

# **Quick Start Guide**

### The geneXplain platform

### https://platform.genexplain.com/bioumlweb/

An inside look at how to create your personal research space on the geneXplain platform.

### What can I learn from this Quick Start Guide?

Whether you have just started to use the geneXplain platform or you have already experienced it, this Quick start guide will help you with some more detailed questions you may have.

You may wonder: How can I get started with my data? How to upload? What can I analyze? How to save my results?

This Quick start guide will answer those questions and more. We are going to take you deep in to the platform.

- $\times$  How do I get started?
- $\times$  How is the geneXplain platform organized?
- $\times$  How to work with the geneXplain platform?
- × How do I upload my data?
- $\times$  How to open/delete a file?
- × Which types of analyses are supported?
- × What is a workflow?
- $\times$  Where are the workflows on the platform?
- × How to start a workflow?
- $\times$  Are there examples on the platform?
- × How to organize my project and my data?
- × How can I get information about my account?



## How do I get started?

### Log in to the geneXplain platform

After registration you will receive an email from BioStore with your password (subject is "BioStore: registration").

For the log in to the platform you need your email address and the password.

Please, log in here to the platform:

https://platform.genexplain.com/bioumlweb/

Login	× ////////////////////////////////////
Platform:	geneXplain web edition 4.3
Enter e-mai	I and password:
E-mail:	MisterX@google.com
Password:	12345678
Register	Demo

When you login into the geneXplain platform for the first time, a window opens that contains the following areas A-E:

121 🗠 🔊 🖉 🔚				
6	Start page			
Databases Data Analyses	Α			
Users		Upstream ana	lysis	
Comparison of the second	RNA-seq	Proteomics Epigenomics	ChIP-seq Sequence analysis	miRNA
B	Microarrays	Drug targets	Pathways	NGS
	Genomic variants	Popular Gene or functions protein list	Complete list of workflows	Metabolism
Search Info ID: <u>Projects</u> Size: 1497 Complete name: data/Projects	Default V	User description is not avail	ription Graph search Script	Clipboard Tasks

**A** - The **Work Space** is the main part of the window. The Start page presents a couple of predefined workflows and methods.

**B**- In the **Tree Area** you find the collection of Databases, the uploaded data files and the available analyses methods under the corresponding tabs.

**C** - The **Info Box** will give you information about the data file or analysis method that you select with a single click in the Tree Area. You can also select the data resource to search in.

**D** - The **Operations Field** provides additional analysis options under the different tabs in a context-dependent manner.

**E** - The general **Control Panel** (tool bar), on top of the different areas, shows a context-dependent set of icons for the available operations.

When you enter your account for the first time, you can see the following three folders in the Tree Area (B) under the **Data** tab (red box): Examples, Projects and Public.



Each of these folders can be expanded (opened) by clicking on the small triangle on the left side of the folder name and symbol.

You can find **your** created **project** (red box; your project name from the registration) after opening the folder Projects.



Your project contains the subfolder **Data** (blue arrow), where all your uploaded data files and analyses results reside.

## How do I upload my data?

Three ways to **import** your **data**:

1. Go to your Data folder (marked blue) and right click on it; select "Import element" and a window "Import file" will open:

Select one of the different upload options: Computer, WebFTP, Repository or Raw.

Press the "Import" button to start loading data.

Image: Solution of the selected format         Image: Solution of the selected format         Image: Solution of the selected format		Import file	//////////////////////////////////////
Databases       Data       Analyses       Users         Import file from:       Computer       Web/FTP       Repository       Raw         Import file from:       Import file from:       Computer       Trept folder:       Gate       Trept folder:       Trept f		Target folder	Jata/Projects/Jeannette_Training/Data
Format: addetect       Import fall     Import fall	Databases Data Analyses Users	Import file from: Computer	Web/FTP Repository Raw
Cancel Import	▲ Jata         ▲ Projects         Projects         ■ DEMO         ● Be GerontoShield         ▲ Beanette_Training         ↓ Deta         ▶ Beanette_Training         ↓ Deta         ▶ New VS Script         ▶ Beanette_Training         ▶ Put ← New diagram         ▶ Put ← New optimization         ▲ Import element         ▲ Export element	Cancel	Import file       ×         Target folder. dataProject3/Jeannette Koschmann project0bta       Upbedreg fla GSM1992269_AR1264_Offrew.txt
			Cancel Import

- 2. Click the Import button of the tool bar [ ], to open the window "Import file".
- 3. Click on any button of the start page. The first list entry is always "Load Data" [ → Load data ]. Clicking on it will lead you to the window "Import file".

## How to open/delete a file?

All analysis tools, and likewise all prepared workflows, require input data from a file in the Tree Area.

Double-clicking on a file will **open** it under a new tab in the Work Space.

Files can also be opened by right-clicking on them and selecting the "Open table/track" option.

