# **RESEARCH**



# The evaluation of transcription factor binding site prediction tools in human and Arabidopsis genomes



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# **Abstract**

**Background:** The precise prediction of transcription factor binding sites (TFBSs) is pivotal for unraveling the gene regulatory networks underlying biological processes. While numerous tools have emerged for in silico TFBS prediction in recent years, the evolving landscape of computational biology necessitates thorough assessments of tool performance to ensure accuracy and reliability. Only a limited number of studies have been conducted to evaluate the performance of TFBS prediction tools comprehensively. Thus, the present study focused on assessing twelve widely used TFBS prediction tools and four de novo motif discovery tools using a benchmark dataset comprising real, generic, Markov, and negative sequences. TFBSs of *Arabidopsis thaliana* and *Homo sapiens* genomes downloaded from the JASPAR database were implanted in these sequences and the performance of tools was evaluated using several statistical parameters at diferent overlap percentages between the lengths of known and predicted binding sites.

**Results:** Overall, the Multiple Cluster Alignment and Search Tool (MCAST) emerged as the best TFBS prediction tool, followed by Find Individual Motif Occurrences (FIMO) and MOtif Occurrence Detection Suite (MOODS). In addition, MotEvo and Dinucleotide Weight Tensor Toolbox (DWT-toolbox) demonstrated the highest sensitivity in identifying TFBSs at 90% and 80% overlap. Further, MCAST and DWT-toolbox managed to demonstrate the highest sensitivity across all three data types real, generic, and Markov. Among the de novo motif discovery tools, the Multiple Em for Motif Elicitation (MEME) emerged as the best performer. An analysis of the promoter regions of genes involved in the anthocyanin biosynthesis pathway in plants and the pentose phosphate pathway in humans, using the three best-performing tools, revealed considerable variation among the top 20 motifs identifed by these tools.

**Conclusion:** The fndings of this study lay a robust groundwork for selecting optimal TFBS prediction tools for future research. Given the variability observed in tool performance, employing multiple tools for identifying TFBSs in a set of sequences is highly recommended. In addition, further studies are recommended to develop an integrated toolbox that incorporates TFBS prediction or motif discovery tools, aiming to streamline result precision and accuracy.

**Keywords:** Transcription factor binding sites, Bioinformatics tools, Performance evaluation



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#### **Background**

The regulation of gene expression is governed by various factors, including transcription factors (TFs). Each cell type or tissue, at a specifc developmental stage or in response to extracellular signals, exhibits a distinctive pattern of activated TFs. This repertoire of TFs is a key determinant for defning cellular identity and function. Studies have identifed 274,633 TF-encoding genes from 183 animal genomes [[1\]](#page-21-0), including >1600 TFs from the human genome [[1\]](#page-21-0) and 320,370 TFs from the genomes of 165 plant species [\[2](#page-21-1)]; more than 1500 TFs have been reported from the Arabidopsis genome [[3\]](#page-21-2).

TFs modulate gene expression by binding to specifc regulatory regions of the genes (upstream and basal promoter elements, enhancers, and silencers) [\[4,](#page-21-3) [5](#page-21-4)]. An important step in unraveling regulatory mechanisms of gene expression is the identifcation of these binding sites in DNA for TFs termed TF binding sites (TFBSs) [[6,](#page-21-5) [7](#page-21-6)]. Tese are typically short and often degenerate stretches, ranging from 5 to 20 base pairs (bp) in length [[8\]](#page-21-7). TFs possess specifc DNA binding domains that recognize and bind to particular sequences within TFBSs. Tese DNA binding domains allow TFs to directly interact with the DNA molecule, facilitating the regulation of gene expression. Alternatively, TFs can bind to DNA indirectly by interacting with another TF [\[9](#page-21-8)].

There are many in vitro and in vivo experimental approaches to determine the interaction of TFs and their potential binding sites on genomic sequences. These methods include the Electro-Mobility Shift Assay [\[10](#page-21-9)], DNase I footprinting/protection assay [\[11](#page-21-10)], Systematic Evolution of Ligands by EXponential enrichment (SELEX) [\[12](#page-21-11)], and Chromatin ImmunoPrecipitation (ChIP) Assay [\[13](#page-21-12)]. Advances in these experimental technologies, particularly ChIP followed by sequencing (ChIP-seq), have led to the generation of large-scale datasets of TFBSs across the genome. These comprehensive datasets provide a valuable resource for scientists to develop and refne computational approaches for predicting TFBSs. Experimental validation of TFBSs is a complex and resource-intensive process. Consequently, in recent years, in silico prediction of TFBSs has emerged as an efficient alternative to these time-consuming experimental methods  $[5, 14-17]$  $[5, 14-17]$  $[5, 14-17]$ .

Numerous algorithms have been developed to predict TFBSs. Many of the methods originally used for predicting TFBSs from sequences were based on position weight matrices (PWMs), also known as position-specific scoring matrices (PSSMs) [\[18\]](#page-21-15). These are the most widely used and well-established mathematical models for predicting TFBSs in DNA sequences [\[14,](#page-21-13) [19](#page-21-16)[–22](#page-22-0)]. PWMs are computed from the multiple sequence alignment of an experimentally validated set of TFBSs, quantitatively scoring binding motifs based on the frequency and positions of the nucleotides within the binding sites for the corresponding TF [\[19](#page-21-16), [23\]](#page-22-1). Several databases, such as JASPAR [[24\]](#page-22-2) and TRANS-FAC [[25\]](#page-22-3), maintain comprehensive collections of TF-binding profles as PWMs or position frequency matrices (PFMs).

