NucVoter: A Voting Algorithm for Reliable Nucleosome Prediction Using Next-Generation Sequencing Data

Boseon Byeon

Institute of Molecular Medicine and Genetics, Georgia Regents University, Augusta, GA 30912, USA

Correspondence should be addressed to Boseon Byeon; bbyeon@gmail.com

Received 17 July 2013; Accepted 8 September 2013

Academic Editors: A. Bolshoy, B. Haubold, S. Liuni, A. Pulvirenti, and C.-A. Tsai

Copyright © 2013 Boseon Byeon. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Nucleosomes, which consist of DNA wrapped around histone octamers, are dynamic, and their structure, including their location, size, and occupancy, can be transformed. Nucleosomes can regulate gene expression by controlling the DNA accessibility of proteins. Using next-generation sequencing techniques along with such laboratory methods as micrococcal nuclease digestion, predicting the genomic locations of nucleosomes is possible. However, the true locations of nucleosomes are unknown, and it is difficult to determine their exact locations using next-generation sequencing data. This paper proposes a novel voting algorithm, NucVoter, for the reliable prediction of nucleosome locations. Multiple models verify the consensus areas in which nucleosomes are placed by the model with the highest priority. NucVoter significantly improves the performance of nucleosome prediction.

1. Introduction

Genes within DNA are transcribed into an RNA product [1]. To be transcribed, the DNA region encoding a gene must be accessible to proteins such as transcription factors and RNA polymerase [2]. As shown in Figure 1, a nucleosome is composed of a DNA sequence wrapped 1.65 times around a histone octamer [3]. If the DNA region is wrapped compactly to prevent proteins from binding to the DNA, the corresponding gene is not transcribed [4]. Therefore, nucleosomes can regulate gene expression by restricting or facilitating the DNA accessibility of proteins.

Figure 2 shows the profile of typical nucleosomes around the transcription start sites (TSSs) of yeast genes. The most prevalent size of nucleosomes is approximately 147 base pairs (bp), and the length of linker DNA between nucleosomes is approximately 18bp [3]. The occupancy of a nucleosome represents the possibility that a nucleosome resides at a particular genomic location. The so-called −1 nucleosome is the first nucleosome upstream of the TSS. The area downstream of the −1 nucleosome is the nucleosome-free region (NFR) which shows very low nucleosome occupancies over approximately 150 bp on average [5]. The NFR contains transcription factor binding sites and is therefore important in transcription regulation [6]. The first nucleosome downstream of the NFR is the +1 nucleosome, followed by the +2, +3, and +4 nucleosomes. While the +1 nucleosome is stable, its upstream and downstream nucleosomes show declines in occupancy and stability and become fuzzy.

For proteins to bind the DNA region associated with compact nucleosomes to initiate transcription, the nucleosome structure needs to be changed. Interestingly, nucleosomes are dynamic, and their structure can be transformed [6, 7]. ATP-dependent remodeling can slide histone octamers a short distance along DNA or remove them temporarily from DNA. Additionally, chemical modifications of histones or histone replacement with histone variants can alter the structure of nucleosomes.

Nucleosome prediction refers to nucleosome positioning in the genome. Using MNase-seq or Chip-seq, it is possible to identify the genomic locations of nucleosomes [4, 5, 8]. Micrococcal nuclease (MNase) is a restriction enzyme that digests DNA that is not wrapped around histones. By treating cells with MNase, linker DNA is removed, and nucleosomes can be extracted (Figures 3(a) and 3(b)). Immunoprecipitation (Chip) with histone antibodies can be further employed to select nucleosomes with regard to specific histone modifications. DNA fragments are subsequently purified from
Figure 1: Organization of nucleosomes and linkers, and DNA. A nucleosome is composed of DNA wrapped around a histone octamer. H indicates a histone octamer. Nucleosomes are connected by linker DNA. DNA is double stranded; the forward strand is in the 5' to 3' direction, while the reverse strand is in the opposite direction.

Figure 2: Profile of typical nucleosomes around TSSs of yeast genes.

