CoopTFD: a repository for predicted yeast cooperative transcription factor pairs

Wei-Sheng Wu*, Fu-Jou Lai, Bor-Wen Tu and Darby Tien-Hao Chang

Department of Electrical Engineering, National Cheng Kung University, Tainan 70101, Taiwan

*Corresponding author: Tel: +886-6-2757575 ext. 62426; Fax: +886-6-2345482; E-mail: wessonwu@mail.ncku.edu.tw

Citation details: Wu, W.-S., Lai, F.-J., Tu, B.-W. et al. CoopTFD: a repository for predicted yeast cooperative transcription factor pairs. Database (2016) Vol. 2016: article ID baw092; doi:10.1093/database/baw092

Received 30 January 2016; Revised 1 May 2016; Accepted 9 May 2016

Abstract

In eukaryotic cells, transcriptional regulation of gene expression is usually accomplished by cooperative Transcription Factors (TFs). Therefore, knowing cooperative TFs is helpful for uncovering the mechanisms of transcriptional regulation. In yeast, many cooperative TF pairs have been predicted by various algorithms in the literature. However, until now, there is still no database which collects the predicted yeast cooperative TFs from existing algorithms. This prompts us to construct Cooperative Transcription Factors Database (CoopTFD), which has a comprehensive collection of 2622 predicted cooperative TF pairs (PCTFPs) in yeast from 17 existing algorithms. For each PCTFP, our database also provides five types of validation information: (i) the algorithms which predict this PCTFP, (ii) the publications which experimentally show that this PCTFP has physical or genetic interactions, (iii) the publications which experimentally study the biological roles of both TFs of this PCTFP, (iv) the common Gene Ontology (GO) terms of this PCTFP and (v) the common target genes of this PCTFP. Based on the provided validation information, users can judge the biological plausibility of a PCTFP of interest. We believe that CoopTFD will be a valuable resource for yeast biologists to study the combinatorial regulation of gene expression controlled by cooperative TFs.

Database URL: http://cosbi.ee.ncku.edu.tw/CoopTFD/ or http://cosbi2.ee.ncku.edu.tw/CoopTFD/

Introduction

Transcriptional regulation of gene expression is one of the major mechanisms for cells to respond to environmental and physiological changes (1, 2). This kind of regulation is usually accomplished by cooperative transcription factors (3–5). For example, the expression of NeuroD1, an essential pancreatic islet gene, is known to be regulated by two cooperative transcription factors Nkx2.2 and Ngn3 (3). Two transcription factors YY1 and E2F1 are known to cooperatively regulate the expression of p73, a protein which plays an important role in tumorigenesis (4). The cooperativity among transcription factors (TFs) enables cells to use a relatively small number of TFs in establishing the complex spatial and temporal patterns of gene expression.
Therefore, identifying cooperative TFs is helpful for uncovering the mechanisms of transcriptional regulation.

With the advent of many high-throughput experimental technologies (e.g. DNA sequencing, microarrays, ChIP-chips, TF knockout experiments and protein arrays), important information of a cell can be obtained. For example, DNA sequencing can provide the DNA sequences of gene promoters. Microarrays can provide gene expression levels. ChIP-chips can provide the binding targets of a specific TF. TF knockout experiments can provide the genes affected by the knockout of a specific TF. Protein arrays can provide protein pairs which have physical interactions. The measurements from different high-throughput experimental technologies are valuable data which can be utilized to computationally identify cooperative TF pairs. Therefore, many computational algorithms have been developed to predict cooperative TF pairs by using one data source or integrating multiple data sources generated by high-throughput experimental technologies. Some algorithms used only gene expression data (6) or ChIP-chip data (7, 8). Several other algorithms integrated ChIP-chip data with gene expression data (9–13), promoter sequence data (14–16), protein–protein interaction data (17) or TF knockout data (18). Another several algorithms integrated more than two high-throughput data sources (19–23).

Previous studies (24, 25) have shown that the performance of an algorithm is varied under different evaluation criteria such as the existence of physical/genetic interaction and the overlap with the benchmark set of known cooperative TF pairs.