JASPAR is an open-access resource that provides access to a manually curated, non-redundant set of TF-binding profiles in the form of PFMs. These PFMs are derived from experimentally defined TFBSs and can be converted into PWMs, which can then be used to scan DNA sequences to predict TFBSs. TRANSFAC, on the other hand, is a commercial database that includes PWMs derived from published collections of experimentally validated binding sites. In addition to these databases, several other resources provide access to collections of PWMs, such as HOmo sapiens COmprehensive MOdel Collection (HOCOMOCO) [[26](#page-22-4)], UniBind [[27](#page-22-5)], UniPROBE [[28](#page-22-6)], and Factorbook [\[29\]](#page-22-7). Several tools are available for predicting TFBSs by scanning for PWMs against DNA sequences. These include Find Individual Motif Occurrences (FIMO) [[30](#page-22-8)], MATCH [[31](#page-22-9)], MOtif Occurrence Detection Suite (MOODS) [[32](#page-22-10)], Ciiider [[33](#page-22-11)], Hypergeometric optimization of motif enrichment (HOMER) [[34](#page-22-12)], INSECT [[35](#page-22-13)], Matrix-scan [[36\]](#page-22-14), Morpheus [\[37\]](#page-22-15), PWMscan [[38\]](#page-22-16), TFBStools [[39](#page-22-17)], Contra V3 [[40](#page-22-18)], TFBS-Discovery Tool Hub (TDThub [[41\]](#page-22-19)), TFBIND [\[42\]](#page-22-20), PROMO [[43](#page-22-21)], Cluster-Buster [[44\]](#page-22-22), rVISTA [[45](#page-22-23)], LASAGNA-search [\[46\]](#page-22-24), and PscanChIP [[47](#page-22-25)]. Furthermore, several freely available de novo motif discovery tools have been developed to handle the large volumes of data generated from high-throughput technologies. These de novo discovery tools, such as AlignAce [\[48\]](#page-22-26), Weeder [[49](#page-22-27)], Improbizer [[50](#page-22-28)], MotifSampler [\[51\]](#page-22-29), rGADEM [[52](#page-22-30)], Sensitive, Thorough, Rapid, Enriched Motif Elicitation (STREME) [\[53\]](#page-22-31), and Multiple Em for Motif Elicitation (MEME) [[54\]](#page-22-32), are capable of identifying novel TFBS motifs without prior knowledge of the binding sites.

In addition to PWM-based models, more complex models have recently been proposed for modeling TFBSs with increased accuracy [[15\]](#page-21-17), in particular, hidden Markov models (HMMs) [\[23,](#page-22-1) [55](#page-22-33)[–58](#page-23-0)], hierarchical mixture models [[59\]](#page-23-1), Bayesian networkbased methods  $[60]$  $[60]$  and deep learning based neural networks  $[61-64]$  $[61-64]$  $[61-64]$ . Tools such as Motif cluster alignment and search tool (MCAST) [[65\]](#page-23-5), TFBSpred [\[17](#page-21-14)], and FABIANvariant [[66](#page-23-6)] utilize HMM-based methods while VOMABT [[67](#page-23-7)], and MotEvo [[68](#page-23-8)], tools use methods based on Bayesian network models. Tools based on deep learning methods include DeepBind [[69](#page-23-9)], DeeperBind [[70\]](#page-23-10), DeepGRN [[61\]](#page-23-3), DeepSNR [[71](#page-23-11)], DeepSTF [[62\]](#page-23-12), Desso [[72\]](#page-23-13), MaResNet [[73\]](#page-23-14), TAMC [\[74](#page-23-15)] and TFImpute [[75](#page-23-16)]. While a variety of computational tools are available for predicting TFBSs, as shown in Fig. [1](#page-3-0), PWM-based methods remain widely used for their simplicity in construction from a set of sequences, and the availability of several curated databases of PWMs applicable to several species.

Accurate prediction of TFBSs in silico is important for understanding gene regulation. However, only a limited number of independent assessments have been conducted to evaluate the performance of TFBS prediction and motif discovery tools [\[14](#page-21-13), [76–](#page-23-17)[81\]](#page-23-18), with the latest comprehensive analysis dating back to 2016. As the feld of computational biology continues to evolve and new tools are developed, assessment of tool performance is essential to ensure accurate and reliable predictions. The findings of such studies will guide the researchers to choose the most appropriate tool(s) for their analyses. Therefore, the present study aimed to evaluate the performance of several widely used TFBS prediction and de novo motif discovery tools, including both previously evaluated and untested tools. The specific objectives of this study were to (1) create a benchmark dataset to evaluate the performance of tools, (2) assess the performance quality of tools using numerous statistics to identify the best tool(s), and (3) conduct case studies involving known genes from key biological pathways in Arabidopsis and humans, using the identifed best tool(s) to predict TFBSs.



<span id="page-3-0"></span>**Fig. 1** Tools available for modeling transcription factor binding sites

<span id="page-3-1"></span>**Table 1** Transcription factors used in the present study

Organism	<b>Transcription factor name</b>	<b>JASPAR ID</b>
Arabidopsis thaliana	APETALA3 (AP3)	MA0556.1
	G-box binding factor 3 (GBF3)	MA1351.1
Homo sapiens	Transcription factor AP-2 alpha (TFAP2A)	MA0003.2
	Paired box 5 (PAX5)	MA0014.2

## **Methods**

## **Development of a benchmark dataset**

A benchmark dataset was generated following the recommendations of Tompa et al. 2005 [[77\]](#page-23-19), to assess the performance of TFBS prediction tools. This dataset comprised four types of sequences: real, generic, Markov, and negative. Real sequences, containing experimentally validated binding sites, were collected from the JASPAR database [[24](#page-22-2), [82](#page-23-20)] for the TFs listed in Table [1](#page-3-1) for both *Arabidopsis thaliana* and *Homo sapiens*. Five sequences of 115 bp length were randomly selected per TF, resulting in a total of 20 real sequences. The choice of 115 bp was based on the typical length of known binding sites from the JASPAR database, which are derived from ChIP-seq experiments and generally fall within this range. Generic sequences were generated by randomly selecting promoter sequences from the genomes of *A. thaliana* and *H. sapiens*. Initially, gf3 fles of *H. sapiens* and *A. thaliana* reference genome sequences were obtained from GENCODE [[83\]](#page-23-21) and NCBI [[84\]](#page-23-22), respectively. Ten genes were randomly extracted per organism, and

115 bp long promoter sequences (upstream from the transcription start site (TSS)) were obtained for each gene using the Ensembl genome browser [\[85](#page-23-23)], resulting in 20 generic sequences. Markov (or artifcial) sequences (20 sequences) were generated using a thirdorder Markov model. Experimentally validated binding sites obtained from the JASPAR database were then implanted into the generic sequences and Markov sequences. Negative sequences, lacking binding sites for the selected four TFs, were also acquired from the JASPAR database; sequences containing similar sites were removed from the dataset, resulting in 20 negative sequences. The real, generic, Markov, and negative sequences, totaling 80 sequences compiled into a single fle in FASTA format to create the benchmark dataset (Additional fle [1](#page-20-0)).

## **Selection of tools for performance evaluation**

As shown in Fig. [1,](#page-3-0) several tools have been released since 2019. However, the currently available deep learning approaches for predicting TFBSs have primarily been trained on human datasets  $[61, 62, 72-74]$  $[61, 62, 72-74]$  $[61, 62, 72-74]$  $[61, 62, 72-74]$  $[61, 62, 72-74]$  $[61, 62, 72-74]$ . Consequently, their direct applicability and performance on plant-specifc data remain uncertain without signifcant retraining and validation  $[86–88]$  $[86–88]$ . Therefore, in the current study, we opted to exclude deep learning models from our evaluation.

