PAIR: the predicted Arabidopsis interactome resource

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ABSTRACT

The predicted Arabidopsis interactome resource (PAIR, http://www.cls.zju.edu.cn/pair/), comprised of 5990 experimentally reported molecular interactions in Arabidopsis thaliana together with 145494 predicted interactions, is currently the most comprehensive data set of the Arabidopsis interactome with high reliability. PAIR predicts interactions by a fine-tuned support vector machine model that integrates indirect evidences for interaction, such as gene co-expressions, domain interactions, shared GO annotations, co-localizations, phylogenetic profile similarities and homologous interactions in other organisms (interologs). These predictions were expected to cover 24% of the entire Arabidopsis interactome, and their reliability was estimated to be 44%. Two independent example data sets were used to rigorously validate the prediction accuracy. PAIR features a user-friendly query interface, providing rich annotation on the relationships between two proteins. A graphical interaction network browser has also been integrated into the PAIR web interface to facilitate mining of specific pathways.

INTRODUCTION

Protein–protein interactions (PPIs) are the major components of many fundamental cellular processes. Identification of the interactions involving a protein is often a key step toward understanding its functions in the cellular context. Many efforts have been made to chart PPI maps in several model organisms. In Saccharomyces cerevisiae (1,2), Drosophila melanogaster (3), Caenorhabditis elegans (4) and Homo sapiens (5,6), genome-wide yeast two-hybrid screens and large-scale affinity purification/mass spectrometry studies have been conducted to map their interactomes. Meanwhile, a number of databases, such as IntAct (7), BioGRID (8), BIND (9) have been established as repositories for interaction data. The STRING database (10) collected a large set of known and predicted protein interactions. The quality of experimentally reported interactions has been rigorously assessed (11). However, to date, no large-scale experiment aiming to map a plant interactome has been reported (12). The number of plant PPIs in these databases remains very limited.

In the plant kingdom, Arabidopsis thaliana is arguably the most important model organism. Even for this best-studied model, less than 6000 experimentally reported interactions can be found in major data repositories. Therefore, the need for predicted interactions has been recognized by several groups and led to a series of efforts to predict the Arabidopsis interactome. Geisler-Lee et al. (13) predicted approximately 20 000 Arabidopsis interactions from homologous interactions in other species (the ‘interolog’ approach). A recent study went one step further to use functional association data to improve prediction reliability, which resulted in approximately 18 000 filtered predictions (14). Though useful, these interolog-based approaches are limited to detecting evolutionarily conserved protein interactions, whereas a significant number of A. thaliana proteins do not have orthologs in other model organisms with rich interactome information. Another work, the AtPID database (15), predicted approximately 23 000 interactions from multiple indirect evidences using a Naïve Bayesian approach. This work represented a conceptual advance in interaction prediction, yet the number of predicted interactions seemed to be too small to represent a comprehensive interactome. In addition, the AtPIN database (16) integrated experimentally reported interactions from the major data repositories and the predicted interactions from the Geisler-Lee data set and the AtPID database.

However, in all the prediction efforts, accuracies were not rigorously assessed with external benchmark data sets, nor could they give a reasonable estimation of the Arabidopsis interactome size. It has been suggested that the yeast interactome includes approximately 18 000 PPIs involving approximately 6000 genes (17). Assuming the same rate of interaction between genes, approximately 200 000 Arabidopsis PPIs would be expected between the...
approximately 20,000 Arabidopsis genes. Therefore, the number of experimentally reported interactions and the sizes of available predicted interactomes are approximately an order of magnitude less than the expected size of Arabidopsis interactome.

Here, we present the predicted Arabidopsis interactome resource (PAIR), which contains the most comprehensive data set of the Arabidopsis interactome to date. These interactions are expected to cover 24% of the entire Arabidopsis interactome with a reasonably high reliability (confidence of each predicted interaction) of 44%. PAIR features an information-rich and user-friendly interface and an integrated, graphical interaction network browser to facilitate mining of specific pathways.

**THE DATABASE**

**Overview**

The PAIR project started as a simple effort to infer Arabidopsis interactions by homology mapping. In 3 years, it gradually evolved into a dedicated effort aiming to provide the most accurate interactome predicted by the state-of-the-art machine learning approach. The current version of PAIR (V3.3) contains 5990 experimentally reported molecular interactions together with 145,494 predicted interactions in A. thaliana.

