Annotating Genes of Known and Unknown Function by Large-Scale Co-Expression Analysis

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Running Head

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Abstract

About 40% of the proteins encoded in eukaryotic genomes are proteins of unknown function (PUFs). Their functional characterization remains one of the main challenges in modern biology. In this study we identified the PUF encoding genes from Arabidopsis thaliana using a combination of sequence similarity, domain-based and empirical approaches. Large-scale gene expression analyses of 1310 publicly available Affymetrix chips were performed to associate the identified PUF genes with regulatory networks and biological processes of known function. To generate quality results, the study was restricted to expression sets with replicated samples. First, genome-wide clustering and gene function enrichment analysis of clusters allowed us to associate 1,541 PUF genes with tightly co-expressed genes for proteins of known function (PKFs). Over 70% of them could be assigned to more specific Biological Process annotations than the ones available in the current Gene Ontology release. The most highly over-represented functional categories in the obtained clusters were ribosome assembly, photosynthesis and cell wall pathways. Interestingly, the majority of the PUF genes appeared to be controlled by the same regulatory networks as most PKF genes, because clusters enriched in PUF genes were extremely rare. Second, large-scale analysis of differentially expressed genes (DEGs) was applied to identify a comprehensive set of abiotic stress response genes. This analysis resulted in the identification of 269 PKF and 104 PUF genes that responded to a wide variety of abiotic stresses, while 608 PKF and 206 PUF genes responded predominantly to specific stress treatments. The provided co-expression and DEG data represent an important resource for guiding future functional characterization experiments of PUF and PKF genes. Finally, the public Plant Gene Expression Database (PED, URL: http://bioweb.ucr.edu/PED) was developed as part of this project to provide efficient access and mining tools for the vast gene expression data of this study.
Introduction

Only a small percentage of the proteins encoded in animal or plant genomes are sufficiently characterized with regard to their cellular functions. The functions for the majority of these proteins remain either completely unknown (40%) or only partially understood (Gollery et al., 2006, 2007). In light of this significant knowledge deficit, our understanding about existing molecular functions appears to be fundamentally incomplete. This is even more evident when we assume that the vast space of unexplored molecular and biological functions is composed of proteins with at least comparable or even greater diversity and importance for cellular processes than the known space. Efforts to narrow this knowledge gap will provide a wide spectrum of opportunities for advancing our understanding about plant and non-plant systems.

Two major methods are in use for defining proteins of unknown functions (PUFs) in model organisms. The widely used similarity approach considers all proteins as PUFs that show no detectable sequence or structural similarities to functionally characterized proteins in reference databases (Leinonen et al., 2004; Boeckmann et al., 2003). In contrast to this, the more conservative empirical approach defines as PUFs all proteins that lack direct experimental evidence as support for a specific function. Conceptually, the empirical approach incorporates most PUFs identified by the similarity approach, as well as functionally uncharacterized sequences that share sequence similarities with proteins of known function (PKFs). Sequence families and ortholog clusters are particularly affected by this fundamental difference between the two unknown definitions. For instance, when a group of related sequences contains one or more members of known function, then the similarity approach tends to assign all of them to the known space, whereas the empirical approach distinguishes between functionally characterized and uncharacterized candidates within groups of related sequences. As a result of this difference, most similarity-based PUFs of a given genome are either singletons or members of families that consist exclusively of uncharacterized sequences. These performance characteristics of the similarity concept result in an underestimation of the number of PUFs, because many genes in eukaryotic organisms are members of poorly characterized gene families (Horan et al., 2005). To illustrate this, all members of large families, like protein kinases or cytochrome P450s, will be assigned by the similarity approach to the known protein space, even though most of their members remain functionally uncharacterized (Horan et al., 2005; Nelson et al., 2004; Wang et al., 2003).

Dividing gene products into only two categories of known and unknown sequences is an oversimplification of a complex knowledge system with incremental and multifaceted differences. Consequently, every definition for drawing a strict separation line remains artificial and controversial. While acknowledging these difficulties, this study will adopt this two-class system mainly for practical reasons.