Figure 3: Nucleosome prediction process. (a) Nucleosomes connected by linker DNA. (b) Nucleosomes following MNase digestion. (c) Purified DNA fragments. Arrowed marks indicate the 5' ends, which are sequenced via NGS. (d) Short sequence tags generated by NGS. (e) Distribution of tags mapped to the reference genome. F and R indicate the forward and reverse strands, respectively. (f) Tag smoothing and nucleosome prediction. Ellipses represent predicted nucleosomes. (g) Example of tag distributions, which are not well separated.

2. Methodology

2.1. Nucleosome Prediction Methods. GeneTrack shifts sequence tags toward the 3' direction by half of the user-defined nucleosome size, smooths the tag distributions using a Gaussian smoothing procedure, and then positions nucleosomes by setting the local maxima to the centers of nucleosome [15]. Nucleosomes can be defined separately on the forward and reverse strands or on the composite strand derived from the sum of the tag frequencies on the forward and reverse strands.

NSeq generates probabilistic distributions of nucleosome centers based on sequence tags and determines significant nucleosomes using triangle statistics, N statistics, and false discovery rates [16]. Nucleosomes predicted by NSeq also have a predetermined size.

TemplateFilter uses templates representing diverse patterns of distributions [17]. TemplateFilter identifies the genomic locations where its templates, including a normal-shaped template, correlate with tag distributions. Then, the locations of optimal templates correlated with the forward and reverse tag distributions determine the left and right boundaries of nucleosomes, respectively. Therefore, TemplateFilter defines nucleosomes of various sizes differently from GeneTrack and NSeq.

2.2. Voting Algorithm. In machine learning, one approach for finding a reliable solution to a difficult problem is the ensemble method, which combines the outcomes of different models [18]. The simplest ensemble technique is voting, in which multiple models take votes, and the majority outcome is adopted as the solution.
In NucVoter, a nucleosome prediction method is regarded as a voter, and the three voters described above locate nucleosomes across the genome. If two or more of the voters position nucleosomes around a genomic location, NucVoter defines those nucleosomes as consensus nucleosomes and their nucleosomal locations as consensus areas where true nucleosomes are likely to reside (Figure 4). Then, NucVoter assigns a priority to each voter as described below and chooses the nucleosome predicted by the voter with the highest priority in each consensus area. If only one voter places a nucleosome in a particular genomic location, NucVoter concludes that the site does not contain a nucleosome.

In a preliminary process, NucVoter normalizes the occupancies of nucleosomes from each voter so that the values show a zero mean and 1 standard deviation. Then, NucVoter further makes the entire occupancies positive by subtracting the global minimum occupancy, which is a negative value, from the normalized occupancies.

To establish the priority of each voter, NucVoter uses the consensus nucleosomes receiving the consent of all three voters. Then, it is hypothesized that if the consensus nucleosomes in a consensus area are nearer, they display higher occupancies than those in other consensus areas, meaning that if voters predict nucleosomes within a short distance of each other, the possibility that a nucleosome resides in that region is high. The consensus nucleosomes with the consent of all three voters are extracted, and their averaged center distance is computed in each consensus area. Then, the correlation between the occupancies of the consensus nucleosomes of each voter and their corresponding averaged center distances is calculated. NucVoter regards the voter showing the higher negative correlation as more accurate and assigns the higher priority to that voter.

Figure 4 depicts the consensus nucleosomes and areas. In NucVoter, the distances between the centers of the consensus nucleosomes in a consensus area should be ≤73 bp (half of the prevalent nucleosome size). The two globally nearest nucleosomes predicted by two different voters create a consensus area, and the third consensus nucleosome predicted by the other voter is added so that the distances between the centers of all consensus nucleosomes are minimized. This process is iterated among the remaining nucleosomes until no more consensus nucleosomes exist. Then, NucVoter determines the priority of each voter using the consensus nucleosomes receiving the consent of all three voters, as described above, and chooses the nucleosome predicted by the voter with the highest priority in each consensus area where two or more voters consented.

When two or more datasets need to be compared (see Section 3.2), NucVoter sets the global voting priorities on the basis of the averaged correlation of each voter across the datasets. Then, the global priorities are used to choose nucleosomes consistently in all datasets.

2.3. Software Availability. NucVoter is available on request.

3. Results

3.1. Synthetic Data. Because the exact locations of nucleosomes are unknown, synthetic datasets were generated to measure the performance of NucVoter. The data generation procedure was carried out in a manner similar to the function of "syntheticNucMap" embedded in the R package of nucleR [19].

