

TRANSFAC Plugin

USER MANUAL

User manual for TRANSFAC TFBS Plugin 1.3

Windows, Mac OS X and Linux

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This software is for research purposes only.

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Chapter 1

Introduction to the TRANSFAC TFBS Plugin

This manual is for the commercial (non-free) version of the TRANSFAC® plugin.¹ The differences from the free version of the plugin are that it bundles all data necessary and should work out of the box without any additional configuration.

The TRANSFAC plugin can be used to search for putative transcription factor binding sites in DNA sequences. The binding site predictions are done by the Match™ tool, which uses the positional weight matrix library from TRANSFAC to analyze your sequences. For each of these matrices, Match™ contains optimized parameters, for minimization of error rates (false negative rate and false positive rate). As a special feature, Match™ provides the option to use profiles that are specific subsets of matrices with optimized cut-offs. These profiles allow users to adapt Match™ to their specific interests. For example, promoters of a certain tissue can be searched with tissue-specific profiles.

In order to be able to use this TRANSFAC plugin you need to install the TRANSFAC plugin into your Workbench.

We cover the above topics first in this manual, and following that, provide information on carrying out searches of TRANSFAC via the CLC Workbench.

¹TRANSFAC® is a registered trademark of BIOBASE

Chapter 2

Installation of the TRANSFAC TFBS Plugin

The TRANSFAC TFBS Plugin is installed using the plugin manager. In order to install plugins, the Workbench must be run with administrator privileges.

Help in the Menu Bar | Plugins... ()

or **Plugins () in the Toolbar**

The plugin manager has three tabs at the top:

- **Manage Plugins.** This is an overview of plugins that are installed.
- **Download Plugins.** This is an overview of available plugins on CLC bio's server.

Install the plugin by clicking the **Download Plugins** button at the top of the dialog. This will open a dialog where you can browse for the plugin. The plugin is called "TRANSFAC TFBS".

When you close the dialog, you will be asked whether you wish to restart the CLC Workbench. The plugin will not be ready for use before you have restarted.

Note that you may wish to click on the button labeled "no" and restart the Workbench yourself. If you choose "yes" at this point, the Workbench restarts using the administrator privileges you were using so that you could install the plugin.

Chapter 3

Searching for transcription factor binding sites

Once the plugin is installed you can start searching for transcription factor binding sites. Depending on the workbench product you have installed:

Toolbox | Epigenomics Analyses (📁) | Transfac TFBS (🔺)

1

This opens the wizard, the first step shown in figure 3.1 and explained below.

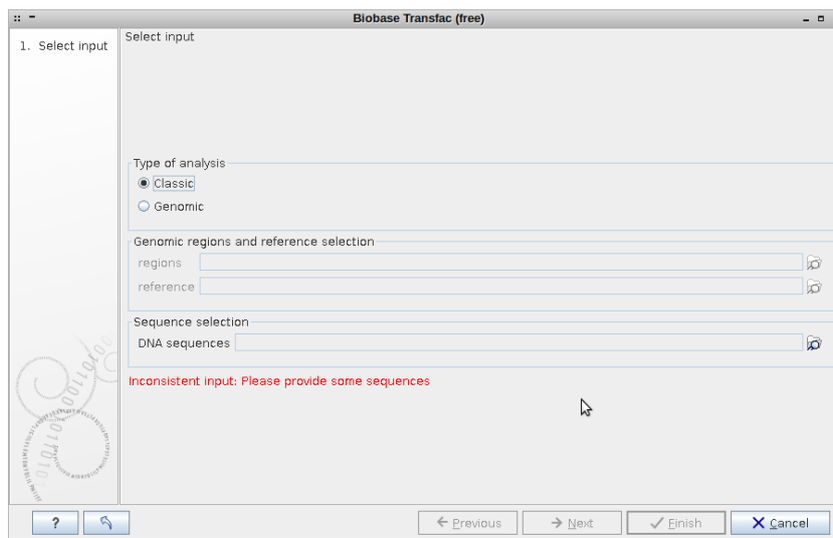


Figure 3.1: Select type of analysis and input data to search for transcription factor binding sites.