Most existing cooperative TFs identification algorithms were applied to the model organism Saccharomyces cerevisiae. Different algorithms predicted different number of cooperative TF pairs ranging from a dozen to more than three thousands. These predicted cooperative TF pairs (PCTFPs) are valuable resources and provide testable hypotheses for future experimental investigation. Unfortunately, these PCTFPs were scattered in different papers and there is still no database that collects these PCTFPs from existing algorithms. This prompted us to construct Cooperative Transcription Factors Database (CoopTFD), which has a comprehensive collection of 2622 PCTFPs in yeast from 17 existing algorithms.

To help users judge the biological plausibility of a PCTFP, CoopTFD provides five types of validation information: (i) the algorithms which predict this PCTFP, (ii) the publications which experimentally show that this PCTFP has physical or genetic interactions, (iii) the publications which experimentally study the biological roles of both TFs of this PCTFP, (iv) the common Gene Ontology (GO) terms of this PCTFP and (v) the common target genes of this PCTFP. Having these five types of validation information could help biologists pick up biologically plausible PCTFPs for further experimental investigation.

Well-established databases such as SGD (26), BioGRID (27) and IntAct (28) can provide protein pairs with physical/genetic interactions, suggesting plausible cooperative TF pairs. In contrast, CoopTFD can provide computationally predicted cooperative TF pairs which may or may not have physical/genetic interactions. Therefore, CoopTFD can generate alternative working hypotheses of the cooperative transcriptional regulation. We believe that CoopTFD will be a valuable resource for yeast biologists to study the combinatorial regulation of gene expression controlled by cooperative TFs.

Construction and contents

Collection of predicted cooperative TF pairs from 17 existing algorithms in the literature

In yeast, many cooperative TF pairs have been predicted by various algorithms in the literature. We collected 3755 non-redundant PCTFPs from 17 existing algorithms (see Table 1 for details). However, we found that many collected PCTFPs are not really TF pairs. Therefore, we removed the PCTFPs whose proteins are not TFs. In CoopTFD, a protein is regarded as a TF if it is annotated as a TF (activator/repressor) or a transcription co-factor in the regulation page of SGD (26). After the data processing, we obtained 2622 PCTFPs among 143 TFs (see Supplementary material, Figure S1 for a distribution of numbers of PCTFPs against number of algorithms which predict a PCTFP of interest).

Construction of five types of validation information for each PCTFP

To help users judge the biological plausibility of a PCTFP, we provide five types of validation information using various data sources, all of which were downloaded in February 2016. First, the number of algorithms which predicted this PCTFP is given. A PCTFP predicted by many algorithms has a low chance to be predicted by random. Therefore, the higher the number is, the higher the statistical confidence of this PCTFP is. Second, the number of publications which experimentally show that this PCTFP has physical or genetic interactions is given. The publications were retrieved from BioGRID database (27). Having physical or genetic interactions strengthens the confidence of the biological plausibility of this PCTFP. Third, the number of publications which experimentally study the biological roles of both TFs of this PCTFP is given. The publications were retrieved from SGD database (26). If a PCTFP is of biological significance, both TFs may well be
studied in the same publication. Therefore, the higher the number is the more biological plausibility of this PCTFP is. Fourth, the common Gene Ontology (GO) terms of this PCTFP are given. The GO terms of a TF were retrieved from SGD database (26). Level of the GO term was calculated from the GO SQL file (29). If this calculation results in multiple levels, the level closest to the root is chosen. Having common GO terms provides users with strengthened evidence of the biological plausibility of this PCTFP. Finally, the common target genes of this PCTFP are provided. The target genes of a TF were retrieved from the YEASTRACT database (30). The regulatory associations between a TF and its target genes are validated by TF binding evidence, which means the experimental evidence (from band-shift, foot-printing or ChIP assay) showing that the TF binds to the promoters of its target genes. Since the biological role of a cooperative TF pair is to co-regulate the expression of a set of genes, knowing the common target genes of the two TFs of a PCTFP helps users evaluate the biological plausibility of a PCTFP.