A total of 1000 stable nucleosomes were generated periodically on the basis of a nucleosome size of 146 bp and a linker length of 20 bp, and 50 nucleosomes were randomly removed. Then, the F number of fuzzy nucleosomes was added at random. The forward/reverse tags for both stable and fuzzy nucleosomes were randomly generated in the range of ±1 to the C coverage (i.e., the number of tags) at the starting/ending locations of nucleosomes. Then, stable tags were randomly shifted in the range of +/-20 bp and fuzzy tags in the range of +/-50 bp. Finally, 49000 true nucleosomes in 40 synthetic datasets were generated based on the combination of various numbers of fuzzy nucleosomes (F: 50 to 500 with an interval of 50) and different coverage (C: 50 to 200 with an interval of 50). Note that most of the parameter values used for this process were the default values of the "syntheticNucMap" function.

GeneTrack, with a nucleosome size 146 bp on the composite strand, as well as NSeq and TemplateFilter, was initially executed using their default parameter values to predict synthetic nucleosomes. However, NSeq and TemplateFilter positioned too few nucleosomes. Therefore, to generate a reasonable number of nucleosomes, the "-f 1" option was used for NSeq, and the options "-overlap 1.0" and "-corr_bound 0.3" were used for TemplateFilter. Note that the parameter values were not optimized because the goal of implementation was not to compare the performance of existing methods.
but to examine how the proposed voting algorithm improves the prediction capability given the outputs of individual methods.

Table 1 shows the prediction of true nucleosomes in the synthetic datasets. If the distance between the centers of a true nucleosome and the nearest predicted nucleosome is ≤73 bp, the true nucleosome is regarded as correctly predicted. The true positives indicate the correctly positioned true nucleosomes, and the false positives represent the incorrectly positioned false nucleosomes. The false negatives are the missed true nucleosomes. Although TemplateFilter predicted the greatest number of true positives, it also positioned many false negatives. NSeq placed the least false positives but missed many true positives. NucVoter and GeneTrack showed reasonably good performance.

Using the true positives, the cumulative frequency ratio was plotted as a function of the distance between the centers of the true and predicted nucleosomes (Figure 5(a)). While NucVoter and NSeq predicted nucleosomes accurately, GeneTrack and TemplateFilter did so less accurately. However, these results might have been produced because different true positives nearest to the true nucleosomes was extracted from each dataset, and the plot of their cumulative frequency was drawn (Figure 5(b)). As expected, the performances of GeneTrack and TemplateFilter were higher. Furthermore, NucVoter outperformed the other methods significantly (P values of the Kolmogorov-Smirnov test < 2.2e −16). The centers of 90% of the NucVoter nucleosomes were within 5 bp from the centers of true nucleosomes. This result supports the hypothesis that the proposed voting algorithm improves the accuracy of nucleosome prediction.

To further analyze the prediction improvement achieved by NucVoter, linker analysis was performed. Figure 6 shows the linker length frequencies obtained. As noted above, the length of true linkers in the synthetic data was 20 bp, and the plot therefore displays a particularly high frequency peak at 20 bp (Figure 6(a)). Additionally, because 5% of nucleosomes were randomly removed, there was a high frequency peak at 186 bp (i.e., 146 bp + 20 bp + 20 bp). Although there were various linker lengths of less than 186 bp observed due to the presence of fuzzy nucleosomes, there were no linkers longer than 186 bp. In the region below 200 bp, NucVoter and NSeq produced frequencies that were more similar to the real linker frequency compared to GeneTrack and TemplateFilter (Figures 6(b) to 6(e)), which means that NucVoter and NSeq predicted nucleosomes more accurately. However, NSeq and TemplateFilter exhibited relatively high
Figure 6: Frequency of linker lengths. (a) Frequency of the lengths of true linkers. Frequencies of the lengths of linkers predicted by (b) NucVoter, (c) GeneTrack, (d) NSeq, and (e) TemplateFilter.

