

Core promoter prediction in eukaryote

genomes

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Despite many recent efforts, in silico identification of promoter regions is still in its infancy. Accurate identification and delineation of promoter regions is important for several reasons, such as improving genome annotation and devising experiments to study and understand transcriptional regulation. Here, we present a novel approach that requires no training for predicting promoters in whole genome sequences by using large-scale structural properties of DNA. We compared our approach to the state-of-the-art in promoter prediction.



- Genome (3Gbp) \rightarrow genes \rightarrow promoter \rightarrow core promoter (50 bp)
- Many genomes (>50)
- No sequence preservation in core promoter







negative strand genes are shown in blue.

16 organisms. The profile is the average over a large number of promoters with the TSS on position zero.

Comparison with the state-of-the-art

Species	F
P. falciparum	0.17
O. pacifica	0.71
O. tauri	0.66
A. thaliana	0.37
O. sativa	0.53
P. trichocarpa	0.46
S. cerevisiae	0.42
S. pombe	0.31
C. elegans	0.26
D. melanogaster	0.19
T. nigroviridis	0.23
M. musculus	0.46
H. sapiens	0.45

→ Right: comparison of EP3 with state-state-of-the-art PPPs. The programs are ranked according to their F-measure on the CAGE data.

 $\leftarrow \textbf{Left:}$ comparison of the performance of EP3 on different eukaryotic genomes. Performance between species differs, but in general the performance is in line with the performance on human. Only EP3 was tested on all genomes because the other programs require training to work on those genomes.

Reference:

Abeel, T., Saeys, Y., Bonnet, E., Rouzé, P., Van de Peer, Y. (2008) Generic eukaryotic core promoter prediction using structural features of DNA. Genome Research 18, 310-23.

Program CAGE			
	Recall	Prec.	F
ARTS	0.38	0.74	0.50
EP3	0.34	0.66	0.45
DragonGSF	0.31	0.75	0.44
PromoterInspector	0.29	0.81	0.43
FirstEF	0.41	0.42	0.42
Eponine	0.28	0.75	0.41
CpgProD	0.34	0.41	0.37
PromoterExplorer	0.39	0.3	0.34
McPromoter (0.0)	0.17	0.68	0.28
PromFD	0.44	0.16	0.23
DragonPF	0.51	0.11	0.18
Promoter2.0	0.63	0.04	0.08
NNPP 2.2 (0.99)	0.03	0.03	0.03