Table 1. The list of 17 computational studies, which developed distinct algorithms to predict cooperative TF pairs by integrating multiple data sources

| Authors of the algorithms | Published year | Data sources integrated | The number of identified predicted cooperative TF pairs (PCTFPs) |
|---------------------------|---------------|-------------------------|---------------------------------------------------------------|
| Banerjee and Zhang [9]    | 2003          | ChIP-chip data and gene expression data | 31 |
| Harbison et al. [14]      | 2004          | ChIP-chip data and promoter sequence data | 94 |
| Nagamine et al. [17]      | 2005          | ChIP-chip data and PPI data | 24 |
| Tsai et al. [10]          | 2005          | ChIP-chip data and gene expression data | 18 |
| Balaji et al. [7]         | 2006          | ChIP-chip data           | 3459 |
| Chang et al. [11]         | 2006          | ChIP-chip data and gene expression data | 55 |
| He et al. [12]            | 2006          | ChIP-chip data and gene expression data | 30 |
| Wang [19]                 | 2006          | ChIP-chip data, gene expression data and promoter sequence data | 14 |
| Yu et al. [15]            | 2006          | ChIP-chip data and promoter sequence data | 300 |
| Elati et al. [6]          | 2007          | Gene expression data     | 20 |
| Datta and Zhao [8]        | 2008          | ChIP-chip data           | 25 |
| Chuang et al. [20]        | 2009          | ChIP-chip data, gene expression data and promoter sequence data | 13 |
| Wang et al. [21]          | 2009          | ChIP-chip data, gene expression data, promoter sequence data, PPI data, TF-gene documented regulation data and comparative genomic data | 159 |
| Yang et al. [18]          | 2010          | ChIP-chip data and TF knockout data | 186 |
| Chen et al. [16]          | 2012          | ChIP-chip data and promoter sequence data | 221 |
| Lai et al. [22]           | 2014          | TF-gene documented regulation data, TFBS data and nucleosome occupancy data | 27 |
| Wu and Lai [23]           | 2015          | TF-gene binding data and TF-gene regulation data | 50 |

In summary, CoopTFD provides five types of evidence (Algorithm Evidence, Physical/Genetic Interaction Evidence, Co-citation Evidence, Common GO Terms Evidence and Common Target Genes Evidence) to help users judge the biological plausibility of a PCTFP. Among them, Physical/Genetic Interaction Evidence and Common Target Genes Evidence are more informative than the other three types of evidence.

Implementation of CoopTFD website

CoopTFD was built using a scripting language PHP and CodeIgniter framework. Python was used to do raw data processing. The processed data was stored in MySQL. The graphics of cooperative TF networks are generated using Cytoscape Web (31).

Utility and discussion

Database interface

CoopTFD provides two search modes and a browse mode. In the first search mode, users can input a list of TFs of interest and specify the lowest number of algorithms that should predict a PCTFP (Figure 1). Then CoopTFD returns
Figure 2. The results of the first search mode. (a) After submission, CoopTFD returns a figure showing a cooperative TF network containing all PCTFPs among the input TFs. (b) A table is given listing five types of validation information of each PCTFP in the cooperative TF network. (c) When clicking on the number in the column of 'Algorithm Evidence', it opens a webpage showing the details of the algorithms. (d) When clicking on the number in the column of '# of common GO terms', it opens a webpage showing the names of the common GO terms. (e) When clicking on the number in the column of '# of common target genes defined by TFB', it opens a webpage showing the names of the common target genes and the numbers of the TF binding (TFB) evidence that experimentally validate the TF-target gene relationship. (f) When clicking on the number in the column of '# of TFB evidence', it opens a webpage showing the publications which provide the TFB evidence.
a figure showing a cooperative TF network containing all PCTFPs among the input TFs (Figure 2a). Moreover, a table is given listing five types of validation information of each PCTFP in the cooperative TF network (Figure 2b). The first three types are the number of algorithms which predict this PCTFP, the number of publications which experimentally show that this PCTFP has physical or genetic interactions, and the number of publications which experimentally study the biological roles of both TFs of this PCTFP. When clicking on the number, it opens a webpage showing the details (e.g. the authors, titles, journals and dates) of the publications (Figure 2c). The abstract of each publication in Pubmed can also be seen by clicking on the title of the publication. The fourth type of validation information is the number of common GO terms of this PCTFP. When clicking on the number, it opens a webpage showing the names
of the common GO terms (Figure 2d). By clicking on the names, users will be redirected to SGD database (26) to see the details of these GO terms. The last type is the number of common target genes of this PCTFP. When clicking on the number, it opens a webpage showing the names of the common target genes and the numbers of the TF binding evidence that validate the TF-target gene relationship (Figure 2e). The publications which provide the TF binding evidence can also be shown by clicking on the number (Figure 2f). In the second search mode, users can input a TF of interest and specify the lowest number of algorithms that should predict a PCTFP (Figure 3a). Then CoopTFD returns
a table listing all possible PCTFPs that are related to the input TF and satisfied the specification (Figure 3b).