The first wizard step allows you to specify the input data.

Type of analysis First you need to select between the two modes of the analysis, each explained below.

- **Classic** This is the legacy mode of analysis where you have raw sequences without genomic information and wish to search for transcription factor binding sites in these

¹In the CLC Main Workbench, the Transfac TFBS tool is found here:

Toolbox in the Menu Bar | Nucleotide Sequence Analyses (📁) | Transfac TFBS (🔺)

sequences.

- **Genomic** Use this mode if genomic information is available in the form of genomic regions for which you wish to search for transcription factor binding sites².

Genomic regions and reference selection: This section will be enabled in the **Genomic** mode of analysis. Data is selected by clicking the corresponding folder icon (📁) in the right-hand side of the wizard.

- **regions:** The regions of interest as a track of annotations.
- **reference:** The reference genome as a track of symbols.

Sequence selection: This section will be enabled in the **Classic** mode of analysis. Data is selected by clicking the corresponding folder icon (📁) in the right-hand side of the wizard.

- **DNA sequences:** The raw DNA sequences of interest.

Clicking **Next** now opens the wizard step shown in figure 3.2.

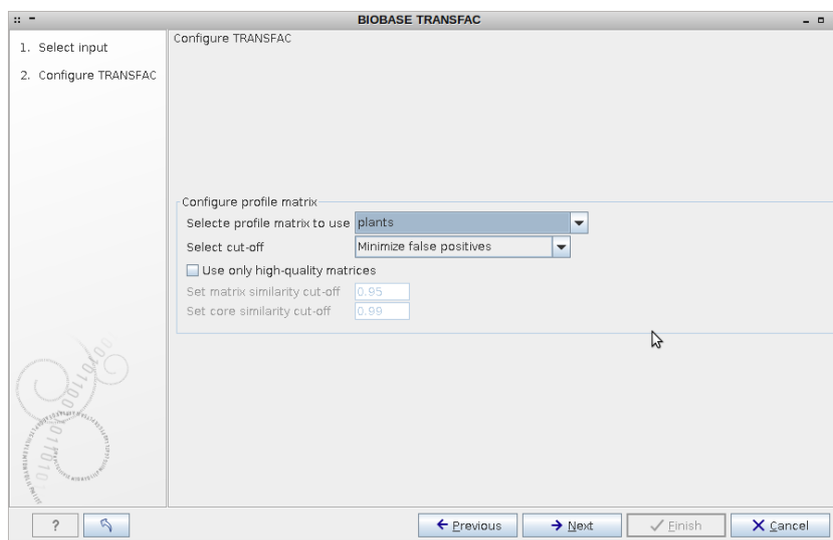


Figure 3.2: Selecting profiles and settings.

This wizard window allows you to configure the profile matrix parameters

Select profile matrix to use: From the dropdown menu, you specify the profile (set of position weight matrices) for a taxonomic group, tissue, etc. with which you want to scan your sequences/regions for putative transcription factor binding sites.

Select cut-off: This menu lets you configure the cut-off to use for the selected set of matrices, the options are:

- **Minimize false positives (minFP).** The minFP cut-off can be used to reduce the number of false positives. The false positive rate is estimated by applying the MatchTM algorithm to upstream sequences. The minFP cut-off is defined as the score that gives one percent of hits in the used sequences relative to the number of hits received at the minFN cut-off.

²For example, if you have done peak detection from a ChIP-Seq analysis, the regions where peaks are found then are the genomic regions you wish to investigate further.

