	IntAct, BioGRID	11 486 365
Protein complexes	node	complexes	EBI Complex Portal [15]	1687
cCREs	node	genomic_elements	ENCODE [16]	2 348 854
Accessible elements	node	genomic_elements	E2G elements	81 532 493
Accessible elements	node	genomic_elements	caQTLs	24 545
Tested elements	node	genomic_elements	ENCODE/IGVF MPRA CRISPR	165 978
Ontology terms	node	ontology_terms	<i>Various</i>	620 729
Gene annotations	edge	gene_products_terms	GO	2 518 911
eQTL	edge	variants_genes	eQTL catalogue (30 studies) [17]	13 681 389
Splice–QTL	edge	variants_genes	eQTL catalogue (30 studies)	5 244 210
CRISPR variant effects	edge	variants_genes	IGVF Variant-EFFECTS	737
caQTL	edge	variants_genomic_elements	regulomeDB [18]	5597
Variant effect predictions	edge	variants_genomic_elements	BlueSTARR [19]	7 553 977
New caQTLs	edge	variants_genomic_elements	AFGR [20], EBI catalogue	460 196
Variant effects	edge	variants_genomic_elements	IGVF MPRA (5 datasets)	24 287
Element–gene	edge	variants_biosamples	STARR-seq [21]	36 481 085
LD (4 ancestries)	edge	variants_variants	topLD [22]	5 939 629 733
Gene–transcript	edge	genes_transcripts	GENCODE (v43)	551 368
Protein–transcript	edge	transcripts_proteins	GENCODE + UniProtKB	188 086
Ontology connections	edge	ontology_terms_ontology_terms	all ontologies	2 300 978
Element effect on Gex	edge	genomic_elements_biosamples	element level MPRA	10 814
Element–gene	edge	genomic_elements_genes	ENCODE-E2G [23]	117 039 661
Element–gene	edge	genomic_elements_genes	CRISPR derived for training	10 412
Element–gene	edge	genomic_elements_genes	Perturb-Seq	1 748 304
Coding variant prediction	edge	coding_variants_phenotypes	MutPred2 [24]	641 276 378
Coding variant prediction	edge	coding_variants_phenotypes	ESM1v	454 527 037
Coding variant prediction	edge	coding_variants_phenotypes	VAMP-seq (inc Multi) [25]	183 696
Coding variant effect prediction	edge	variants_phenotypes_coding_variants	SGE [26]	15 143
Functional effect prediction	edge	variants_phenotypes	cV2F [27]	2 046 460
GWAS	edge	variants_phenotypes_studies	OpenTargets	315 159
GWAS studies	node	studies	OpenTargets	22 690
eQTL studies	node	studies	EBI QTL catalogue	33
Pathways	node	pathways	Reactome [28]	2711
Pathway hierarchy	edge	pathways_pathways	Reactome	2730
Gene–pathway	edge	genes_pathways	Reactome	146 697
Motifs	node	motifs	HOCOMOCO v11/SEMP1 [29]	624
Motif–proteins	edge	motifs_proteins	HOCOMOCO v11/SEMP1	2 282
TF allele SB	edge	variants_proteins	AdAstra [30]	1 147 815
TF allele SB	edge	variants_proteins	GVATdb [31]	4 288 875
UKBB plasma pQTLs	edge	variants_proteins	PPP [32]	42 154
Predicted TF allele SB	edge	variants_proteins	SEMP1 [33]	357 795 432
Co expression	edge	genes_genes	CoXPresdb [34]	3 538 462
Genetic interactions	edge	genes_genes	BioGrid	15 085
Drugs	node	drugs	PharmGKB [35]	4613
Drug–variant	edge	variants_drugs_genes	PharmGKB	24 695
Disease–gene	edge	diseases_genes	Orphanet [36]	8218
Disease–gene–variant	edge	variants_diseases_genes	ClinGen [37]	3820

An entity refers to a specific type of data entry, where an “element” denotes a genomic region. Type indicates whether the entry is represented as a node or an edge in the graph database, with nodes corresponding to individual data entries and edges representing connections between nodes. Arango collection specifies the identifier of the collection in the Arango database, analogous to a table in a relational database. Source denotes the origin of the data, with IGVF-generated datasets highlighted in bold. The final column reports the total counts of each data entity.

Frontend user interface

Unified search bar

The catalog features a unified search bar designed to let users “search by anything” (Fig. 1A). It supports both free-text names and structured identifiers across diverse entities such as genes, proteins, diseases, and studies. The search system provides real-time autocomplete with type-aware suggestions, and a deterministic parser recognizes standard identifiers—including rsIDs, genomic coordinates, SPDI strings [38],

PMIDs, and more—directly routing users to the appropriate page. Users can further refine queries with slash commands (e.g. /genes) to confine results to a specific entity type or enter a lexical mode for full-text searching. A persistent search history enables quick re-querying, and the interface showcases scrollable example queries to guide exploration.

