TFBSPred: A functional transcription factor binding site prediction webtool for humans and mice

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Abstract. Transcription factors (TFs) play a major role in the regulation of gene expression. Discovering the TFs which bind to the regulatory regions of each gene has been long-term focus of research. Since the experimental verification of TF binding sites (TFBSs) is a complex process, webtools that perform predictions have been developed. However, the majority of the tools do not provide a user-friendly environment for data input and a number of these tools produce a large number of false-positive results. The present study introduces TFBSPred, a TFBS prediction webtool that utilises hidden Markov model-based TF flexible models (TFFM) for predicting binding sites while providing an automated and minimal input user interface. TFBSPred uses DNAse I hypersensitivity data from ENCODE to identify open chromatin regions and takes advantage of the conservation between Homo sapiens and Mus musculus, by using Ensembl Compara pairwise alignments, to increase the true positive rate of the prediction. The users input a gene name or genomic location of a human or mouse genome, select the cell types of interest and TFBSPred outputs the conserved open chromatin region of the selected

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Abbreviations: DH, DNase I hypersensitivity; GHMM, general hidden Markov model; GTF, general transcription factor; HMM, hidden Markov model; MAF, multiple alignment format; PSSM, position-specific scoring matrix; PWM, position weight matrix; TF, transcription factor; TFBS, transcription factor binding site; TFFM, transcription factor flexible model; TSS, transcription start site

Key words: transcription factor binding sites, webtool, prediction, DNase I hypersensitivity, conservation, TFFM

regulatory sequences and cell types as a pairwise alignment and displays the predicted TFBSs. The present study benchmarked TFBSPred and several similar functioning webtools using experimentally verified TFBSs. TFBSPred exhibited the optimal trade-off between sensitivity and specificity in the case of the well-studied IFNB1 enhanceosome, while outperforming the other web tools in subsequent use-cases. TFBSPred may thus prove to be a valuable tool for TFBS prediction and for the provision of hypotheses for experimental validation. TFBSPred also has the potential for further improvement with future updates of TFFM data. TFBSPred is freely available at: https://www.michalopoulos.net/tfbspred/.

Introduction

RNA polymerase II binds to the promoter of a gene and it assembles the transcription mechanism by gathering general transcription factors (GTFs), creating the pre-initiation complex to initiate transcription. Transcription is regulated by cis-regulatory elements, including the promoter. Distal elements can exert a positive effect on transcription (termed enhancers) or a negative effect (termed silencers) (1-3). Transcription factors (TFs) are proteins that bind to such regulatory regions of genes recognising multiple specific DNA sequence motifs, termed TF binding sites (TFBSs) (4). There are >1,600 human TFs catalogued (5), controlling processes of cell type specification, developmental patterning (6), as well as specific biological pathways (7). The binding affinity of a TF is dependent on its DNA-binding domain and the specific sequence of nucleotides which is targeted (8). Potential binding sites are predicted based on matches to a consensus sequence, often allowing for certain mismatches. The methods originally used for searching genome sequences to predict TFBSs were based on position weight matrices (PWMs), also known as position-specific scoring matrices (PSSMs) (9), derived from the multiple sequence alignment of experimentally verified DNA target motifs. A weight matrix is a two dimensional array of values that represent the score for finding each of the four nucleotides at each position in the

DNA sequence (10). The main databases for the collection of PWMs are TRANSFAC (11) and JASPAR (12). The majority of algorithms used for searching for PWMs in genomic sequences are MATCH (13) and FIMO (14). To simulate TFBS motif intricacies, more complex models have been proposed in recent years (15-18) and, in particular, hidden Markov models (HMMs) (19) have been applied successfully for TFBS prediction (20,21). To account further for TFBS length variability and interactions between nucleotides, HMM-based TF flexible models (TFFMs) (22) were developed, by consulting already existing JASPAR models. They can be graphically represented with a sequence logo, in a manner similar to PWMs (23).

Comparing the relative order of gene orthologs in the human and mouse genomes has revealed that a long-range sequence organisation has been preserved to a large extent from their last common ancestor (24). Approximately 80% of the common genes can be matched between the two organisms with the addition of a high rate of conservation of nucleotides (25). Genes that share close evolutionary associations are likely to possess similar functions and, likewise, functionally similar *cis*-regulatory elements have been shown to be conserved between species (1,26). Therefore, it can be considered that the majority of functional TFBS sequences are conserved between human and mouse.