To advance our knowledge beyond a roadmap of knowing what we don’t know, it is important to develop and apply approaches for predicting putative functions for PUFs. Bioinformatic techniques provide here a wide spectrum of opportunities. For instance, PUFs can be associated with remotely related PKFs by using sensitive sequence and structure similarity search strategies (Altschul et al., 1997; Eddy, 1996). The detected similarities can reveal important clues for testing their functions experimentally. Additionally, one can predict functional features from their sequences, such as subcellular targeting signals, secondary structures and membrane domains (Schwacke et al., 2003; Gollery et al., 2006). Proteomics and protein interaction technologies provide additional important functional links (Johnson and Liu, 2006). However, for plants the required proteome resources are not yet available on a genome-wide level. One of the most promising and readily available information resources for systematic functional assignment studies of PUF genes represent large-scale gene expression data from public microarray databases. These data sets over vast opportunities for associating PUF genes with molecular functions and cellular processes of co-regulated PKF genes.

In this study we identified and analyzed the genome-wide PUF encoding genes from
Arabidopsis using both, empirical and similarity strategies. Large-scale analysis of publicly available gene expression array data allowed us to associate PUF with PKF genes based on similarities of their expression and treatment response profiles. For this, cluster analysis was used to identify groups of co-regulated PUF and PKF genes based on the similarity of their expression profiles across a wide range of tissue and treatment samples. Subsequently, enrichment analysis of Gene Ontology terms was applied to annotate the obtained clusters by over-represented gene functions. Second, statistical analysis of differentially expressed genes (DEGs) allowed us to identify PUFs that exhibit generic and specific expression changes in response to a large number of different abiotic stress treatments. Finally, the Plant Gene Expression Database was developed to provide to the public efficient data mining utilities for the complex differential expression and clustering data of this project.
Results and Discussion

Identification of PUFs
To obtain for this study a comprehensive set of PUFs from Arabidopsis, we compared three profoundly different PUF identification methods. The three approaches are based on Gene Ontology annotations, sequence similarities and protein domain searches.

First, we mined the Gene Ontology (GO) annotations to estimate the number of PKFs and PUFs from a manually curated knowledge system that combines empirical and computational methods for assigning gene functions (Berardini et al., 2004; Falcon and Gentleman, 2007). Alternative pathway annotation systems from KEGG and AraCyc could have been used for the same purpose (Kanehisa et al., 2006; Mueller et al., 2003). However, due to the limited number of Arabidopsis genes (<40%) assigned to pathways, the GO system, with close to 95% genome coverage, appears to be currently the more efficient resource for identifying nearly complete PUF sets. This number includes the direct assignments to the root term of each ontology which are the new GO annotations for sequences of unknown function (see Material and Methods for more details).

The evidence codes of the GO annotations specify which functional assignments are supported by experimental evidence data from the public domain and which annotations are solely based on computational prediction methods (Ashburner et al., 2000). To gain insight into the nature of the annotations with regard to the evidence type for assigning members to the known and unknown space, we combined in Table I the current set of thirteen evidence codes into four custom categories. The category with the highest level of functional support (Empirical) is based on direct evidence from traditional single sample experiments, the second one is based on large-scale screening data (Large-Scale), the third one on computational predictions (Sequence), and the fourth one are the GO-based PUF entries that lack functional support from experiments or in silico analyses. The detailed assignment schema of the evidence codes to the four categories is provided in the legend of Table I.

