FAANGMine Documentation

Release 1.6

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FAANGMine is a data mining resource that integrates reference genome assemblies for cattle, horse, pig, sheep, chicken, cat, dog and water buffalo with many other biological data sets. Powered by InterMine, this platform provides access to a number of datasets from a variety of source. It also provides customized bioinformatics tools that researchers can use to create their own custom datasets. FAANGMine is part of FAANGMine.org. The FAANG (Functional Annotation of ANimal Genomes) Consortium is "a coordinated international action to accelerate genome to phenome" and aims to generate comprehensive maps of functional elements in genomes of domesticated animals. FAANGMine will integrate data generated by the FAANG Consortium for animal researchers with or without bioinformatic programming skills to use in their own research projects.

	GMine 🗸								help relea
Home	A MyMine	Templates	Lists	QueryBuilder	Regions	Data Sources	Help	API	Contact Us Log in

Main site: http://faangmine.org/faangmine

Link to the available datasets in FAANGMine: http://128.206.116.18:8080/faangmine/dataCategories.do

FAANGMine.org is based upon work supported by the National Science Foundation under Award Number 0054449. Any opinions, findings, and conclusions or recommendations expressed in this material are those of the author(s) and do not necessarily reflect the views of the National Science Foundation. FAANGMine is developed and hosted at the University of Missouri. If you have comments or if you wish to report a problem, please contact the Database Administrator.

CHAPTER 1

Overview of FAANGMine

This section provides a brief overview of the layout for FAANGMine.



The navigation panel highlights different functionalities of FAANGMine.

Home - The home page for FAANGMine

MyMine - The MyMine serves as a portal for account management. When logged in to FAANGMine Users can access their saved templates, most recent queries and saved lists.

Templates - List of templates that users can select from based on the nature of their query. Each template is a predefined query with a simple form containing a description of what input is expected and the type of output that will be generated.

Lists - Allows users to upload lists of genes on which they can perform enrichment analyses and export the results. Users that log in to FAANGMine can save their lists for future use.

QueryBuilder - A flexible interface that allows users to create their own custom query template while browing the FAANGMine data models. Queries can be exported in a variety of formats to share with other users.

Regions - The Genomic Region Search tool where users can enter a series of genomic coordinates, specify flanking regions and fetch all features that fall within the given interval. The result can be exported or saved as a list for further analyses.

Data Sources - Provides a summary of all the data loaded into FAANGMine including their sources, associated publications and links to source sites.

Help - Links to the FAANGMine help docs and tutorials

API - Describes the InterMine API that allows users to programmatically access FAANGMine.

CHAPTER 2

Navigation and Searching in FAANGMine

There are several ways for users to query FAANGMine.

2.1 Quick Search

Quick Search allows users to search keywords from any of the FAANGMine datasets. There is a Quick Search box on the FAANGMine home page or a smaller search box in the upper right corner of all pages.

We are currently working to fix our Quick Search tool and hope to have it running again soon.

Quick Search
Search FAANGMine. Enter names, identifiers or keywords for genes, proteins, ontology terms, authors, etc. (e.g. RCAN1, COX2, NM_001040473.2, NP_001076063.1, 282087, ENSFCAG00000031270, rs385910977, IL13)
e.g. IGF2, ENSBTAG0000013066
SEARCH

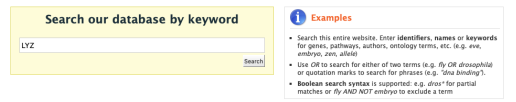
Fig. 1: Quick Search from home page



Fig. 2: Quick Search from any page

Quick Searches can be conducted with a number of identifiers including gene names, transcripts, pathways, gene identifiers or organisms. The wildcard character * can be used to retrieve all results that match a particular search query.

For an example, we will use the lysozyme gene *LYZ* as an example. Enter LYZ in the search box and click **Search**. The results page is tabulated and displays a summary about your query, as shown below.



Search results 1 to 40 out of 40 for LYZ

	Туре	Details	
Categories		ENSG0000090	0382 + - + LYZ
lits by Category		Source:	Ensembl95
 Gene: 17 	Gene	Length:	
 mRNA: 14 Protein: 7 		Chromosome Location:	[unknown]
 Publication: 2 		Organism:	H. sapiens
Hits by Organism H. sapiens: 6		777776 - I Source:	LYZ RefSeg
 B. taurus: 4 C. lupus familiaris: 4 E. caballus: 4 		Description: Length:	lysozyme 8807 FASTA
 F. catus: 4 G. gallus: 4 	Gene	Chromosome Location:	5: 44506988-44515794
 O. aries: 4 		Organism:	B. taurus
 S. scrofa: 4 R. norvegicus: 2 		Assembly:	ARS-UCD1.2
 M. musculus: 1 		474442 -	LYZ
		Source:	RefSeq
		Description:	lysozyme
	Gene	Length:	4140 FASTA
	dene	Chromosome Location:	10: 11346500-11350639
		Organism:	C. lupus familiaris
		Assembly:	CanFam3.1

Fig. 3: Results table for Quick Search with the gene LYZ

Hits are summarized in the box to the left of the results tables and can be filtered based on **Category** and **Organism**. Clicking on any of them will filter by the selected category. Note that for results with sequence data available, the sequence can be downloaded in FASTA format by clicking on the FASTA box within the hit Details box. The score column in the result table indicates the similarity of your query to each of the hits. The results page can also be converted to a list (and saved if users are logged in). To enable this feature click on **Gene** in the **Hits by Category** then click on *C. lupus familiaris* in **Hits by Organism**.

After the table has been filtered for gene and organism, checkboxes will be available for users to select genes they would like to add to their list. Once the genes are selected, click on **CREATE LIST**. See the lists section for more detail on creating and saving lists.

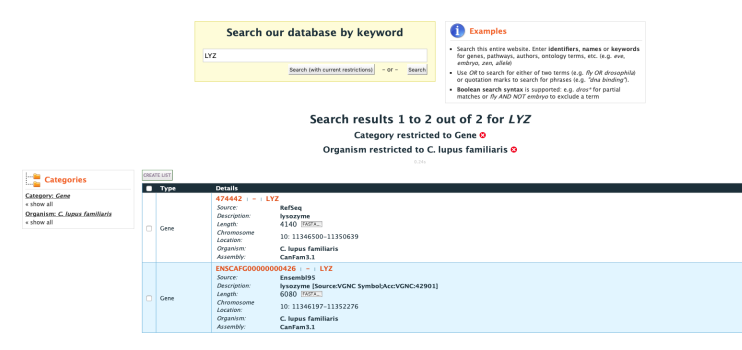


Fig. 4: Results table for Quick Search with the gene LYZ filtered by gene then by organism ..

2.2 Templates

Templates or predefined queries are another search method within FAANGMine. Popular templates are displayed on the home page, grouped by category (e.g., Genes, Proteins, Interactions) and the complete list can be seen by clicking the **Templates** menu tab.

As an example, the **Gene -> Homologues** template queries FAANGMine to retrieves all homologue for a given gene. Here, we will do a search for the gene *GSTM1*.

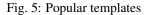
The results page displays all of the homologues for that query gene. When logged in to FAANGMine, users can save their results as a list for further analyses by clicking on the **Save as List** button above the results table then choosing columns to save in their list. See the lists section for more detail on creating and saving lists. Note the "Trail: Query" text at the upper left of the results table. Clicking on the "Query" link will bring you back to the query that generated the table to allow for edits without having to start with a new template.

2.2.1 Generate query code

The code for each template query can be retrieved by clicking on the arrow next to **Generate Python Code** and choosing the desired language from the pull-down menu. The language options are Python, Perl, Java, Ruby, JavaScript, and XML.

Please contact us if you would like any additional template queries or if you have a concern about a query not completing.