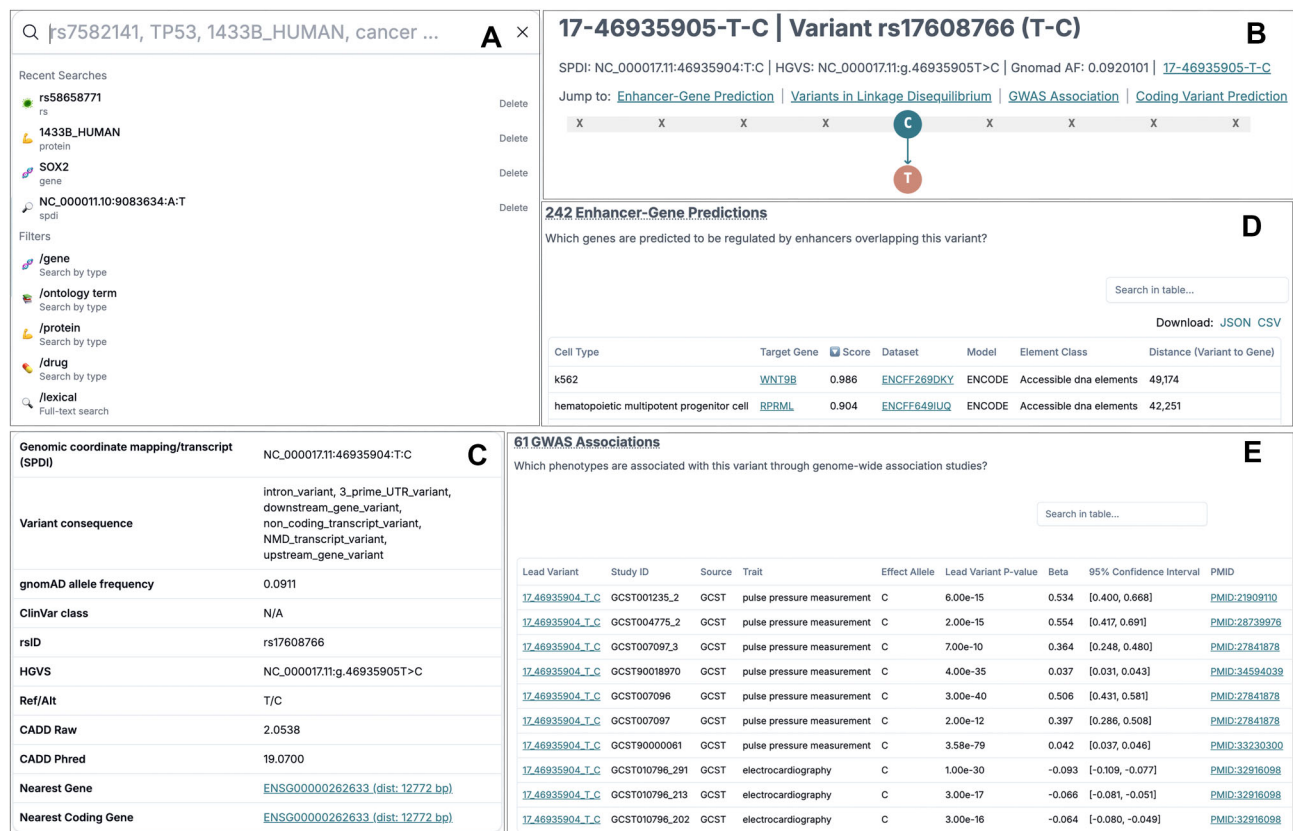


Figure 1. Unified search interface and table-based variant data visualization in the IGV Catalog. **(A)** The unified search bar on the Catalog homepage provides a centralized entry point for querying any data stored in the Catalog backend database, supporting both variant- and gene-centric searches. **(B)** Start of a variant page, which begins with basic descriptive information about the variant, followed by a navigation panel linking to all associated data tables on the page. **(C)** Example of a variant detail table. **(D)** The table displayed is the enhancer–gene predictions table, which summarizes predicted regulatory interactions. **(E)** The GWAS associations table, which displays which phenotypes are associated with this variant from genome-wide association studies.