- **Minimize false negatives (minFN).** The minFN cut-off can be used to reduce the number of false negatives. The false negative rate is measured, as far as available, on known genomic binding sites for the transcription factors. In case not sufficient (less than 10) genomic binding sites are available, SELEX sites or sets of oligonucleotides based on the nucleotide distribution in the weight matrix are used for estimating the minFN cut-off. The minFN cut-off is defined as that score at which at least 90% of the positive test set are recognized, i.e. it equals a false negative rate of 10%.
- **Minimize the sum of both error rates (minSUM).** The minSUM cut-off can be used to minimize the sum of both error rates. The sum of corresponding percentages for false positives and false negatives is computed for every cut-off ranging from minFN to minFP, whereby the false positive rate at minFN (10% false negative rate) is defined as 100%. The score at which this sum is minimal is used for the minSUM cut-off.
- **Set Matrix similarity cut-off.** Here it is possible to define similarity score to use in the search (see [Kel et al., 2003] for details). The range is from 0.0 to 1.0 where a score of 1.0 is given to the exact match and 0.0 will match anything.
- **Use cutoffs from selected profile.** The cutoffs that were defined in the profile will be used.

Use only high quality matrices: When enabled this will exclude highly abundant matrices which produce at minSUM (see below) more than 10 hits (false positives) per 1000 nucleotides³.

³"high quality matrices" are defined as matrices producing less than 10 hits (FP) per 1000 nucleotides (in sequences, 10.000 to 5.000 nucleotides upstream of the transcription start sites) at minSUM. About 5% of the current matrices producing higher FP rate, can be excluded as "highly abundant" / "low quality"; these 5% of matrices produce about 50% of all FP hits.

Chapter 4

Output of TRANSFAC TFBS Plugin

Clicking **Next** allows you to specify the output options as shown in figure 4.1.

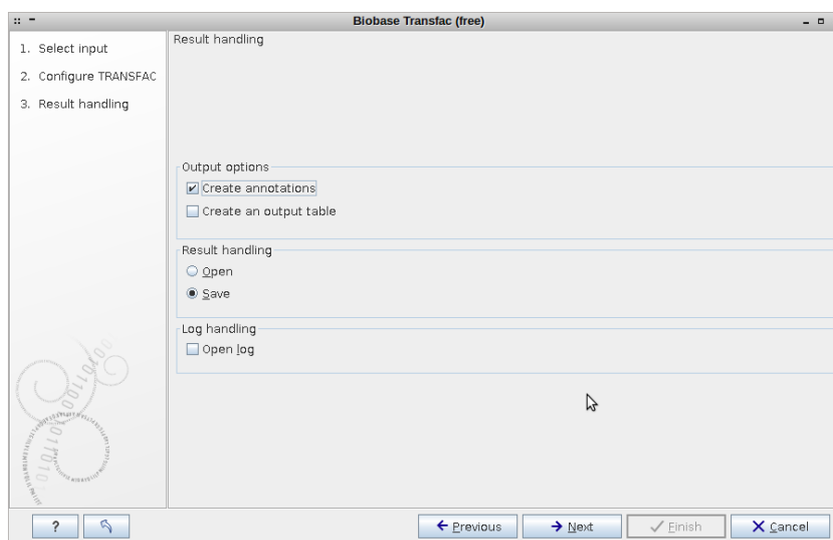


Figure 4.1: Selecting profiles and settings.

Based on the settings in figure 3.2, the Workbench uses the Match™ program from BIOBASE to search for transcription factor binding sites. The binding sites can be reported as annotations that are either added to the input sequences or reported as a track of annotations (depending on the analysis type, Classic or Genomic, see Chapter 3) and/or as an output table as shown in figure 4.4.

4.1 Output table

The table includes the following information:

- **Matrix ID.** Identifier for the matrix with which the putative binding site was found.
- **Factor name.** Name of the binding factor, represented by the matrix. If a group of factors is assigned to a matrix, only representative factor(s) are given. For a complete list of linked binding factors, please see the Matrix Report in TRANSFAC.