GENES	PROTEINS	FUNCTION	HOMOLOGY	INTERACTIONS	VARIATION	ENTIRE GENE SET	ALIAS AND DBXREF				
The source of gene annotations in FAANGMine are NCBI (RefSeq) and Ensembl.											
Query for genes:											
• Ge	ne 📫 Chromo	somal location									
• Ge	ne 🔶 GO term	15									
• Ch	romosomal loca	ation 🔶 genes									
• Ge	ne 🔶 Publicat	ions									
• Ch	romosome 📫	genes									
• Ge	ne 📫 Transcri	pts + Exons					5				
• Ge	ne Source and O	Organism 📦 G	enes and Chromo	somal location			eile				
• Or	ganism 📦 Tra	nscripts + Gene	es + Database Cro	oss Reference IDs			CH ^R				
							aula.				
» More	queries						popular queites				
	queries	1									



2.2.2 Download results

The search results from a template query can be downloaded by clicking the **Export** button above the table and choosing the desired format from the pull-down menu to the right of the File name field. Available formats are tabseparated values, comma-separated values, XML, and JSON. When the results contain genomic features, they may also be downloaded in FASTA, GFF3, or BED format. Other options may be specified in the submenu to the left of the download box. By default, all rows and all columns are downloaded, but individual columns may be included or excluded by clicking on the toggles next to the column headers in the **All Columns** submenu. The number of rows and row offset are set in the **All Rows** submenu. Download the results as a compressed file by choosing GZIP or ZIP format in the **Compression** submenu (default is **No Compression**). Column headers are not added by default but may be included under the **Column Headers** submenu. Finally, the **Preview** submenu displays the first three rows of the file to be downloaded so that the desired format and options may be finalized before beginning the download. When ready, click the **Download file** button to download the results.

2.2.3 Customize output and manage columns

To customize the results table layout, click the **Manage Columns** button. This allows users to rearrange, remove or order columns. Filters can be edited by clicking the **Manage Filters** button. To specify the entity relationships within the query and change the way the results are presented in the table, click **Manage Relationships**. Clicking on the **Manage Relationships** option bring also brings up a blue information panel ("What does this do?") that provides more detailed information.

To further manage column data, each column has a set of icons in its header. Mousing over any icon will reveal what it does. The sort icon resembles two triangles and allows you to sort a column in either direction (ascending/descending; a ->z, z ->a). To delete any column from your table, click on the "x" icon. Alternatively, the three dots "…" can be selected to "Toggle column visibility" and hide the column. Once hidden, the icon turns into a double arrow that can be selected to expand the column into its original form. The next icon resembling a funnel can be selected to edit or remove any currently active filters. A summary of the data within a column can be viewed by clicking on the icon that resembles a graph. The data within each summary can be selected for further filtering or downloading.

Templ	nplates lates are predefined queries, each has a simple form and a description. Click on a template to run it, you can search for templates by keyword Iter them by category.
ote: Please conta	ct us if you would like any additional template queries or if you have a concern about a query not completing.
ïlter:	Filter: all categories 📀 Reset
Actions: Export :	elected Options: 🕑 Show descriptions 🗌 Show Tags
	You are not logged in. Log in to mark items as favourites 😭.
	omosomal location e id or gene symbol, retrieve the chromosomal coordinates.
Gene> GO Given a gen	terms e id or symbol, retrieve GO terms. Be sure to view the Qualifier column for genes annotated as "NOT".
	location> genes omosome id and coordinates, retrieve genes.
	Chromosome Location 2 rsID, retrieve chromosome location.
Gene> Path Given a gen	way e id or symbol, retrieve pathways. The pathway data sources retrieved may be affected by the input id (RefSeq, Ensembl or gene symbol).
SNP rsID> (Given a SNP	Gene PrsID and organism, retrieve gene. No rows will be returned if the SNP is not within a gene.
Gene> Pub	lications e id or symbol, retrieve publications.

□ Gene ID --> Variant Consequences

Given an Ensembl gene id, retrieve variants and their consequences.

Fig. 6: Full list of templates on Templates page

	Given a gene id or symbol in the selected organism, retrieve homologues. Optionally select homologue type.									
	Gene									
	LOOKUP: GSTM1		0							
	🗆 constrain to be 🛛 🔊 saved	Gene list B. taurus Ensembl95 All Genes	۵							
optional ON OFF	Organism > Short Name = 🗘 B. taurus	0								
	Homologue > Type									
optional ON OFF	= O (orthologue O)									
	Show Results			Edit Que	ry					
	📾 web service URL	Perl Python Ruby Java [help]		export XML						

Fig. 7: Example: Gene -> Homologue

🗆 Manage	Columns T Manag	e Filters	< Manage Relat	ionships	່ວ Undo	•		Generate Python	code 👻 🖹 Export
Showing 1 to	25 of 155 rows				Row	s per page: 25 🔹	← «-	← page 1 → → →	
≎ x ···· ▼ Gene Organism	≎ × ··· ▼ Gene Gene ID	≎ × ···· ▼ ddl Gene Source	≑ x ··· ▼ <u>III</u> Homologues Type	¢ × ···· ▼ <u>III</u> Homologues Homologue . Primary I	Identifier	≑ x ···· ▼ dd Data Sets Name	♦ X ···· Y <u>Idd</u> Data Source Name	≎ x ··· र जा Homologues Homologue . Organism . Short Name	[▲] x ···· Y <u>III</u> Homologues Last Common Ancestor
B. taurus	ENSBTAG00000017765	Ensembl95	orthologue	ENSGALG00000032922		EnsemblCompara data set	Ensembl	G. gallus	Amniota
B. taurus	ENSBTAG0000037673	Ensembl95	orthologue	ENSGALG00000016328		EnsemblCompara data set	Ensembl	G. gallus	Bilateria
B. taurus	ENSBTAG0000037673	Ensembl95	orthologue	ENSG00000134201		EnsemblCompara data set	Ensembl	H. sapiens	Boreoeutheria
B. taurus	ENSBTAG0000037673	Ensembl95	orthologue	ENSG00000213366		EnsemblCompara data set	Ensembl	H. sapiens	Boreoeutheria
B. taurus	ENSBTAG0000037673	Ensembl95	orthologue	ENSMUSG0000004035		EnsemblCompara data set	Ensembl	M. musculus	Boreoeutheria
B. taurus	ENSBTAG0000037673	Ensembl95	orthologue	ENSRNOG0000018937		EnsemblCompara data set	Ensembl	R. norvegicus	Boreoeutheria
B. taurus	ENSBTAG00000017765	Ensembl95	orthologue	ENSCHIG0000023592		EnsemblCompara data set	Ensembl	C. hircus	Bovidae
B. taurus	ENSBTAG00000017765	Ensembl95	orthologue	ENSCHIG0000024688		EnsemblCompara data set	Ensembl	C. hircus	Bovidae
B. taurus	ENSBTAG0000017765	Ensembl95	orthologue	ENSOARG00000019297		EnsemblCompara data set	Ensembl	O. aries	Bovidae

Fig. 8: Example: Gene -> Homologue template search results, identifier for Gene GSTM1

🔓 Save	e as List -	ⓓ Generate Python code ■ Export				
		Python				
	page 1 –	Perl	1			
		Java				
×		Ruby	≑ × ··· ▼ <u>IIII</u>			
ata ource	Homologue Homologue	JavaScript	Homologues Last Common			
ame	Name	XML	Ancestor			

Fig. 9: Options for generating code from template query

Download TSV file	File name	
	Gene_to_orthologues	.tsv
All Columns		x Tab separated values.
All Rows	A flat file format suitable for spreadsheet programmes	Comma separated values.
No Compression	Destination	XML
No Column Headers	 Download file 	{}JSON
Dura di auto	 Send to Galaxy 	FASTA sequence.
Preview	 Upload to Genomespace 	GFF3 features.
		BED locations.

Fig. 10: Options for downloading results from template query



Fig. 11: Options for customizing the results of a template query.

≑ x T		≑ ×…▼		≑ × ··· ▼ 📶		≑ × ··· τ <u>Iai</u>
ald	↑×…▼ <u>Iall</u>	dil	≑ × … ▼ 📶	Homologues	¢ × ···· ▼ 📶	Data
Gene	Gene	Gene	Homologues	Homologue . Primary	Data Sets	Source
Organism	Gene ID	Source	Туре	Identifier	Name	Name

Fig. 12: Column header icons available to edit the results of a template query.