Table-based data display

To effectively organize and present the highly interconnected information within the IGV Catalog graph database, we employ table-based views as the primary mechanism for displaying data associated with individual database items, such as variants (Fig. 1B–E) or genes (Fig. 2). Tables were selected as the default representation because each item in the underlying graph database is typically linked to many other items, and a tabular format provides a structured, scalable, and user-friendly way to summarize these relationships. For example, on a variant detail page, the Catalog displays enhancer–gene predictions associated with that variant in a table format (Fig. 1D). Each row corresponds to a prediction within a specific cell type, while the columns capture key attributes, including the predicted target gene(s), prediction score, and links to the relevant supporting datasets. The table is further enhanced with interactive features such as grouping predictions by cell type, sorting rows by prediction score, and filtering entries by attributes, enabling users to explore the data dynamically. Multiple other types of information related to a variant are also presented in tabular form. For instance, more information about this variant (Fig. 1B and C), GWAS associations (Fig. 1E), coding variant functional predictions, and allelic effects on transcription factor (TF) binding, among others. Each of these tables provides a concise yet comprehensive view of the associated evidence, allowing users to seamlessly navigate between data categories. To facilitate access, in-page naviga-

tion controls are provided, enabling users to jump directly to the table of interest without excessive scrolling. By adopting this consistent, table-centric display strategy, the Catalog ensures that diverse and heterogeneous datasets can be explored within a unified framework, improving both accessibility for casual exploration and efficiency for hypothesis-driven research.

Figure 2 illustrates examples of table-based displays within the Gene page of the Catalog. Like the Variant page, in-page navigation is provided to facilitate quick access to different tables containing gene-associated information. At the top of the page, summary tables present coding variants linked to the selected gene, including predicted functional impacts and computational scores (Fig. 2A). Downstream tables provide additional layers of annotation: predicted enhancer–gene links derived from the ENCODE-rE2G model [23], with each row corresponding to a predicted enhancer and its relevant cell type (Fig. 2B); and variants associated with the gene through multiple lines of evidence, such as eQTL, splicing QTL (sQTL), and other regulatory relationships (Fig. 2C). Additional resources, including associated pathways, gene–gene interactions, and co-expression patterns, are also available, ensuring a comprehensive view of the gene’s functional and regulatory landscape.