The 5990 experimentally reported interactions were collected from three interaction repositories, i.e., IntAct (7), BioGrid (8) and BIND (9), as of 23 July 2010, and the interaction data set compiled by TAIR curators (18). All these major interaction data repositories manually curate data from literature. However, the Arabidopsis interactions collected by these repositories showed a small overlap (19). In our compilation, only 131 interactions were shared by all repositories. Therefore, integration of their data is necessary and helpful. In this effort, PPIs in these repositories with experimental evidence were extracted. Protein identifiers in respective repositories were mapped to the Arabidopsis gene loci according to the conversion tables provided by TAIR (Version 9) (18). Interactions involving proteins that could not be unambiguously mapped were discarded. This resulted in 5990 experimentally reported PPIs involving 2824 proteins.

The 145,494 interaction predictions were made at an earlier time, 1 February 2010, as part of the PAIR V3.0 major release. Due to the heavy computational requirements, interaction predictions were only updated with each major data release. These interactions were predicted by a support vector machine (SVM) model that integrates several indirect evidences for interaction, such as gene co-expressions, domain interactions, shared GO annotations, co-localizations, phylogenetic profile similarities and interologs (20). The SVM model was trained using a set of example interactions known as the Gold Standard Positives (GSPs), which is a collection of interactions in the major repositories as of 15 June 2009. The prediction accuracy was validated using two external benchmark data sets, containing interaction examples that were not available at the time when we trained our prediction model (detailed below). The algorithmic details on how the indirect evidences were computed, how the prediction models were trained, and how the prediction accuracies were evaluated can be found in the Help/FAQ page of the PAIR website. Altogether, 145,494 interactions involving 9480 proteins were predicted by the PAIR V3 prediction model. These predicted interactions were expected to cover 24% of the entire Arabidopsis interactome, and the reliability of each predicted interaction was estimated to be 44%. These predicted interactions had 1584 (26.44%) overlap with the 5990 experimentally reported interactions mentioned above.

Altogether, the PAIR 3.3 release contains 149,900 interactions involving 10,380 proteins. They can be queried through a user-friendly web interface, downloaded in a number of widely-used data formats or mined with a graphical interaction network browser integrated within the PAIR website.

**The quality of predicted interactions**

Two external benchmark data sets were used to verify the accuracy of predicted interactions. Before showing the assessment results, it needs to be clarified that the interaction data set searchable at the PAIR website (PAIR V3.3) is essentially a compilation including the PAIR V3 predictions and the experimentally reported interactions deposited in the major interaction databases before 23 July 2010 (update V3.3). In the accuracy assessments below, only the PAIR V3 predictions without the additional experimentally-reported interactions were evaluated.

The first benchmark data set contained newly reported interactions that were not included in the major interaction databases at the time (15 June 2009) our positive interaction examples were assembled. This independent evaluation set was retrieved from an update of the BioGRID database (8) (as of 27 December 2009), which included 448 new interactions that had been double-checked to avoid any overlap with our GSPs used in model training. As shown in Table 1, 115 (26%) of these new interactions were successfully recognized by our prediction model. This sensitivity (26%) was comparable to the expected sensitivity (24%). In contrast, these new interactions showed much less overlap with other predicted interactomes. Only 50 interactions could be found in the Geisler-Lee data set (the Interologs data set) (13), 20 in the De Bodt data set (the Filtered Interologs data set) (14), 16 in the AtPIN database and 57 in the predicted interactions in the AtPIN database (16). Detailed results are provided in Supplementary Table S1.