2.2.4 Optional filters

Some templates have optional filters that are disabled by default. For the Gene -> Homologue query, there is an optional filter to specify the organism used in the query. To enable the filter, click **ON** below the **Organism > Short Name** label.

	Gene Homologues Given a gene id or symbol in the selec	ted organism, retrieve homologues. Optionally sel	ect homologue type.
	Gene		
	LOOKUP: RCAN1		0
	🗆 constrain to be 🛛 🔊 saved	Gene list B. taurus Ensembl95 All Genes	
optional ON OFF	= B taurus	0	
	Homologue > Type		
optional ON OFF			
	Show Results		Edit Que
	📾 web service URL	Perl Python Ruby Java [help]	export XML

Fig. 13: Using the optional organism filter in the Gene -> Homologue query template.

2.3 QueryBuilder

The provided templates are suitable for many different types of searches, new queries may be built from scratch using the **QueryBuilder**. The possibilities of queries using the QueryBuilder are endless. The output may be formatted exactly as desired, and the query constraints may be chosen to perform complex search operations.

Advanced users can use a flexible query interface to construct their own data mining queries. The QueryBuilder lets you view the data model, apply constraints and select output. You can also export queries to share them with others. Browse data model O Import query from XML O View saved queries O	Alias Name Author BRENDA Term Bio-Entity CDS CMO Term CRISPR
--	--

To begin, select a **Data Type**. For example, select **Gene** as a Data Type then click the **Select** button to be taken to the Model browser.

Click on a class name for a starting at that class	description or double click on a class name to create a new query
Gene Protein Alias Name Author BRENDA Term Bio-Entity CDS CMO Term CRISPR	Gene: "Gene" is an abstract term used to describe a collection of transcripts and related regulatory features.

Fig. 14: Selecting data type as **Gene** in QueryBuilder.

2.3.1 Model browser

After selecting a data type, the **Model builder** appears displaying the attributes for the chosen feature class **Gene**.

First lets select Gene as a Data Type in the QueryBuilder. Then click on Select. This will take you to a Model browser where you can select the attributes for the feature class 'Gene', which you would want to be shown in your results.

2.3.2 QueryBuilder Examples

The following three examples provide details as to how to use the QueryBuilder using "Gene" as the selected data type.

2.3.2.1 Example 1: Querying for protein coding genes

In the Model browser, click **Show** next to **Biotype**, **Gene ID** and **Symbol**, which will add these fields to the query. Notice that these two fields appear below the data type **Gene** in the Query Overview section.

Then click **Constrain** next to **Biotype**. The first drop-down menu defaults to = (equals sign). In the second drop-down menu, select **Protein Coding**, then click the **Add to query** button. This adds a constraint to the query to search only for protein coding genes. Notice that the Query Overview section now shows "Biotype = Protein Coding". Also, two types of icons appear next to the attributes. Clicking on the red "X" icon next to an attribute will remove that field or constraint from the query. Clicking on the blue pencil icon next to a constraint brings up the constraint editing window where changes may be made to the query filters.

Model browser Query Overview Browse through the classes and attributes. Click on [UIIIARY] links to add summary of fields to the results table or on [NIIIIARY] links to add individual fields to the results. Use [OIIITRAIN] constrain a value in the query. no fields constrained Gene [I] [UIIIARY] [OINITRAIN] - Biotype [NIIIII [OIIIITRAIN] - Description [RIIIII] [OIIIITRAIN] - Description [RIIIIII] [OIIIIIIIIIIIIIIIIIIIIIIIIIIIIII		
Inks to add summary of fields to the results table or on SHOW+ no fields constrained Inks to add individual fields to the results. Use CONSTRAIN+ Inks Inks to add individual fields to the results. Use CONSTRAIN+ Inks Cene SUMMARY + CONSTRAIN+ Inks Biotype SHOW+ CONSTRAIN+ - Description SHOW+ CONSTRAIN+ - Length * Integer SHOW+ CONSTRAIN+ - Name * SHOW+ CONSTRAIN+ - Secondary Identifier * SHOW+ CONSTRAIN+ - Source (SHOW+ CONSTRAIN+) - Status SHOW+ CONSTRAIN+ - Status SHOW+ CONSTRAIN+ - H CDSs CDS * SUMMARY + CONSTRAIN+ - H Chromosome Chromosome SUMMARY + CONSTRAIN+ - H Data Sets Data Set SUMMARY + CONSTRAIN+ <	Model browser	Query Overview
	Browse through the classes and attributes. Click on SUMMARY + links to add summary of fields to the results table or on SHOW+ links to add individual fields to the results. Use CONSTRAIN+ links to add individual fields to the results. Use CONSTRAIN+ links to add individual fields to the results. Use CONSTRAIN+ links to add individual fields to the results. Use CONSTRAIN+ links to add individual fields to the results. Use CONSTRAIN+ Inks to add individual fields to the results. Use CONSTRAIN+ Biotype SHOW+ CONSTRAIN+ Description SHOW+ CONSTRAIN+ Name SHOW+ CONSTRAIN+ Secondary Identifier SHOW+ CONSTRAIN+ Secondary Identifier SHOW+ CONSTRAIN+ Status SHOW+ CONSTRAIN+ Symbol SHOW+ CONSTRAIN+ - Status SHOW+ CONSTRAIN+ - CDSs CDS SHOW+ CONSTRAIN+ - Chromosome Choromosome ButmARY + CONSTRAIN+ - Chromosome Location Location B SUMMARY + CONSTRAIN+ - Cross References Cross Reference B SUMMARY + CONSTRAIN+	no fields constrained Constraint logic:
	Fields selected for output	

Columns to Display

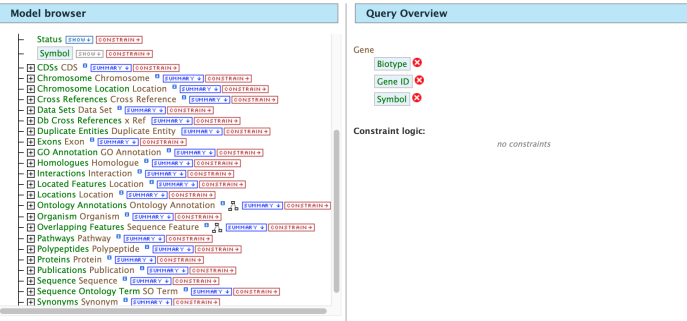
Use the SHOW+ or SUMMARY + links to add fields to the results table.

No fields selected for output

Show results

Show results

Fig. 15: Model browser with "Gene" selected as the data type.



Show empty fields

Fig. 16: Model browser with "Gene" selected as the data type. Example 1, Step 1: Select fileds to be added to the query

Lastly, click on **Show Results** above the Model Browser. The resulting table contains all protein coding genes in the database, with Gene ID, Gene Symbol and Gene Biotype as the table columns. Because the Biotypes should all be the same (protein coding), that column can be deleted by clicking the "x" above it. Alternatively, the three dots "…" can be selected to "Toggle column visibility" and hide the column. See the *Customize output and manage columns* section to review ways to manage column data.

2.3.2.2 Example 2: Querying for protein coding genes on a particuler chromosome

Users can customize the previously run query by adding another constraint for **Chromosome**. Note the "Trail: Query" text at the upper left of the results table. Clicking on the "Query" link will bring you back to the query that generated the table to allow for edits without having to build a new query. In the Model browser, click on the "+" (plus sign) next to the Chromosome feature class to display its attributes.

Next click on **Constrain** next to the **Chromosome ID** attribute and in the text box of the pop-up window select "=" (equals). Enter **18** for chromosome number 18then click on **Add to Query**, which adds the additional constraint to the query. Just clicking chromosome 18 will bring up that chromosome in multiple organisms so we will add another organism constraint. Click on the **Organism** to reveal its attributes then click on **constrain** next to Genus. In the resulting pop-up window, slect "=" (equals) then "Equus" for the chromosome 18 of horse.

