agriGO v2.0: a GO analysis toolkit for the agricultural community, 2017 update

Tian Tian†, Yue Liu†, Hengyu Yan†, Qi You, Xin Yi, Zhou Du, Wenyiing Xu* and Zhen Su*

State Key Laboratory of Plant Physiology and Biochemistry, College of Biological Sciences, China Agricultural University, Beijing 100193, China

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ABSTRACT

The agriGO platform, which has been serving the scientific community for >10 years, specifically focuses on gene ontology (GO) enrichment analyses of plant and agricultural species. We continuously maintain and update the databases and accommodate the various requests of our global users. Here, we present our updated agriGO that has a largely expanded number of supporting species (394) and datatypes (865). In addition, a larger number of species have been classified into groups covering crops, vegetables, fish, birds and insects closely related to the agricultural community. We further improved the computational efficiency, including the batch analysis and P-value distribution (PVD), and the user-friendliness of the web pages. More visualization features were added to the platform, including SEACOMPARE (cross comparison of singular enrichment analysis), direct acyclic graph (DAG) and Scatter Plots, which can be merged by choosing any significant GO term. The updated platform agriGO v2.0 is now publicly accessible at http://systemsbiology.cau.edu.cn/agriGOv2/.

INTRODUCTION

An enrichment analysis is an efficient and fast method to determine the functions associated with large gene lists and to increase the likelihood of interpreting biological processes (1). Information on biological processes (BP), molecular functions (MF) and cell components (CC) is organized into a structured vocabulary, known as gene ontology (GO), that could describe almost all organisms. The majority of emerging enrichment tools (2–7) employ GO as their annotation resource. However, these resources generally address human or model species but are limited when concerning agricultural species. For examples, the Database for Annotation, Visualization and Integrated Discovery (DAVID) provides a comprehensive platform of functional annotations and analyses based on any given gene list mainly focusing on humans and human diseases. The Gene Ontology enrichment analysis and visualization tool (GORILLA) is used for model species enrichment analyses. Our laboratory developed two GO-based tools for the agricultural community, easyGO (2) and the ensuing agriGO (8), and has continuously maintained them. With the rapid development of high-throughput technologies, these web servers cannot include the enormous amount of sequencing data in the agricultural field (9–14).

Additionally, the GO database is released monthly in several versions, while the definitions of terms and the parent-child relationship between terms changes owing to the integration of information from different sources (15). Many sources and methods induce irregular annotation levels and gene identities (IDs) (15–19). Relatively more comprehensive and accurate versions are being produced based on the evidence codes and/or comparative genomics to reduce the substantial number of false positives (20). For example, a rice gene or gene product’s GO annotation contains versions 5.0, 6.1 and 7.0 from TIGR (21), Gramene release 50 (22), Phytozome v11 (18) and others. Thus, it is necessary to increase the GO annotation, as well as the corresponding GO terms under constantly expanding knowledge.

Up to January 2017, Google Scholar showed that agriGO was referenced 982 times with >80 000 analysis requests since it went online. At the same time, users have put forward a series of requirements for accurate analytical results with reliable backgrounds. To satisfy the ever-increasing number of sequenced species, maintain an up-to-date GO background (15) and meet user requirements, we have updated agriGO with new features and better support. The major updates are supporting more species and datatypes, and adding new visualization tools.

METHODS

AgriGO v2.0 mainly contains analysis tools for processing with the agriGO v2.0-provided background and custom analyses, including search, singular enrichment analysis,
ysis (SEA), parametric analysis of gene set enrichment (PAGE), SEACOMPARE, Batch SEA, DAG drawer and Scatter Plots (Figure 1). The former four were derived from agriGO and updated with broad species and datatypes, while the DAG and Scatter Plots were newly developed for visualization. The difference between SEA and Batch SEA is that the latter can tackle multiple input datasets simultaneously based on one reference (Figure 2). SEACOMPARE is also convenient to compare the significant GO terms based on the false discovery rates (FDRs) from the Batch SEA results (Figure 3).