Matrix ID	Factor name	Region	Strand	Match sequence	Core similarity	Matrix similarity	Matrix report
V\$GRE_C	GR	complement(18..33)	minus	cAGGACaaagcagcct	0.978	0.720	M00205
V\$RFX1_01	RFX1	34..50	plus	ttggagcctGGAAcctt	0.982	0.930	M00280
V\$MUSCLEINI_B	Muscle initiator	93..113	plus	gtggagcagCACCCagcgcct	1.000	0.873	M00321
V\$DBP_06	DBP	26..32	plus	AGCGAGcc	0.969	0.925	M00624
V\$DBP_06	DBP	67..73	plus	AGCAGct	0.965	0.899	M00624
V\$DBP_06	DBP	70..76	plus	AGCTctc	0.899	0.878	M00624
V\$DBP_06	DBP	100..106	plus	AGCAGcct	0.965	0.942	M00624
V\$FAX_06	Pax	42..52	plus	CTGGAGatttaa	1.000	0.803	M00808
V\$FAX_06	Pax	complement(51..61)	minus	aagatGTCCAG	1.000	0.705	M00808
V\$FAX_06	Pax	66..76	plus	CAGAGctctc	0.675	0.672	M00808
V\$FAX_06	Pax	complement(79..89)	minus	ctcagggCCAG	0.900	0.711	M00808
V\$FAX_06	Pax	94..104	plus	ctGGAGcagcagc	1.000	0.724	M00808
V\$FAX_06	Pax	complement(110..120)	minus	gcctctTCCAG	1.000	0.667	M00808
V\$FAX_06	Pax	complement(145..155)	minus	ttaactTTAG	0.818	0.754	M00808
V\$FAX_06	Pax	157..167	plus	CCGGGaaetcc	0.735	0.764	M00808
V\$FAX_06	Pax	complement(159..169)	minus	gggaatCTCTG	0.575	0.671	M00808
V\$FAX_06	Pax	complement(170..180)	minus	tgaagtCCTAG	0.660	0.666	M00808
V\$DR4_02	LXR, FXR, CAR, COUP, RAR	complement(118..134)	minus	caGGTCTctctgggtcc	0.754	0.708	M00965
V\$CHC1_01	Churchill	158..163	plus	CGGGaa	0.999	0.995	M00986
V\$HELOSA_02	Helios A	16..26	plus	ccqAGGACaaa	0.916	0.915	M01004
V\$HELOSA_02	Helios A	complement(112..122)	minus	ctcCTCCAggt	0.877	0.790	M01004
V\$HELOSA_02	Helios A	complement(121..131)	minus	gtcCTCTggg	0.812	0.811	M01004
V\$AP2ALPHA_03	AP-2alphaA	27..41	plus	gcAGCTTggagcc	0.990	0.771	M01047
V\$AP2ALPHA_03	AP-2alphaA	complement(27..41)	minus	gcagccttTGGAGcc	0.608	0.771	M01047
V\$AP2ALPHA_03	AP-2alphaA	68..82	plus	gcAGCTctcggtcca	0.609	0.721	M01047
V\$AP2ALPHA_03	AP-2alphaA	complement(68..82)	minus	gcagctctCGGCTca	0.891	0.721	M01047
V\$AP2ALPHA_03	AP-2alphaA	74..89	plus	ctCGCTcaagccca	0.719	0.812	M01047
V\$AP2ALPHA_03	AP-2alphaA	complement(74..89)	minus	ctcggctAGGCCca	0.954	0.812	M01047
V\$AP2ALPHA_03	AP-2alphaA	80..84	plus	tcAGGCCcaagcagc	0.609	0.765	M01047
V\$AP2ALPHA_03	AP-2alphaA	complement(80..84)	minus	tcagcccAGGCAgc	0.966	0.765	M01047
V\$CTCF_01	CTCF	complement(106..125)	minus	cagagCTCTctccaggtcct	0.917	0.899	M01240
V\$BEN_01	BEN	complement(13..20)	minus	gtcCGAG	0.889	0.884	M01240
V\$BEN_01	BEN	108..113	plus	CAGCGctc	1.000	0.914	M01240
V\$MAN06_02	Manng	14..33	plus	tccagagCAAGcagcct	1.000	0.927	M01317
V\$SIK1_01	SiK-1	complement(47..63)	minus	acttaGATGtccagag	0.659	0.659	M01313
V\$SIK1_01	SiK-1	56..72	plus	gtccagaGATCagcagc	0.818	0.736	M01313
V\$SIK1_01	SiK-1	155..171	plus	gcccgggAATCTctgtg	0.659	0.714	M01313
V\$HDX1_01	HDX1	complement(44..59)	minus	ggagcCTTAagatgctc	0.967	0.681	M01317
V\$HDX1_01	Hdx	18..34	plus	caggacaAAGCAGcctt	0.769	0.723	M01333
V\$HDX1_01	Hdx	complement(47..63)	minus	acttaGATGtccagag	0.674	0.703	M01333
V\$HDX1_01	Hdx	56..72	plus	gtccagaGATCagcagc	0.852	0.837	M01333
V\$HDX1_01	Hdx	155..171	plus	gcccgggAATCTctgtg	0.832	0.755	M01333
V\$HDX1_01	Hdx	complement(165..181)	minus	ctctGTGAAGtctagc	0.630	0.686	M01333
V\$RHOK1_01	Rhok1	48..64	plus	cttaaGATGtccagaga	0.904	0.755	M01347
V\$RHOK1_01	Rhok1	68..104	plus	aggcaGCTGGAGcagc	0.756	0.701	M01347