According to the above strategy, 32-38% of the Arabidopsis genes are currently annotated by the GO system as PUF encoding genes (Table I). This is largely in agreement with the estimates from previous studies (Wortman et al., 2003; Gollery et al., 2006). Interestingly, only 7% of all entries are functionally characterized by traditional one-gene-at-a-time experiments in the Molecular Function (MF) ontology and 14% in the Biological Process (BP) ontology, while 34% and 18% have functional support from high-throughput experiments, respectively. This means that 93% of the genes from Arabidopsis code for poorly characterized proteins or PUFs when the most conservative empirical criteria are applied within the MF ontology. The relative amount of PUFs for the combined empirical and large-scale categories is 59% in the MF ontology and 68% in the BP ontology. The Cellular Component (CC) ontology contains by far the largest number of entries with sequence-based annotations and the lowest for the empirical categories. This trend is due to the majority of the CC annotations presently being based on computational ab initio predictions of sub-cellular localizations, whereas annotations with experimental support are much less frequent in this category than in the other two ontologies. The subsequent analysis steps of this study utilize the standard PUF set from the MF ontology containing 8,665 members. These genes are exclusively assigned to the root term of the MF ontology (GO:0003674) and they carry the evidence code ND (no biological data available). The MF category was selected here, because protein functions are most profoundly described at the mechanistic molecular level, whereas the other two ontologies, BP and CC, provide rather indirect information in this regard.

To compare the results obtained from the MF ontology with alternative PUF identification methods, we also used one sequence similarity and one domain-based approach using Hidden Markov models. First, all predicted Arabidopsis proteins were searched against the Swiss-Prot database with the BLASTP program (Altschul et al., 1990; Wu et al., 2006). Protein sequences that showed no similarities to functionally characterized proteins in the Swiss-Prot database were classified as PUFs
using an expectation value (E-value) of $10^{-6}$ as cutoff. Second, the same protein set was used to search the Pfam database with the HMMPFAM program (Eddy, 1996; Bateman et al., 2004). Likewise, sequences without similarities to protein domains of known function (E-value $\geq 10^{-2}$) or those matching exclusively domains of unknown function (DUF) were considered PUFs. Due to different calculation methods, the E-values of the two search algorithms are not directly comparable. Therefore, we chose for both methods conservative cutoff values that are commonly used for sensitive sequence similarity searching with low false positive detection rates (e.g. Gollery et al., 2006; Horan et al., 2005; Girke et al., 2004). Table II provides a comparison of the results from the three different PUF identification approaches. Based on the chosen confidence thresholds, all three approaches identified PUF sets of comparable sizes with 8,272-8,681 members, while 5,456-6,260 PUFs are common among two and 4,667 among all three methods. The corresponding gene lists for the three methods are provided in Supplement S1.

To simplify the description of the subsequent functional analysis steps of this study, the remaining text is restricted to the PUF set obtained from the MF ontology, while the data for the remaining PUF identification methods are included in the corresponding Supplements S1, S3, S5 and S7. The GO PUF set was given preference, because of the high quality of the manually curated GO annotation system and its broad acceptance in the scientific community.

**Relative Amount of Expressed Genes**

To functionally associate PUF with PKF encoding genes based on the similarity of their mRNA expression profiles, large-scale gene expression analysis of publicly available Affymetrix GeneChip microarrays was performed. Only experiment sets containing at least two replicate samples were used for this analysis to enable statistical analysis of differentially expressed genes (DEG) and to increase the confidence of the obtained results. In total, the study included the raw expression data from 1,310 Affymetrix chips from the AtGenExpress and GEO sites (Schmid et al., 2005; Barrett et al., 2006). Table III provides a summary of the chosen experiment sets that covers a wide spectrum of treatment series and tissue samples. The complete list of the analyzed data is available in Supplement S2.

The relative amount of expressed genes can be expected to be lower in the PUF than in the PKF category, because many predicted PUF genes may be the result of genome annotation artifacts or may represent untranscribed pseudogenes. In addition, a certain fraction of PUF genes may be expressed below the detection limit of the GeneChip microarray technology. To estimate the extent of these limitations, the amount of detectable genes across all experiment categories was compared between the PUF and PKF sets. The present call information of the non-parametric Wilcoxon signed rank test of the MAS5 algorithm provides for this purpose relatively reliable estimates (Liu et al., 2002; Schmid et al., 2005; McClintick and Edenberg, 2006). According to this test, the amount of detectable genes between the PUF and PKF sets differs 0.5-8% within the five frequency intervals plotted in Figure 1. The detailed data set of this analysis is available in Supplement Table S3. Based on these rather small relative differences, it is likely that the majority of the PUF genes are expressed at high enough levels to obtain for them meaningful data in the downstream cluster and differential gene expression analyses of this study.