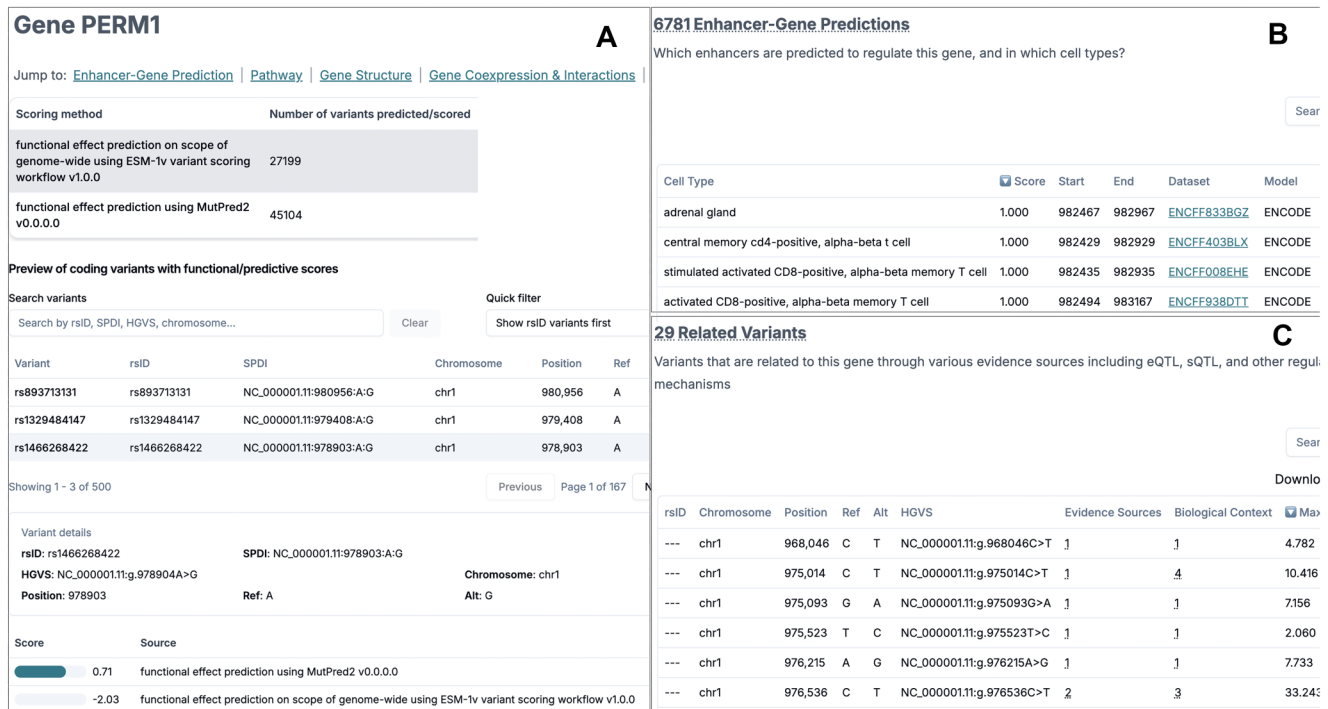


Figure 2. Table-based displays in the Gene page of the IGVF Catalog. **(A)** Summary of coding variants associated with the selected gene, including functional predictions and annotations, with most data generated directly by the IGVF Consortium. Two methods, esm-1v-workflow [19] and MutPred2 [24], are used to show the number of variants predicted/scored. **(B)** Predicted enhancer–gene links from the ENCODE-rE2G model, where each row represents a predicted enhancer and its corresponding cell type. **(C)** Variants linked to the gene through diverse evidence sources, including eQTL, sQTL, and other regulatory relationships.

Other visualization components

The IGVF Catalog provides a diverse suite of interactive, HTML canvas-based visualizations designed to facilitate intuitive exploration of genomic data (Fig. 3A–D). All visualizations retrieve data dynamically from the IGVF API and render responsively to the user’s screen, allowing real-time interaction with large datasets. Complementing the tabular displays of gene- and variant-related information (Figs 1 and 2), these visual modules provide an integrated, graphical perspective that highlights functional and mechanistic relationships. On variant pages, users can view an animated edit diagram that depicts nucleotide substitutions, along with a population allele frequency bar chart enhanced with hover-enabled meta-data for each population (Fig. 3A). A linkage disequilibrium (LD) [39] heatmap is also available (Fig. 3B), enabling users to assess correlation structures between variants within the same genomic region. On gene pages, multiple complementary visualization modules are provided. These include distribution plots of functional and predictive scores across diverse annotation sources, a fully zoomable and interactive gene-to-gene interaction network that captures co-expression and regulatory relationships (Fig. 3C), and a tree-like plot highlighting pathway enrichments (Fig. 3D).

Feedback and glossary system

The Catalog includes a glossary that provides clear contextual definitions for scientific terms used throughout the application. Users can access additional details via hover-enabled tooltips, and each glossary entry includes a “Suggest an Edit” option that opens a prepopulated form, allowing users to propose updated definitions, add notes, and share contact

information. Submissions are stored in a database for review, and their status can be tracked as feedback is processed (Supplementary Fig. S1). A password-protected Glossary Editor enables approved maintainers to incorporate updates directly, ensuring that definitions remain current and community-informed. This system not only enhances user comprehension but also serves as a collaborative platform, engaging sister consortia and researchers worldwide in refining and standardizing scientific terminology within the Catalog.

API service and information

The preferred way to access the data in the backend Graph Database is via the API. The IGVF Catalog API exposes access to the data from the Graph Database using internal optimized queries. We provide an HTTP access and a TRPC-based access. The HTTP access is made public using the OpenAPI specification, allowing REST calls from several clients. The TRPC-based access is made available using the tRPC (typescript remote protocol call) protocol written in Typescript. This allows direct RPC calls using the Typescript language as a client, a widely used language to build front-end applications, speeding up development and ingestion of a fully type-safe API. Direct access to the backend Graph Database is supported through the Arango query language (AQL), with a demonstration provided in a Jupyter notebook (Supplementary File 1).

In addition to backend APIs, the IGVF Catalog’s front-end includes an “X-ray” feature that exposes the underlying API calls used to generate each table, providing users with full transparency into how data are retrieved and displayed (Fig. 3E). For every supported table, users can toggle an X-

