# PanelAppRex aggregates disease gene panels and facilitates sophisticated search

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#### Abstract

Motivation: Gene panel data provides critical insights into disease-gene correlations. However, aggregating and interrogating this diverse dataset can be challenging. PanelAppRex addresses this by first preparing a machine-readable aggregate and second by offering a sophisticated natural search interface that streamlines data exploration for both clinical and research applications.

Results: PanelAppRex aggregates gene panel data from source including Clin-Var, UniProt, and Genomics England's PanelApp, including the approved panels used in the NHS National Genomic Test Directory and the 100,000 Genomes Project. It enables users to execute complex queries by gene names, phenotypes, disease groups and more, returning integrated datasets in multiple downloadable formats. Benchmarking demonstrates 93% - 100% accuracy, effectively simplifying variant discovery and interpretation to enhance workflow efficiency. The greatest benefit is the analysis-ready format for bioinformatic integration.

Availability: https://switzerlandomics.ch/services/panelAppRexAi/ (A standalone webpage will be substituted for publication version). The source code and data are accessible at https://github.com/DylanLawless/PanelAppRex. PanelAppRex is available under the MIT licence. The dataset is maintained for a minimum of two years following publication.

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#### Acronyms

API Application Programming Interface	2
CSV comma-separated values	5
GE Genomics England	2
HGNC Human genome organisation Gene Nomenclature Committee	
HTML HyperText Markup Language	3
JACI Journal of Allergy and Clinical Immunology	
MOI Mode of Inheritance	4
OMIM Online Mendelian Inheritance in Man	4
PDF Portable Document Format	5
PID Primary immunodeficiency	4
RAG Retrieval-augmented generation	2

#### 1 Introduction

Disease-gene panels are pivotal for the diagnosis and interpretation of genetic disorders. Sources like Genomics England (GE)'s PanelApp and PanelApp Australia host comprehensive panels that support genomic testing (1). For instance, these are integral in the NHS and research projects such as the 100,000 Genomes Project(1). Despite its utility, manual panel selection and data aggregation remain labour intensive. PanelAppRex was developed to simplify this process by automating data retrieval from Application Programming Interface (API) sources and integrating the data into machine and user-friendly formats. We include the use of GE's PanelApp, ClinVar, and UniProt (1–3). The novel natural language-style search capability further streamlines the discovery of disease gene panels by allowing queries based on gene names, phenotypes, disease groups and additional key attributes which were supplemented with evidence-based Retrieval-augmented generation (RAG).

#### 2 Materials and methods

#### 2.1 Data

The PanelAppRex core model contained 58,592 entries consisting of 52 sets of annotations, including the gene name, disease-gene panel ID, diseases-related features, confidence measurements (4). Data from gnomAD v4 comprised 807,162 individuals, including 730,947 exomes and 76,215 genomes (5). This dataset provided 786,500,648 single nucleotide variants and 122,583,462 InDels, with variant type counts of 9,643,254 synonymous, 16,412,219 missense, 726,924 nonsense, 1,186,588 frameshift and 542,514 canonical splice site variants. ClinVar data were obtained from the variant summary

dataset (this version: 16 March 2025) available from the NCBI FTP site, and included 6,845,091 entries, which were processed into 91,319 gene classification groups and a total of 38,983 gene classifications; for example, the gene A1BG contained four variants classified as likely benign and 102 total entries (2). For our analysis phase we also used dbNSFP which consisted of a number of annotations for 121,832,908 single nucleotide variants (6).

#### 2.2 Implementation

PanelAppRex was implemented in R and integrates data from GE's PanelApp, ClinVar, and UniProt (1–3). It performed credentialed access to the API to retrieve all approved panels, merging them into two formats: a simplified version (Panel ID, Gene) and a complex version (including metadata such as confidence level, mode of inheritance, and disease information), and several metadata summary statistics. In addition, the tool incorporates a search module to execute complex user queries. The search functionality supports queries by gene names, phenotypes, disease names, disease groups, panel names, genomic locations and other identifiers. RAG was used to improve the natural queries in hidden states based on evidence about the disease and gene function by supplementing the data with additional sources including ClinVar, UniProt, etc.

#### 2.3 Usage

On a desktop browser, queries can be executed via the integrated search bar in our HyperText Markup Language (HTML) version, where a JavaScript function splits the query into individual terms and progressively filters rows - retaining only those that match all active terms while ignoring unmatched ones. This enables users to perform complex, partial matching queries (e.g. "paediatric *RAG1* primary immunodeficiency skin disorder") to rapidly identify the panel most closely associated with their hypothesis on primary immunodeficiency and paediatric skin disorders.

