The TRANSFAC® Database

Dating back as early as 1988, when the first data collection of transcription factors (TFs) and their binding sites was published [Wingender, Nucleic Acids Res. 16:1879-1902, 1988], TRANSFAC® is the oldest and most comprehensive database on eukaryotic TFs. Since 1998, it has been taken over for further maintenance by BIOBASE GmbH, and was merged later on with the resources TRANSCompel (a database on composite elements) and TRANSpro (a comprehensive collection of promoters of human and eight other genomes).

TRANSFAC® is now also available under the geneXplain platform, providing the most comprehensive collection of TF DNA-binding profiles available for the state-of-the-art sequence analysis implemented in the platform.

GeneXplain offers a one-stop shopping solution for the platform together with the TRANSFAC® database.

Applications

The most popular application of TRANSFAC contents is the prediction of potential transcription factor binding sites (TFBSs). Its contents can also be used to train own pattern finding algorithms, or to mine the wealth of information about transcription factors.

Further reading


About geneXplain

GeneXplain’s mission is to provide a comprehensive platform for bioinformatic, systems biological and cheminformatic tools. The raison d’être of this platform is to assist translational research in the life sciences, mainly in the context of personalized medicine and pharmacogenomics. We intend to make our expertise available to academic and commercial partners in collaborative research projects.

To achieve this, geneXplain offers:

- The geneXplain platform to provide an integrated and comprehensive workflow management of a large number of “bricks”, each providing a specific function in analyzing biological data
- In Silico Molecular Cloning (IMC) for handling large-scale genome data
- GenomeTraveler (GT) for handling next generation sequencing (NGS) data
- PASS and PharmaExpert for predicting biological activities of compounds qualitatively
- GUSAR for QSAR model building and quantitative activity prediction

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**Key features**

- 34,000+ transcription factor binding site reports containing details from the primary literature for more than 300 species, with a focus on human, mouse, rat, yeast, and plants
- 18,000+ transcription factor reports (including miRNAs), a subset of which provide GO functional assignments, disease associations and expression pattern assignments
- 2,300,000+ ChIP fragment reports that include the best scoring site prediction for the respective factor as well as downloadable sequences and gene lists
- 277,000+ promoter reports including ChIP-chip/Seq based histone modification locations, transcription start sites, and single nucleotide polymorphisms (SNPs)
- A pathway visualization tool for building custom regulatory networks out of experimentally demonstrated factor-DNA and factor-factor interactions

**The structure**

The core of TRANSFAC® comprises contents of two domains: Eukaryotic transcription factors (TFs) and TF binding sites (TFBSs).

Binding sites referring to the same TF are merged into **positional weight matrices (PWM)**. A PWM reflects the frequency with which each nucleotide is found in each position of the known and aligned TFBSs and, thus, the base preference in each position.

Transcription factors are classified based on the general properties of their DNA-binding domains. The most up-to-date and comprehensive TF classification available has been included in the geneXplain platform.

**Encyclopedic use**

TRANSFAC® is the most comprehensive encyclopedia about eukaryotic transcription factors. The structural and functional properties of each factor are documented by extensive manual annotation from the scientific literature by the BIOBASE team.

Individual TFBSs are documented including experimental details and a corresponding quality assessment.

**Overview of high-throughput data**

TRANSFAC® also documents HTP data on TF binding sites in eukaryotic genomes, usually from ChIP-chip or ChIP-seq experiments. These data are carefully selected and interpreted w.r.t. the binding regions and motifs found in the corresponding data sets.

**Availability**

The most up-to-date version of TRANSFAC® can be obtained either

- for downloading as textual flat files, to have the full content locally at your disposal;
- for online use through the geneXplain platform, making full use of the rich functionality of this unique toolbox, or
- for online use of the familiar look-and-feel provided by the BIOBASE server, or any combination of these options.

**Site & promoter analysis**

Using the rich library of 1600+ positional weight matrices of the TRANSFAC® database, DNA sequences can be scanned for potential transcription factor binding sites. One option for this is the proven tool Match™, which comes along with a standard TRANSFAC® license, or to use one of the new sophisticated tools that are additionally provided by the geneXplain platform.