You can also **delete** files ("Remove"; default value is "No") this way.

The content of the table is sorted according to the values in one of its columns. Being opened for the first time, a default column is defined for sorting, usually the **ID** column. This default column is indicated by a blue arrowhead (red box).

Start page 🛃 Upre	egulated Ensembl ge 🕽						
				Edit	Apply Car	icel Selec	t all Select page
First Previous Page	e 1 of 11 Next Last	Showing 1 to 50 of 503 entries				Show	50 • entries
ID 🔺	Affymetrix ID 👙	Gene description	Gene symbol	\$	Species 🖕	-log(P- value) 🔶	LogFoldChange
ENSG0000001167	204108 at	alpha,nuclear transcription factor Y	NFYA		Homo sapiens	3.77848	1.34116
ENSG0000002586	<u>201028 s at,</u> 201029 s at	CD99 molecule	CD99		Homo sapiens	3.26959	2.9574
ENSG0000004455	<u>208967 s at</u>	adenylate kinase 2	AK2,RP4-592A1.2		Homo sapiens	3.1983	0.94765
ENSG0000011304	202189 x at, 211270 x at, 211271 x at, 212015 x at, 216306 x at	polypyrimidine tract binding protein 1	PTBP1		Homo sapiens	4.02757	1.21257
ENSG0000014123	<u>212633 at</u>	UFM1-specific ligase 1	UFL1		Homo sapiens	3.53308	1.22277

You may **sort** the table according to the values of any column in ascending or descending order by clicking on the up- or downwards pointing arrowhead on top of this column, respectively.

On top of the table, you can **navigate** between the individual pages of the table; it is also shown on which page of the table you are, and in the right top corner, the page size in terms of number of entries (rows) is shown and can be adjusted.

In the Tree Area under the tab Analyses, you find all **Methods** (red box) you may apply using the geneXplain platform.



Upon double-clicking on a method name in the Tree Area the input mask for this method will be opened in the work space.

The **input form** of one method looks like this:

Start page		
Experiment	[?]	(select element)
Species	[?]	Human (Homo sapiens)
Annotation source	[?]	databases/Ensembl/Data/gene
Annotation columns	[?]	(no selection)
Cutput table	[?]	(select element)

Show expert options >> Run

For starting a method just drag and drop you input file, verify your settings and press the **Run** button.

As a **result** of this analysis, a new table will be created and automatically stored in your data folder.

## What is a workflow?

**Workflows** on the geneXplain platform are linked chains of methods to perform a bioinformatic analysis with one click.



The picture shows a small workflow for the gene set enrichment analysis (GSEA) using four different ontologies: Gene Ontology (GO) Biological Process, GO Cellular Compartment, GO Molecular Function, and Reactome's functional assignments (see right side).

The larger light **blue boxes** are analysis functions (program modules, "Bricks"). **Green boxes** stand for input files, especially user-defined inputs. **Yellow boxes** represent automatic delivery or output files.

The platform provides workflows for different topics e.g.

- × Analyze ChIP-seq/RNA-seq data
- × Analyze Microarray data
- × Proteomics/Epigenomics
- × Discover functional enrichment (Gene Ontology)
- X Analyze networks
- imes Identification of regulatory regions
- 🔀 Finding drug targets
- 🔀 Analyze genomic variants
- × Expression and binding sites of miRNAs
- X Working with NGS data

If you have valid licenses for e.g. **TRANSFAC**<sup>®</sup> or **TRANSPATH**<sup>®</sup> databases the number of applications is higher.

## Where are the workflows on the platform?

 In the **Tree Area** under the tab Analyses, you find all workflows you may apply using the geneXplain platform. Upon double-clicking on a workflow name in the Tree Area (A) the method will be opened in the work space (B). Additional information is shown in the Info box (C). A schematic overview of the workflow is presented in the operation field (D).