**Cluster Analysis**

Since many dynamic cellular processes are tightly associated with coordinated transcriptional changes, cluster analysis of gene expression profiles can be used to identify candidate sets of co-regulated genes that are directly or indirectly involved in related processes (Steinhauser et al., 2004a; Gachon et al., 2005; Toufighi et al., 2005; Haberer et al., 2006; Jen et al., 2006; Vandepoele et al., 2006; Wei et al., 2006; Gutierrez et al., 2007). For instance, if a group of genes exhibits correlated expression profiles and it is significantly enriched in genes involved in a specific process then it is reasonable to assume
that some of the PUF members of this cluster may share overlapping functions with its functionally characterized members. This association-based approach was applied here on a genome-wide level to systematically assign PUF to PKF genes based on the similarity of their expression profiles. Despite the great potential of this approach, it is important to keep in mind that correlation does not prove causal relationships. It only provides useful leads for establishing hypotheses and causal links in downstream investigations. Accordingly, the results of this study need to be interpreted as preliminary computer predictions that offer useful information for guiding future gene characterization experiments. Final evidence about gene and protein functions cannot be inferred directly from this data. Alternative network modeling approaches were not considered for this study, because of the lack of efficient statistical methods to efficiently represent, score and interpret the resulting network architectures on a genome-wide scale (e.g. Wolfe et al., 2005; Gutierrez et al., 2007; Ma et al., 2007). At this point, the traditional clustering approach appears to be more practical for the goals of this study.

To generate reliable and biologically relevant gene clusters from expression data, we evaluated several available clustering algorithms (e.g. K-means, SOM) and selected agglomerative hierarchical clustering as the method of choice (Murtagh, 1985; Eisen et al., 1998; de Hoon et al., 2004; R Development Core Team, 2006). The hierarchical clustering method was chosen because of three main advantages: (1) the method requires no prior knowledge about the optimum number of the final clusters, (2) it is extremely robust in joining highly similar items into proper similarity groups and (3) it provides an information-rich data output that represents the relative distances between all clustered items in a dendrogram (Becker et al., 1988). The main disadvantages of the approach are the complexity of its data output, the lack of predefined boundaries between clusters and its weaker performance in identifying local expression similarities in a small subset of the samples (Prelic et al., 2006). However, most of these challenges can be overcome by applying efficient post-processing methods of the obtained dendrograms, such as tree cutting methods (e.g. Gutierrez et al., 2007). Popular fuzzy clustering approaches (Krishnapuram et al., 2001) that allow memberships in several clusters - as opposed to strict clustering with unique memberships - were not considered for this study, because of the difficulty to efficiently prioritize and mine the complex cluster memberships from these methods in the downstream functional analysis steps. As an implementation of the hierarchical clustering algorithm, we used the hclust function (Murtagh, 1985) from the statistical programming environment R (R Development Core Team, 2006). As distance measurement we used correlation coefficients and as cluster joining method complete linkage (see Material and Methods for more details). To obtain discrete clusters from the resulting dendrograms, we developed for this study a novel hierarchical threshold clustering (HTC) method. The corresponding R script is available in Supplement S10. This method selects clusters in hierarchical clustering dendrograms based on a maximum tolerable distance between cluster members by applying an all-against-all distance test on all possible sub-trees, while maintaining unique cluster memberships. As threshold we chose for this step a minimum correlation coefficient of 0.6. This relatively conservative HTC setting ensures that all members of any given cluster share with all other members of the same cluster correlation coefficients between the selected cutoff of 0.6 and the highest possible value of 1.0. The exact cutoff value of 0.6 was chosen because it resulted in the highest enrichment of functionally related genes compared to alternative cutoff settings (Supplement S4). Additionally, other gene expression correlation studies have used the same or very similar cutoff values (Haberer et al., 2006; Wei et al., 2006).