Bioinformatically, users can import the provided, ready-for-use, datasets in TSV or Rds formats. A typical use case might involve users merging with their own omic data based on gene or protein ID. The following code snippet, available in minimal\_example.R, demonstrates how to load the data in R:

```
# TSV format
path_data <- "../data"
core_path <- paste0(path_data, "/PanelAppData_combined_core")
minimal_path <- paste0(path_data, "/PanelAppData_combined_minimal")

df_core <- read.table(
    file= paste0(core_path, ".tsv"),
    sep = "\t", header = TRUE)</pre>
```

```
df_minimal <- read.table(
    file= pasteO(minimal_path, ".tsv"),
    sep = "\t", header = TRUE)

# Rds format
rds_path <- pasteO(path_data, "/PanelAppData_combined_Rds")
df core <- readRDS(file= rds path)</pre>
```

#### 2.4 Validation for completeness in core data

To ensure the reliability and completeness of the core dataset, we systematically assessed whether key gene-level fields were present for each entry after merging the core dataset. Specifically, we checked for non-missing values in gene symbols, associated publications, disease panel names, Mode of Inheritance (MOI), and Online Mendelian Inheritance in Man (OMIM) gene IDs. Where Human genome organisation Gene Nomenclature Committee (HGNC) or Ensembl gene IDs were missing, we used programmatic queries to the Ensembl database via the biomart R package to recover the identifiers using the available HGNC symbol as input (7). This validation step was applied to the full integrated PanelAppRex dataset.

#### 2.5 Benchmarking for manual queries

To mimic a clinician diagnosing a new disease such as Primary immunodeficiency (PID), we began by systematically selecting genetic diagnosis case studies from the current online catalogue from the Journal of Allergy and Clinical Immunology (JACI), using the first five results (8–12). Although the method presented here is generalisable to genetic diagnosis across disease areas, we chose to validate it using PID case studies, as this is the primary focus of our own research.

The clinical background from these studies was used to construct keyword queries from patient features, simulating a naïve starting position for a clinician. We then tested whether our PanelAppRex tool could successfully retrieve panels that included the final causal gene reported in each case study. The sources, queries, and results are shown in **Table 1**.

### 3 Results

#### 3.1 Core dataset

PanelAppRex successfully aggregated data, currently from 451 panels, and several genomics databases to offer a user-friendly search functionality (**Figure 1**). Users

can retrieve results filtered by gene names, phenotypes, disease groups and other criteria. The system returned a table view with panel details and provided options for exporting results in comma-separated values (CSV), Excel, or Portable Document Format (PDF) formats. Bioinformatic uses may include generating virtual panels, constructing prior odds, or supporting formal reporting in qualifying variant protocols.

#### 3.2 Validation shows completeness in the core dataset

The raw dataset exhibited near-complete coverage in most fields after merging the core dataset. Specifically, 99.9915% of gene entries had HGNC symbols and 99.2315% had Ensembl gene IDs. All entries included at least one publication, a disease panel name, a mode of inheritance, and an OMIM gene ID (100% each). After recovery of missing identifiers, completeness reached 100% across all these core fields. This validated dataset ensures that users can confidently build on a consistent foundation, using standardised and stable identifiers to link or enrich the core data with external resources (**Figure 2**).

# 3.3 Benchmarking for manual queries confirms accurate retrieval of causal gene

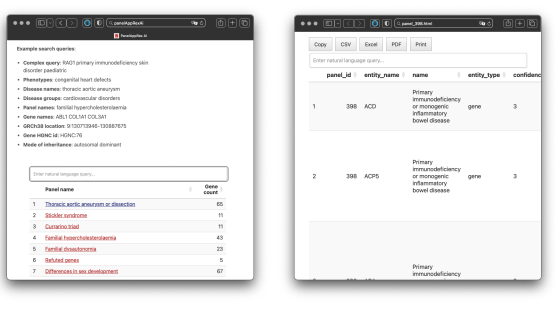
To evaluate the practical utility of our system on a desktop browser, we applied the benchmarking approach described in methods using five published case studies of PID (8–12). The tool's query process retrieved gene panels using natural language-style input. For example, in the first case study on Hereditary Angioedema, the authors reported suspecting that the condition was linked to "SERPING1, Factor XII, and edema" which we used as the query terms. Although the true causal gene term F12 (using its official HGNC name) was not mentioned, the inclusion of the related disease terms enabled the system to correctly retrieve the panel containing F12 based on the hidden search knowledge.