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032	Start page 🔊 Compute differentially ex X
Databases Data Analyses Users	Compute differentially expressed genes (Affymetrix probes)
⊿ Janalyses	- C Experiment normalized
	- Control normalized
Methods	Human (Homo saniens)
A Morkflows	Results folder
🖌 🐖 Common	
- A BAM to Counts	Run workflow Edit workflow
- As Compute differentially expressed genes (Affymetrix probes)	
- Pb Compute differentially expressed genes (Agilent probes)	
<ul> <li>- Ph Compute differentially expressed genes (Illumina probes)</li> </ul>	
<ul> <li>- Pb Compute differentially expressed genes using Hypergeometric test (Affymetrix probes)</li> </ul>	
<ul> <li> <sup>a</sup>b Compute differentially expressed genes using Hypergeometric test (Agilent probes)     </li> </ul>	
- 2% Compute dimensitive expressed genes using hypergeometric test (numina proces)	
- Ph Find common effectors in networks (GeneWays)	
<ul> <li>         -          -         </li></ul>	
<ul> <li>- Phi Find master regulators in networks (GeneWays)</li> </ul>	
- A Gene set enrichment analysis (Affymetrix probes)	
- 2% Gene set enrichment analysis (Agrent proces)	
- A Gene set enrichment analysis (Ilumina probes)	
- Ph Gene set enrichment analysis - select a classification (Gene table)	
<ul> <li>- Ph Mapping to ontologies (Gene table)</li> </ul>	
<ul> <li>- A Mapping to ontologies for multiple gene sets</li> </ul>	
Comparison of RNA-seq with Cuttinks for multiple BAM files	
ROTEOME	
TRANSFAC	
E RANSPATH	
Search Info Default	Overview My description Graph search Script Clipboard Tasks
ID: Compute differentially expressed genes (Affymetrix probes)	
Title: Compute differentially expressed genes (Affymetrix probes)	
Size: 129	
Complete name: analyses/Workflows/Common/Compute differentially expressed genes (Affymetrix probes)	Experiment normalized
Description:	
This workflow is designed to identify up-regulated, down-regulated and non-changed genes for experimental data	
with three and more data points for each experiment and control.	Results blder)
As input, normalized data with Affvmetrix probeset IDs can be submitted. Such normalized files are the output of	
the "Normalize data" procedure.	
1	Control correstration

 The new platform **Start page** arranges workflows according to major areas of interest. Select one button with your favorite analysis topic. You can expand the list of available workflows by clicking. The workflows from the start page are marked as blue links.



Open the workflow input form from the Tree Area or the Start page (see e.g. red box) by clicking.



For starting a workflow just drag and drop your input file(s), verify your settings and press the  ${f Run}$  button.

lentify enriched motifs in promoters (TRANSFAC(R))	
🗅 Input Yes gene set	(select element)
🗅 Input No gene set	📑 sets/Data/Housekeeping genes (Human) 300
🗅 Profile	ଲ013.4/Data/profiles/vertebrate_human_p0.001
Species	Human (Homo sapiens)
Filter by TFBS enrichment fold	1.0
Start promoter	-1000
D End promoter	100
Result folder	(select element)

In the **Tree Area** under the tab Data, you find all stored output examples from different input data sets. Examples include data from different microarray platforms (Affymetrix, Agilent, Illumina), ChIP-seq-data, FASTQ, BAM and BED files, SNP data or sequence collections.



Each example comprises several analyses results, sorted in different folders. You are welcome to open the output files and folders.



Upon one mouse click on the folder Data, as in the picture below, you can apply the button from the top control menu (red box) to create a new folder within the selected one.

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Databases Data Analy	vses Users
🔺 🚛 data	
Examples	
🛛 🚛 Projects	
🔺 📷 Olga Kel projec	t
👂 🌆 Data	
👂 🤛 Journal	Create folder X
👂 🌆 tmp	
🕨 🚛 Public	Folder name:
	Cancel Ok

In this way, you can define the **hierarchical organization** of your folders and subfolders within your project, for example as shown below. Every time you run a workflow, you need to specify a location of the results folder, and you can specify any particular location within your project area.

Databases	Data	Analyses	Users	
🔺 🌆 data				
🕴 🌆 Exa	amples			
🖉 🤞 🚛 Pro	jects			
<b>™</b>	Olga Ke	l project		
4	📗 Data	8		
	k 🔎	GSE9187 an	alysis	
	4 🔊	Research1		
	Þ	Pathway	analysis	
	⊳	Promoter	analysis	
	- k 🔊	Research2		

Your first password for the geneXplain platform is generated automatically and sent to you by email. Once you enter the platform you can change your password and also have an option to edit your personal information.

To get information about your account, select the **Account info** button (red box) on the top menu control panel.

Account info	
Databases Data Analyses Users	Start page Account info X
▲ 🚛 data	
Examples	Username: jeannette.koschmann@genexplain.cor
Projects	Password: Change
Public	Expiration:
I	
	Courtesy: Dr.
	First name: Jeannette
	Last name: Koschmann
	Work phone:
	Home phone:
	Cell phone:
	Country: Germany
	Edit account info

To check your totally available work **space**, click on your project name, so that it is highlighted in blue. In the Info Box you can see information about this project including disk quota (red box). This is the space available for you. If you plan to upload large files, please make sure that you have enough work space available.



# Let's start

- 1. <u>Log in to the platform.</u>
- 2. Import your own datasets.
- 3. Start your workflows.

For any further questions/suggestions /problems please contact:

info@genexplain.com

jeannette.koschmann@genexplain.com (customer support)

Useful links:

www.genexplain.com

http://genexplain.com/genexplain-platform/

http://genexplain.com/wp-content/uploads/2016/09/User-Guide-4.0.pdf