Applying the above strategy, we calculated four separate clustering data sets using both the Pearson and Spearman correlation coefficients, in their signed and absolute forms as distance measures. The following text will refer to the four methods as PCC, SCC, PCCa and SCCa, respectively (Supplement S5). All four data sets were generated, because of their complementary performance characteristics. The clustering with absolute correlation values allows the identification of positively and negatively correlated gene expressions, whereas the sign-specific approach joins only positively correlated items into similarity groups. The rank-based Spearman approach is limited to identifying
global similarities in expression profiles, while the Pearson approach is very sensitive in detecting both, global and local similarities. In particular, the latter detects local similarities with wide amplitude changes relative to the background, which can result in extreme cases in co-clustering of outliers. A consensus approach between several or all methods was not considered, because such a strategy would artificially deflate the cluster sizes and compromise the transparency of the results.

The distributions of the obtained numbers of clusters including their sizes from the four clustering methods are summarized in Figure 2. Because the sign removal increases the potential pool sizes of gene pairs with correlation values above a given cutoff, one would expect larger cluster sizes for the data sets with absolute correlation values compared to their signed counterparts. This trend can be observed in the many individual clusters in Supplement S5, but the effect is not very pronounced in the global representation of Figure 2. These relative increases in cluster sizes are not as frequent as expected, because of two main reasons. First, the number of highly negatively correlated gene pairs is much smaller than the number of positively correlated gene pairs (data not shown, compare Haberer et al., 2006). Second, the assignment of a negatively correlated gene to a cluster at an earlier stage of the hierarchical clustering process can prevent other potential members from joining the same cluster at a given cutoff level, if they do not share the required degree of correlation with the existing members. This is particularly the case in combination with a complete linkage joining method, that was chosen for this study to minimize the number of false positive members in the generated clusters.

The most obvious differences among the four clustering data sets in Figure 2 are the numbers of singlet genes that do not join any clusters in the different methods. There are about 2000 fewer singlet genes in the Pearson than in the Spearman data sets. This is expected because the latter method tends to generate slightly lower correlation values on gene expression data. Due to space restrictions, the subsequent text focuses on the clustering results from the distance method with the signed Pearson correlation coefficients (PCC), whereas the results for the other three methods are included in Supplement S5. In addition, the clustering data for individual genes are available in the associated public database of this study (see below).

Functional Categorization of Gene Expression Clusters
Gene expression clusters with highly enriched functions provide more conclusive information about the potential roles of their PUF encoding members than clusters with very heterogeneous compositions. To functionally annotate the obtained clusters and to select the most informative gene sets with over-represented gene functions, we performed enrichment analysis of Gene Ontology terms using the hypergeometric distribution as a statistical test (Falcon and Gentleman, 2007). This method computes the enrichment test for all ~18,000 GO nodes of the three ontology networks and ranks the results by p-values (see Material and Methods, and Supplement S9). The results of this method are more comprehensive and informative than generalized functional categorization systems, like GO slim or high-level pathway classification systems. Clusters with fewer than 5 members were excluded from this analysis, because the predictive value of extremely small clusters is rather limited. The complete result set of this enrichment analysis is available in Supplement S6. It contains the data for 916 clusters composed of a total of 11,077 genes. To prioritize the clusters based on the obtained enrichment data, we applied two selection filters. First, each cluster of interest needed to contain at least one over-represented GO term in one of three ontologies (enrichment filter). Second, at least 20% of the cluster members had to be associated with this GO term in order to select clusters with relatively homogeneous compositions (uniformity filter). An overview of the number of clusters that meet these filter criteria is provided in Table IV. It contains the results for four different p-value cutoffs of the GO term enrichment filter ranging from 0.05 to $10^{-6}$. The corresponding GO annotations for the prioritized cluster set, that passed the most stringent selection criteria of $10^{-6}$, are listed in Table V. For space and readability reasons, the table presents only the highest ranking GO term for each of the three
ontologies. The full set of GO annotations can be found in Supplement S6. The following discussion of selected clusters is restricted to this most conservative data set (Table V). It contains 66 clusters with a total of 1,279 genes that include 277 PUF genes derived from 53 clusters (see Table IV). Our focus on these clusters does not indicate that the other clusters of this study are biologically less important. This selection is mainly based on the assumption that clusters with uniform GO annotations are particularly informative for functionally associating PUF with PKF genes.