Click on **Show results** and the query will result in all protein-coding genes on the Chromosome with constrained ID and organism. Note that the number of results has been reduced with the addition of the constraint.

Home 💄 MyMine Templates Lists Qu	Other	ata Sources Help	API Contact Us best	goat@fangpicking.edu Lo
	Processed Pseudogene Protein Coding			ch: e.g ATP synthase, AT5G2
	Pseudogene	_		e.g ATP synthase, AT5G2_
	RNase MRP RNA			
	SRP RNA			Show results
	TR C gene			
	TR V gene			
Model browser	TR J gene	Query Overvie	ew	
	V segment			
	antisense RNA			
links to add individual fields to the resul	quide RNA	Gene		
to constrain a value in the query.	lincRNA	Biotype 😣		
	IncRNA	Biotype		
	miRNA	Gene ID 🙆		-
Biotype SHOW CONSTRAIL Constraint for Gen	misc RNA			x
— Description SHOW ↓ CONST	rRNA			
- Length Integer SHOW + Constrain	ribozyme			
Hune Brown Constants	sRNA			
Gene ID SHOW CONSTR	scaRNA			
- Secondary Identifier	snRNA			
- Source SHOW+) CONSTRAIN-	snoRNA	ecific value		
- Status SHOW CONSTRAIN	tRNA	Jecific value		
- Symbol SHOW + CONSTRAIN Gene > Biotyp	telomerase RNA			
	,			
- + Chromosome Chromosom OR Contained	in list: IN OB	taurus Ensembl95 All Genes	Add to query	
- F Chromosome Location Lo				
- I Cross References Cross R				
🗕 🛨 Data Sets Data Set 🎙 💷 🔽 Filter query resu	Its on this field having a	any value or not ———		
– 🛨 Db Cross References x Re 💿 Has no value	Has a value Add to d			
🗕 🛨 Duplicate Entities Duplica	Add to t	query		
- + Exons Exon B SUMMARY 4			,	
- I GO Annotation GO Annotation				
Homologues Homologue				
<u>, </u>				
Show empty fields				
Fields selected for output				

Fig. 17: Example 1, Step 2 Adding a constraint to the query on Biotype.

Trail: Query > Results						
Manage Columns	▼ Manage Filters	Kanage Relationships		Save	e as List • 🕼 Generate Python code 🔹	
Showing 1 to 25 of 346,	084 rows		Rows per page: 25 ♣	· «- «	$\vdash \dots \rightarrow \rightarrow \rightarrow \rightarrow $	
▲ x ··· YI Gene Biotype		Ger	····▼ lini ne ne ID		≎ x ··· ▼ .ht Gene Symbol	
Protein Coding			008585	TICAM1		
Protein Coding		100	009677	HK2		
Protein Coding		100	009678	FMOD		
Protein Coding		100	009679	HSPA8		
Protein Coding		100	100009680		HTR1B	
Protein Coding		100	100009681		LUM	
Protein Coding		100	009682		EEF1A1	
Protein Coding		100	009683		HTR1A	

Fig. 18: Example 1, Step 3 Display query results.

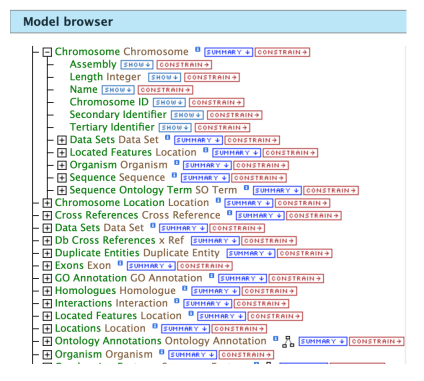


Fig. 19: Example 2, Step 1 View attributes of Chromosome feature class

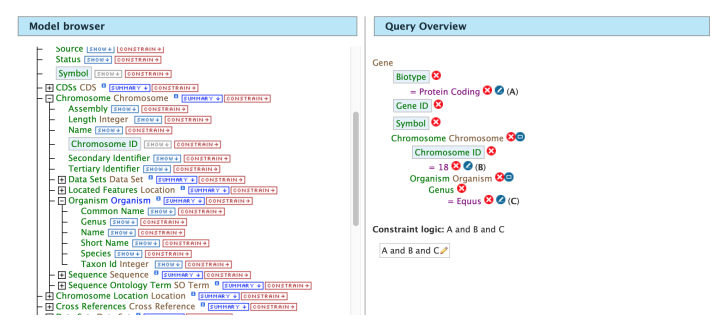


Fig. 20: Example 2, Step 2 Adding constraints to the Chromosome ID and Organism

Trail: Query > Results			
	rs		
S Manage Relationships		Save as	s List → 🕼 Generate Python code 🖌 🗈 Export
Showing rows 1 to 25 of 803	Rows per p	age: 25 ♣ + ← pag	$pe 1 \rightarrow \rightarrow \rightarrow \rightarrow$
<mark>^ x … Y lill</mark> Gene Biotype	≎ x ··· ▼ ևll Gene Gene ID	≎x…▼ <u>ldd</u> Gene Symbol	◆ x ··· ▼ Int Chromosome Chromosome ID
Protein Coding	100033832	MSTN	18
Protein Coding	100033909	ACTR3	18
Protein Coding	100034068	TNFAIP6	18
Protein Coding	100049822	DBI	18
Protein Coding	100049823	ACVR2A	18
Protein Coding	100049893	C1QL2	18
Protein Coding	100049894	MBD5	18
Protein Coding	100049957	INSIG2	18
Protein Coding	100049958	EPC2	18

Fig. 21: Example 2, Step 3 Result table after constraining by Chromosome ID and Organism

2.3.2.3 Example 3: Querying for Protein Coding genes on a particular chromosome and their exons

This last example builds upon the previous queries to display all exons for each gene on a particular chromosome. Again, note the "Trail: Query" text at the upper left of the results table. Clicking on the "Query" link will bring you back to the query that generated the table to allow for edits without having to build a new query. Scroll down to **Exon** feature class and select the "+" (plus sign) next to Exons to display its attributes. Click **Show** next to **Length** and **Exon Identifier**.

The Query Overview shows the query in progress with the selected fields. Also notice that a third type of icon, a blue square, appears next to some attributes. Clicking on a blue square icon brings up a window where the query Join Style may be modified. When adding a constraint, you can decide whether you want to show only those results with the information (genes with exons) or all results and the constrained feature if it exists (e.g., all genes and indicate exons if they exist). Click on the blue square icon next to **Exon collection** to bring up the Switch Join Style window. The default option is to show only Genes if they have a exon (inner join). Change this to **Show all Genes and show Exons if they are present** (outer join) then click **Add to query**.

Then click "Show results" to run the new query.

The results table now lists a new column **Gene Exons**, which we added to the query. If we look at the second row in the table, it lists 12 exons. Click on the **12 exons** link to expand that entry. That column now has additional rows containing the **Exon identifier** and **Length** for each of the 12 exons.