Figure 4.2: Result table.

- **Region.** Position of the matrix match (putative binding site) within the analyzed sequence.
- **Strand.** Plus/minus. The strand on which the putative site was found depends on the orientation in which the matrix is given in TRANSFAC.
- **Match sequence.** Shows the matching sequence. Capital letters indicate the positions in the sequence that match with the core sequence of the matrix, while the lower case letters refer to positions that match to the remaining part of the matrix.
- **Core similarity.** The core similarity score for the matrix match. (The matrix core is defined as the five consecutive most conserved nucleotides within the matrix.)
- **Matrix similarity.** The matrix similarity score for the matrix match. The Match score can vary from 0 to 1, with 0 for the lowest similarity and 1 for the highest similarity of the match to the matrix. Only those matches are listed in the result, for which the core and matrix similarity are higher than the chosen cut-offs.
- **Matrix Report.** Matrix accession with a hyperlink to a detailed build-in matrix report that will open in a default browser on your system. See Section 4.2 for a detailed description of what the matrix report contains.

You can **Export** (📄) the table in csv or Excel format.

4.2 Matrix Reports

Matrix reports can be accessed by clicking the link in an entry in the Matrix Report column of the output table. It will load a html page in the default web browser of your system (see figure 4.3).

In the following we will describe the sections that exists in a Matrix Report.



Figure 4.3: A Matrix Report in a browser window.

Positional Weight Matrix Describes the characteristics of the matrix.

Matrix overview

- **Sequence logo:** A matrix logo, which displays the consensus sequence graphically, is shown. The reverse complement consensus sequence is displayed to the right.



Figure 4.4: Example matrix logo.

- **Nucleotide position frequency:** Displays the nucleotide frequency matrix. Read down to identify the binding site and across to identify the nucleotide frequency at that particular binding site position. The derived IUPAC consensus is provided in the last column. Click here for more information about the Matrix block. The nucleotide frequency matrix for the reverse complement is displayed to the right.
- **Transcription factors:** Lists the transcription factors from which binding sites were used for matrix construction, hyperlinked to corresponding Locus Reports. The count of binding sites contributed to the calculation of the positional weight matrix by each factor is displayed in the bar graph.
- **Aligned Binding Sites:** Lists the transcription factor binding sites used to generate the matrix, with links to the corresponding Site Reports. Includes the part of the sequence within the matrix window, the start of the matrix window (and the displayed sequence) relative to the complete sequence as given in the Site Report. The aligned binding sites are not given for all matrices. (This depends on the availability of the sites, e.g. not for all SELEX matrices the sites have been published. Also, for ChIP-based matrices no site alignments are given.)

The following information is provided for each site, when available: the transcription factor hyperlinked to the corresponding Locus Report, the gene from which the binding site was derived hyperlinked to the corresponding Locus Report, a graphical summary of the

experimental evidence supporting the TF-DNA binding interaction hyperlinked to the detailed Site Report, the experimental source, and the references supporting the TF-DNA binding interaction.