Given that this study involves TFs from both human and Arabidopsis genomes, we focused our evaluation on tools mainly based on PWMs and HMMs. However, we did not include the recently released FABIAN-variant tool [[66\]](#page-23-6) in our analysis. Although it uses transcription factor fexible models (an extension of HMMs) and PWMs, FABIANvariant is specifcally designed to predict the efects of DNA variants on human TFs [\[66](#page-23-6)]. Additionally, we excluded two PWM-based tools developed after 2019, TDTub and TFBSPred. TDTub is a web server that relies on a set of pre-computed TFBS in plant species [\[41](#page-22-19)], and TFBSPred is designed for human and mouse TFs [\[17\]](#page-21-14), limiting their relevance to our study, which includes both human and Arabidopsis TFs.

In our study, we evaluated the performance of 12 TFBS prediction tools and four de novo motif discovery tools commonly used in research. Tis focused evaluation ensures that our fndings are robust and relevant to both human and plant TFs. Table [2](#page-5-0) provides an overview of the tools employed in this study. PWMs for the selected TFs were obtained from the JASPAR database in JASPAR, MEME, and TRANSFAC matrix formats; the list of PWMs used in this study, along with related information is provided in Additional fle [2](#page-20-0). Default parameters were applied to all tools to enable comparisons among them.

## **Performance metrics**

Several statistical parameters were used to assess the performance quality of the tools. For each tool, the number of true positives (nTPs), false positives (nFPs), and false negatives (nFNs) were calculated for the benchmark dataset, considering diferent overlap percentages between the lengths of predicted and known binding sites or motifs (i.e., 80, 90, and 100%). True positives (TPs) denote the number of nucleotide positions in both known and predicted binding sites, false positives (FPs) represent the number of nucleotide positions not in known binding sites but in predicted binding sites, and false negatives (FNs) indicate the number of nucleotide positions in known binding sites



<span id="page-5-0"></span>

but not in predicted binding sites [[14,](#page-21-13) [77](#page-23-19)]. Employing these values, statistical parameters were calculated for each tool, including sensitivity (Sn) (Eq. ([1\)](#page-5-1)), positive predictive value (PPV) (Eq.  $(2)$  $(2)$ ), performance coefficient (PC) (Eq.  $(3)$  $(3)$ ), average site performance (ASP) (Eq.  $(4)$  $(4)$ ), and geometric accuracy (ACCg) (Eq.  $(5)$  $(5)$ ). The tools were individually ranked using Excel's RANK function, considering each of the above-mentioned statistical parameters. Subsequently, the average rank for each tool across all statistical parameters was computed using Excel's AVERAGE function.

$$
Sensitivity (Sn) = (nTPs)/(nTPs + nFNs)
$$
\n(1)

<span id="page-5-2"></span><span id="page-5-1"></span>Positive predictive value (PPV) = 
$$
(nTPs)/(nTPs + nFPs)
$$
 (2)

<span id="page-5-3"></span>Performance coefficient (PC) = 
$$
(nTPs)/(nTPs + nFNs + nFP)
$$
 (3)

<span id="page-6-0"></span>Average site performance (ASP) = 
$$
(Sn + PPV)/2
$$
 (4)

<span id="page-6-1"></span>Geometric accuracy (ACCg) = 
$$
\sqrt{\text{(Sn * PPV)}}
$$
 (5)

#### **Evaluation of the best‑performing tools**

To assess the best-performing tools, we randomly selected 50 genes from both the *A. thaliana* and *H. sapiens* genomes (See Additional fle [3](#page-20-1) for the gene list). Promoter regions of these genes, defned as the 1000 bp upstream from the TSS, were retrieved from the Eukaryotic Promoter Database (EPD) [\[107\]](#page-24-7). These sequences were scanned for 879 vertebrate PWMs and 1,407 plant PWMs obtained from the JASPAR database (Additional fle [2\)](#page-20-0) using the three best-performing tools: MCAST, FIMO, and MOODS. All the predicted motifs were ranked by frequency, and the top 20 motifs from each tool were compared to evaluate the consistency of their prediction.

# *Case study 1: transcription factor binding sites in genes associated with the anthocyanin biosynthesis pathway in plants*

A total of 149 genes involved in regulating anthocyanin biosynthesis in *A. thaliana* were obtained from Grau et al. 2022 [[41\]](#page-22-19) (Additional fle [3\)](#page-20-1). Additionally, we downloaded genes associated with anthocyanin biosynthesis in *Zea mays*, *Oryza sativa*, and *Glycine max* using TDThub [\[41](#page-22-19)] (Additional file [3](#page-20-1)). The promoter regions of these genes (1000 bp upstream from the TSS) were extracted from the EPD and NCBI databases [[105,](#page-24-8) [107](#page-24-7)]. These sequences were then scanned against PWMs of all plant TFBSs available in the JASPAR database, using the three best-performing TFBS prediction tools identifed in this study.

# *Case study 2: transcription factor binding sites in genes associated with the pentose phosphate pathway in humans*

Tirty-one genes involved in the pentose phosphate pathway were retrieved from the Kyoto Encyclopedia of Genes and Genomes (KEGG) database (Additional fle [3\)](#page-20-1). Promoter regions (1000 bp upstream from the TSS) were extracted from the EPD and scanned against 879 vertebrate PWMs obtained from the JASPAR database using the three best-performing tools.

## **Results**

## **Performance evaluation of diferent TFBS prediction tools**

The benchmark dataset comprised a total of 60 TPs, indicating that the maximum nTPs that any tool could predict in this study was limited to 60. Among the 12 TFBS prediction tools examined, none of them achieved 100% accuracy in predicting all 60 binding sites. However, both MotEvo and DWT-toolbox successfully identifed all TPs at both 80% and 90% overlap (Table [3](#page-7-0)). Interestingly, while the nTPs remained consistent at 80% and 90% overlap with a few exceptions, there was a notable decrease in the nTPs identifed at 100% overlap across all TFBS prediction tools, resulting in higher nFNs (Table [3\)](#page-7-0). Particularly, TFBStools predicted fewer TPs at 80% overlap compared to other tools and failed to predict any binding sites at 90% and 100%