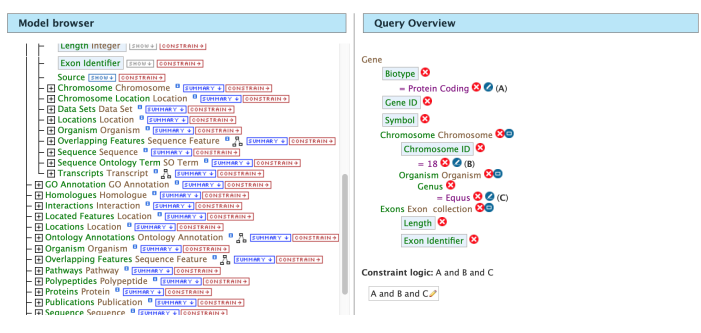


Fig. 22: Example 3, Step 1 Expand exon attributes and add fields to the query

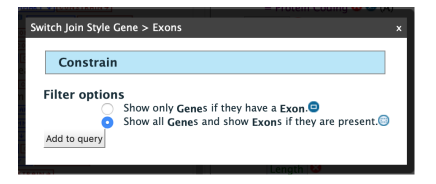


Fig. 23: Example 3, Step 2 Select join style for exons

Trail: Query > Results				
Manage Columns Manage	age Filters			
Manage Relationships			lob Save as List - I I I I I I I I I I I I I I I I I I	Python code
Showing rows 1 to 25 of 803		Rows per page: 25 ♣	- ← ← page 1 → → →	
[▲] x ···· ▼ Gene Biotype	≎ x ··· ▼ ini Gene Gene ID	¢x…▼lini Gene Symbol	★ × ··· ▼ <u>Int</u> Chromosome Chromosome ID	i≣ x ··· ▼ 네 Gene Exons
Protein Coding	100033832	MSTN	18	I 3 Exons
Protein Coding	100033909	ACTR3	18	I2 Exons
Protein Coding	100034068	TNFAIP6	18	I 6 Exons
Protein Coding	100049822	DBI	18	I 14 Exons
Protein Coding	100049823	ACVR2A	18	II Exons≣
Protein Coding	100049893	C1QL2	18	III 2 Exons
Protein Coding	100049894	MBD5	18	I 65 Exons
Protein Coding	100049957	INSIG2	18	I 49 Exons
Protein Coding	100049958	EPC2	18	I 14 Exons
Protein Coding	100050022	CCDC93	18	I 140 Exons

Fig. 24: Example 3, Step 3 Query results with exon constraint

Manage Columns	▼ Manage Filters		
K Manage Relationship	ips		Save as List ◄ 🕼 Generate Python cod
showing rows 1 to 25 o	of 803	Rows per page:	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$
[▲] ★ ···· ▼ <u>.ltl</u> Gene Biotype	≎ x ··· ▼ <u>lill</u> Gene Gene ID	≎ x ··· Y <u>lill</u> Gene Symbol	× ··· ▼ Int Chromosome Gene Chromosome ID Exons
Protein Coding	100033832	MSTN	18 ≣ 3 Exons
Protein Coding	100033909	ACTR3	18 🗮 12 Exons
			● Length ● Exor
			311 exon3
			56 exon3
			125 exon3
			111 exon3
			96 exon3
			108 exon3
			144 exon3
			174 exon3
			93 exon3
			126 exon3
			84 exon3
			1230 exon3
Protein Coding	100034068	TNFAIP6	18 🔚 6 Exons

Fig. 25: Example 3, Step 4 Query results with exon column expanded

In changing the join style to an outer join, the exons have been grouped together by gene making it easier to determine how many exons are contained in each gene. If the same query is run with the default join (outer join) of **Show only Genes if they have an Exon**, the results table adds a new row for each new exon rather than grouping exons by gene.

Trail: Query > Results					
Manage Columns	▼ Manage Filters				
K Manage Relationships			Save	e as List 🗸 🕼 Generate I	Python code
Showing rows 1 to 25 of 51	1,208	Rows p	per page: 25 ♦ I← ← ←	→ → →]
▲ x ···· Y lılıl Gene Biotype	≎ × ··· ▼ Gene Gene ID	≎ x ··· ▼I Gene Symbol	≎ x ···· ▼ III Chromosome Chromosome ID	≑ × ··· ▼ <u>lill</u> Exons Length	\$ X ··· ▼ ⊡ Exons Exon Identifier
Protein Coding	100033832	MSTN	18	373	exon370698
Protein Coding	100033832	MSTN	18	374	exon370699
Protein Coding	100033832	MSTN	18	381	exon370700
Protein Coding	100033909	ACTR3	18	56	exon335560
Protein Coding	100033909	ACTR3	18	84	exon335569
Protein Coding	100033909	ACTR3	18	93	exon335567
Protein Coding	100033909	ACTR3	18	96	exon335563
Protein Coding	100033909	ACTR3	18	108	exon335564
Protein Coding	100033909	ACTR3	18	111	exon335562
Protein Coding	100033909	ACTR3	18	125	exon335561

Fig. 26: Example 3, Step 5 Query results with default join style run for genes containing exons

CHAPTER 3

Report Pages

All objects in FAANGMine (e.g., gene, protein, transcript, publication) have report pages that can viewed after running a query. It allows users to view all available information for that object while providing links to related objects. As an example, we can revisit the templates example. In the list of templates under the **Templates** tab on the FAANGMine home page, select Gene -> Homologues to query FAANGMine to retrieve all homologues for a given gene. Enter "GSTM1" into the the LOOKUP search box then click **Show Results**. In the results table, note that every entry is contains a link. You can mouse over any link to bring up a summary of that object. If we hover over the first Gene ID, we can see a summary box that includes

Clicking on that same item will bring up its report page that includes a comprehensive for gene GSTM1. The report page header shows the Gene ID and its Biotype, for this example, protein coding. The tabs at the top of the page in the Quick Links menu bar quickly bring you to the data listed. The column on the right side of the report page displays external links to other Mines and databases.

The content of the report page is divided into categories based on the type of information provided for that particular object. Clicking on links within each category bring up more details about the objects of interest.

3.1 Summary

The **Summary** section near the top of the report provides information on the gene such as its length, chromosome location, and strand information. Users can also get the complete FASTA sequence of the gene by clicking on the FASTA tab.

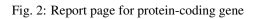
3.2 Transcripts

The **Transcripts** section contains information about the gene model, such as transcripts and exons. Links to FASTA files are included where applicable.

🖽 Manage Columns		▼ Manage F	▼ Manage Filters		nage Relationships	
Showing row	/s 1 to 25	Gene				p
		Biotype	Pr	otein Cod	ing	
≎ × ··· ▼ 🛄 Gene	^ x ··· ▼ Gene	Chromosome	AF	RS-UCD1.	2	
Organism	Gene ID	Description	glutathione S-transferase M1			fie
B. taurus	<u>327709</u>	Gene ID		327709		
B. taurus	327709	Length	10	,766		
D. taurus	521105	Organism	В.	B. taurus		
B. taurus	327709	Source	Source RefSeq			
B. taurus	327709	Symbol	G	STM1		
B. taurus	327709					
B. taurus	327709	RefSeq ort		orthologue 100156976		
B. taurus	327709	RefSeq or		orthologue 100169963		
B. taurus	327709	RefSeq orth		thologue 100514231		
B. taurus	327709	RefSeq	RefSeg ortholog		100861222	

Fig. 1: Summary for gene entry in query results table

Gene : GSTN	1 B. taurus				
Gene ID 🏾	327709	So	urce	RefSeq	
Description	glutathione S-transfera	ase M1 Bio	otype	Protein Coding	
SHARE					
Quick Links:	Summa	ary Transcripts Pro	teins Function Hor	nology Interactions Public	ation Other
Genome feature					Lists
Region:	gene 🖁	Length:	10766	FASTA	This Gene is in one list:
Location:	3:33690883-3	3701648 reverse strand			B. taurus RefSeq All Genes (30580)
1 Db Cross Referen	ices				Links to other Mines
Manage Columns	▼ Manage Filters			ⓓ Generate Python code ▼	Export HumanMine H. sapiens
K Manage Relations	hips			🕰 Sav	e as List - CSTM4 [™] , GSTM5 [™] , CSTM2 [™] , GSTM3 [™] , CSTM1 [™]
Showing rows 1 to 1 o	of 1				MouseMine M. musculus Gstm4 ^[A] , Gstm1 ^[A] , Gstm6 ^[A] , Gstm3 ^[A] ,
x ···· ▼ Inl Db Cross References Identifier			x ···· ▼ [.ii] Db Cross References Source		Gstm7 [™] , Gstm5 [™] , Gstm2 [™] YeastMine RatMine
ENSBTAG00000017765			Ensembl95		R. norvegicus Gstm4 ^A , Gstm1 ^A , Gstm7 ^A , Gstm3 ^A ,



Genome feature			
Region:	gene ^B	Length:	10766 FASTA
Location:	3:33690883-33701648 rev	verse strand	

Transcripts

All Transcripts for Gene - Transcripts: 1 Exons: 8 Coding		
Transcript	Exons	Coding Sequence
	exon632091	100 [FASTA]
	exon632092	76 FASTA
	exon632093	65 FASTA
GSTM1 NM_175825.3	exon632094	82 FASTA NM_175825.3-CDS
1141 FASTA	exon632095	101 FASTA 657 FASTA
	exon632096	96 FASTA
	exon632097	111 FASTA
	exon632098	510 FASTA

3.3 Proteins

The **Proteins** section provides information about the protein product of the gene. The comments section gives a brief description about the protein along with the UniProt accession and links to any outside data sets.

