

TRANSFAC® release 2017.3

The TRANSFAC[®] database on transcription factors, their genomic binding sites and DNA-binding motifs (PWMs), contains these new data features:

• Annotation of transcription factor binding sites based on sequence conservation

Known transcription factor binding sites located in human, mouse or rat genomes were extracted from TRANSFAC® and highly conserved sites were retained. Given high conservation as a prerequisite, binding sites were annotated for the two other species in respective genomic location if not more than one mismatch was observed in the sequence alignment with the primary species. This resulted in 1,565 new binding site entries.

• ChIP-Seq experiment browse pages

New browse page for 161 human DNase hypersensitivity ChIP-Seq experiments imported from ENCODE. The genomic intervals for each data set can be downloaded in .BED format.

The TFBS and DNase ChIP-Seq experiment browse pages can be accessed from the tools menu. The BED download buttons have also been added to the TFBS ChIP-Seq experiment browse page, providing easier access for the now 2,032 data sets.

113 new transcription factor binding site ChIP-Seq experiments released by the <u>ENCODE phase 3</u> project between February 2017 and May 2017. The data sets comprise 1,329,758 fragments bound by 98 distinct transcription factors, of which 66 factors were not yet covered by ChIP-Seq data. For 71 of the sets, an existing positional weight matrix for the respective transcription factor was used together with the MATCH tool to predict altogether 816,574 best binding sites inside the fragments. Predicted best binding sites as well as complete fragments are available in FASTA and BED format via the ChIP Experiment Reports, as are lists of genes in a distance range to the fragments as specified by the user.

• Reorganization of the in vivo transcription factor bound fragment section on a Locus Report

To improve clarity, only those fragments are listed that overlap with one of the promoter sequences of the entry. As new information fields, the table contains the relative position of the fragment to the transcription start site (TSS) of promoter, as well as the sequence of the predicted best binding site for the transcription factor inside the fragment.

• HOCOMOCO v10 matrix library integration

134 mononucleotide position weight matrices based on ChIP-Seq experiments have been incorporated from HOCOMOC0 v10 (<u>http://hocomoco.autosome.ru/</u>).

• Enhanced human SNP content

The new March 2017 dbSNP Build 150 data for human has been integrated and increases the number of SNPs mapped to human promoter sequences more than two-fold from 34,839,288 in the last release to 73,423,232.

• Ensembl version update

Genomic information for genes, promoters, and ChIP fragments for the species human, mouse, rat, macaque, and Arabidopsis is now based on Ensembl release 89.