Tool	Overlap percentage*									
	80%			90%			100%			
	<b>nTPs</b>	nFNs	<b>nFPs</b>	<b>nTPs</b>	nFNs	<b>nFPs</b>	<b>nTPs</b>	<b>nFNs</b>	<b>nFPs</b>	
Ciiider	43	17	10	43	17	10	34	26	9	
DWT-toolbox	60	$\mathbf{0}$	65	60	0	65	45	15	65	
<b>FIMO</b>	57	3	17	57	3	17	45	15	17	
<b>HOMER</b>	58	2	1002	58	$\overline{2}$	1002	43	17	1002	
<b>INSECT 2.0</b>	45	15	12	45	15	12	36	24	12	
Matrix scan	50	10	13	50	10	13	41	19	13	
<b>MCAST</b>	51	9	$\Omega$	51	9	0	42	18	$\Omega$	
<b>MOODS</b>	52	8	12	52	8	12	43	17	12	
Morpheus	54	6	142	51	9	142	46	14	142	
MotEvo	60	$\mathbf{0}$	35	60	0	35	45	15	35	
PWMScan	57	3	20	57	3	20	45	15	20	
TFBStools	9	51	$\overline{2}$	0	60	$\overline{2}$	0	60	$\overline{2}$	

<span id="page-7-0"></span>**Table 3** Number of true positives (nTPs), false negatives (nFNs), and false positives (nFPs) detected by diferent TFBS prediction tools

\* Overlap percentage: the percentage overlap between the length of predicted and known binding sites



<span id="page-7-1"></span>**Fig. 2** Sensitivity of TFBS prediction tools at different overlap percentages

overlap. Additionally, the nFPs identifed by the HOMER tool was relatively high compared to the other TFBS prediction tools examined (Table [3\)](#page-7-0).

At both 80% and 90% overlap, Sn was generally higher compared to 100% for all the tools except for TFBStools (Fig. [2\)](#page-7-1). Although Morpheus showed the highest sensitivity at 100% overlap with a sensitivity value of 0.767, DWT-toolbox, FIMO, HOMER, MotEvo, and PWMScan surpassed Morpheus at both 80% and 90% overlap. Moreover, TFBStools, which was unable to predict any binding sites at both 90% and 100% overlap, was able to predict binding sites at 80% overlap. However, the Sn was very low (0.150). Overall, it was evident that DWT-toolbox, MotEvo, Morpheus, PWM-Scan, and FIMO consistently delivered  $Sn \geq 0.750$  across all three overlap percentages.



<span id="page-8-0"></span>**Fig. 3** Positive predictive values of TFBS prediction tools at three diferent overlap percentages



<span id="page-8-1"></span>**Fig. 4** Performance coefficient of TFBS prediction tools at three different overlap percentages

When examining the PPVs obtained for the three diferent overlap percentages, it was observed that the tools Ciiider, INSECT 2.0, Matrix-scan, MCAST, and MOODS consistently provided the best PPVs (0.750 or higher) throughout all three overlap percentages; of these, MCAST tool exhibited the highest PPV (1.000) at all overlap percentages (Fig. [3](#page-8-0)). Although the PPV of TFBStools was null at 100% and 90% overlap, a signifcant increase was observed at 80% overlap, making it the second highest (Fig. [3\)](#page-8-0). Additionally, the PPV of the DWT-toolbox, HOMER and Morpheus tools was relatively low compared to other tools examined, at all three overlap percentages (PPV<0.500).

Assessing the performance of TFBS tools based on the PC values revealed that the MCAST tool exhibits the best performance across all three overlap percentages (Fig. [4](#page-8-1)). Following MCAST, the tools MOODS and FIMO, secured the second and third-highest PC values, respectively, at 100% overlap (Fig. [4\)](#page-8-1). However, at 90% and 80% overlap, FIMO emerged as the tool with the second-highest PC value, while MOODS exhibited the third-highest PC value (Fig. [4](#page-8-1)). Similar to statistical parameters: Sn, and PPVs, the TFBStools showed zero PC at 100% and 90% overlap but exhibited a value of 0.145 at



<span id="page-9-0"></span>**Fig. 5** Average site performance of TFBS prediction tools at three diferent overlap percentages



<span id="page-9-1"></span>**Fig. 6** Geometric accuracy of TFBS prediction tools at three different overlap percentages

80% overlap (Fig. [4\)](#page-8-1). Additionally, the PC of DWT-toolbox, HOMER, and Morpheus tools were considerably low across all three overlap percentages (PC<0.500).

As shown in Fig. [5](#page-9-0), MCAST demonstrated the best performance across all three overlap percentages based on the estimated ASP measures. Similar to the tool performances based on PCs, the MOODS tool exhibited the second-highest ASP and the FIMO tool showed the third-highest ASP at 100% overlap (Fig. [5\)](#page-9-0). Furthermore, at 90% and 80% overlap, the second-best and third-best ASPs were shown by FIMO and MOODS tools, respectively. It was also noted that TFBStools showed ASP value only at 80% overlap. Similar to PCs and ASP measures, a consistent trend in tool performance was observed when considering the ACCg of the TFBS prediction tools (Fig. [6\)](#page-9-1). The MCAST demonstrated the highest ACCg across all overlap percentages, followed by FIMO and MOODS tools. HOMER and Morpheus tools exhibited relatively low ACCg compared to other tools examined, at all three overlap percentages ( $ACCg < 0.500$ ). Taking into account the PC (Fig. [4](#page-8-1)), ASP (Fig. [5\)](#page-9-0), and ACCg (Fig. [6](#page-9-1)), it became evident that the best-performing tools were MCAST, MOODS, and FIMO.

The best TFBS prediction tool/tools could not be solely identified from the values of statistical parameters, as various tools dominated diferent parameters. Hence, to determine the best-performing tool/tools across all the statistical parameters considered in the present analysis, the tools were ranked under each parameter, and the average rank was calculated. The average ranks obtained at different overlap percentages are provided in Additional fle [4](#page-20-2). It was revealed that the MCAST tool outperformed others at 100% overlap, while MOODS and FIMO tools secured the second and third positions, respectively. The average ranks obtained at  $90\%$  overlap (Additional file [4](#page-20-2)), further showed that the MCAST tool performed the best at 90% overlap, following FIMO and MOODS tools. Consistent with the fndings at 100% and 90% overlaps, it was revealed that the MCAST tool exhibited the best performance across all statistical parameters at 80% overlap. Additionally, the FIMO and MOODS tools were observed to secure the secondbest and third-best rankings, respectively at 80% overlap (Additional fle [4\)](#page-20-2).

Furthermore, each data type of the benchmark dataset was analyzed independently, irrespective of species, to assess the performance of TFBS prediction tools with an 80% overlap between the length of predicted and known binding sites. DWT-toolbox and MotEvo identifed the maximum nTPs in all three datasets. In addition to these tools, HOMER and PWMScan identifed all 20 binding sites present in the real dataset (Table [4\)](#page-10-0). However, HOMER resulted in a considerably higher number of FPs across all data types. Furthermore, TFBStools resulted in a notably lower number of TPs across all data types.

