S1 appendix

Reference graphs

All graphs used in the experiments can be downloaded from:

- Arabidopsis thaliana:
 hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/arabidopsis_thaliana.tar.gz
- Drosophila melanogaster:
 hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/drosophila_me
 lanogaster.tar.gz
- Human:
 hyperbrowser.uio.no/graph-peak-caller/static/graph peak caller data/human.tar.gz

ChiP-seq reads and motifs used

For all transcription factors on *Arabidopsis thaliana*, raw ChIP-seq reads were fetched from the NCBI Sequence Read Archive using the following accession numbers:

ERF115: SRR931836
SEP3: SRR1042995
AP1: SRX387187
SOC1: SRR1044950
PI: SRR502859

For the validation on human and *Drosophila melanogaster*, raw ChIP-seq reads were collected from experiments on ENCODE having the following experiment IDs. In all cases, raw fastq reads from replicate 1 were used:

Human CTCF: ENCSR000DUBHuman SRF: ENCSR000BIV

D. melanogaster JRA: ENCSR471GSA
 D. melanogaster SQZ: ENCSR923VWW
 D. melanogaster ANTP: ENCSR082RBU
 D. melanogaster JIM: ENCSR978WED

The motifs were collected from the Jaspar database [1] (for human and *A. thaliana*) or from OnTheFly [2]:

- *A. thaliana* ERF115*:

 https://hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/motifs/A
 RABIDOPSIS ERF115.meme
- A. thaliana SEP3: http://jaspar.genereg.net/api/v1/matrix/MA0563.1.meme
- A. thaliana AP1: http://jaspar.genereg.net/api/v1/matrix/MA0940.1.meme
- A. thaliana SOC1: http://jaspar.genereg.net/api/v1/matrix/MA0554.1.meme
- A. thaliana PI: http://jaspar.genereg.net/api/v1/matrix/MA0559.1.meme
- Human CTCF: http://jaspar.genereg.net/api/v1/matrix/MA0139.1.meme

- Human SRF: http://jaspar.genereg.net/api/v1/matrix/MA0083.2.meme
- D. melanogaster JRA:
 https://hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/motifs/D
 M JRA.meme
- D. melanogaster SQZ:
 https://hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/motifs/D
 M SQZ.meme
- D. melanogaster ANTP:
 https://hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/motifs/D
 M_ANTP.meme
- D. melanogaster JIM:
 https://hyperbrowser.uio.no/graph-peak-caller/static/graph_peak_caller_data/motifs/D
 M JIM.meme
- * Since we were not able to find a motif for A. thaliana ERF115 in the Jaspar database, we used the following motif from the Plant Cistrome Database [3]: http://neomorph.salk.edu/dap_web/pages/browse_table_ai.php?AGI=ERF115

References

- [1] Khan, A. et al. JASPAR 2018: update of the open-access database of transcription factor binding profiles and its web framework. Nucleic Acids Res. 2018; 46:D260–D266, doi:10.1093/nar/gkx1126
- [2] Shazman, Shula, Hunjoong Lee, Yakov Socol, Richard S. Mann, and Barry Honig. "OnTheFly: a database of Drosophila melanogaster transcription factors and their binding sites." Nucleic acids research 42, no. D1 (2013): D167-D171.
- [3] Ronan C. O'Malley, Shao-shan Carol Huang, Liang Song, Mathew G. Lewsey, Anna Bartlett, Joseph R. Nery, Mary Galli, Andrea Gallavotti, and Joseph R. Ecker (2016). Cistrome and Epicistrome Features Shape the Regulatory DNA Landscape, Cell. 2016 May 19;165(5):1280-92. doi: 10.1016/j.cell.2016.04.038.