DNase I is an endonuclease that cleaves DNA adjacent to pyrimidine nucleotides. In order for DNase I to cleave a DNA strand, it needs to be able to access it. Within cells, TFs displace histone octamers, unwinding the tightly packed chromatin structure. Through the technique of DNase I hypersensitivity (DH) assays (27) and its evolutions (28), DH sites, which denote the open and accessible areas that DNase I can operate on, are discovered. Analysing the whole genome accessibility landscape using DNase I, yields DH maps that denote parts of the genome that are probably transcriptionally active. From these data, it is possible to discover potential cell-type specific TFBSs, as the DH areas originate from open chromatin regions that contain regulatory elements of active genes (29-31).

Predicting TFBSs in regulatory sequences of a gene of interest requires some of the aforementioned computational tools. The present study introduces TFBSPred, a TFBS prediction webtool which eliminates false discoveries by using novel TFFM searches on cell type-specific open chromatin regions, which are conserved between *Homo sapiens* and *Mus musculus*.

Data and methods

Data collection. Using in-house PHP scripts, the present study downloaded and processed various *Homo sapiens* and *Mus musculus* genomic data, such as gene symbols and annotations from HGNC (32) and MGI (33), as well as gene stable IDs, transcript stable IDs and transcription start site (TSS) information from Ensembl (34) through BioMart (35). To determine the open chromatin regions of each biological replicate of each cell line and/or treatment, BroadPeak (36) DH data (31) for human hg19 and mouse mm9 genomes were downloaded from the UCSC Genome Browser database (37). The LiftOver tool (38) was used to update the DH data coordinates to those of the latest versions of the two genomes (hg38)

for human and mm10 for mouse), removing unmapped or duplicate regions. DH data for each replicate, cover, on average, 3.1 and 3.6% of the human and mouse genomes, respectively (Table SI).

Furthermore, ENCODE (39) common cell type details from UCSC Genome Browser were collected and parsed. To determine the conserved sequences between Homo sapiens and Mus musculus, Ensembl Compara (40) pairwise alignments in multiple alignment format (MAF) for the latest versions of the two genomes were downloaded, which were constructed using the LASTZ (41) alignment program. When parsing all MAF files, all pairwise alignments were converted into FASTA format and their genomic coordinates were extracted. The alignments cover, on average, 32.8 and 35.6% of the human and mouse genomes, respectively (Table SII). All aforementioned data were stored in a relational database using the MySQL management system (https://www.mysql. com/) on a Linux Ubuntu 64-bit, 16-core (Intel(R) Xeon(R) CPU E5-2650 v3), 64 GB memory virtual machine, which is provided by GRNET (https://okeanos-knossos.grnet.gr/).

The TF and TFFM data were procured from the latest version of the JASPAR database (http://jaspar.genereg.net/). From the TFFMs available, only the detailed trained ones that belonged to vertebrates were retained. A total of 462 TFFMs are included.

Web interface. The website is hosted on an Apache2 HTTP server. It was developed in HTML5, which was validated by HTML validator addon (https://www.gueury.com/mozilla/). Bootstrap styling library (https://getbootstrap.com/) was used for the website design. JavaScript was used to implement the selection checkboxes and autocompletion fields. All scripts were written in PHP scripting language.

To perform a TFBS search on TFBSPred, the users are guided through a number of online steps, using a web wizard. As input to TFBSPred, the users initially select an organism between human or mouse on the main page (Fig. 1A) and consequently, they type an autocompleted HGNC or MGI gene symbol, respectively, or an Ensembl Gene or Transcript stable ID, or an exact hg38 or mm10 genomic location in the form of Chromosome: Location(Strand), e.g., 4:102501329(+). In the case a gene symbol or Ensembl Gene ID is provided, the users are redirected to the TSS selection page of the gene of interest (Fig. 1B). The TSS selection page provides links to the TSSs of the input gene and to related Ensembl pages. If an Ensembl Transcript ID is used as input, a single link to a TSS will appear on the TSS selection page. By clicking on one of the links to a TSS, the users proceed to the cell type filtering page. The users can directly reach this page from the main page, if they use an exact genomic coordinate input, to perform a TFBS search on a region containing a specific genomic location, instead of a region of a gene promoter.