When assessing the Sn of tools in predicting binding sites, it was observed that generally, all tools except TFBStools exhibited Sn greater than 0.500 across all data types (Fig. [7a](#page-11-0)). Among these, two tools (DWT-toolbox and MotEvo) exhibited maximum Sn across all three data types. Based on the results obtained for the PPVs of the tools, it was observed that only the MCAST tool consistently performed well across all three data types (Fig. [7b](#page-11-0)). All tools, except HOMER and Morpheus, exhibited PPVs greater

Tool	Data type									
	Real			Generic			<b>Markov</b>			
	<b>nTPs</b>	nFNs	<b>nFPs</b>	<b>nTPs</b>	<b>nFNs</b>	<b>nFPs</b>	<b>nTPs</b>	nFNs	<b>nFPs</b>	
Ciiider	15	5	$\overline{4}$	14	6	$\mathbf{0}$	14	6	1	
DWT toolbox	20	$\mathbf{0}$	10	20	$\mathbf{0}$	$\overline{4}$	20	$\mathbf 0$	11	
<b>FIMO</b>	16	$\overline{4}$	$\overline{4}$	17	3	$\mathbf{0}$	17	3	$\mathbf{0}$	
<b>HOMER</b>	20	$\mathbf{0}$	267	19	1	262	19	1	264	
<b>INSECT 2.0</b>	15	5	$\overline{4}$	15	5	$\mathbf{0}$	15	5	$\Omega$	
Matrix scan	16	$\overline{4}$	$\overline{4}$	17	3	$\mathbf{0}$	17	3	$\Omega$	
<b>MCAST</b>	17	3	0	17	3	$\Omega$	17	3	$\cap$	
<b>MOODS</b>	17	3	$\overline{4}$	18	$\overline{2}$	$\Omega$	17	3	$\Omega$	
Morpheus	18	2	33	18	$\overline{2}$	30	18	2	29	
MotEvo	20	$\mathbf 0$	$\overline{4}$	20	$\mathbf{0}$	2	20	0	$\overline{2}$	
PWMScan	20	$\overline{0}$	3	19	1	1	18	$\overline{2}$	3	
TFBStools	3	17		3	17	$\mathbf{0}$	3	17	$\mathbf{0}$	

<span id="page-10-0"></span>**Table 4** Number of true positives (nTPs), false negatives (nFNs), and false positives (nFPs) detected by each TFBS prediction tool, categorized by data type with an 80% overlap



<span id="page-11-0"></span>**Fig. 7** Tool performance matrices across diferent data types. **a** Sensitivity, **b** Positive predictive values, and **c** Performance coefficient

than 0.500 for all data types. Similarly, the comparison of PC values of the tools across the three data types revealed that the MCAST tool performed equally well across all data types (Fig. [7c](#page-11-0)). In addition, MotEvo showed greater performance across all data types. It was also observed that the TFBStools, Morpheus, and HOMER tools exhibited comparatively lower performance (PC< 0.500).

Tool	nTPs	nFNs	nFPs
MEME	59	01	20
MotifSampler	39	21	18
<b>STREME</b>	39	21	14
rGADEM	19	41	08

<span id="page-12-0"></span>**Table 5** Number of true positives (nTPs), false negatives (nFNs), and false positives (nFPs) detected by de novo motif discovery tools at 70% overlap



<span id="page-12-1"></span>**Fig. 8** Distribution of the top 20 motifs predicted by MCAST, FIMO, and MOODS tools for randomly selected promoter sequences. **a** *Arabidopsis thaliana*, **b** *Homo sapiens*

## *Performance evaluation of* **de novo** *motif discovery tools*

The results of our independent assessment of four commonly used de novo motif discovery tools provide valuable insights into their performance. The evaluation was conducted as described by Jayaram et al.  $[14]$  $[14]$ , using a 70% overlap between the known binding site and the binding site identifed by the tool. Among the tools assessed, MEME emerged as the top performer, identifying 59 binding sites of the 60 binding sites in the benchmark dataset. MotifSampler and STREME both identifed 39 binding sites, indicating a comparable level of performance between these two tools (Table [5](#page-12-0)). When evaluating the tools based on Sn, PC, ASP, and ACCg values, MEME consistently demonstrated the highest performance across these metrics.

## **Evaluation of the best‑performing tools**

To further evaluate the performance of the top three TFBS prediction tools identifed in the present study—MCAST, FIMO, and MOODS, a comparative analysis was conducted using the top 20 motifs identifed from randomly selected promoter regions of *A. thaliana* and *H. sapiens*. In the *A. thaliana* sequences, none of the predicted motifs were shared across the three tools (Fig. [8](#page-12-1)a). However, 12 motifs were commonly identifed by both MCAST and FIMO, while MOODS identifed a distinct set of motifs within its top 20 results (Fig. [8a](#page-12-1)). In the human promoter sequences, three motifs were predicted by all three tools. Additionally, six motifs were commonly identifed by both MCAST and FIMO, while five motifs were shared between FIMO and MOODS (Fig. [8](#page-12-1)b).

# *Case study 1: assessing transcription factor binding sites in the promoters of the genes involved in the anthocyanin biosynthesis pathway*

The anthocyanin biosynthesis is one of the most extensively studied pathways in plants, making it an ideal target for analyzing TFBSs in the promoter regions of related genes. In this study, we analyzed the promoter regions of 149 *A. thaliana*, 129 *G. max*, 74 *O. sativa*, and 70 *Z. mays* genes associated with anthocyanin biosynthesis using the three best-performing TFBS prediction tools: MCAST, FIMO, and MOODS. The analysis revealed that none of the top 20 motifs were shared across all three tools in any of the species analyzed. However, a notable overlap was observed between the motifs identifed by MCAST and FIMO. Specifcally, these two tools shared eight motifs in *A. thaliana*, ten motifs in *G. max*, ten motifs in *O. sativa*, and eight motifs in *Z. mays*, respectively (Additional fle [5](#page-20-3)).

MYB (v-Myb myeloblastosis viral oncogene homolog) TFs are key regulators of anthocyanin biosynthesis in plants [\[108–](#page-24-9)[112](#page-24-10)]. In our analysis, MCAST identifed only a few MYB binding sites among the top 20 motifs for *A. thaliana* (10%), *G. max* (5%), and *O. sativa* (5%) (Additional fle [5](#page-20-3)). In contrast, neither FIMO nor MOODS detected any MYB motifs in the top 20 motifs for these species. Moreover, in the promoter sequences of *Z. mays*, none of the tools detected MYB or MYB-related motifs among the top 20 motifs. As the number of analyzed motifs increased, the detection of MYB and/or MYBrelated motifs increased. In *A. thaliana*, both FIMO and MOODS began detecting MYB and/or MYB-related bindings sites within the top 40 motifs (Fig. [9a](#page-14-0)). Similarly, in *G. max*, FIMO detected MYB motifs within the top 60 motifs, while MOODS detected MYB-related motifs starting at the top 40 (Fig. [9](#page-14-0)b). For *O. sativa* sequences*,* MOODS failed to identify any MYB binding sites even within the top 100 motifs (Fig. [9c](#page-14-0)). However, for *Z. mays*, MYB binding sites were detected by MCAST, FIMO and MOODS starting from the top 40, 80 and 60, respectively (Fig. [9](#page-14-0)d).