Statistical test method
The main default statistical method for Fisher’s tests and z-scores in PAGE (23) can be calculated using the following formulae:

\[ P = \frac{n}{k} \left( \frac{N-n}{K-k} \right) \]  

\[ Z = \frac{\bar{x}_n - \mu}{\sigma/\sqrt{n}} \]

where \( N \) represents the total number of one organism’s genes annotated with GO or of the user-provided background, \( n \) represents the number of genes mapped to the background in the query list, \( K \) represents the total number of genes in one GO term and \( k \) represents the counts of overlapped genes. Additionally, \( \bar{x}_n \) represents the mean of sample \( n \), \( \mu \) represents the average for all samples, \( \sigma \) represents the standard deviation and \( n \) represents the sample counts. We also provided other statistical test methods, such as Hypergeometric and Chi-square tests. The Fisher’s exact and Hypergeometric tests are more suitable for situation where the reference lists covers the query list. The Chi-square is appropriate for fewer intersections between the target and reference lists.

The P-value distribution (PVD) can display the distribution of \( P \)-values of significant GO terms. We also use the same formulas as in the SEA function to compute the \( P \)-values for another 99 lists of randomly selected genes of the same number as the observed query lists. Every GO term has a \( P \)-value distribution with 100 values, and the real \( P \)-value’s position is recorded (Figure 2H).

Multi-test adjustment method
The multiple hypothesis adjustment can control the type I error effectively when there is an increase in the number of hypotheses tests. Several multiple hypotheses tests, including Holm (24), Hochberg (25), Hommel (26), Benjamini & Hochberg (BH) (27) and Benjamini & Yekutieli (BY) (28), are available. The first four methods have a greater control of the family-wise error rate. But, the ‘BH’ and ‘BY’ methods use FDRs under less stringent conditions than those of family-wise error rates, therefore, these methods are more powerful than the others.

Batch SEA and SEACOMPARE
With the development of high-throughput sequencing technologies, various platforms and the generation of multiple omics datasets across genomes, transcriptomes, epigenomes, proteomes and others, have been developed. Researchers usually focus on one key biology problem from different points of view (29). However, during transcriptome analysis, some studies on diurnal cycles (30), time-course (31), and the correlations of different abiotic or biotic stresses (31) often produce differentially expressed gene lists with intrinsic connections. Therefore, we developed the Batch SEA function to address requests to analyze multiple samples simultaneously. Batch SEA can cycle analyze the input gene lists with the same background, parameters and cut-offs, which accelerates the speed of processing and improves the efficiency.

SEACOMPARE was produced for the subsequent comparison of Batch SEA results. Two or more jobs can be selected for a comparative analysis with heatmaps to display the common or specific significant terms. The color in the tables indicates the \( P \)-values or FDRs.

The new developed visualization tools
The direct acyclic graph (DAG) drawer and Scatter Plots are two visualization tools with different formats to illustrate the significant GO terms. The DAG, based on the nature of the GO structure, can indicate submitted terms and the inter-relationships between terms, as in agriGO (8). Scatter Plots can display the significant GO terms after a multi-step filtering process. We first remove the GO terms above the \( P \)-value cut-off, then select one of the GO categories and filter the similar GO terms based on the semantic similarity measurement method (32). Scatter Plots remove similar GO vocabularies and can show the core GO terms.

The Semantic Similarity Measurement (32) estimates the functional similarities of gene products based on GO vocabularies (33) using the formula:

\[ \text{Similarity} (A, B) = \frac{\sum_{t \in T_i \cap T_j} S_A(t) + S_B(t)}{SV(A) + SV(B)} \]

Wang (33) defined the contribution of the GO term \( t \) (closer to term A) to the semantic GO term A as the S-value of GO term \( t \) related to term A, where \( S_A(t) \) is the S-value of GO term \( t \) related to term A and \( S_B(t) \) is the S-value of GO term \( t \) related to term B. Multidimensional Scaling for Java (http://www.inf.uni-konstanz.de/algo/software/mdsj/) can translate the dissimilarity matrices to coordinate information.