In the browse mode, users can browse CoopTFD by a TF name. In total, 2622 PCTFPs among 143 TFs are deposited in CoopTFD (Figure 3c). When users click the number in the column of ‘# of PCTFPs related to the TF of interest’, our database returns a table listing five types of validation information of each PCTFP that is related to the TF of interest (Figure 3d). This is actually the same result when users select the second search mode and specify one as the lowest number of algorithms that should predict a PCTFP.

Three scenarios of using CoopTFD

Here, we introduce three scenarios of using CoopTFD. The first scenario is as follows. If researchers have a TF of interest (e.g. Mbp1) and want to know which TFs may have cooperativity with Mbp1, they can (i) select the second search function, (ii) input Mbp1 and (iii) require that each PCTFP must be predicted by at least four algorithms (Figure 3a). After submission, CoopTFD returns seven PCTFPs, suggesting that seven TFs (Ace2, Fkh1, Fkh2, Skn7, Stb1, Swi4 and Swi6) may have cooperativity with Mbp1 (Figure 3b). Among them, five PCTFPs (Mbp1-Fkh1, Mbp1-Skn7, Mbp1-Stb1, Mbp1-Swi4 and Mbp1-Swi6) are highly biologically plausible since they are supported by five types of validation information. The other two PCTFPs (Mbp1-Ace2 and Mbp1-Fkh2) are moderately biologically plausible since they are supported by four types of validation information.

The second scenario is as follows. Researchers often have a set of genes of interest (e.g. differentially expressed genes under a specific biological condition) from microarrays. They then use existing algorithms or tools to identify TFs that regulate this set of genes (32–34). If they also want to know possible PCTFPs among the identified TFs, they can use CoopTFD to do this task. For example, researchers can have a list of 17 predicted cell cycle TFs (Ace2, Ash1, Cin5, Cst6, Fkh1, Fkh2, Mbp1, Mcm1, Ndd1, Rlm1, Stb1, Ste12, Stp1, Swi4, Swi5, Swi6 and Tec1) from an existing algorithm (32). Now if they (i) select the first search function, (ii) input the list of 17 TFs and (iii) require that each PCTFP must be predicted by at least four algorithms, CoopTFD returns 34 PCTFPs (Figure 4). Among them, 18 PCTFPs are highly biologically plausible since they are supported by five types of validation information. The other 16 PCTFPs are moderately biologically plausible since they are supported by four types of validation information.

The third scenario is as follows. If researchers have already known key TFs of a specific biological process and would like to know the cooperative network of these TFs, they can use CoopTFD to do this task. Here, we use yeast oxidative stress response as an example. Yeast cells are constantly challenged by the oxidative stress which is induced by the reactive oxygen species (ROS). If not eliminated properly, ROS can damage all cellular components. Yeast cells respond to the oxidative stress by activating many genes involved in the oxidant defence mechanisms. Msn2, Msn4, Skn7 and Yap1 are key TFs which cooperatively regulate the expression the antioxidant genes (1). If users input these four TFs and require that each PCTFP must be predicted by at least two algorithms, then CoopTFD returns a densely connected cooperative TF network (consisting of six PCTFPs) among the four input TFs (Figure 2a and b). The provided cooperative TF network is likely to be biologically relevant since (i) all PCTFPs are predicted by at least two algorithms, (ii) 67% (4/6) PCTFPs have experimental evidence of having physical or genetic interactions, (iii) the two TFs of each of the six PCTFPs are studied in the same publications, (iv) all PCTFPs have common GO terms and (v) all PCTFPs have common target genes.