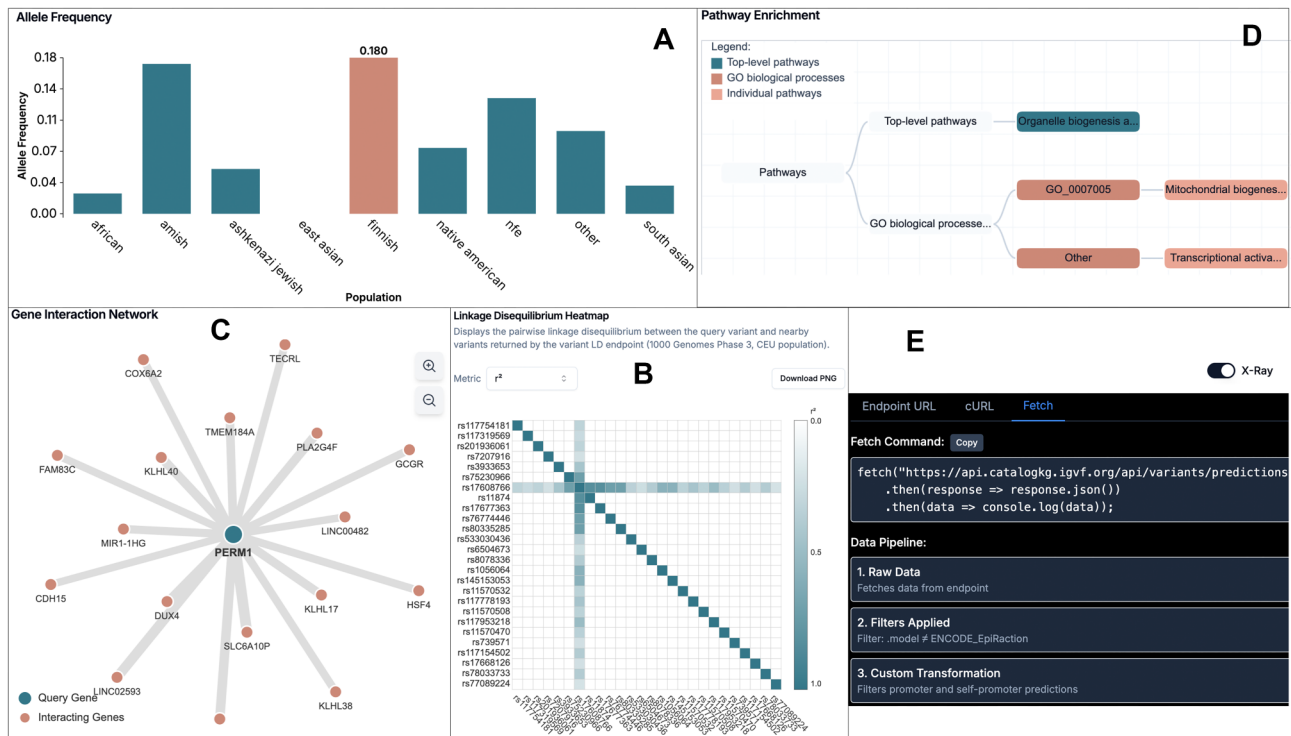


Figure 3. Interactive visualization components and the X-Ray feature in the IGVF Catalog. **(A)** Dynamic population allele frequency bar chart, where hovering over each bar displays detailed metadata for the corresponding population. **(B)** LD heatmap, with a metrics dropdown menu that enables users to select the measure (r^2 or D^2) used to generate the plot. **(C)** Zoomable gene interaction network, allowing users to explore co-expression and regulatory relationships among genes. **(D)** A tree-like plot illustrating pathway enrichment for a selected gene, providing a hierarchical view of functional annotation. **(E)** X-Ray feature, which exposes the underlying code and query pipeline used to fetch data from the Catalog API, including API endpoints, parameters, and optional client-side transformations, enabling reproducibility and transparency.

ray panel to view the fully resolved query URL (including parameters, pagination, and filters), alongside ready-to-use examples such as a cURL command and a JavaScript “fetch()” snippet for reproducing the query externally. When client-side filtering or transformations are applied, the panel further documents the processing pipeline, showing how raw data are mapped, filtered, enriched, or otherwise transformed before display. Because the table data fetcher and X-ray panel share the same data source, consistency is ensured between what is shown and what is retrieved. This design makes IGVF not only a data exploration platform but also a developer- and researcher-friendly resource, lowering barriers to reusing and integrating its data into external workflows, something rarely offered by other genomic databases.

For users working in R, the *rigvf* package (<https://igvf.github.io/rigvf/>) offers a convenient interface to explore and analyze Catalog data directly within R/Bioconductor. Users can query the Catalog by variants, genomic elements, and genes using several accepted identifiers, and retrieve the other nodes that are connected in the graph database, with variants, elements, and genes represented using standard Bioconductor classes such as *GenomicRanges*. Genomic elements, which tend not to have stable identifiers as do variants or genes, may be queried by specifying an extent of the genome.

We have also implemented an experimental “free text” query endpoint within the API. This system employs an LLM-RAG (LLM with RAG) framework via LangChain (<https://www.langchain.com/>). Specifically, input text is first classified to infer a reduced graph database schema, which is then

passed as additional tokens to the ArangoGraphQChain. This process generates AQL queries that are executed on the database, with the resulting output returned to the LLM to produce a plain-text response (in addition to a data table). The current implementation uses OpenAI’s GPT-4.1 (<https://openai.com/>) to generate both the graph queries and the final output. While results can be further enhanced by employing larger models or incorporating a broader representation of the graph schema, these approaches require substantially greater computational resources. To facilitate easier navigation of the IGVF Catalog resources, we implemented a chatbot (<https://catalog.igvf.org/chat>). The chatbot attempts to translate users’ questions into AQL queries whenever possible and returns results directly from our knowledge graph database. This provides a more intuitive interface for users who may not be familiar with query languages. An example of such a query and its corresponding output is shown in [Supplementary Fig. S2](#).