Notably, in the promoter regions of *A. thaliana* and *G. max*, MCAST primarily identifed cystein-rich polycomb-like protein (CPP) motifs as the most abundant among the top 20 motifs, followed by DNA-binding with one fnger (DOF) motifs. In contrast, FIMO predicted DOF motifs as the most abundant, followed by CPP motifs (Additional fle [6\)](#page-20-4). Furthermore, all of the top 20 motifs identifed by MOODS in both species were DOF binding sites (Additional fle [6\)](#page-20-4). In the monocot species *O. sativa* and *Z. mays*, MCAST and FIMO primarily detected dehydration-responsive element-binding protein (DREB) binding sites within the top 20 motifs, while MOODS predominately identifed DOF binding sites, followed by DREB binding sites (Additional fle [6\)](#page-20-4).

These findings highlight the variability in motif detection across different tools, even when analyzing the same gene list within a species. This underscores the importance of carefully selecting the appropriate tools for TFBS discovery.

# *Case study 2: assessing transcription factor binding sites in the promoters of the genes involved in the pentose phosphate pathway in Homo sapiens*

The pentose phosphate pathway (PPP) is a fundamental component of cellular metabolism and plays a signifcant role in various human diseases [[113\]](#page-24-11). An analysis of the promoter regions of 31 genes involved in the PPP using the TFBS prediction tools MCAST,



<span id="page-14-0"></span>**Fig. 9** Frequency of motifs detected in promoter sequences of genes involved in the anthocyanin biosynthesis pathway in plants. **a** *Arabidopsis thaliana*, **b** *Glycine max*, **c** *Oryza sativa*, **d** *Zea mays*

FIMO, and MOODS revealed that six motifs were consistently detected among the top 20 motifs by all three tools (Fig. [10](#page-15-0)). Additionally, FIMO and MOODS shared eight motifs, while MCAST and FIMO shared fve motifs, highlighting both the overlap and variation in motif detection across these tools. Notably, most of the top 20 motifs identifed by all three tools corresponded to binding sites for TFs belonging to the C2H2 zinc fnger family (C2H2-ZF), highlighting the signifcance of this TF family in PPP regulation (Additional fle [6](#page-20-4)).

## **Discussion**

Advances in bioinformatics have revolutionized biological research by providing scientists with a range of computer tools for retrieval, analysis, and visualization of omics data, thereby uncovering the complexities of living organisms. However, this also presents a challenge for researchers, as the fndings and conclusions may



<span id="page-15-0"></span>**Fig. 10** Distribution of the top 20 motifs identifed by MCAST, FIMO, and MOODS across the promoter regions of the genes involved in the pentose phosphate pathway in *Homo sapiens*

signifcantly depend on the choice of tools used for the analysis. To address this, benchmarking studies are often conducted to rigorously compare the performance of diferent tools, identify their strengths, and provide recommendations on the most suitable tools for specifc analyses [[114\]](#page-24-12). Designing a robust performance evaluation of computational tools is challenging due to various factors, such as selecting appropriate benchmark datasets, determining optimal tool parameters, and choosing suitable statistical measures for performance evaluation. Therefore, it is essential to design benchmarking studies carefully to ensure accurate, unbiased, and informative results [[114](#page-24-12)].

To date, only a limited number of assessments have been conducted to evaluate the performance quality of TFBS prediction tools. For instance, Tompa et al. [\[77](#page-23-19)], provide a comparative analysis of several TFBS discovery tools, including AlignACE, ANN-Spec, Consensus, GLAM, Improbizer, MEME, MITRA, MotifSampler, oligo/ dyad-analysis, QuickScore, SesiMCMC, Weeder, and YMF. Additionally, Jayaram et al. [[14\]](#page-21-13), evaluate a set of TFBS prediction tools (i.e., MCast, Baycis, Cister, Cluster-Buster, Comet, FIMO, Clover, Matrix-Scan, Patser and Possum-Search), as well as de novo motif discovery tools (i.e., MEME-ChIP, HOMER, ChIPMunk and rGADEM). Among the 16 tools assessed in the present study, the performance of INSECT 2.0, Morpheus, Ciiider, PWMScan, MOODS, MotEvo, DWT-toolbox, and TFBStool has not been previously evaluated. Our study was conducted from a general user's perspective, employing default settings wherever possible or the recommended settings provided in the corresponding user manuals. If no recommendations were available, settings were chosen based on personal preference.

The selection of the reference dataset plays a crucial role in benchmarking computational methods [[114\]](#page-24-12). Previous studies evaluating TFBS prediction tools often relied on reference datasets consisting of preprocessed ChIP-seq peaks or annotated TFBSs from the TRANSFAC database (14, 77, 81]. However, in our study, we opted to utilize the information stored in the JASPAR database which provides annotated TFBSs alongside sequences obtained from ChIP-seq experiments and is freely accessible, unlike TRANSFAC. To construct the reference or benchmark dataset, we incorporated four types of data: real, generic, Markov, and negative sequences. This approach aimed to minimize biases that could potentially favor certain tools over others [\[77](#page-23-19), [114\]](#page-24-12).

Establishing criteria for defning an actual binding site (true positive) is crucial for comparing diferent TFBS or motif prediction tools. Employing a 100% overlap percentage between the length of actual and predicted binding sites to classify them as TPs may introduce bias, given the difficulty in precisely predicting TFBSs by many tools. In the study conducted by Jayaram et al.  $[14]$  $[14]$  $[14]$ , an overlap percentage of 70% was used, while Tompa et al. [\[77\]](#page-23-19), considered an overlap by at least one-quarter of the length of the known site as a TP. Consequently, in this study, diferent overlap percentages were applied: 100%, 90%, and 80% for TFBS prediction tools. No signifcant changes in results were observed beyond an 80% overlap, and thus, lower overlap percentages were not considered. Additionally, for de novo motif discovery tools, a 70% overlap was employed, following the approach described by Jayaram et al. [\[14\]](#page-21-13).