Conclusion

In this article, we present CoopTFD which provides 2622 predicted cooperative TF pairs (PCTFPs) among 143 yeast TFs from 17 existing algorithms. By integrating multiple data sources, we also provide five types of validation information for each PCTFP to help users judge the biological plausibility of a PCTFP. The information includes the algorithms which predict a PCTFP, the publications which experimentally show that a PCTFP has physical or genetic interactions, the publications which experimentally study the biological roles of both TFs of a PCTFP, the common GO terms of a PCTFP and the common target genes of a PCTFP. Using three scenarios, we show that CoopTFD can return biologically plausible PCTFPs of a TF or a biologically relevant cooperative network of a list of TFs.

CoopTFD has an easy-to-use interface for biologists to search or browse for the PCTFPs of the TFs of interest. CoopTFD will be regularly updated based on the newly published literature and the latest releases of the BioGRID, SGD and YEASTRACT databases. We believe that the PCTFPs deposited in CoopTFD will be a very useful resource for yeast biologists to study the combinatorial regulation of gene expression by cooperative TFs.

Supplementary Data

Supplementary data are available at Database Online.
Acknowledgements

The physical or genetic interaction data, co-citation papers, co-annotated GO terms and the co-regulatory target genes were retrieved from BioGRID, SGD and YEASTRACT on 24 February 2016. We greatly appreciate the effort of these research teams in collecting and curating such valuable data from the literature.

Funding

This work was supported by National Cheng Kung University and Ministry of Science and Technology of Taiwan [MOST-103-2221-E-006-174-MY2]. Funding for open access charge: National Cheng Kung University and Ministry of Science and Technology of Taiwan.

Conflict of interest. None declared.