Table 2: Accuracy of the nucleosomal and linker locations predicted by the indicated methods. The number in parentheses is the \( P \) value of the paired \( t \)-test, where the alternative hypothesis is that the accuracy of NucVoter is greater than that of the indicated method.

| Method         | Sensitivity      | Specificity       | Overall accuracy |
|----------------|------------------|-------------------|------------------|
| NucVoter       | 94.25%           | 94.00%            | 94.22%           |
| GeneTrack      | 92.95% (5.25e-09)| 75.34% (<2.2e-16)| 90.73% (<2.2e-16)|
| NSeq           | 87.89% (6.51e-12)| 94.61% (1.0)     | 88.73% (1.117e-11)|
| TemplateFilter | 84.12% (<2.2e-16)| 79.58% (<2.2e-16)| 83.55% (<2.2e-16)|

To measure the overall performance of NucVoter, individual locations in synthetic genomes were scanned, and a confusion matrix of the locations of nucleosomal and linker DNA was generated [18, 20]. True positives (TP) and true negatives (TN) represent correct predictions of nucleosomal and linker locations, respectively. While a false positive (FP) occurs when a linker location is incorrectly predicted as a nucleosomal location, a false negative (FN) is observed when a nucleosomal location is incorrectly predicted as a linker location. The sensitivity is TP divided by TP + FN and measures the ability to correctly predict true nucleosomal locations. The specificity, which is TN divided by TN + FP, measures the ability to correctly predict true linker locations. The overall accuracy is defined as the number of correct predictions divided by the total number of predictions (i.e., TP + TN over TP + TN + FP + FN). As observed in Table 2, NucVoter achieved the best performance in terms of both sensitivity and overall accuracy. Regarding specificity, NucVoter displayed a performance similar to that of NSeq, which exhibited the best performance. These results confirm that the proposed voting algorithm significantly improves the nucleosome prediction capability.

3.2. Real Data. NucVoter was applied to published MNase-seq datasets generated from normal and heat-shocked yeast cells [14]. To achieve consistent voting between the two datasets, the global priorities for nucleosome selection were set in the order of TemplateFilter, NSeq, and GeneTrack on the basis of the averaged correlation values across the two datasets, as described above. To make the occupancy levels of two datasets comparable, the occupancy values of

frequencies, approximately 352 bp and in the area of 500 bp or longer (Figures 6(d) and 6(e)), which indicated missing consecutive true nucleosomes.
nucleosomes predicted by NucVoter were further normalized via the quantile method.

Additionally, GeneTrack and NSeq were executed with the “-f 1” option, and TemplateFilter was run with its default options to generate a reasonable number of nucleosomes. Figure 7 shows the similarity of the predicted nucleosomes to published nucleosomes. Assuming the published nucleosomes to be true nucleosomes, if the distance between the centers of a published nucleosome and the nearest predicted nucleosome was ≤ d, the published nucleosome was regarded as correctly predicted. Then, the similarity was defined as the number of correctly predicted nucleosomes divided by the total number of predicted nucleosomes (i.e., TP over TP + FP + FN). Refer to Table 1 description for the definitions of TP, FP, and FN. Interestingly the nucleosomes predicted by NucVoter showed the highest similarity to the published nucleosomes overall.

Figure 8 depicts the averaged frequency of nucleosomes predicted by NucVoter as a function of the distance from TSSs. The number of nucleosomes in the heat-shocked cells is greater than the number in normal cells, which indicates that nucleosomes are gained under heat shock. While the frequencies of the nucleosomes downstream of the +1 nucleosome decrease slightly, the numbers of the nucleosomes upstream of the −1 nucleosome increase to a small degree.

Figure 9 is the averaged occupancy plot for the nucleosomes predicted by NucVoter as a function of the distance from TSSs. It can be observed that the occupancy of nucleosomes in heat-shocked cells is higher than in normal cells. Note that this is identical to the published result [14]. The +1 nucleosome displays strong occupancy, and its downstream nucleosomes show substantially decreased occupancy. Although the −1 nucleosome exhibits a lower frequency than its upstream nucleosomes, as shown in Figure 8, its occupancy is higher. Additionally, the length of the NFR is slightly shorter in the heat-shocked compared to the normal cells.

Figures 10(a) and 10(b) provide plots of the frequencies of the lengths of linkers predicted by NucVoter in the range of +/−1000 bp from the TSSs of genes. The linkers are shorter overall in the heat-shocked cells than in the normal cells, and the small peak observed at approximately 180 bp in the normal cells, which reflects the NFR, disappears following heat shock.