The experimental evidence categories are organized as:

- CI - chromatin immunoprecipitation
- DM - DNA modification (methylation, etc)
- FA - functional analysis
- FO - footprinting
- GS - simple gel shift/gel retardation
- IP - immunoprecipitation
- SE - SELEX
- SS - supershift/competitive gel shift
- OT - other

All experimental methods that do not fall into one of the other categories will be assigned to the OT (other) category.

- **Matrix type:** Specifies whether a matrix is specific for a factor, or more generally representative of a family of related factors.
- **Matrix classification:** For matrices built from vertebrate transcription factors, specifies the class that the matrix has been assigned to based on a matrix clustering algorithm submitted for publication. As of the 2012.3 release there are 44 classes: AP2-EREBP, ARID, ATHOOK, BHLH, BSH, BZIP, CHCH, CU-FIST, DM, E2, E2F, ETS, FORKHEAD, GCM, GENINI, GRAINY, HISTONE, HMG, HOX, HSF, IRF, MADS, MYB, NAM, P53, REL, RFX, RUNT, SAND, SMAD, STAT, SWI4, TBP, TBX, TCP, TEA, WRKY, ZFC2H2, ZFC4-NR, ZFC6, ZFDOF, ZFGATA, ZFPD, ZFRING. Matrices that do not fall into one of these classes are classified as "unclassified".
- **Matrix category:** Describes which method was used to create the matrix. Possible methods include matrix compiled from individual genomic sites, SELEX (CASTing, SAAB, Target Detection Assay), and more.
- **Application details:** Provides additional information about the experimental source of the binding sites used for matrix construction, the experimental approach applied to obtain this set, etc.
- **Number of sequences used:** Provides information about the number of binding sites used for matrix construction.
- **Additional transcription factors linked to the matrix:** Lists those transcription factors which did not contribute binding sites to the construction of the matrix but which are linked to the matrix, usually through homology.
- **Profile Membership:** Displays information about the profiles that the matrix is a part of
- **Profiles which include this matrix:** When relevant, lists whether the matrix of the page is part of the vertebrate non-redundant (VNR) or other profile for use in Match analysis.
- **Other vertebrate non-redundant (VNR) matrices that this matrix represents:** When relevant, lists the group of matrices that the matrix of the page represents in the vertebrate non-redundant profile (VNR) for use in Match analysis.

Related matrices Displays related matrices

- **This matrix is an older version of:** When relevant, lists any newer versions of this matrix, such as a version that was created after additional binding sites became available.
- **This matrix is a newer version of:** When relevant, lists any older versions of this matrix, such as a version that was created before additional binding sites became available.
- **Related family matrices:** Lists related matrices that are generally applicable to a family of transcription factors to which the factors used to construct this positional weight matrix belong.
- **Related factor-specific matrices:** Lists other matrices that have been constructed using the factors used to construct this positional weight matrix.

Identifiers Displays the identifiers associated with the matrix.

- **BKL Accessions:** Lists the identifier for the matrix of the page.
- **Similar matrix in:** Lists closely related matrices in public databases including JASPAR and UnIPROBE.

References Lists references from which whole sets of sequences or complete matrix was derived, including for the individual binding site sequences. Matrices constructed by us, typically have a link to a "TRANSFAC Report", which describes the way we used for site alignment and matrix construction.

This block gives the full citations, with titles, that correspond to the PubMed identifiers displayed. When a PubMed identifier is not available, the Medline identifier is displayed. When neither a PubMed nor a Medline identifier is available, a BIOBASE-specific number is assigned preceded by a "P". Nearly all of the reference numbers are hyperlinked to the PubMed database where the abstracts may be read. All references cited in the annotations and properties section are listed, and other references known to contain information about the protein may also be listed. The first five references are shown, click on the button labeled [more ...] to view all references.

Bibliography

[Kel et al., 2003] Kel, A. E., Ling, E. G., Reuter, I., Cheremushkin, E., Kel-Margoulis, O. V., and Wingender, E. (2003). MATCHM: a tool for searching transcription factor binding sites in DNA sequences. *Nucleic Acids Research*, 31(13):3576–3579.