While an expert might readily recognise "Factor XII" as referring to F12, this correct query result demonstrates that the system can make such connections automatically, supporting users with varying levels of genetic expertise.

In our evaluation, PanelAppRex returned panels in which the causal gene was present in 93% of all returned panels, with the subjective user-selected "best panel" achieving a perfect accuracy of 100% (**Figure 1**). These metrics were derived by comparing the causal gene identified from the case study abstracts to the panels returned by our query, and by applying a subjective relevance measure to mimic intuitive panel selection - acknowledging that certain panels, such as the established PID gene panel, are inherently more reliable than broader, less specific panels. Overall, the results confirm that our approach accurately identifies the most relevant panels and effectively supports clinical decision-making in complex diagnostic scenarios.

Table 1: Summary of case study queries and PanelAppRex results. \*Case study 5 had five individual cases and patient information was significantly longer that other studies. It was therefore converted to keyword query automatically by the OpenAI model o3-mini to remove subjective bias and align with the other queries.

Case study (Ref)	Source title	Query	Causal gene	Result panels	Sub- jec- tive best pan- els	Pan- els with causal gene	Sub- jec- tive rele- vance ratio		Result ID, panelName, geneCount
1 (8)	Genetic Analysis As a Practical Tool to Di- agnose Hereditary An- gioedema with Normal C1 Inhibitor: A Case Report	SERPING1 Factor XII edema	F12	3	1	3	0.3	1	64 COVID-19 research 695; 192 Primary immunodefi- ciency or monogenic inflam- matory bowel disease 572; 311 Research panel - Severe Paediatric Disorders 2691
2 (9)	Severe dermatitis, multiple allergies, and metabolic wasting syndrome caused by a novel mutation in the N-terminal plakin domain of desmoplakin	SAM syndrome DSG1 dermatitis metabolic wasting	DSP	3	3	3	1	1	205 Fetal anomalies 2185; 210 DDG2P 2422; 211 Pae- diatric disorders 3903
3 (10)	Hematopoietic stem cell transplantation in a patient with proteasome-associated autoinflammatory syndrome (PRAAS)	resistant cutaneous vasculitis SH2D1A	PSMB4	3	1	2	0.3	0.7	64 COVID-19 research 695; 192 Primary immunodefi- ciency or monogenic inflam- matory bowel disease 572; 311 Research panel - Severe Paediatric Disorders 2691
4 (11)	Autoimmune lympho- proliferative syndrome caused by a homozy- gous null FAS ligand (FASLG) mutation	Autoimmune lympho- proliferative syndrome ALPS lymphoprolifera- tion hypergammaglob- ulinemia autoimmune cytopenia	FASLG	2	1	2	0.5	1	64 COVID-19 research 695; 192 Primary immunodefi- ciency or monogenic inflam- matory bowel disease 572
5* (12)	Fatal combined immunodeficiency associated with heterozygous mutation in STAT1	primary immunod- eficiency recurrent pneumonia chronic diarrhea oral thrush bronchiectasis lym- phadenopathy hep- atosplenomegaly autoimmune hepatitis Addison	STAT1	1	1	1	1	1	192 Primary immunodefi- ciency or monogenic inflam- matory bowel disease 572



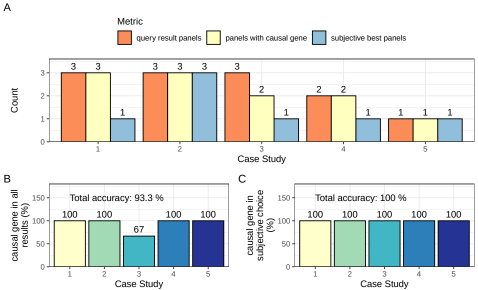


Figure 1: PanelAppRex interface displaying search results for a complex query. Top: screenshots showing the search interface, with the left panel displaying the full database before filtering and the right panel showing detailed results for a selected panel. Bottom: benchmark metrics are presented as follows: (A) for each of the 5 case studies, the total number of panels returned, the number of panels that included the causal gene, and the single best panel selected (always 1 by default); (B) the percentage of all returned panels that included the true causal gene; (C) the percentage of the single best panels that contained the true causal gene.