To analyze the NFRs predicted by NucVoter, the longest linkers in the range of TSS − 250 bp to TSS + 50 bp were defined as NFRs, and the 264 genes whose NFR lengths were reduced by ≥150 bp following heat shock were extracted. Figure 11 provides the nucleosome profile of these genes aligned based on their TSSs. Then, Gene Ontology (GO) analysis of the genes was performed using FunSpec [21, 22]. GO provides annotation of genes based on biological knowledge [2]. Table 3 describes the GO biological processes of the 264 genes. A total of 29 and 28 genes significantly contribute to translation and metabolic processes, respectively. This result supports the notion that the NFR plays an important role in regulating gene expression.

4. Conclusions

Although nucleosomes are critical for regulating gene expression, the prediction of nucleosome locations is difficult. This paper proposed the NucVoter algorithm, which is a novel
Figure 8: Averaged frequency of the nucleosomes predicted by NucVoter as a function of the distance from the TSSs of yeast genes.

Figure 9: Averaged occupancy of the nucleosomes predicted by NucVoter as a function of the distance from the TSSs of yeast genes.

Figure 10: Relative frequency of the lengths of linkers predicted by NucVoter in the range of +/-1000 bp from the TSSs of yeast genes. (a) Linkers from normal cells. (b) Linkers from heat-shocked cells.
The plot was smoothed by the Lowess method [2].

Contributed to translation and metabolic processes. NFR lengths were considerably reduced following heat shock on analysis, it was further noted that genes whose heat-shocked cells compared to the normal cells. Based in normal cells. Additionally, the linkers became shorter in nucleosomes in the heat-shocked cells was higher than observed to be gained under heat shock, and the occupancy to various nucleosome analyses. Many nucleosomes were significantly improved the performance of nucleosome prediction. NucVoter achieved the best prediction accuracy and significantly improved the accuracy of the other methods. Furthermore, it was confirmed that linkers in a manner that was more similar to actual linkers than most in this analysis. In addition, NucVoter produced linkers in linkers in true nucleosomes than any other method examined in this analysis. In addition, NucVoter produced linkers in a manner that was more similar to actual linkers than most of the other methods. Furthermore, it was confirmed that NucVoter achieved the best prediction accuracy and significantly improved the performance of nucleosome prediction.

Using published datasets from normal and heat-shocked yeast cells, it was shown that NucVoter could be applied to various nucleosome analyses. Many nucleosomes were observed to be gained under heat shock, and the occupancy of nucleosomes in the heat-shocked cells was higher than in normal cells. Additionally, the linkers became shorter in the heat-shocked cells compared to the normal cells. Based on GO analysis, it was further noted that genes whose NFR lengths were considerably reduced following heat shock contributed to translation and metabolic processes.

**Conflict of Interests**

The author declares that there is no conflict of interests regarding the publication of this paper.

**Table 3: GO analysis of the 264 genes whose NFR length was reduced by ≥150 bp following heat shock.**

| Biological process                                           | P value     | Number of genes |
|-------------------------------------------------------------|-------------|-----------------|
| Translation [GO:0006412]                                     | 2.31e−05    | 29              |
| Nucleobase, nucleoside, nucleotide, and nucleic acid transport [GO:0015931] | 0.000660    | 4               |
| Tryptophan transport [GO:0015827]                            | 0.001593    | 2               |
| Ergosterol biosynthetic process [GO:0006696]                 | 0.001830    | 5               |
| Steroid biosynthetic process [GO:0006694]                    | 0.002706    | 5               |
| Metabolic process [GO:0008152]                               | 0.005827    | 28              |
| Nucleobase transport [GO:0015851]                            | 0.006152    | 3               |
| Group I intron splicing [GO:0000372]                         | 0.006152    | 3               |

**References**

[1] J. C. Rajapakse and S. L. Ho, “Markov/neural model for Eukaryotic promoter recognition,” in *Machine Learning in Bioinformatics*, Y. Zhang and J. C. Rajapakse, Eds., pp. 283–299, John Wiley & Sons, New York, NY, USA, 2009.

[2] S. Draghici, *Statistics and Data Analysis for Microarrays Using R and Bioconductor*, CRC Press, New York, NY, USA, 2012.

[3] C. Jiang and B. F. Pugh, “Nucleosome positioning and gene regulation: advances through genomics,” *Nature Reviews Genetics*, vol. 10, no. 3, pp. 161–172, 2009.