On the cell type filtering page (Fig. 2A), the users filter the cell types whose open chromatin region will be used for analysis, by checking at least one of the boxes of each of four categories: Lineage, Tissue, Sex and Karyotype. By submitting these selections, the users proceed to the cell line selection page (Fig. 2B), where only filtered cell lines are displayed, along with relevant details from Encode. The users check at least one of the boxes of the cell lines. Alternatively, if the users wish to

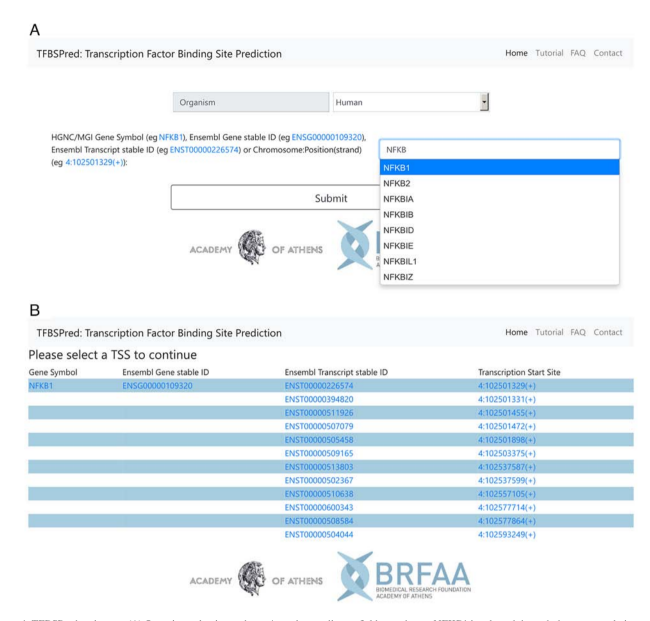


Figure 1. TFBSPred main page (A) Organism selection and gene/genetic coordinates fields are shown. NFKB1 is selected through the autocompletion results. (B) The TSS selection screen of the NFKB1 input gene. The first TSS in the table is selected. TFBS, transcription factor binding site; TSS, transcription start site.

identify all TFs which bind to the promoter region of their gene of interest, irrespective of the tissue context, they need to check all boxes of the cell type filtering and cell line selection pages. To conduct TFFM searches, a TFFM threshold must be specified at the bottom of the cell line selection page. The threshold value ranges from 0.60 to 1, and the higher the number, the stricter the TFBS search is. By submitting the selected cell lines and cut-off value, the TFBS prediction page appears, after a few minutes. A conserved open chromatin region whose extent depends on the cell lines selected, is depicted as a pairwise alignment between human and mouse, with the top sequence belonging to the organism that was initially selected (Fig. 3A). The pairwise alignment follows the 'pair' format (http://emboss.sourceforge.net/docs/themes/AlignFormats. html#pair) and can also be downloaded in FASTA format, to be used as input for downstream analyses. The TSS or specified genomic location is marked with an arrow highlighted red, indicating its orientation of transcription. The predicted TFBSs are displayed above their corresponding location in each pairwise alignment with consecutive arrows indicating their binding orientation. Detailed TFBS prediction results can be shown, located below (Fig. 3B). The predicted TFs are sorted alphabetically by their gene symbol, which links to its corresponding JASPAR entry. Each TF contains one or more discovered binding sites presented as a human and mouse pairwise alignment in 'pair' format, indicating the actual genomic coordinates of each binding site.

TFFM search. TFBSPred TFBS prediction is based on TFFM searches which are executed in the webtool backend, after the users have submitted their data. To this end, the TFFM framework (22), as well as its prerequisites, the general hidden Markov model (GHMM) (42) and Biopython (43) libraries, were installed.