The other benchmark data set was reported by a recent (April 2010) article in Plant Cell (21), published 2 months after PAIR V3 was released. In this report, 917 protein pairs involving 58 core cell-cycle proteins were tested by two complementary interaction assays, bimolecular fluorescence complementation and high-confidence yeast-two-hybrid, resulting in 357 interactions, of which 293 had not been reported before (21). Among the 357 reported interactions, PAIR predicted 170 (48%) of them. Among the 293 newly reported interactions, 140 (48%) were correctly predicted. Again, in this data set, PAIR predicted many
more interactions than other predicted interactomes. As shown in Table 1, the sensitivity of PAIR V3 was over four times higher than others. Detailed results are provided in Supplementary Table S2. On the other hand, PAIR predicted 338 interactions from the 917 experimentally tested protein pairs, of which 170 were confirmed. Therefore, the reliability of PAIR predictions reached 50%. In other words, over half of the predicted interactions were real in this test. However, it should be noted that the coverage and reliability observed with this data set (48/50%) were higher than our estimations (24/44%). This might have happened because the proteins tested in this experiment were all well-studied core cell-cycle proteins. Well-studied proteins tend to have more comprehensive and accurate supporting data to compute the indirect evidences based on which our predictions are made. Therefore, the high coverage and reliability observed with this data set may not apply to the entire predicted interactome. Even so, these results showed that with the ever-growing volume of protein characteristics data, PAIR has the potential to predict the Arabidopsis interactome at higher levels of coverage and reliability.

It is also worth noting that the size, estimated coverage and reliability of the predicted Arabidopsis interactome implied an estimation of the Arabidopsis interactome size, $\sim 2.58 \times 10^5$. In other words, 1 out of 893 random protein pairs was expected to interact. This ratio is similar to the experimentally observed ratio in yeast (1/775) (17). Considering a smaller fraction of the genome is usually expressed at the same time in higher organisms as compared to unicellular species, this estimated ratio of interacting protein pairs seemed to make sense. Details of the above results are provided at the PAIR website.

In addition, a recent Plant Cell article by another group (19) showed that the previous version of PAIR (PAIR V2) already had the highest coverage among all available predicted interactomes. Using the newly curated interactions in the IntAct database (7) as a benchmark, this article reported that the coverage of PAIR V2 predictions was more than double the coverage of the second-best predicted interactome (19 versus 9%).

The high coverage of our predicted interactions is supported by multiple assessments. However, the high reliability of these predicted interactions is not validated with external experimental data. This is because that most negative results, pairs of proteins that do not interact, are never reported in literature. Consequently there is no reliable data source of non-interactions that is large enough to support an accurate estimation of our prediction reliability. But given that the estimated coverage and reliability led to a reasonable estimation of the interactome size, the estimated reliability (44%) should be roughly accurate.

### THE WEB INTERFACE

#### Search methods

In PAIR, interactions can be searched by specifying one or both of their component proteins or by specifying a homologous interaction (with both component proteins) in one of four other model organisms: *H. sapiens*, *S. cerevisiae*, *C. elegans* and *D. melanogaster*. Proteins may be specified by their identifiers, such as AGI codes (gene loci), UniProt accessions and RefSeq identifiers, or by keywords in their annotation texts. Alternatively, users can perform a BLAST sequence search to retrieve interactions involving a particular gene family or protein domain. In addition, PAIR supports gene set search, which allows a user to enter a number of AGI codes (gene loci). According to user option, PAIR can return interactions between the specified proteins or all interactions involving the specified proteins. This function is most useful to extract interaction sub-networks related to a specific cellular process.

#### Interaction information

For every PPI, PAIR provides rich annotation on the relationships between the two proteins involved. Taking the interaction between DMC1 (At3g22880) and RAD51 (At5g20850) as an example, we show in Figure 1a typical Interaction Information page. This page contains three sections. The first section shows a summary of the interaction and its component proteins (Figure 1a). If the interaction has been experimentally reported, the related experimental evidences will be shown. The second section shows the indirect evidences supporting this interaction, including gene co-expressions, domain interactions, shared GO annotations, co-localizations, phylogenetic profile similarities and homologous interactions in other organisms (interologs) (Figure 1b). For domain interactions, domains in both proteins are retrieved from Pfam (22). Known interactions between the domains were collected from the DOMINE database (23), which contains multiple domain interaction data sets reported by different approaches. The homologous interactions in four

| Table 1. Assessments of prediction accuracy using independent benchmark data sets |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|
| Independent example data set    | PAIR$^a$        | AtPID$^b$       | Interologs$^c$  | Filtered Interologs$^d$ |
| New interactions in BioGrid (448)$^f$ | 115             | 16              | 50              | 20              | 57              |
| All cell-cycle protein interactions (357)$^g$ | 170             | 22              | 27              | 11              | 37              |
| New cell-cycle protein interactions (293)$^h$ | 140             | 16              | 16              | 2               | 26              |

$^a$The PAIR V3 predicted interactions.
$^b$The predicted interactions in the AtPID database (15).
$^c$The homologous interactions in the AtPID database (15).
$^d$The filtered interolog-based predicted interactions by De Bodt et al. (14).
$^e$The interactions reported in a recent Plant Cell article by another group (19) based on homology to known interactions in other organisms (interologs).
$^f$The newly curated interactions in the BioGRID database (8).
$^g$The 375 interactions between 58 cell-cycle proteins reported in a recent Plant Cell article (21).
$^h$The 293 interactions reported in (21) that had not been reported before.