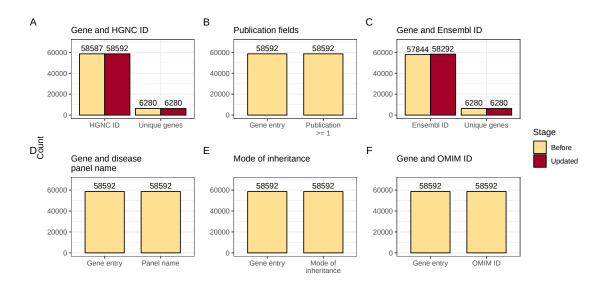


Figure 2: Validation and recovery of core annotation fields in the PanelAppRex dataset. (A) Unique genes and HGNC IDs before and after retrieving missing HGNC entries via Ensembl. (B) Availability of publication annotations for genes. (C) Unique genes and Ensembl gene IDs before and after biomart-based recovery of missing Ensembl IDs. (D) Gene entries with associated disease panel names. (E) Gene entries with annotated mode of inheritance. (F) Gene entries linked to an OMIM gene ID. Each bar shows the count of entries with non-missing values for the respective field. Updated fields (HGNC and Ensembl ID) reflect values recovered via external lookup using HGNC symbols.

#### 4 Discussion and conclusion

PanelAppRex provides an accessible and validated platform for querying, aggregating, and exporting curated disease-gene panel data. Designed for both clinical and research use, it simplifies study planning and variant interpretation through a user-friendly interface, while also offering a harmonised dataset for programmatic workflows. The core model integrates over 58,000 panel-gene associations, with full coverage of key fields including HGNC symbols, Ensembl IDs, OMIM annotations, MOI, and supporting publications.

Beyond immediate usability, PanelAppRex lays a foundation for genome-wide statistical modelling of disease risk. With annotations for gene-disease relationships, the dataset can support rigorous estimation of the prior probability of observing variant classifications (e.g. benign, pathogenic) under different MOI. This helps address a persistent challenge in clinical genetics: the lack of principled priors for variant interpretation that account not only for known pathogenic variants (true positives), but also for unobserved variants and negative evidence (false negative causal pathogenic variant and true negative absence of causal variants) (13; 14). By integrating high-resolution allele frequencies from gnomAD (5), curated classifications from ClinVar (2), and structured gene-disease associations (1), PanelAppRex can enable the derivation of calibrated genome-wide priors for Bayesian models of genetic risk.

These capabilities go beyond panel lookup to support evidence-aware diagnostics. Probabilistic models incorporating both observed and unobserved variation can quantify uncertainty, resolve ambiguous findings, and refine variant prioritisation, scaling naturally to support complex, multi-gene disorders. The dataset also provides a substrate for AI-driven approaches to variant interpretation, including probabilistic inference, reinforcement learning, and deep annotation pipelines (15; 16).

Several limitations should be acknowledged. Not all known coding genes are currently linked to a disease panel; we prioritised high-confidence, traceable annotations over broad but less reliable coverage. Some genes are over-represented across multiple panels due to historical research biases, leading to non-uniform panel enrichment. Additionally, not all panels returned by queries may be equally informative. For example, during benchmarking, the COVID-19 research panel frequently appeared alongside the PID panel due to overlapping genes. While technically accurate, such panels may be less relevant to users focused on clinical diagnosis of immunodeficiency. In bioinformatic workflows, these choices can be refined systematically or excluded using filters or downstream logic.

PanelAppRex bridges expert-curated panel knowledge with genome-scale statistical reasoning. It offers a robust tool for interactive search, a validated dataset for programmatic access, and a scalable framework for quantitative genomic modelling. Future work will focus on expanding supported queries, integrating additional variant types and annotations, and enabling more sophisticated applications in variant interpretation and risk estimation.

#### 5 Conclusion

PanelAppRex offers a robust solution for aggregating and querying gene panel data. Its sophisticated search feature simplifies data exploration and enhances variant interpretation.

## Acknowledgements

We acknowledge GE for providing public access to the PanelApp data. The use of data from GE panelapp was licensed under the Apache License 2.0. The use of data from UniProt was licensed under Creative Commons Attribution 4.0 International (CC BY 4.0). ClinVar asks its users who distribute or copy data to provide attribution to them as a data source in publications and websites (2).

# Competing interest

We declare no competing interest.

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