References

1. Hohmann, S. and Mager, W.H. (2003) Yeast Stress Responses. Springer-Verlag, Berlin.
2. Wu, W.S. and Li, W.H. (2008) Identifying gene regulatory modules of heat shock response in yeast. BMC Genomics, 9, 439.
3. Anderson, K.R., Torres, C.A., Solomon, K. et al. (2009) Cooperative transcriptional regulation of the essential pancreatic islet gene NeuroD1 (beta2) by Nkx2.2 and neurogenin 3. Biol. Chem., 284, 31326–31324.
4. Wu, S., Muri, S., Kataoka, K. et al. (2008) Yin Yang 1 induces transcriptional activity of p73 through cooperation with E2F1. Biochem. Biophys. Res. Commun., 365, 75–81.
5. Kim, H.Y., Lee, S.B., Kang, H.S. et al. (2014) Two distinct domains of Flo8 activator mediates its role in transcriptional activation and the physical interaction with Mss11. Biochem. Biophys. Res. Commun., 449, 202–207.
6. Elati, M., Neuvial, P., Bolotin-Fukuhara, M. et al. (2007) LICORN: learning cooperative regulation networks from gene expression data. Bioinformatics, 23, 2407–2414.
7. Balaji, S., Babu, M.M., Iyer, L.M. et al. (2006) Comprehensive analysis of combinatorial regulation using the transcriptional regulatory network of yeast. J. Mol. Biol., 360, 213–227.
8. Datta, D. and Zhao, H. (2008) Statistical methods to infer cooperative binding among transcription factors in Saccharomyces cerevisiae. Bioinformatics, 24, 545–552.
9. Banerjee, N. and Zhang, M.Q. (2003) Identifying cooperativity among transcription factors controlling the cell cycle in yeast. Nucleic Acids Res., 31, 7024–7031.
10. Tsai, H.K., Lu, H.H.S. and Li, W.H. (2005) Statistical methods for identifying yeast cell cycle transcription factors. Proc. Natl. Acad. Sci. USA, 102, 13532–13537.
11. Chang, Y.H., Wang, Y.C. and Chen, B.S. (2006) Identification of transcription factor cooperativity via stochastic system model. Bioinformatics, 22, 2276–2282.
12. He, D., Zhou, D. and Zhou, Y. (2006) Identifying synergistic transcriptional factors involved in the yeast cell cycle using Microarray and ChIP-chip data. In Proceedings of the Fifth International Conference on Grid and Cooperative Computing Workshops (GCCW’06), Hunan, China, pp. 357–360.
13. Wu, W.S., Li, W.H. and Chen, B.S. (2006) Computational reconstruction of transcriptional regulatory modules of the yeast cell cycle. BMC Bioinformatics, 7, 421.
14. Harbison, C.T., Gordon, D.B., Lee, T.I. et al. (2004) Transcriptional regulatory code of a eukaryotic genome. Nature, 431, 99–104.
15. Yu, X., Lin, J., Masuda, T. et al. (2006) Genome-wide prediction and characterization of interactions between transcription factors in Saccharomyces cerevisiae. Nucleic Acids Res., 34, 917–927.
16. Chen, M.J., Chou, L.C., Hsieh, T.T. et al. (2012) De novo motif discovery facilitates identification of interactions between transcription factors in Saccharomyces cerevisiae. Bioinformatics, 28, 701–708.
17. Nagamine, N., Kawada, Y. and Sakakibara, Y. (2005) Identifying cooperative transcriptional regulations using protein-protein interactions. Nucleic Acids Res., 33, 4828–4837.
18. Yang, Y., Zhang, Z., Li, Y. et al. (2010) Identifying cooperative transcription factors by combining ChIP-chip data and knockout data. Cell Res., 20, 1276–1278.
19. Wang, J. (2006) A new framework for identifying combinatorial regulation of transcription factors: a case study of the yeast cell cycle. J. Biomed. Informatics, 40, 707–725.
20. Chuang, C.L., Hung, K., Chen, C.M. et al. (2009) Uncovering transcriptional interactions via an adaptive fuzzy logic approach. BMC Bioinformatics, 10, 400.
21. Wang, Y., Zhang, X.S. and Xia, Y. (2009) Predicting eukaryotic transcriptional cooperativity by Bayesian network integration of genome-wide data. Nucleic Acids Res., 37, 5943–5958.
22. Lai, F.J., Jhu, M.H., Chiu, C.C. et al. (2014) Identifying cooperative transcription factors in yeast using multiple data sources. BMC Syst. Biol., 8(Suppl 5), S2.
23. Wu, W.S. and Lai, F.J. (2015) Properly defining the targets of a transcription factor significantly improves the computational identification of cooperative transcription factor pairs in yeast. BMC Genomics, 16(Suppl 12), S10.
24. Lai, F.J., Chang, H.T., Huang, Y.M. et al. (2014) A comprehensive performance evaluation on the prediction results of existing cooperative transcription factors identification algorithms. BMC Syst. Biol., 8(Suppl 4), S9.
25. Lai, F.J., Chang, H.T. and Wu, W.S. (2015) PCTFPeval: a web tool for benchmarking newly developed algorithms for predicting cooperative transcription factor pairs in yeast. BMC Bioinformatics, 16(Suppl 18), S2.
26. Costanzo, M., Engel, S.R., Wong, E.D. et al. (2014) Saccharomyces Genome Database provides new regulation data. Nucleic Acids Res., 42, 717–725.
27. Charlery-Aryamothri, A., Breitkreutz, B.J., Heinicke, S. et al. (2013) The BioGRID interaction database: 2013 update. Nucleic Acids Res., 41, D816–D823.
28. Orchard, S., Ammari, M., Aranda, B. et al. (2014) The MIntAct project–IntAct as a common curation platform for 11 molecular interaction databases. Nucleic Acids Res., 42, D358–D363.
29. Ashburner, M., Ball, C.A., Blake, J.A. et al. (2000) Gene ontology: tool for the unification of biology. The Gene Ontology Consortium. Nat. Genet., 25, 25–29.
30. Teixeira, M.C., Monteiro, P.T., Guerreiro, J.F. et al. (2014) The YEASTRACT database: an upgraded information system for the analysis of gene and genomic transcription regulation in Saccharomyces cerevisiae. Nucleic Acids Res., 42, 161–166.
31. Lopes, C.T., Franz, M., Kazi, F. et al. (2010) Cytoscape Web: an interactive web-based network browser. *Bioinformatics*, 26, 2347–2348.

32. Wu, W.S. and Li, W.H. (2008) Systematic identification of yeast cell cycle transcription factors using multiple data sources. *BMC Bioinformatics*, 9, 522.

33. Wang, H., Wang, Y.H. and Wu, W.S. (2011) Yeast cell cycle transcription factors identification by variable selection criteria. *Gene*, 485, 172–176.

34. Chang, D.T., Li, W.S., Bai, Y.H. et al. (2013) YGA: identifying distinct biological features between yeast gene sets. *Gene*, 518, 26–34.