The processing flow for predicting the GO annotation
Glycyrrhiza uralensis was used as an example to introduce the comparative genomic approach to predict the GO annotation. We used BLAST-2.2.19 (-e 1e-3 -m 8) to compare the protein sequences of G. uralensis (34) and Arabidopsis thaliana (35). Then, the orthologous pairs with A. thaliana were determined, as was the corresponding GO annotation. Additionally, Blast2GO has a comprehensive bioinformatics platform that can automatically predict GO annotation.
Figure 1. The analysis framework for agriGO v2.0. AgriGO v2.0 provides three analysis entrances and six major tools. For all query lists, P-values are computed and adjusted by multiple testing corrections or permutation tests. Then, significant GO terms can be further analyzed for similarity or parent–child hierarchical structures. Four visualization methods can help users understand the results. The orange color represents the new function in agriGO v2.0.

RESULTS
The updates in agriGO v2.0 are mainly embodied in three parts, adding a large number of species and datatypes, newly developed tools and more flexible visualization methods. These can increase the range, quality and strength of the statistical analyses, as well as present a convenient and intuitive graphical interface with new features.

Integrative collection of GO annotations
Reference files with higher whole genome coverage levels, as well as bigger ratios of GO terms (relative to all GO entries) are key points in producing reliable analysis results. We collected additional GO annotation files from different methods, including published databases (18,19,21,22,35,39), publications and those computed by comparative genomics (detailed in METHODS) (11,34,40–42).

Identifying the correct GO structure is another challenge (Supplementary Figure S1). We determined the structure for each GO annotation source by comparing its similarity with the GO files from the 2010 and 2016 versions. The background of version 2010 was retained to be consistent with agriGO v1, ensuring that former study results had a stable continuity. The 2016 version was corresponded to the new datatypes. We plan to update the older version and manually add into our web server later. Then, we ranked the datatype at the front based on their larger coverage of GO terms and genes, reliable data sources and newer annotated versions. The most recommended datatype was ranked first, and users are free to choose the appropriate datatype (Supplementary Table S1). In summary, a great deal of useful information regarding GO annotations was gained, including 394 species containing 865 datatypes.

Species classification
The previous display method for the organisms is not suitable for more species and datatypes become available. We categorized the species into plant, animal and fungi categories (Table 1). The plant category was further divided into Brassicaceae, Poaceae, Malvaceae, Fabaceae, Solanaceae, Medicinal plants, etc. Of those, the Brassicaceae family includes both vegetable and oilseed crops (43); the Poaceae (grass) family includes major cereal crops (Supplementary Table S1); the medicinal plants have attracted much atten-
Figure 2. Batch SEA for up-regulated genes after cold treatments (0.5, 1, 3, 6, 12 and 24 h) in *Arabidopsis thaliana*. Input the query lists into the expanded text box with clicking on the ‘add’ button (A). All successful job IDs (B) corresponding to query lists can be selected for the SEACOMPARE analysis (C). A brief summary of one job for 24-h cold stress (D). Users can find the basic information on the reference ‘Affymetrix ATH1 Genome Array’ that contains 22,479 datatypes better matches the GO version 2010, and 34 significant GO terms were obtained with 165 annotated genes in the query list. The detailed information about the BP category was displayed in a tabular format, including significant GO terms with descriptions, gene numbers in input and background lists, P-values, FDR by multi-test adjustments, and P-value distribution (E). The DAG (F) and Scatter Plots (G) illustrates the BP category. A few GO terms were not in the top 1% and they were filtered out by multi-test adjustments, like the GO term ‘response to hormone stimulus’ (GO:0009725) (H).

Batch SEA

Several new functions have only been added to the single species analysis pages. We have now provided a new function, Batch SEA, to tackle multiple input datasets simultaneously based on one reference. The Batch SEA method often works best for time-course or continuous datasets, providing not only increased calculation speeds, but also benefits for the subsequent comparative analysis. SEACOMPARE is applied to compare the significant GO terms from the results of selected datasets to effectively identify GO terms.
Figure 3. SEACOMPARE analysis based on multiple hypothesis tests (FDRs) and P-values of the up-regulated gene list after a 6-h cold treatment of Arabidopsis thaliana. The detailed information in each column, from left to right, represent the significant GO terms, GO category (BP, CC and MF), GO annotation, a heatmap (columns 4–10) for FDRs or P-values (columns 11–17) from different computational methods, and the last column is the top percent based on the P-value distribution as assessed by the permutation test. The colored blocks in the heatmap represent the level of significance in each term. The yellow-to-red indicates the level from low to high, and grey indicates not significant.