Depending on the stringency of the applied prioritization filters listed in Table IV, our combined clustering and GO term enrichment strategy associated 277-1,541 PUF genes to overrepresented GO annotations. In comparison to the GO annotations currently available for these PUF genes, our method associated 216-1,050 of them to more specific GO terms in the MF category, 225-1,089 in the BP category and 239-1,096 in the CC category (Supplement S6). The large number of PUF genes associated to functionally informative annotations demonstrates the great potential of our approach for guiding future experimental studies on these genes.

Based on enrichment p-values, the most highly over-represented functional categories in the obtained cluster set are the biological processes: ribosome assembly, photosynthesis pathways and cell wall metabolism (Table V). This finding is largely in agreement with related gene co-regulation studies in Arabidopsis (Haberer et al., 2006; Wei et al., 2006). With regard to ribosome assembly, 124 of the 410 GO annotated genes for cytosolic, plastidial and mitochondrial ribosome components appear in seven clusters (see Table V, cluster IDs: 23, 32, 37, 39, 182, 239 and 299); and 272 ribosomal genes appear in clusters with ≥5 members of the non-prioritized data set. While cluster 23 consists exclusively of genes annotated as ribosomal genes (GO:0005840, p-value: 1.2*10^{-64}), the other six clusters are highly enriched in ribosomal genes, and they contain among others 16 PUF genes. Equally interesting is the observation that photosynthesis-related annotations are highly over-represented in five large clusters (cluster IDs: 4, 9, 45, 110 and 304). These clusters represent 51 of all the 121 genes that are currently annotated by the GO system as photosynthesis components (GO:0015979). Because both processes, photosynthesis as well as ribosomal activities, require the coordinated assembly of many proteins to large complexes and protein-protein interaction networks, it is not unexpected that their corresponding genes are tightly co-regulated. In alignment with the association hypothesis of this study, several of the PUF members in these functionally extremely uniform clusters may be involved in processes that are connected to the enzymatic or regulatory networks of photosynthesis and ribosomal activities.

Interestingly, our method also identified a cluster (ID 77) that is highly enriched in cell wall-related annotations (e.g. GO:0009834, p-value: 4.2*10^{-15}), such as cellulase synthase genes. A very similar cluster of genes was recently described and experimentally verified by two groups (Persson et al., 2005; Brown et al., 2005) who specifically mined public expression data for genes that are co-regulated with the cellulose synthase genes CESA4, 7 and 8. In addition, comparable results were described by Jen et al. (2006). This example demonstrates that our genome-wide expression clustering approach generates biologically meaningful data. An additional interesting cell wall-related cluster is cluster 349 that contains eight genes for proline-rich extensin domain proteins.

The majority of the clusters in our data set contain one or more PUF genes (Table IV), but only a few of the larger clusters consist predominantly of PUF genes. Cluster 17 represents an exception to this rule. The 43 members of this cluster contain 26 PUF genes, and its characterized members show no clear enrichment of specific functions. Based on the high abundance of PUF genes in the entire data set (~32%), PUF gene enriched clusters occur much less frequent than those enriched in PKF genes; and clusters consisting exclusively of PUF genes are entirely absent (Table IV). One explanation for this difference could be that the expression of most PUF genes is controlled by the same regulatory networks as many PKF genes. If this is the case, PUF genes are more likely to appear in expression clusters together with PKF genes than without them.
Our method also identified clusters that are enriched in abiotic stress response annotations. For instance, clusters 85 and 912 are highly enriched in heat stress-related genes (GO:0009408, p-values: $6.7 \times 10^{-18}$, $1.8 \times 10^{-6}$). Interestingly, 10 of the 23 members in the cluster 85 were identified by the subsequent DEG analysis of this study, as genes that respond specifically to heat stress and to a much lesser extent to other types of abiotic stresses (see Supplement S7). Based on the available co-expression data, the 9 PUF genes of this cluster are now excellent candidates for discovering novel gene functions involved in heat stress response pathways. Additionally, this example illustrates that the two chosen approaches of this study, expression clustering and DEG analysis, complement and confirm each other. The hypoxia cluster 203 is another interesting abiotic stress cluster (Supplement S6; Fukao and Bailey-Serres, 2004). This cluster does not appear in the most stringently prioritized data set (Table V), because it did not pass the applied uniformity filter. Nevertheless, it is enriched in hypoxia-responsive genes (cluster ID 203, GO:0001666, p-value: $2.0 \times 10^{-5}$), and it contains several members that are involved in cellular respiration processes, such as genes for the alcohol dehydrogenase ADH1 (AT1G77120), a pyruvate dehydrogenase (AT4G33070) and a hemoglobin-like oxygen binding protein that affects ATP levels under hypoxia (AT2G16060, Hebelstrup et al., 2007). Whether the five PUF genes of this cluster are also involved in hypoxia-response processes, can be addressed in experimental studies.