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well-studied organisms (human, fruit fly, yeast and worm) are listed in the Homologous Interactions table. The sub-cellular localizations of Arabidopsis proteins were collected from the SUBA database (24), which also contains multiple sub-cellular localization data sets reported by different approaches. The co-localization table summarizes the shared localizations in each data set, which are distinctly colored for clarity. In the GO annotations table, the relationship between the annotation terms of two proteins that share a significant parent term is present in a graphical manner. The expression profiles for two proteins are illustrated by charts in the co-expression table, so users may develop an intuitive idea on how they correlate under different perturbations (e.g. light, development or abiostress). Co-publication (two interacting proteins appearing in the same articles) information was not used for interaction prediction, but is also presented here to facilitate literature mining. (c) The confidence scores for this interaction. The SVM score is the overall confidence score.
positive prediction. However, for experimentally reported interactions, this score may be negative as our prediction model may not correctly predict all real interactions. The confidence score derived from each indirect evidence is also presented. A detailed account of these scores is provided in the Help/FAQ page.

**Graphical interaction network browser and ‘My collection’**

Each PAIR web page showing interactions includes an integrated interaction network browser, developed from the Adobe Flash™-based program Cytoscape Web (http://cytoscapeweb.cytoscape.org). Figure 2 shows an example network graph obtained by searching PAIR with a set of genes in the sulfur metabolism pathway. The query proteins are displayed as triangle nodes. Other proteins are shown as circle nodes. All nodes are colored by their molecular function annotations. The color scheme is provided in the Help/FAQ page. Interactions are presented as edges between nodes. Experimentally reported interactions are in red. Predicted interactions that are homologous to known interactions in other organisms are in blue. Other predicted interactions are colored grey. The layout of the

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**Figure 2.** The graphical interaction network browser showing a network of interactions between proteins in the sulfur metabolism pathway. Several core sulfur metabolism proteins (triangle nodes) are used to search PAIR for interactions between them. This network graph is then expanded to include additional interactions. Double click on a node will bring up a window showing all interactions involving this protein and allow selection of some interactions to be added into this graph. Newly added proteins are shown as circle nodes. Nodes are colored according to their protein functions, as detailed in the Help/FAQ page. Experimentally reported interactions (edges) are in red. Predicted interactions that are homologous to known interactions in other organisms are in blue. Other predicted interactions are in grey.
network graph can be changed by applying a number of layout algorithms or by simply dragging the nodes. Double clicking on a node will bring up a window showing all interactions involving this protein. Users can select some of these interactions to be added into the network graph. Once a desired network has been created, it is possible to save it as an image or export it in various data formats, including the Microsoft Excel, Cytoscape SIF, GraphML, PSI 2.5 XML and PAIR XML formats.

We also support the use of ‘My collection’ where interactions of interest may be stored as browser cookies and later retrieved in the ‘My collection’ page. This feature will facilitate mining of specific biological pathways. In many PAIR web pages, there is a button to add selected interactions to ‘My collection’. Right-clicking on an edge in the interaction network browser will also bring up an option to add this interaction to ‘My collection’.

SUMMARY

The PAIR database contains experimentally reported interactions integrated from major interaction repositories and the most comprehensive prediction of the Arabidopsis interactome with a high reliability. These predictions were expected to cover 24% of the entire Arabidopsis interactome, and their reliability was estimated to be 44%. PAIR features a user-friendly query interface, providing rich annotation on the relationships between two proteins. A graphical interaction network browser has also been integrated into the web interface to facilitate mining of specific pathways. PAIR is a resource not only for large-scale mining of Arabidopsis interactome networks but is also an exploratory tool for cell/molecular biologists to understand more about the relationships between the proteins in specific cellular processes.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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