To define the extent of the open chromatin region of the genome of the organism of interest, which includes the selected TSS or specific genomic location, the borders of the Please select all or at least one of each category

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Figure 2. TFBSPred cell type selection (A) The cell type filtering page. Categories corresponding to normal blood-type cells of all sexes are selected. (B) The filtered cell type selection page. Only CD4⁺ cell types are selected. TFBS, transcription factor binding site.

open chromatin region broad peaks of all selected cell lines, which contain the specified genomic coordinate are retrieved from the database and merged together using BEDTools (44), defining the borders of a genomic region which represents the union of all open chromatin regions. The borders of the conserved genomic region which contains the specified genomic coordinate are also retrieved. The conserved open chromatin region is the cross section of the merged open chromatin region and the conserved region, and its borders are calculated using BEDTools. From the pairwise alignment of the conserved open chromatin sequence, the corresponding sequence of the other organism is determined. Both the human and mouse genomic sequences are then searched upon with all TFFMs available, using an in-house python script which outputs the matches above the user designated cut-off value. TFBSPred parses all search results and only displays as pairwise alignments the predicted TFBSs, which are conserved between the human and mouse sequences.

Benchmarking TFBS prediction webtools. TFBS predictions on various human genes were performed using TFBSPred and other webtools, mostly using the default settings. Gene names were submitted to TFBSPred. FASTA-formatted gene regulatory sequences were submitted to AliBaba2.1 (45),

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mm10	135691505 G-CAGGAAGAGGAGGTTTCGCCACCCGAGCA	CCCGGCTGCGCGCTGACG	135691553						

Figure 3. TFBSPred NFKB1 results (A) A part of the conserved open chromatin region of NFKB1 with its predicted TFBSs, using data from the selected cell types. (B) A part of the detailed results of predicted TFs and their TFBSs on the conserved open chromatin region of NFKB1. TFBS, transcription factor binding site.

TFBIND (46), SITECON (47), PROMO (48), MATCH (13) and STAMP (49). Both human and their corresponding mouse sequences were submitted to ConSite (50), FOOTER (51) and rVista 2.0 (52). To execute rVista 2.0, the integrated zPicture (53) alignment program was used to align the two sequences. Gene names were used as input and both human and mouse gene promoter sequences were selected, along with JASPAR Core Matrices with all available TFs, in LASAGNA-Search 2.0 (54). Gene names were used as input, the exploration function was executed followed by the visualisation function using the discovered TFs, in ConTra v3 (55). Finally, the motif features discovered in the regulatory regions of genes in Ensembl regulatory build (56) were also identified.

Results

Interferon beta 1 (IFNB1) enhanceosome. To assess the predictive capabilities of TFBSPred and other TFBS prediction webtools, the IFNB1 enhanceosome (57) was used. The enhanceosome is an ~50 bp enhancer sequence of the IFNB1 gene and requires the coordinate activation and DNA binding of the TFs ATF2/Jun, IRF3 and IRF7 and NF- κ B (57,58), effectively dividing the enhanceosome into four positive regulatory domains (PRDI to PRDIV). The exact binding sites of each of these TFs have been experimentally confirmed (57,59). PRDM1 has also been shown to bind specifically to the PRDI domain of IFNB1 enhanceosome, with a negative effect on transcription (60), while additionally playing a role in recruiting co-repressor complexes required to silence gene expression (61). With the default 0.90 TFFM threshold, TFBSPred discovered

PRDM1, NF-KB (RELA, RELB and NFKB1), IRF1, SPIB and TEAD2 TFBSs in the enhanceosome region. ATF1 was also discovered among other TFs, when the threshold was decreased to 0.62 (Fig. 4); however, 26 false-positive TFBSs were also predicted (data not shown, as the default cut-off value is not 0.62). The ConTra v3 exploration function predicted only two TFs, NF-κB and PRDM1. AliBaba2.1 discovered five total TFBSs, including IRF8 and ATF2 sites but not NF-κB. LASAGNA-Search 2.0 discovered NF-KB and STAT1, MATCH discovered NF-kB, IRF1 and IKZF1, and the single sequence option of ConSite discovered NF-KB, TEAD1, IRF1 and IRF2. The pairwise alignment option of ConSite only discovered the IRF2 site. PROMO, TFBIND, rVista2.0 and SITECON discovered a large amount of TFBSs. PROMO found TFBSs for all verified TFs; however, it displayed 23 false-positive TFBSs. TFBIND discovered IRF-family TFs, NF-κB and JUN, with 10 false-positive sites. rVista2.0 discovered NF-kB and IRF-family TFs (including IRF7), also having 10 false-positives, and SITECON only found IRF-family TFs, while having 29 false-positive TFBSs. Although STAMP predicted less TFBSs than those of the four previous tools, it only found NF-κB from the verified TFs, presenting 7 false positive sites. Finally, FOOTER did not succeed in predicting any TFBSs. Although the majority of the experimentally verified TFBSs (including NFKB1, NFKB2, IRF3, IRF7, JUN, ATF7 and PRDM1) were computationally predicted by the Ensembl Regulatory Build of IFNB1 (ENSR00001147395), 53 false-positive TFBSs were also found (https://www.michalopoulos.net/tfbspred/erb.html). Nevertheless, none of the putative TFBSs was marked as 'experimentally verified' in the Ensembl genome browser (all