[4] F. Zambelli and G. Pavesi, “Algorithmic issues in the analysis of Chip-seq data,” in *Algorithms in Computational Molecular Biology*, M. Elloumi and A. Y. Zomaya, Eds., pp. 425–448, John Wiley & Sons, New York, NY, USA, 2011.

[5] L. Bai and A. V. Morozov, “Gene regulation by nucleosome positioning,” *Trends in Genetics*, vol. 26, no. 11, pp. 476–483, 2010.

[6] O. Bell, V. K. Tiwari, N. H. Thomá, and D. Schübeler, “Determinants and dynamics of genome accessibility,” *Nature Reviews Genetics*, vol. 12, no. 8, pp. 554–564, 2011.

[7] G. Felsenfeld and M. Groudine, “Controlling the double helix,” *Nature*, vol. 421, no. 6921, pp. 448–453, 2003.

[8] K. Cui and K. Zhao, “Genome-wide approaches to determining nucleosome occupancy in metazoans using MNase-Seq,” *Methods in Molecular Biology*, vol. 833, pp. 413–419, 2012.

[9] W. J. Ansorge, “Next-generation DNA sequencing techniques,” *New Biotechnology*, vol. 25, no. 4, pp. 195–203, 2009.

[10] X. Zhou, L. Ren, Q. Meng, Y. Li, Y. Yu, and J. Yu, “The next-generation sequencing technology and application,” *Protein and Cell*, vol. 1, no. 6, pp. 520–536, 2010.

[11] M. L. Eaton, K. Galani, S. Kang, S. P. Bell, and D. M. MacAlpine, “Conserved nucleosome positioning defines replication origins,” *Genes and Development*, vol. 24, no. 8, pp. 748–753, 2010.

[12] R. M. Fraser, D. Keszenman-Pereyra, M. W. Simmen, and J. Allan, “High-resolution mapping of sequence-directed nucleosome positioning on genomic DNA,” *Journal of Molecular Biology*, vol. 390, no. 2, pp. 292–305, 2009.

[13] N. Ponts, E. Y. Harris, J. Prudhomme et al., “Nucleosome landscape and control of transcription in the human malaria parasite,” *Genome Research*, vol. 20, no. 2, pp. 228–238, 2010.

[14] S. Shivaswamy, A. Bhinge, Y. Zhao, S. Jones, M. Hirist, and V. R. Iyer, “Dynamic remodeling of individual nucleosomes across a eukaryotic genome in response to transcriptional perturbation,” *PloS Biology*, vol. 6, no. 3, article e65, 2008.

[15] I. Albert, S. Wachi, C. Jiang, and B. F. Pugh, “GeneTrack—a genomic data processing and visualization framework,” *Bioinformatics*, vol. 24, no. 10, pp. 1305–1306, 2008.
[16] A. Nellore, K. Bobkov, E. Howe, A. Pankov, A. Diaz, and J. S. Song, "NSeq: a multithreaded Java application for finding positioned nucleosomes from sequencing data," *Frontiers in Genetics*, vol. 3, article 320, 2013.

[17] A. Weiner, A. Hughes, M. Yassour, O. J. Rando, and N. Friedman, "High-resolution nucleosome mapping reveals transcription-dependent promoter packaging," *Genome Research*, vol. 20, no. 1, pp. 90–100, 2010.

[18] I. H. Witten and E. Frank, *Data Mining*, Morgan Kaufmann Publishers, 2005.

[19] O. Flores and M. Orozco, "nucleR: a package for non-parametric nucleosome positioning," *Bioinformatics*, vol. 27, no. 15, Article ID btr345, pp. 2149–2150, 2011.

[20] J. Pevsner, *Bioinformatics and Functional Genomics*, John Wiley & Sons, New York, NY, USA, 2009.

[21] D. W. Huang, B. T. Sherman, and R. A. Lempicki, "Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists," *Nucleic Acids Research*, vol. 37, no. 1, pp. 1–13, 2009.

[22] M. D. Robinson, J. Grigull, N. Mohammad, and T. R. Hughes, "FunSpec: a web-based cluster interpreter for yeast," *BMC Bioinformatics*, vol. 3, article 35, 2002.