Table 1. Species classifications and statistics

| Category         | Classification | Species counts |
|------------------|----------------|----------------|
| Plant            | Brassicaceae   | 12             |
|                  | Poaceae        | 29             |
|                  | Malvaceae      | 6              |
|                  | Fabaceae       | 16             |
|                  | Solanaceae     | 12             |
|                  | Rosaceae       | 5              |
|                  | Medicinal plant| 12             |
|                  | Tree           | 29             |
|                  | Algae          | 20             |
| Animal           | Fish           | 20             |
|                  | Bird           | 11             |
|                  | Amphibia       | 3              |
|                  | Insecta        | 56             |
|                  | Mammalia       | 58             |
|                  | Sordariomycetes| 5              |

We took six groups of up-regulated Arabidopsis genes under different cold treatment time intervals as inputs for a batch enrichment analysis at the A. thaliana SEA analysis webpage (8). There were 61, 64, 105, 150, 165 and 165 up-regulated genes corresponding to cold stress after 0.5, 1, 3, 6, 12 and 24 h (Supplementary Table S2), and these were inputted into the expanded text box after clicking on the ‘add’ button (Figure 2A). The detailed enrichment results, as well as the next visual analysis, with the significant GO terms are shown in Figure 2.

Job IDs are listed in the first analysis (Figure 2B), and the SEACOMPARE analysis can be performed using selected job IDs (Figure 2C). Thus, the GO terms were reproducible and a significant elevation occurred with the longer cold treatments. GO terms concerning ‘respond to cold’ (GO:0009409) were significantly represented throughout all of the stages, and ‘cold acclimation’ (GO:0009631) appeared after 24 h. Then, a brief summary about the job of 24-h cold stress is available. Based on the selected reference ‘Affymetrix ATH1 Genome Array’ (with GO version 2010), all 165 inputted genes were present in the 22 479 background genes and 34 significant GO terms were identified (Figure 2D). The detailed significant GO information in a tabular format ranked by FDR contains the GO term’s description, P-value, FDR and PVD (Figure 2E). A DAG (Figure 2F) and Scatter Plots (Figure 2G) can be generated for each GO category (BP, CC and MF). The GO terms (GO:0009469, ‘response to cold’ and GO:0009414, ‘response to water deprivation’) were located at the lower level of the hierarchy but had higher significance levels (Figure 2F). Several GO terms (GO:0006950, ‘response to stress’ and GO:0009311, ‘oligosaccharide metabolic process’) that had too many or too few relationships with the rest of the GO terms were filtered out by Scatter Plot drawer (Figure 2G).

The PVD is represented by a line chart that can display the distribution of P-values of significant GO terms of observed query and random genes. It can decrease the false positive rate through the permutation test. The line chart can illustrate the location and percentage among 99 other random results to help users distinguish the real significant GO terms owing to their background bias or integrity. The PVD for each significant GO term can be illustrated when clicking on the line chart icon with a short description of the P-value position in the last column (Figure 2H). The P-value of the GO term concerning ‘response to hormone stimulus’ (GO:0009725) was not ranked first and was removed after a multi-test adjustment.

Additional tools for customized analyses

The accumulation of species in agriGO cannot catch up with the rate of genomic data generation or GO annotation.

Reference: 'Affymetrix ATH1 Genome Array' (with GO version 2010).