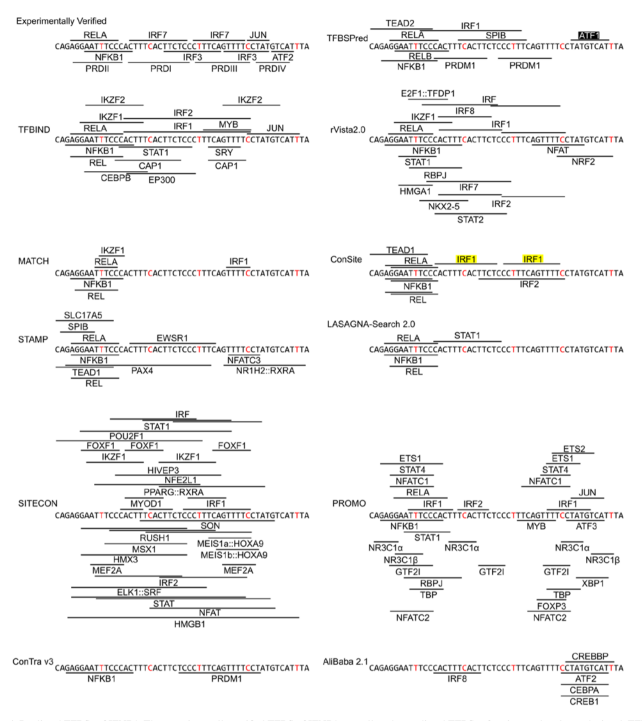


Figure 4. Predicted TFBSs of IFNB1. The experimentally verified TFBS of IFNB1, as well as the predicted TFBSs of various webtools are depicted. TFBSs are depicted as continuous lines. When a TFBS is depicted above the nucleotide sequence, the name of the TF corresponding to it is then located above it. Respectively, if a TFBS is depicted below the sequence, the name of the TF is then located below it. The TF name is displayed once for multiple adjacent converging TFBS lines for the same TF. ATF1 (highlighted in black) was discovered with a TFFM cut-off value of 0.62. In ConSite, IRF1 (highlighted in yellow) was also found using the pairwise alignment option. TFBS, transcription factor binding site; TF, transcription factor.

TFBSs are coloured in grey), indicating that none of the TFBS hits was supported by ChIP-Seq peak evidence.

FGA gene promoter. In its basal state, Fibrinogen alpha chain (FGA) is regulated by HNF1 (-59 to -47 nucleotides from the TSS) and CEBP (-142 to -134 nucleotides from the TSS) family proteins. In an acute state, STAT3 additionally binds to the IL-6RE region, upstream of FGA TSS (62). To examine the tissue-specificity of TFBSPred, two analyses on FGA

were conducted: In the first, only endodermal normal cell lines were used, while in the second analysis, only the HepG2 hepatoblastoma cell line was selected. In both cases, the FGA gene was used as input. In the first analysis, TFBSPred discovered CEBP-family factors (CEBPE, CEBPA, CEBPD), HNF1 (HNF1A, HNF1B) and VDR. In the second analysis, a larger conserved open chromatin region was displayed, due to cancer cells having an abnormal regulation pattern. Multiple TFs of the STAT (STAT3, STAT4, STAT5A, STAT5B) and