3.4 Function

The Function section displays Gene Ontology annotations for a gene. Annotations are divided into three categories:

- · Cellular Component
- Molecular Function
- Biological Process

The GO terms are displayed along with the evidence code indicating how the annotations were derived. A results with Pathway information is also displayed if applicable.

3.5 Homology

The **Homology** section provides information for all homologues. The first portion displays a summarized view of the homologues reported in different organisms. The next portion provides more detailed information about the homologue, the type of homologue and from which dataset the information was obtained all displayed in a results table.

3.6 Interactions

The **Interactions** section provides interaction information. For GSTM1 there are no interaction information available but for genes that do have interaction information, a network is displayed showing all interactors for the current gene.

Proteins

Curated comments from UniProt					
Туре	Comment				
function	Conjugation of reduced glutathione to a wide number of exogenous and endogenous hydrophobic electrophiles. Protects aga mediated metal-catalysed oxidative inactivation of enzymes.	inst the thiol-			
similarity	Belongs to the GST superfamily. Mu family.				

1 Proteins				
II Manage Columns	▼ Manage Filters	团 Generate Python code	•	Export
Kanage Relationship	DS		🗅 Sa	ave as List -

Showing rows 1 to 1 of 1

 ★ x ··· ▼ [.iii] Proteins DB identifier 	★ x ··· ▼ III	x ···· ▼ <u>III</u>	★ x ···· ▼	x ···· ▼ III
	Proteins	Proteins	Proteins	Proteins
	Primary Accession	Organism . Name	Length	datasets
GSTM1_BOVIN	Q9N0V4	Bos taurus	218	I≣ 3 Data Set datasets

Function

Gene Ontology	
cellular component cytoplasm	ECO:0000501
molecular function glutathione transferase activity ^B	⑧ ECO:0000501 □
biological process	
No terms in this category.	

17 Pathways							
Manage Columns	▼ Manage Filters	ⓓ Generate Python code ▮ Export					
Kanage Relationshi	os	▲ Save as List -					

Showing rows 1 to 17 of 17

★ ★ ··· ▼ <u>iiil</u> Pathways Identifier

R-BTA-1430728 R-BTA-156580 R-BTA-156590 R-BTA-211859

1 to 17	of 17	Rows per page: 10 🛊
	¢ x ··· ▼ <u>Iıll</u> Pathways Name	x ··· ▼ .inl Pathways Dataset
1	Metabolism	i≣ 1 Data Set Dataset
	Phase II - Conjugation of compounds	I Data Set Dataset
	Glutathione conjugation	I Data Set Dataset
	Biological oxidations	i≣ 1 Data Set Dataset
		Reactome pathways data set

Homology

B. bubalis	C. hircus	C. lupus familiaris	E. caballus	F. catus	G. gallus	H. sapiens	M. mus	culus	O. aries	R. norvegie	cus	S. scrofa
LOC102396303 LOC102398085 GSTM3 LOC102397004	GSTM3 LOC106503993 LOC108633298 LOC100861222 LOC102185621 LOC102185621 LOC102189813	LOC479911 LOC479912	GSTM3 LOC100061761 LOC100058290 LOC100058329	LOC101100824 GSTM3	GST2L GSTM2	GSTM2 GSTM3 GSTM4 GSTM1 GSTM5	Gstm Gstm Gstm Gstm Gstm Gstm	1 5 4 3 2	GSTM3 LOC101107831 LOC101108092 LOC101108705 LOC101107401	Gstm6		LOC10651020 LOC11026035 LOC11026035 LOC780435 LOC10015697 LOC11026034 GSTM3
🌸 Gene>	Homologues	for report p	age (59 row	s) 🏠								
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« Manage R	elationships									6	Sa	ve as List -
Showing rows	1 to 25 of 59											
	1 10 25 01 55			Rows p	er page	25	\$	I←	* ←	page 1		· → →I
	1 10 23 01 39			Rows p	er page	25	\$	←	* ←	page 1		* →I
▲ x ···· Y III Data Sets Name	¢ x ···· ▼ Homologue Type	 ★ x ··· ▼ III Homologue Last Comm Ancestor 	es non	Rows p ✦ x … Y Homologues Homologue . Pr Identifier		23	\$ × ···	▼ <u>III</u> logue	 ★ ★ ··· ▼ Intervention ↓ ★ ··· ▼ Intervention ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓ ↓	<u>l</u> les		
Data Sets	≑ × … ▼ ⊡ ll Homologue	Homologue s Last Comm	non	≑ × ··· ▼ ⊡ Homologues Homologue . Pr		23	‡ x … Homo	T ⊡II blogue ol	¢ x ···· Υ ⊡ Homologu Homologu	<u>l</u> les		

Interactions

🌸 Gene --> Interacting genes (0 rows) 🖙

3.7 Publications

The **Publications** section displays a table of publications related to the gene with links to full citations.

Publication

🜸 Publications (3 rows) 🏠						
Manage Columns	▼ Manage Filters		Generate Python code	•	Export	
K Manage Relationship	os			🔒 Sa	ave as List -	

Showing rows 1 to 3 of 3

→ X ··· ▼ Int	★ ★ ···· ▼ Int	◆ x ··· ▼ Int	★ ★ ···· ▼ Int	★ X ··· Y <u>ldd</u>	★ X ··· ▼ <u>Idd</u>	★ ···· ▼
Publications	Publications	Publications	Publications	Publications	Publications	Publications
Year	First Author	Title	Journal	Volume	Pages	PubMed ID
2015	Hering D M	Missense mutation in glutathione-S-transferase M1 gene is associated with sperm motility and ATP content in frozen- thawed semen of Holstein-Friesian bulls.	Anim. Reprod. Sci.	159	94-7	26091956

3.8 Other

This last section provides miscellaneous information that do not fit into any of the above categories. This example lists protein coding annotations and their sources.

Other

5 Data Sets

KEGG pathways data set, Bos taurus RefSeq Protein Coding Genes, OrthoDB data set, Swiss-Prot data set, PubMed to gene mapping

CHAPTER 4

Genomic Regions Search

The **Genomic Regions Search** is a tool to fetch features that are within a given set of genomic coordinates or are within a given number of bases flanking the coordinates.

To begin this type of search, click the **Regions** tab on the menu bar. A form will appear asking for the search parameters (organism, feature types, genomic coordinates, etc.)

The coordinates must have one of three formats:

- 1. chromosome_number:start..end
- 2. chromosome_number:start-end
- 3. chromosome_number start end (tab delimited)

Click on the input examples above the text input box (number 4) to view a representative set of coordinates in each format. Click the **Genome coordinates help** link near the top of the form for more detailed information on the input format requirements.

During a search, regions may be extended on either side of the genomic coordinates using the slider or by entering text in the field to the left of the slide bar. There is also the option to perform a strand-specific region search using the checkbox at the bottom of the form (number 6)

As an example, select *B. taurus* from the Select Organism drop-down, and ARS-UCD1.2 as the Assembly. Slick the box next to Select Feature Types to uncheck all of the boxes, then check the box next to Gene, and enter the following coordinates into the genomic regions search text field:

14:2000000..2800000

Click the search box to conduct the genomic regions search. If there are no overlaps within your search coordinates, the search can be done again with the search region extended using the slide bar or entering text into the search box (e.g., 10k).