Relationship to the IGVF data portal

The IGVF Data Portal (<https://data.igvf.org/>) hosts the raw and processed datasets generated by the IGVF consortium, while the IGVF Catalog provides a user-facing interface for querying, visualizing, and integrating the analyzed results produced by consortium working groups alongside public annotations from external databases. IGVF data on the portal is organized primarily around Files and FileSets (representing raw experimental data, processed results, analytical outputs, predictions, and models) covered by extensive metadata.

Release files from AnalysisSets, along with data from public sources, are the raw material that is loaded into the Catalog Knowledge Graph. Metadata is explored via a faceted search interface where results, resources, and metadata can be viewed and files downloaded. Results are presented in a configurable, paginated table with adjustable columns, ontology-linked references, and export options in JSON or CSV. In addition to serving as the basis for the IGVF Catalog, the IGVF data portal is a web application that enables the exploration of raw data, integrated annotations, and streamlines access to the diverse experimental outputs generated by IGVF.

Comparison with other similar databases

Most widely used genomic resources, such as gnomAD [1] (population-scale variant frequencies), GWAS Catalog [3] (variant–trait associations), GTEx [2] (expression QTLs across tissues), ClinVar [4] (clinical variant annotations), and Open Targets Platform [5, 40] (integration of genetic associations with functional genomics to prioritize causal variants and genes), focus on different sets of data or user groups. These databases provide critical insights about naturally occurring human genetic variants and their associations with molecular phenotypes or disease.

The IGVF Catalog builds on some of these resources and incorporates unique experimental datasets and predictive models generated by the IGVF Consortium. These datasets include genomic perturbation experiments that build on observational resources to directly probe causal mechanisms of genomic variation.

While MaveDB [41] focuses on results from multiplexed assays of variant effects, which experimentally assess the functional consequences of thousands of variants in parallel, IGVF expands on this concept with a broader suite of experimental strategies—including CRISPR-based perturbations, single-cell transcriptomics, and chromatin accessibility profiling. By integrating these complementary methods, IGVF is uniquely positioned to move beyond single-variant effects and uncover the regulatory architecture and gene networks that shape phenotypic outcomes.

All this information is organized into a unified, graph-based framework that connects functional, genetic, and genomic resources, enabling cross-linked exploration across data types. In this way, the IGVF Catalog enables linking genomic variation to molecular and cellular phenotypes through both experimental and predictive insights.

Limitations and outlook

The IGVF Catalog is an actively evolving resource, and several limitations remain. Data coverage is still incomplete, with ongoing integration of multiple datasets (e.g. Biobank OR results and MaveDB-derived assays). Some interactive features and autocomplete functionality, are being refined to improve performance and search accuracy. Clearer labeling of IGVF-generated versus public data is also needed to enhance interpretability. Finally, documentation and user guidance are under active development and revision, and feedback mechanisms have only recently been expanded to accommodate broader community participation.

The DACCs are continually enhancing both the backend database and frontend interface of the IGVF Catalog; thus, future updates will address these limitations by incorporating

additional datasets (including both new datasets generated by IGVF consortium and other resources from the community), optimizing query and visualization performance, implementing explicit data source labels, embedding contextual documentation, and refining feedback and glossary systems. Regular updates and subsequent publications will ensure the Catalog remains an up-to-date, comprehensive, and user-friendly resource.

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Supervision: M.I.L., J.M.E., B.C.H., T.W.

Funding acquisition: B.C.H., T.W.

Supplementary data

Supplementary data is available at NAR online.

Conflict of interest

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Data availability

The IGVF Catalog is freely available at <https://catalog.igvf.org/>, with detailed documentation and user guidance provided at <https://docs.catalog.igvf.org>. The Catalog's underlying data can also be accessed programmatically through its API service at <https://api.catalogkg.igvf.org/>, which includes comprehen-

sive reference materials. All of the code for the IGVF catalog backend, including database and API, is available at Zenodo (10.5281/zenodo.17468372).