In conclusion, the combined clustering and gene function enrichment strategy allowed us to associate a considerable fraction of the PUF encoding gene pool with tightly coexpressed gene sets of known function. Depending on the chosen stringency settings, the approach allowed us to assign 277-1,541 PUF genes (Table IV) to more specific GO terms than those available in the latest GO annotation release for Arabidopsis.

### Analysis of Differentially Expressed Genes (DEGs)

DEG analysis can identify groups of genes that exhibit expression changes in response to specific treatments or cellular changes. Because this information is not easily obtainable from clustering of global expression profiles, DEG analysis of publicly available expression data complements the previous approach by associating PUF with PKF encoding genes based on common differential expression responses to environmental changes, such as abiotic stresses. If a group of genes shares similar expression patterns across a wide spectrum of treatments then it is likely that certain members are involved in similar or connected response pathways to these perturbations. The association of genes with these response mechanisms can provide valuable information for future functional characterization experiments of PUF or PKF genes.

One of the main challenges of performing systematic DEG analyses on large and diverse gene expression data sets from public sources is the identification of the given design parameters to determine for each experiment set its biologically most meaningful analysis strategy. This step is extremely crucial, because every analysis needs to focus on the specific treatment factors of an experiment. The alternative of performing simply all possible comparisons will provide meaningless results for many experimental designs, because it would generate a large number of illegitimate contrasts between biologically incomparable samples. In order to define reasonable analysis strategies for public GeneChip microarray expression data sets, all their replicates and the most useful sample comparisons need to be determined manually to provide the proper experimental design parameters to the downstream statistical methods for identifying DEGs. The MIAME and MGED Ontology annotations (Brazma et al., 2001; Whetzel et al., 2006) of the public microarray depositories provide the essential information about the experiments, but efficient facilities to completely automate the DEG analyses on a large scale are not available at this point.

To perform large-scale DEG analysis of public expression data, we chose for this study a human-supervised analysis strategy, in which we determined for each experiment set its optimum
analysis parameters. The goal of this analysis was to identify all PUF and PKF genes that respond to specific or a wide range of conditions by enumerating their significant expression modulations in the corresponding experiment classes. For this, the available experiment annotations were manually evaluated and the most reasonable set of sample comparisons were recorded in an experiment definition table that contained all the required input parameters to control the downstream statistical DEG analysis in an automated manner (Supplement S2). Typically, we chose for each experiment set a design strategy that focused the analysis on the primary treatment as the main experimental factor. Multifactorial analysis strategies were avoided as much as possible. For instance, when an experiment contained a stress treatment as the primary experimental factor and time or different tissue types as secondary factors, then we compared only samples from identical tissues that were collected at the same time points. Additionally, comparisons between different experiment sets were not considered to exclude unknown variables, such as sample handling differences between laboratories (Hong et al., 2006). It is important to stress here, that depending on the design of a given experiment and its available annotations, it is often difficult to select a single most meaningful analysis strategy. Thus, our chosen strategy may not provide a perfect solution for every experiment set, but it represents a practical and reasonable compromise for performing systematic DEG analyses on large expression data sets from public databases.