The search results page presents a list of features present within the genomic interval that was searched. In this case, the feature type was limited to Gene. The results may be exported as tab-separated or comma-separated values. If they contain genomic features, there is also the option to saved the results in GFF3 or BED format. The FASTA sequences

Search for chromoson numbers fo Z,W for chi chromoson "ENTIRE GE	features that overlap a list ne identifier system for th or autosomes. Cat uses A1 cken. The mitochondrial c ne identifiers, you can use	of genome coor e selected specie A3, B1-B4, C1- hromosome is M	dinates you ent s, and be sure t C2, D1-D4, E1 T for all species	er or upload, e.g. to select both the -E3 and F1–F2 for 5. Scaffolds use Re	6:500001 species and r autosomes efSeq ids ("N	omic Region 00000. Be sure to use the the assembly. All species . Sex chromosomes are X, W") for all species. To re found on the FAANGMine	correct except cat use / for mammals and etrieve a map of
1.	Select Organism: B. bub	alis 🗘					
2.	Assembly: ARS-UCD1.2	0					
3.	✓ Select Feature Types:						
	CDS B		🗹 mRNA 🛛			🛃 tRNA 🗉	
	✓ Exon ^B ✓ Gene ^B		✓ Polypeptide ✓ rRNA [□]	e 8		🗹 Transcript 🏾	
	(example for input forma (example for tab delimite		,		6		
	or Upload genomic regio Choose File no file selec		2				
5.	Extend your regions at b	oth sides:					
		1k	10k	100k	1M	10M	
	0						
6.	Check this box to perference of the perfect of the perference of t	orm a strand-spe	cific region sea	rch (search + str	and if region	n start <end; search="" stran<="" td="" –=""><td>Reset Search</td></end;>	Reset Search

Fig. 1: Genomic Regions search form

Overlap features set Search for features that overlap a list of chromosome identifier system for the s numbers for autosomes. Cat uses A1-A Z,W for chicken. The mitochondrial chru chromosome identifiers, you can use th "ENTIRE GENE SET" Genome coordinates help 1. Select Organism: B. taurus 2. Assembly: ARS-UCD1.2	f genome coordinates you er elected species, and be sure \3, B1-B4, C1-C2, D1-D4, E omosome is MT for all specie le template query "Genome A	iter or upload, e.g. 6:500 to select both the specie 1–E3 and F1–F2 for autos es. Scaffolds use RefSeq io	00100000. Be sure to use the co s and the assembly. All species ex omes. Sex chromosomes are X,Y f ds ("NW") for all species. To retr	rrect cept cat use or mammals and ieve a map of
3. 🗹 Select Feature Types:				
 CDS ^B Exon ^B Gene ^B Polypeptide ^B Primary Transcript ^B Indel ^B Lnc RNA ^B mRNA ^B SNP ^B 4. Type/Paste in genomic regission (example for input formation (example for input formation (example for tab delimited) 14:20000002800000 	thr:11000)▼ thr:1-1000)▼	e 0 ot 0	 Sequence Alteration ^B Substitution ^B tRNA ^B Tandem Repeat ^B 	
or Upload genomic regions Choose File no file selected 5. Extend your regions at both	1			
	1k 10k	100k 1M	10M	
0			I	
 Check this box to perform end<start)< li=""> </start)<>	n a strand-specific region se	arch (search + strand if r	egion start <end; i<="" search="" strand="" td="" –=""><td>f region Reset Search</td></end;>	f region Reset Search

Fig. 2: Genomic Regions search example with Bos taurus

of the features may also be downloaded. Links within the features provide detailed reports. If users are interested in creating a list of particular features from the result page then they can filter based on feature type (if applicable), shown in red box, and click on **Go**.

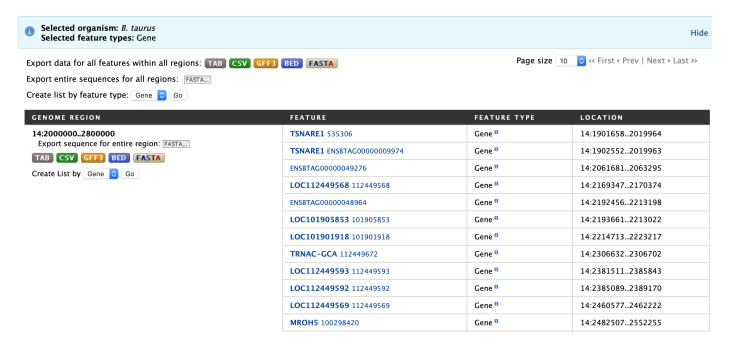


Fig. 3: Genomic Regions search results

Lists

5.1 Creating Lists

Users may create and save lists of features, such as gene IDs, transcript IDs, gene symbols, etc. The list tool searches the database for the list items and attempts to convert each identifier to the selected type. Click on the Lists tab from the menu to access the full list upload form. A short version of the form is also in the Quick List box on the home page.

As an example, enter the following comma-separated identifiers into the Lists upload form under the **Lists** tab. Notice that they do not have to be in the same format. A Summary table is displayed with the results of searching for each of the five identifiers in the list.

CAPN2, ENSCHIG00000014802, BTG1, XDH, 101107826

Leave the **Select Type** drop-down menu to **Gene** and the **Organism** drop-down to **Any**. Click on **Create List**. Note that you can also upload a list from a .txt file.

The summary table provides information regarding those identifiers that had a direct hit without any duplicates. If there are any duplicates, users can decide to add the relevant entries individually by clicking on the **Add** button under the **Action** column or choosing the **Add all** tab. Here we will click **Add all**. Once the selections have been added, the list can be saved by clicking the **Save a list of 66 Genes** button on the top of the summary table. Name the list by entering text into the **Choose a name for the list** box at the top of the results page.

After the list is saved, users are presented with a **List Analysis** page. This page provides users with widgets to perform analyses on gene lists that they have created.

The selection of widgets provided on the List Analysis page depend on the contents of the list. The available widgets for this list example include:

1. Gene Ontology Enrichment



Create a new list

Select the type of list to create and either enter in a list of identifiers or upload identifiers from a file. A search will be performed for all the identifiers in your list.

- Separate identifiers by a comma, space, tab or new line.
- Qualify any identifiers that contain whitespace with double quotes like so: "even skipped".

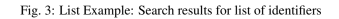
Select Type:	Gene
for Organism:	Any
Type/Paste in identifiers	(click to see an example with different types of IDs)▼ (example to query a list of milk production genes)▼
or Upload identifiers from a .txt file	Choose File no file selected Match on case
	Reset Create List

Fig. 1: List upload form

Enter a list of identifiers. Gene 515523, CAPN2, ENSCHIG00000014802, BTG1, ENSBTAG0000006858, 102398240		
515523, CAPN2, ENSCHIG00000014802,	tifiers.	
	0	Gene
BIG1, ENSBIAG0000006858, 102398240		
	2398240	BIG1, ENSE
advanced	,	
ANALYZE		

Fig. 2: Quick list from FAANGMine home page

ldentifier you provided	Matches	i								
	symbol	organism short name	chromosome assembly	length	description	secondary identifier	source	primary identifier	class	Act i
BTG1	BTG1	H. sapiens					RefSeq	694	Gene	Ad
	Btg1	M. musculus					RefSeq	12226	Gene	Ad
	Btg1	R. norvegicus					RefSeq	29618	Gene	Ad
	BTG1	H. sapiens					Ensembl95	ENSG00000133639	Gene	Ad
	Btg1	M. musculus					Ensembl95	ENSMUSG0000036478	Gene	Ado



Il_organism_gene	_search		(e.g. Sr	nith 2013)						
dd additi	ional mat	ches								
ou entere We found	d: 5 identi d: 2 Genes							Sav	e a list of	66 Genes
hy are the num	bers different?	See below.								
									Add all	Remove a
-			do you want?	? 🔟						
									5 rola	
aye i ol 13 k	2 3 4	5 15 /							0104	/s per page `
lage 1 of 13 4	Matches								0101	Action
ldentifier you	_		chromosome assembly	length	description	secondary identifier	source	primary identifier	class	
ldentifier you provided	Matches	i organism		length	description		source RefSeq	primary identifier		Action
ldentifier you provided	Matches symbol	i organism short name		length	description				class	Action
ldentifier you provided	Matches symbol BTG1	organism short name H. sapiens		length	description		RefSeq	694	class Gene	Action i Remove Remove
	Matches symbol BTG1 Btg1	organism short name H. sapiens M. musculus		length	description		RefSeq RefSeq	694 12226	class Gene Gene	Action

Fig. 4: List Example: Saving list of identifiers

- 2. Publication Enrichment
- 3. Pathway Enrichment
- 4. Orthologues

5.2 Saving Lists

To see your saved lists, click the **View** tab on the **Lists** page. If not logged in, lists will be saved temporarily during your current session. However, you must be logged in to save your lists permanently. Further analyses of lists can be done with the **Actions** links at the top of the list. The links become active once lists are selected for analyses. Saved lists may also be accessed from the **MyMine** menu tab.