References

- Karczewski KJ, Francioli LC, Tiao G *et al.* The mutational constraint spectrum quantified from variation in 141,456 humans. *Nature* 2020;581:434–43. <https://doi.org/10.1038/s41586-020-2308-7>
- Consortium GT. The GTEx Consortium Atlas of genetic regulatory effects across human tissues. *Science* 2020;369:1318–30. <https://doi.org/10.1126/science.aaz1776>
- Cerezo M, Sollis E, Ji Y *et al.* The NHGRI-EBI GWAS Catalog: standards for reusability, sustainability and diversity. *Nucleic Acids Res* 2025;53:D998–D1005. <https://doi.org/10.1093/nar/gkae1070>
- Landrum MJ, Lee JM, Benson M *et al.* ClinVar: improving access to variant interpretations and supporting evidence. *Nucleic Acids Res* 2018;46:D1062–7. <https://doi.org/10.1093/nar/gkx1153>
- Buniello A, Suveges D, Cruz-Castillo C *et al.* Open Targets Platform: facilitating therapeutic hypotheses building in drug discovery. *Nucleic Acids Res* 2025;53:D1467–75. <https://doi.org/10.1093/nar/gkae1128>
- Consortium IGVF. Deciphering the impact of genomic variation on function. *Nature* 2024;633:47–57. <https://doi.org/10.1038/s41586-024-07510-0>
- Oughtred R, Stark C, Breitkreutz BJ *et al.* The BioGRID interaction database: 2019 update. *Nucleic Acids Res* 2019;47:D529–41. <https://doi.org/10.1093/nar/gky1079>
- Orchard S, Ammari M, Aranda B *et al.* The MIntAct project—IntAct as a common curation platform for 11 molecular interaction databases. *Nucl. Acids Res.* 2014;42:D358–63. <https://doi.org/10.1093/nar/gkt1115>
- Zhou H, Arapoglou T, Li X *et al.* FAVOR: functional annotation of variants online resource and annotator for variation across the human genome. *Nucleic Acids Res* 2023;51:D1300–11. <https://doi.org/10.1093/nar/gkac966>
- Sherry ST, Ward MH, Kholodov M *et al.* dbSNP: the NCBI database of genetic variation. *Nucleic Acids Res* 2001;29:308–11. <https://doi.org/10.1093/nar/29.1.308>
- Liu X, Li C, Mou C *et al.* dbNSFP v4: a comprehensive database of transcript-specific functional predictions and annotations for human nonsynonymous and splice-site SNVs. *Genome Med* 2020;12:103. <https://doi.org/10.1186/s13073-020-00803-9>
- Mudge JM, Carbonell-Sala S, Diekhans M *et al.* GENCODE 2025: reference gene annotation for human and mouse. *Nucleic Acids Res* 2025;53:D966–75. <https://doi.org/10.1093/nar/gkae1078>
- Boeckmann B, Bairoch A, Apweiler R *et al.* The SWISS-PROT protein knowledgebase and its supplement TrEMBL in 2003. *Nucleic Acids Res* 2003;31:365–70. <https://doi.org/10.1093/nar/gkg095>
- Boutet E, Lieberherr D, Tognolli M *et al.* UniProtKB/Swiss-Prot. *Methods Mol Biol* 2007;406:89–112.
- Meldal BHM, Perfetto L, Combe C *et al.* Complex Portal 2022: new curation frontiers. *Nucleic Acids Res* 2022;50:D578–86. <https://doi.org/10.1093/nar/gkab991>
- Project Consortium ENCODE. An integrated encyclopedia of DNA elements in the human genome. *Nature* 2012;489:57–74. <https://doi.org/10.1038/nature11247>
- Kerimov N, Tambets R, Hayhurst JD *et al.* eQTL Catalogue 2023: new datasets, X chromosome QTLs, and improved detection and visualisation of transcript-level QTLs. *PLoS Genet* 2023;19:e1010932. <https://doi.org/10.1371/journal.pgen.1010932>
- Dong S, Zhao N, Spragins E *et al.* Annotating and prioritizing human non-coding variants with RegulomeDB v. *Nat Genet* 2023;55:724–6. <https://doi.org/10.1038/s41588-023-01365-3>
- Sverchkov Y. ESM-1v predictions for all AA substitutions in all MANE proteins. February 2025, Zenodo, <https://zenodo.org/doi/10.5281/zenodo.14828609>
- DeGorter MK, Goddard PC, Karakoc E *et al.* Transcriptomics and chromatin accessibility in multiple African population samples. bioRxiv, <https://doi.org/10.1101/2023.11.04.564839>, 6 November 2023, preprint: not peer reviewed.
- Arnold CD, Gerlach D, Stelzer C *et al.* Genome-wide quantitative enhancer activity maps identified by STARR-seq. *Science* 2013;339:1074–7. <https://doi.org/10.1126/science.1232542>