Among the tools evaluated, MCAST stands out for its unique ability to avoid FPs across all three overlap percentages. Tis remarkable performance can be attributed to its algorithm, which is based on HMMs. MCAST is specifcally designed to identify clusters of non-overlapping motifs, commonly referred to as *cis*-regulatory modules (CRMs) [[65](#page-23-5)]. By focusing on clusters of motifs rather than individual occurrences, MCAST efectively reduces the likelihood of FPs, enhancing the accuracy of motif detection. However, while MCAST excels in minimizing FPs, its sensitivity was below that of many of the tools including FIMO and MOODS. Sensitivity is essential for ensuring that true binding sites are not overlooked. In applications where detecting all possible binding sites is crucial, this limitation of MCAST should be taken into account. Lower sensitivity means that some true binding sites may not be identifed, which can be a signifcant drawback in comprehensive motif discovery studies.

The findings of the present study align with those of Jayaram et al.  $[14]$  $[14]$  $[14]$ , where MCAST and FIMO emerged as the best best-performing tools. However, in our study, MOODS outperformed FIMO at a 100% overlap percentage, while FIMO surpassed MOODS at the other two overlap percentages. Both of these tools are designed to identify individual binding sites, unlike MCAST. FIMO offers both web-based and command-line interfaces, with the web tool being particularly user-friendly [\[30\]](#page-22-8). In contrast, MOODS operates solely via the command line on Linux systems, though it boasts a faster runtime compared to FIMO [[32\]](#page-22-10).

Comparing the performance quality of TFBS prediction tools poses a challenge due to their reliance on diferent algorithms for motif discovery. Our benchmark dataset included only one binding site per sequence and therefore, was not afected by these algorithmic diferences. However, in real datasets, multiple binding sites may be present in a given DNA sequence. Hence, it is more reliable to conclude that for an average-end user aiming to identify CRMs with minimal FPs, MCAST stands out as the best option. Conversely, for identifying individual binding sites, FIMO and MOODS emerge as the best options. The choice between FIMO and MOODS may depend on factors such as run-time, user-friendliness, and the expected level of accuracy.

Although MCAST, FIMO, and MOODS outperformed other tools overall, DWTtoolbox and MotEvo excelled in identifying the highest number of TPs at both 80% and 90% overlaps. DWT-toolbox, a command line tool employing a Bayesian network model, efectively addresses a critical issue in TFBS prediction tools: pairwise dependency [[15,](#page-21-17) [115\]](#page-24-13). Unlike the PWM method used in most tools, which assumes independence between positions, the DWT-toolbox considers all possible pairwise dependencies within a rigourous probabilistic framework devoid of tunable parameters, automatically avoiding overftting [[106\]](#page-24-0). MotEvo, another command line tool, employs a Bayesian probabilistic model-based algorithm specifcally designed for analyzing TFBSs across phylogenetically related multiple organisms [[68\]](#page-23-8), but can also handle single species sequences. However, considering its fle requirements and parameters, it is conceivable that MotEvo could have performed even better with phylogenetically related sequences.

Consistent with previous fndings [\[77\]](#page-23-19), our evaluation of TFBS prediction tools across three diferent data types: real, generic, and Markov, further underscores the critical role of experimental design in performance evaluation research. Notably, a considerable increase in the number of FPs was observed in the real dataset compared to the other data types. This suggests the presence of putative binding sites for other TFs beyond those known and utilized in the current experimental design. Consequently, tools that successfully identify these additional binding sites may be unfairly penalized due to the experimental setup. To mitigate this bias, generic and Markov datasets were included in the benchmark, ensuring a more balanced evaluation across diferent tools. MCAST demonstrated consistent performance across all three data types and all statistical parameters assessed, reinforcing its reliability as a TFBS prediction tool.

When evaluating the overall performance of TFBS prediction tools across various overlap percentages and data types, most tools demonstrated satisfactory performance, with some emerging as the best performers. However, both HOMER and TFBStools exhibited notably poor performances. HOMER, a command-line tool, ranked among the lowest performers due to its tendency to identify a high number of FPs, despite its ability to accurately recognize TPs. Tis high FP rate reduces its overall reliability, highlighting the need for more precise algorithms or improved parameter settings. Furthermore, TFBStools, an R package, performed poorly in identifying TPs in our study, especially at 100% and 90% overlaps, with only a few TPs detected at 80% overlap. While TFBStools did not generate many FPs, its limited ability to predict most binding sites, even after parameter adjustments, raises concerns regarding its efectiveness as a TFBS prediction tool. Tis limited predictive capability suggests that TFBStools may require signifcant algorithmic improvements or more sophisticated parameter tuning to enhance its performance.

Among the de novo motif discovery tools examined, the comprehensive dominance of MEME underscores its reliability and efficiency in accurately identifying binding motifs. MEME directly identifes motifs within user input sequences, making it highly efective for straightforward motif discovery tasks. Although both MEME and STREME are part of the MEME suite, they utilize diferent algorithms. STREME compares user sequences with a set of control sequences to detect relatively enriched motifs [\[53\]](#page-22-31). Tis methodological distinction suggests that STREME may ofer superior performance for certain types of analyses, particularly those involving diferential motif enrichment. MotifSampler's performance, while notable, may be impacted by a lack of updates and maintenance. This could affect its usability and reliability. Furthermore, our findings present a contrast to the results reported by Jayaram et al. [[14](#page-21-13)], where rGADEM was shown to outperform tools such as ChIPMunk, HOMER, and MEME-ChIP. In our study, rGA-DEM exhibited the poorest performance across most metrics, except for PPV. Tis discrepancy could be attributed to the parameter settings used during the motif discovery process. While Jayaram et al. [\[14\]](#page-21-13), manually optimized rGADEM's parameters to achieve better results, our study utilized the default settings. This highlights the importance of fine-tuning parameters for motif discovery tools to maximize their efficacy. Default settings, while convenient, may not always capture the nuances required for accurate motif identifcation. Users of rGADEM and similar tools should consider customizing parameters based on the specifc characteristics of their datasets to improve performance outcomes.

To further evaluate the performance of the best-performing tools—MCAST, FIMO, and MOODS, we focused on two distinct biological pathways: the anthocyanin biosynthesis pathway in plants and the PPP in humans. In addition, we analyzed a set of random promoter sequences from *A. thaliana* and *H. sapiens* genomes to assess the consistency in motif detection across the tools. Overall, our comparative analysis revealed considerable variability in motif detection across the tools, highlighting the importance of selecting the appropriate TFBS prediction tool(s) for diferent biological contexts to enhance prediction accuracy.