Supplementary Table S2.
tion updates, especially the de novo data that is user-defined. The custom tools independent of species were available in the previous version but were often overlooked by users. To facilitate user awareness, we highlighted these tools in the navigation bar and also added new tools to agrirGO v2.0. The custom SEA and SEA share the same principle, but the custom tool can be used with any pair of queries and references. The SEACOMPARE tool can compare ID numbers from SEA or a data matrix with GO terms as the first column and P-value as the left column (Figure 2C). Both the DAG drawer (Figure 2F) and Scatter Plots (Figure 2G) are graphics generation tools of different forms based on the significant GO terms and P-values in each of the categories, respectively. These can improve the usage range and compatibility with external data independent of the species and statistical methods. The DAG drawer can reproduce the structure with the significant GO terms based on the relationship between parent and child terms. The Scatter Plots can remove terms with similar definitions.

**DISCUSSION**

With the accumulation of sequencing data and various user requirements related to adding new species or datatypes, operating processes, downloads, GO versions and others (Supplementary Figure S2), we updated agrirGO (v2.0) to contain more selected species and datatypes, a newly added species classification system and different GO structure versions. We maintained a similar interface and provide three analysis entrances: the group species analysis based on the classifications consistency and single species analyses with new Batch SEA and PVD options. Additionally, custom tools independent of species produced in the previous version (SEA and SEACOMPARE), as well new developed tools (DAG drawer and Scatter Plots) were gathered and placed in a striking position (Figure 1).

The GO terms were updated monthly by changing the terms or relationship between terms. The current number of GO entries for BP, CC and MF are far greater than in the old version of agrirGO (Supplementary Figure S1A). Additionally, the same GO term may correspond to different annotations, as well as a relationship between two parent–child terms (Supplementary Figure S1B). To have consistency between analyses from different periods, agrirGO v2.0 must adapt to the changes in GO while maintaining uniformity between the GO annotations for each organism and the ontology structure (Supplementary Table S1).

Although the structure of GO and the coverage of GO annotations are continuously improving, the overall annotation level is low, except for those of some model species (46). Other means of gene functional annotations, like gene families and pathways, can be divided into gene sets for enrichment analyses (3). Furthermore, gene annotations can be more explicitly compared with the increasing number of relationships between genes and GO terms. Thus, we added gene annotations for some model species with high-quality gene annotations, including Arabidopsis, rice, maize, soybean and others.

The PVD option, which was embedded in the ‘optional advanced options’, provides a trend graph to display the P-values of query dataset and 99 random datasets (Figure 2H) (47). The line chart highlights the practical P-value with a red box and is displayed in two forms (normal and logarithmic scales). In SEA, we used Fisher’s exact test and all of the multiple hypothesis tests (48) provided in agrirGO v2.0 to analyze the up-regulated gene list under a 6-h cold stress treatment. The analysis results with six kinds of multiple hypothesis tests were compared by the SEACOMPARE tool displayed in Figure 3. We found that the GO terms selected by the multiple hypothesis test are always located at the top. The GO term concerning ‘calcium ion binding’ (GO:0005509) can be selected by the permutation test but is filtered out by the multi-test adjustment. Ca²⁺ plays an important role in the signal transduction pathways that lead to stress gene expression (49,50). Through the comparison between different statistical testing methods, the PVD can help users avoid missing important GO terms (47).

This was a practical demonstration that the distribution of P-values, to some extent, can distinguish false positive events (Figure 3). From our empirical point of view, lists of several hundred genes worked better (for A. thaliana as example). If there are too few genes on the list, the sensitivity is lost, but with too many, specificity is lacking. Some large input gene lists, encompassing >60–70% of the whole genome might be confused with the analysis results of other similar scale datasets. This is partially due to the incompleteness or preference of the GO terms in the reference background.

We have been constantly developing and enhancing our GO analysis tools and databases to support the agricultural community based on user experiences and feedback. The updated version agrirGO v2.0 has an increased number of species and datatypes available, which have been classified into several groups. SEA, as the central tool, has been improved with a batch analysis to handle multiple input datasets simultaneously. Custom tools, including custom SEA and SEACOMPARE inherited from agrirGO, and some new tools (DAG drawer and Scatter Plots) were placed together and highlighted in the navigation bar. These will provide users with more efficient and systematic analyses. We will continue to maintain our web server and welcome feedback on the need for new datatypes, the usability of existing tools, or suggestions for new features.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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