- Taliun D, Harris DN, Kessler MD *et al.* Sequencing of 53,831 diverse genomes from the NHLBI TOPMed Program. *Nature* 2021;590:290–9. <https://doi.org/10.1038/s41586-021-03205-y>
- Gschwind AR, Mualim KS, Karbalayghareh A *et al.* An encyclopedia of enhancer-gene regulatory interactions in the human genome. bioRxiv, <https://doi.org/10.1101/2023.11.09.563812>, 23 February 2025, preprint: not peer reviewed.
- Pejaver V, Urresti J, Lugo-Martinez J *et al.* Inferring the molecular and phenotypic impact of amino acid variants with MutPred2. *Nat Commun* 2020;11:5918. <https://doi.org/10.1038/s41467-020-19669-x>
- Matreyek KA, Starita LM, Stephany JJ *et al.* Multiplex assessment of protein variant abundance by massively parallel sequencing. *Nat Genet* 2018;50:874–82. <https://doi.org/10.1038/s41588-018-0122-z>
- Findlay GM, Daza RM, Martin B *et al.* Accurate classification of BRCA1 variants with saturation genome editing. *Nature* 2018;562:217–22. <https://doi.org/10.1038/s41586-018-0461-z>
- Fabiha T, Raine I, Kundu S *et al.* A consensus variant-to-function score to functionally prioritize variants for disease. bioRxiv, <https://doi.org/10.1101/2024.11.07.622307>, 10 November 2024, preprint: not peer reviewed.
- Milacic M, Beavers D, Conley P *et al.* The Reactome Pathway Knowledgebase 2024. *Nucleic Acids Res* 2024;52:D672–8. <https://doi.org/10.1093/nar/gkad1025>
- Vorontsov IE, Eliseeva IA, Zinkevich A *et al.* HOCOMOCO in 2024: a rebuild of the curated collection of binding models for human and mouse transcription factors. *Nucleic Acids Res* 2024;52:D154–63. <https://doi.org/10.1093/nar/gkad1077>
- Abramov S, Boytsov A, Bykova D *et al.* Landscape of allele-specific transcription factor binding in the human genome. *Nat Commun* 2021;12:2751. <https://doi.org/10.1038/s41467-021-23007-0>
- Yan J, Qiu Y, Ribeiro Dos Santos AM *et al.* Systematic analysis of binding of transcription factors to noncoding variants. *Nature* 2021;591:147–51. <https://doi.org/10.1038/s41586-021-03211-0>
- Sun BB, Chiou J, Traylor M *et al.* Plasma proteomic associations with genetics and health in the UK Biobank. *Nature* 2023;622:329–38. <https://doi.org/10.1038/s41586-023-06592-6>
- Nishizaki SS, Ng N, Dong S *et al.* Predicting the effects of SNPs on transcription factor binding affinity. *Bioinformatics* 2020;36:364–72. <https://doi.org/10.1093/bioinformatics/btz612>
- Obayashi T, Kodate S, Hibara H *et al.* COXPRESdb v8: an animal gene coexpression database navigating from a global view to detailed investigations. *Nucleic Acids Res* 2023;51:D80–7. <https://doi.org/10.1093/nar/gkac983>
- Barbarino JM, Whirl-Carrillo M, Altman RB *et al.* PharmGKB: a worldwide resource for pharmacogenomic information. *WIREs Mechanisms of Disease* 2018;10:e1417. <https://doi.org/10.1002/wsbm.1417>
- Weinreich SS, Mangon R, Sikkens JJ *et al.* [Orphanet: a European database for rare diseases]. *Ned Tijdschr Geneesk* 2008;152:518–9.
- Rehm HL, Berg JS, Brooks LD *et al.* ClinGen—the Clinical Genome Resource. *N Engl J Med* 2015;372:2235–42. <https://doi.org/10.1056/NEJMs1406261>

38. Holmes JB, Moyer E, Phan L *et al.* SPDI: data model for variants and applications at NCBI. *Bioinformatics* 2020;36:1902–7. <https://doi.org/10.1093/bioinformatics/btz856>
39. Slatkin M. Linkage disequilibrium—understanding the evolutionary past and mapping the medical future. *Nat Rev Genet* 2008;9:477–85. <https://doi.org/10.1038/nrg2361>
40. Ochoa D, Hercules A, Carmona M *et al.* Open Targets Platform: supporting systematic drug-target identification and prioritisation. *Nucleic Acids Res* 2021;49:D1302–10. <https://doi.org/10.1093/nar/gkaa1027>
41. Esposito D, Weile J, Shendure J *et al.* MaveDB: an open-source platform to distribute and interpret data from multiplexed assays of variant effect. *Genome Biol* 2019;20:223. <https://doi.org/10.1186/s13059-019-1845-6>