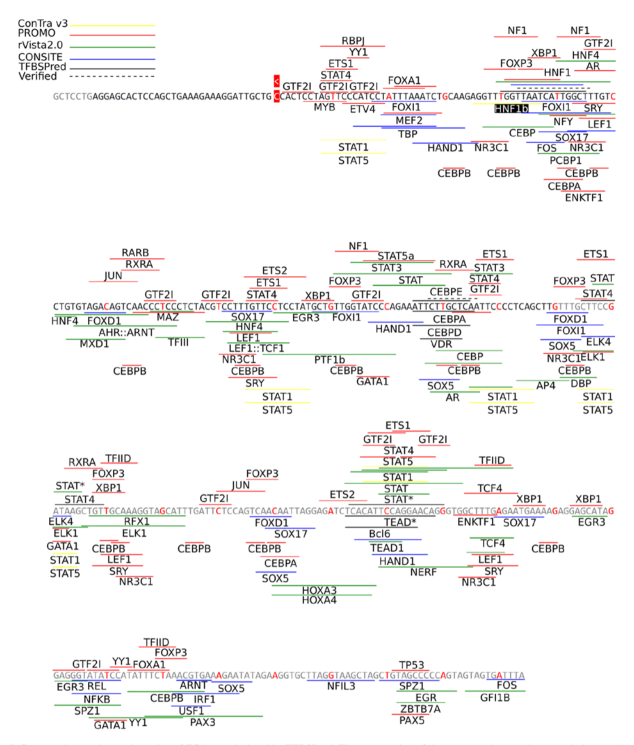


Figure 5. Conserved open chromatin region of FGA as calculated by TFBSPred. The grey section of the sequence denotes the expanded open chromatin region that appears when the HepG2 cell line is selected. The TSS and an arrowhead showing its orientation of transcription are highlighted in red. The experimentally confirmed binding sites are portrayed with dashed lines, while the TFBSs predicted by the selected webtools with coloured continuous lines. When a TFBS is depicted above the nucleotide sequence, the name of the TF corresponding to it is then located above it. Respectively, if a TFBS is depicted below the sequence, the name of the TF is displayed once for multiple adjacent converging TFBS lines for the same TF. STAT* includes STAT1, STAT3, STAT5A and STAT5B factors and TEAD* includes TEAD1, TEAD2, TEAD3 and TEAD4 factors. HNF1b (highlighted in black) was predicted by ConTra v3 in other species, but not in *Homo sapiens*. TFBS, transcription factor binding site.

TEAD (TEAD1, TEAD2, TEAD4) families, as well as Bcl6, were additionally discovered further upstream of the TSS (Fig. 5).

Four other webtools that performed best for IFNB1 (single sequence ConSite, rVista2.0, PROMO and ConTra v3), were also used to predict TFBSs for FGA (Fig. 5). The

FASTA-formatted human sequence which corresponds to the conserved acute-state open chromatin area upstream to the FGA TSS, as calculated by TFBSPred (345 nucleotides length), was used as input. ConSite predicted only the HNF1 site with 35 false-positives. While rVista2.0 and PROMO all predicted the experimentally verified TFs (HNF1, CEBP and STAT3),

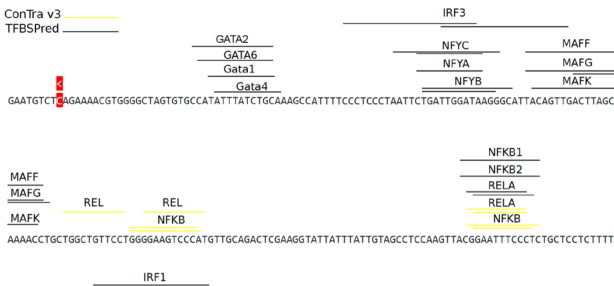




Figure 6. Conserved open chromatin region of CXCL10 as calculated by TFBSPred and the TFBSs predicted by TFBSPred and Contra V3 (coloured yellow). The TSS and an arrowhead showing its orientation of transcription are highlighted in red. Only the region upstream of the promoter where the TFBSs are predicted is shown. TFBS, transcription factor binding site.

they also found 30 and 80 false-positive TFBSs, respectively. ConTra v3 discovered STAT-family TFs and HNF1b, although the latter was not found on the *Homo sapiens* sequence, but rather in other organisms. TFBIND was also tested as a candidate webtool; however, it was not included since it predicted >400 binding sites (Table SIII).