In plants, MYB TFs are one of the key regulators of anthocyanin biosynthesis [[110](#page-24-14), [116](#page-24-15)]. Our analysis revealed inconsistencies in detecting MYB binding sites among diferent tools when analysing the same set of promoter sequences and focusing on the top 20 motifs. Specifcally, MCAST identifed MYB motifs within the top 20 motifs in the promoters of anthocyanin biosynthesis-related genes in *A*. *thaliana*, *G. max*, and *O. sativa*. In contrast, neither FIMO nor MOODS detected MYB motifs within this range for these species. This suggests that MCAST may be more sensitive in detecting motifs based on their frequency within the top 20 motifs. Nevertheless, only a few MYB motifs were identifed among the top 20 motifs: MYB23 and MYB58 binding sites were detected in *A. thaliana*, and MYB31 binding sites were identifed in both *G. max* and *O. sativa*. As this analysis was expanded to include the top 40–100 motifs in increments of 20, both FIMO and MOODS began to detect MYB motifs, indicating that these tools may perform better with a broader search range.

A recent study analyzing TFBSs in Arabidopsis anthocyanin biosynthesis-related genes using FIMO, sorted by a user-defned Signifcance Score (S-Score), reported a prevalence of MYB TFs within the top 20 motifs [\[41\]](#page-22-19). In contrast, our present study, which also analyzed the same Arabidopsis anthocyanin gene list using FIMO but sorted based on motif frequencies, did not detect any MYB binding sites within the top 20 motifs. However, binding sites for MYB51, MYB80, and MYB93, as identifed in Grau and Franco-Zorrilla [\[41\]](#page-22-19), were detected among the top 100 motifs in our analysis. This discrepancy in motif detection may be due to diferences in the TFBS sorting criteria, and/or the set of PWMs employed during promoter scanning. For instance, in the present analysis, we utilized PWMs from the JASPAR database, which do not include certain PWMs such as MYB6, MYB37, and MYB38, which were considered in the analysis by Grau and Franco-Zorrilla [\[41](#page-22-19)].

Furthermore, the PWMs for several important MYB TFs that regulate anthocyanin biosynthesis in plants, specifcally, MYB11, MYB12, MYB75, MYB90, and MYB114 [[117–](#page-24-16)[119](#page-24-17)] were not included in either the JASPAR database or the study by Grau and Franco-Zorrilla  $[41]$  $[41]$ . This could limit the detection of important binding sites in the anthocyanin biosynthesis-related genes of the plant species if those binding sites are present. Therefore, as new TFs are identified and characterized, it is essential to continuously update freely accessible databases like JASPAR to enhance the accuracy of TFBS detection.

In addition to the MYB TF family, several other TF families are known to be involved in anthocyanin biosynthesis [[120](#page-24-18)[–122](#page-24-19)]. However, the role of DREB TFs in this pathway has not been extensively studied. Interestingly, our analysis revealed a higher prevalence of DREB binding sites in the anthocyanin biosynthesis-related genes of monocotyledon species such as *O. sativa* and *Z. mays*, as detected by MCAST and FIMO. Tis fnding suggests that DREB TFs may play a regulatory role in the anthocyanin pathway in monocots. Therefore, future studies exploring the potential involvement of DREB TFs in mediating anthocyanin biosynthesis in monocots could provide valuable insights into the regulation of this important secondary metabolic pathway in plants. Moreover, we observed a predominant presence of C2H2-ZF family binding sites in the genes of the human PPP. This family of TFs represents the largest class of multifunctional regulators in both humans and plants [\[123](#page-24-20), [124\]](#page-24-21). However, there is currently no information on the involvement of these TFs in the human PPP, opening avenues for future research.

## **Conclusion**

As various TFBS prediction tools become available, understanding the performance of diferent tools in terms of accuracy, sensitivity, and specifcity is crucial in genomic research. This study identified MCAST, FIMO, and MOODS as the top-performing TFBS prediction tools, while DWT-toolbox and MotEvo demonstrated increased efficacy in predicting a higher number of true binding sites, indicating superior sensitivity. MEME emerged as the most reliable de novo motif discovery tool.

The observed performance variations in this study emphasize the necessity for continuous benchmarking and independent assessments of TFBS prediction tools. Such evaluations help to identify the strengths and weaknesses of tools, guiding users in selecting the most appropriate tool(s) for their specifc research needs. Moreover, tool developers can leverage these insights to enhance algorithm robustness and usability, ensuring that default settings are more universally efective or providing clear guidance on parameter customization. Further studies should explore the impact of parameter optimization across diferent datasets to provide a more comprehensive understanding of each tool's capabilities and limitations.

It is recommended to further evaluate the performance quality of TFBS predicting tools using diverse benchmark datasets derived from multiple organisms, containing a range of TFBSs. Furthermore, relying solely on a single type of binding site per sequence, as done in the present study, may not provide a comprehensive evaluation of a tool's performance. Utilizing benchmark datasets from various organisms with multiple binding sites per sequence can provide valuable insights into the tool's performance across a broader spectrum of genomic contexts, better refecting the complexity of regulatory regions in biological systems. Additionally, this approach will enhance the accuracy and reliability of the evaluations.

In summary, due to variations in sensitivity and specifcity among TFBS prediction tools, it is recommended to integrate predictions from multiple tools to mitigate the limitations of individual tools, leading to a more thorough analysis of TFBSs. Tus, the development of a comprehensive toolbox that integrates TFBS prediction tools and de novo motif discovery tools would enable users to utilize a diverse array of tools, thereby increasing the likelihood of identifying significant binding sites/motifs. This integrative approach would facilitate a more robust and detailed understanding of transcriptional regulatory mechanisms, ultimately advancing genomic research.

#### **Abbreviations**



#### **Supplementary Information**

The online version contains supplementary material available at [https://doi.org/10.1186/s12859-024-05995-0.](https://doi.org/10.1186/s12859-024-05995-0)

<span id="page-20-4"></span><span id="page-20-3"></span><span id="page-20-2"></span><span id="page-20-1"></span><span id="page-20-0"></span>Additional fle1 (TXT 11 KB) Additional fle2 (XLSX 60 KB) Additional fle3 (XLSX 25 KB) Additional fle4 (XLSX 14 KB) Additional fle5 (PDF 205 KB) Additional fle6 (XLSX 30 KB)

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#### **Author contributions**

A.M.W. and S.V. conceived the study, participated in its design, and assisted in drafting the manuscript. D.V.W. contributed to the study design, performed analysis and interpreted data, and drafted the manuscript. All authors read and approved the fnal manuscript.

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#### **Data availability**

All data generated or analysed during this study are included in this published article [and its supplementary information fles]. No datasets were generated or analysed during the current study.

#### **Declarations**

## **Ethics approval and consent to participate**

Not applicable.

**Consent for publication** Not applicable.

#### **Competing interests**

The authors declare that they have no competing interests.

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