List Analysis for All_organism_gene_search (66 Genes)

Manage Columns Manage Filters			ൾ Gen	erate Pyth	on cod	e		Export
S Manage Relationships						6	Save	as List •
Rows per page: 25	I←	≪	~	page 1	\rightarrow	-*	→I	

Showing 1 to 25 of 51 rows

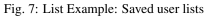
≑ × ··· ▼ <u></u> Gene Gene ID	♦ X ···· ▼ Identifier	≑ x ··· ▼ <u>i.i.i</u> Gene Symbol	♦ × ··· ▼ Gene Source	★ x ···· ▼1 Gene Description	♦ × ··· ▼ Gene Length	♦ X ··· T <u>.lil</u> Gene Organism	★ ★ ···· ▼ <u></u> Gene Assembly
100055184	NO VALUE	CAPN2	RefSeq	calpain 2	58996	E. caballus	EquCab3.0
100101473	NO VALUE	BTG1	RefSeq	BTG anti-proliferation factor 1	2743	S. scrofa	Sscrofa11.1
100127211	NO VALUE	CAPN2	RefSeq	calpain 2, (m/II) large subunit	49823	O. aries	Oar_v3.1
100217400	NO VALUE	BTG1	RefSeq	B-cell translocation gene 1, anti-proliferative	1235	O. aries	Oar_v3.1
100515259	NO VALUE	XDH	RefSeq	xanthine dehydrogenase	64984	S. scrofa	Sscrofa11.1
100629230	NO VALUE	BTG1	RefSeq	BTG anti-proliferation factor 1	2623	E. caballus	EquCab3.0

Fig. 5: List Example: Analysis for gene list

, ,	nent lenes in this list - data fron : list not analysed in this w		Counts	ologues s of genes in list with orthologues. ns in your list have been analysed.	
Test Correction	Max p-value	DataSet			
Holm-Bonferroni	0.05	KEGG pathways d	View	Download Organism.name	Genes
Default Change				Canis lupus familiaris	60
				Equus caballus	60
No enrichment found.				Gallus gallus	60
				Bos taurus	59
				Homo sapiens	59
				Mus musculus	59
				Ovis aries	59
				Sus scrofa	59
				Felis catus	58
				Capra hircus	57
				Bubalus bubalis	33

Fig. 6: List Example: Displayed widgets for list analysis





MyMine

MyMine serves as a portal where logged-in users may manage their lists, queries, templates, and account details.

To access MyMine, click on the MyMine menu tab. A submenu appears with six options:

Lists - Lists saved by the user when logged in.

History - List of most recently run queries.

Queries - List of saved queries.

Templates - Templates created or marked as "favorite" by the user.

Password - Password reset form.

Account Details - User preferences form.

Lists History 🏶 Queries 🌣 Templates Password Account Details								
		DESCRIPTION	Түре	NUMBER OF OBJECTS	DATE CREATED			
	All_organism_gene_search 2 🏠		Gene	66 values	2019-11-26 15:07			
	Chromosome_List_example_Equus a the second s		Chromosome	1 value	2019-11-26 14:33			

Fig. 1: Saved lists found under MyMine. Note that currently saved lists can be selected for analyses to contribute to new lists.

API

An API is available for users who would like to programmatically access FAANGMine.

Perl Web Service Client
The Perl web service client library makes it easy to run queries in FAANGMine directly from Perl programs. You can use these modules to construct any query you could run from web interface and fetch the results in a number of formats, including native Perl data structures and objects, and TSV/CSV strings.
Like all our code, it is open-source, coming licensed under the LGPL. For information on our API, visit our wiki pages.
Prerequisites
You should install the Perl webservice client library module to get started. You can install it directly from CPAN (the comprehensive Perl archive network). This is a public repository of thousands of modules. Using CPAN will ensure you get the latest version, and that your dependencies are automatically managed for you. To install the InterMine Perl client library type the following command into a shell:
 > sudo cpan Webservice::InterMine • Examples of using the downloaded script
On each Template Query page and the QueryBuilder there is a link to get Perl code to run that particular query using the web service API. Just click the link, save the generated Perl script in a file and execute it. You can use the generated code as a starting point for your own programs.
You can run the downloaded script by running the following command in a shell:
> perl path/to/downloaded/script.pl
If you get an error saying Cannot find Webservice::InterMine in @INC, or similar then see 'Prerequisites' above. Feel free to edit the script – these are designed to be spring-boards to help you get where you want to. See the Cookbook, which contains a set of short tutorial 'recipes' that demonstrate particular features of the Perl API, for ways to edit the scripts, and even extend their functionality. For a good reference to writing programs in Perl, see here.

Perl, Python, Ruby, and Java are the languages supported by the InterMine API.

For more detailed information, view the InterMine documentation.

Data Sources

The Data Sources table provides a description of the datasets that are integrated into FAANGMine, along with their download location, version or release, citations wherever applicable, and any additional comments.

Data This page lists all data sources loaded along with the date the data was downloaded and related publication. NOTE: All external links open in a new window.							
Data Category	Organism	Data	Source	PubMed	Note		
	Bos taurus	NCBI Genes (RefSeq)	NCBI Bos taurus Annotation Release 106 (ARS-UCD1.2)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Ovis aries	NCBI Genes (RefSeq)	NCBI Ovis aries Annotation Release 101 (Oar_v3.1)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Capra hircus	NCBI Genes (RefSeq)	NCBI Capra hircus Annotation Release 102 (ARS1)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Sus scrofa	NCBI Genes (RefSeq)	NCBI Sus scrofa Annotation Release 106 (Sscrofa11.1)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Gallus gallus	NCBI Genes (RefSeq)	NCBI Gallus gallus Annotation Release 104 (GRCg6a)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Equus caballus	NCBI Genes (RefSeq)	NCBI Equus caballus Annotation Release 103 (EquCab3.0)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Canis lupus familiaris	NCBI Genes (RefSeq)	NCBI Canis lupus familiaris Annotation Release 105 (CanFam3.1)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Felis catus	NCBI Genes (RefSeq)	NCBI Felis catus Annotation Release 104 (Felis_catus_9.0)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Bubalus bubalis	NCBI Genes (RefSeq)	NCBI Bubalus bubalis Annotation Release 101 (UOA_WB_1)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Homo sapiens	NCBI Genes (RefSeq)	NCBI Homo sapiens Annotation Release 109 (GRCh38.p12)	O'Leary et al – PubMed 26553804	NCBI FTP		
	Mus musculus	NCBI Genes (RefSeq)	NCBI Mus musculus Annotation Release 106 (GRCm38.p6)	O'Leary et al – PubMed 26553804	NCBI FTP		
Gene	Rattus norvegicus	NCBI Genes (RefSeq)	NCBI Rattus norvegicus Annotation Release 106 (Rnor_6.0)	O'Leary et al – PubMed 26553804	NCBI FTP		

Fig. 1: BovineMine Data Sources table

How to cite

FAANGMine is a project supported by the National Science Foundation to address the need for a high performance data mining resource that enables fine-grained querying and integrating the heterogeneous FAANG data with existing information, such as functions of known genes and research datasets.

For more generic examples on how to use InterMine, click here. These are tutorials created by FlyMine that showcase the different features of InterMine.