C-X-C motif chemokine ligand 10 (CXCL10) promoter. CXCL10 is a chemokine that is secreted during the immune response. STAT1, NF-κB, AP1 and heat shock factors bind to an ~230 bp region upstream from the TSS (63). Additionally, IRF3 binds to CXCL10 promoter during hepatitis C infection (64). The CXCL10 gene was used as input to TFBSPred along with all the available cell-lines and default TFFM settings. TFBSPred revealed several factors binding upstream of the gene promoter, including members of the NF-κB (NFKB1, NFKB2 and RELA) family, all the factors forming the NFY complex (NFYA, NFYB and NFYC), as well as IRF1, IRF8, MAFG and STAT2. ConTra v3 exploration analysis discovered only NF-κB family-related TFs (Fig. 6).

Discussion

The most popular approach for TFBS prediction, PWM-based search, considers that the nucleotides of each position exhibit independent participation in the DNA-protein interactions. ConTra v3 (55), a widely used webtool, employs PWM-based technologies which use JASPAR and TransFac PWMs to search a given genomic region for potential TFBSs. Specifically, it uses FIMO for its exploration function and MATCH for its

visualisation function. The majority of other webtools, such as TFBIND, PROMO and AliBaba2, use their own TFBS prediction algorithms, all of which are based on PWM profiles. The Ensembl Regulatory Build is a collection of regulatory features (denoted as ENSR) across the whole genome. PWMs based on TF motifs which were discovered through SELEX, were matched upon those regulatory regions (34), using MOODS (65), a PWM matching program. However, in the case of the IFNB1 enhanceosome, the loose parameters used for the execution of MOODS in Ensembl Regulatory Build, resulted in low specificity, as >50 pseudo-positive TFBSs were predicted. TFFMs, as a complete HMM-based approach, were designed to address the confounding properties of nucleotide composition, interpositional sequence dependence and variable lengths observed in the recently emerging extensive experimental data. They have been shown to perform more effectively than the majority of PWM-based models (22), while also being publicly available. TFBSPred is the only web based TFBS prediction tool which, thus far, employs the HMM-based TFFM algorithm. In the case of IFNB1, TFBSPred predicted an IRF1 site which coincided with the experimentally verified IRF3 and IRF7 sites in PRDI (57), as well as PRDM1, which was also discovered in the same coordinates. In addition, it should be noted that TFBSPred cannot predict the IRF3 and IRF7 binding sites, as there is no TFFM model for these. The TFBSPred search is connected to the available TFFM profiles from JASPAR. As JASPAR is a continuously evolving project, TFBSPred will incorporate future TFFMs, as well as updates to already existing models. Finally, ATF2/c-Jun leucine zipper heterodimer, does not bind

tightly, as the c-Jun half-site is not fully complementary to c-Jun binding and DNA is bent (57). Moreover, the majority of TFFMs for dimer TFs are created for homodimers, rather than heterodimers. Thus, model-based prediction algorithms could not predict this heterodimeric factor. Thus, TFBSPred only predicted ATF1 when the cut-off was lowered, indicating that it is possible to predict difficult TFBS cases, when the TFFM threshold is sufficiently decreased, although this should be used with caution, as it may introduce false-positives.

To reduce the false discovery rate, ConTra v3 takes advantage of the evolutionary conservation of regulatory elements across various related species by expanding the PWM matching on a multiple genome sequence alignment and subsequently visualising the results of the TFBS predictions. However, it is up to the users themselves to evaluate the importance of the conservation of each predicted TFBS among the multitude of species presented. Other webtools, such as rVista2.0 and ConSite, require the user to input two homologous genomic sequences to search for conserved TFBSs by creating a pairwise alignment where conserved TFBSs are automatically determined and displayed. Similar to rVista2.0 and ConSite, TFBSPred searches for conserved TFBSs in a pairwise alignment. However, as opposed to those tools, TFBSPred uses high-quality pairwise alignments from Ensembl Compara, which cover ~1/3 of each genome, while identifying homologous regions between two genomic species is a complex task, which is not easily performed by the average wet lab experimentalists. Another difference is that TFBSPred only compares human and mouse sequences. Although the ability to select between any combination of species and set multiple parameters to search for TFBSs renders rVista2.0 and ConSite more versatile, it markedly increases usage complexity. FOOTER requires the user to input a human and a mouse or rat genomic sequence, and predicts conserved TFBSs between the two; however, its default parameters are not adequate for optimal results. TFBPred requires only one human or mouse genomic sequence as input, and automatically identifies its homologous region of the other species: Mus musculus and Homo sapiens are extensively studied model organisms which belong to the orders of Rodentia and Primates, respectively. These orders which belong to the mammalian superorder of Euarchontoglires, split 85-97 million years ago (66), sharing a high degree of genomic sequence conservation. Thus, predicted TFBSs which are conserved between human and mouse are likely true positives. Searching Compara-aligned human and mouse sequence pairs and automatically discarding TFBSs which are not highly conserved, offers the best possible TFBS results, without any further user participation.

The boundaries of regulatory regions (promoters, enhancers and insulators, as well as TF motifs in open chromatin regions) were defined in Ensembl regulatory build, through a variety of genome-wide experimental data from multiple epigenomic consortia. Predicted TFBSs which are verified by ChIP-Seq can be displayed in the Ensembl genome browser tracks with additional information indicating cell-line/tissue specificity. Although this is an extensive assortment of TFBSs, in the case of the IFNB1 enhanceosome, experimental verification has a low sensitivity, as no experimentally verified TFBSs were identified. As opposed to Ensembl regulatory build, ConTra v3 is unable to define on its own the extent of regulatory regions and has no cell-line/tissue specificity. ConTra v3 requires a user-specified region length to search for TFBSs, with the arbitrary default value being 500 bps. In addition, the majority of webtools perform TFBS prediction solely on a FASTA-formatted genomic sequence the user selects. On the other hand, TFBSPred automatically identifies the boundaries of cell-line/tissue specific regions of open-chromatin which cover around 3-4% of each genome, based on DH data of the selected cell lines, and then searches for conserved TFBSs. Consequently, by selecting certain tissues and cell types, TFBSPred can discover potential cell-type specific TFBSs, as DH areas indicate open chromatin regions that contain regulatory elements of active genes. To examine the basal state of FGA, non-cancer cell lines were used for a TFBSPred analysis: TFBSs for HNF1 and CEBP-family factors were predicted, in accordance with experimentally confirmed sites. To study the acute state of FGA, only Hep2G cancer cell line was used as input for a TFBSPred analysis: Not only was a larger conserved open chromatin region produced, but also STAT family factors were predicted in this expanded open chromatin sequence of the acute state, including experimentally confirmed STAT3. This displays the potential for TFBSPred for tissue-specific differential TFBS predictions, as an experimentally characterised TFBS (STAT3) is predicted in the acute state (Hep2G cancer cell line) and not in the basal state (non-cancer cell lines). R-based BinDNase (67), also incorporates DH data to increase PWM accuracy and has demonstrated the validity of this approach.

ConTra v3 has a long execution period, particularly if a number of PWMs are selected in a visualization analysis. On the contrary, one of the main objectives of TFBSPred was the provision of a rapid and user-friendly interface specifically catering for wet lab biologists. Throughout the website, user interaction is minimal, as the majority of the input fields and selections are provided through TFBSPred database. The TFBSPred web wizard begins with the selection of the TSS of a gene or a chromosomal location in either human or mouse, followed by the filtering and selection of cell types. Finally, the maximum execution time does not exceed a couple of minutes.

To summarise, TFBSPred is a comprehensive and easy-to-use TFBS prediction webtool, based on open chromatin patterns, genomic conservation among human and mouse and HMM-based searches. The present study demonstrated that the predictions of TFBSPred were in accordance with both universal and tissue-specific experimentally verified TFBSs and that in a number of cases, it outperformed existing PWM-based webtools of similar function. TFBSPred is expected to further improve as the TFFM quality and quantity increases. TFBSPred may thus be a useful addition to the experimental biologist community as it provides a working hypothesis that can be experimentally verified.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

IM conceived the study. PGB and DAS were also involved in the conception of the study. DH data were downloaded and processed by CN. Multiple alignment between *Homo sapiens* and *Mus musculus* data were downloaded and processed by KS. VLZ downloaded the TFFM framework and TFFM profiles, and developed the database and Graphical User Interface (GUI). IK assisted in the website front end development. GAL performed the webtool benchmarking. VLZ and IM confirm the authenticity of all raw data. The manuscript was written by VLZ and IM. All authors contributed to the revision of the work and, have read and approved the final manuscript.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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