In total our large-scale DEG analysis survey included 333 comparisons between samples with 2-4 technical or biological replicates from 41 experiment sets of 6 experiment categories. Table III provides an overview of the corresponding sample and experiment sets, and Supplement S2 contains all detailed information including the chosen analysis strategies for these data sets. Since the abiotic stress category is by far the largest data set, containing 524 chip hybridizations of 254 biosamples (Kilian et al., 2007), the following description of our DEG results will be restricted to this most comprehensive treatment category (Table VI). The data for the other categories are provided in the online database of this project (see below). As the statistical method for identifying DEGs with the determined experiment analyses strategies, we used Linear Models for Microarray Data (LIMMA) from Smyth (2004, 2005) using in all cases as confidence threshold a false discovery rate (FDR) of \( \leq 0.01 \) in combination with a minimum fold-change filter of 2.

Applying the above DEG analysis strategy, we were able to identify 269 PKF and 104 PUF genes that showed expression changes in the majority of the ten considered abiotic stress categories (Figure 3, Supplement S7). This set of a total of 373 generic stress DEGs was determined by filtering the generated DEG data set for members that showed one or more significant expression changes in at least 80% of all stress categories. Interestingly, 95% of these DEGs also appear in the generated gene expression clusters of the previous analysis (Supplement S5). The subsequent GO term enrichment analysis revealed that stress-related annotations are highly over-represented in this group of DEGs (see Supplement S8). About 48 of its members (13%) are associated with the GO term “response to stress” from the BP ontology (GO:0006950, p-value: \( 2.0 \times 10^{-13} \)). This enrichment indicates that our strategy has a high selectivity for identifying stress-response genes. Therefore, many PUF encoding genes in this data set may be directly or indirectly involved in generic stress response pathways. Among the different groups of identified stress responsive genes (see below and Figure 3), the generic stress DEG set represents by far the largest group.

Similarly, other studies have shown that stress-regulated genes frequently exhibit expression changes to a wide range of different abiotic stress treatments rather than a refined subset of stresses (Rodriguez and Redman, 2005; Kilian et al., 2007). The group of generic stress DEGs contains 48 genes that are annotated as transcription regulators in the MF ontology (GO:0030528, p-value: \( 2.5 \times 10^{-3} \), Supplement S8). This enrichment emphasizes the central role of transcription factors for the control of many stress response pathways. Moreover, it opens the possibility that several of the 104 PUF genes of this data set may be involved in similar transcription control processes.
We also used the generated abiotic stress DEG data set for identifying genes that respond predominantly to a specific type of stress. These specific stress DEGs were defined as follows. Firstly, they had to show in 25% of all comparisons of a given stress type significant changes. Secondly, they had to exhibit at the same time at least four times as many changes than in the other nine stresses (Figure 3, Supplement S7). This frequency-based filtering approach appeared to be more efficient for associating DEGs with specific stresses than overly strict filtering methods. This is the case because most stress response genes are not highly specific for a single type of stress (Kilian et al., 2007). As a result, strict filtering for genes responding only to a single stress will fail to identify any candidate genes in our comprehensive data sets. It is important to emphasize here that the chosen filtering approach is a practical compromise, but not a perfect solution to the problem of assigning DEGs reliably to different stress types. Therefore, the complete DEG results are provided in Supplement S7 where users can apply their own custom filters and prioritize strategies.

With the chosen frequency filter we were able to identify specific stress DEG sets within six of the ten treatment types (Table VI, Figure 3). The data sets for the stress treatments - light, oxidative and wounding stress - did not contain any genes that meet our filtering criteria, and the drought data set contained only a single member. The lack of specific stress DEGs in these data sets indicates that the genome-wide expression response patterns to these four stresses widely overlap with those from other stresses. For the remaining six treatment categories we identified in total 608 PKF and 206 PUF genes that responded predominantly to single stresses. The functional analysis of these specific stress DEG sets with our GO term enrichment approach showed no outstanding enrichment of specific gene functions. Instead, the results contained mostly moderately enriched GO annotations from a wide spectrum of molecular and biological processes (Supplement S8). Similar to the generic stress data, the different groups of specific stress DEGs included various marker genes that are characteristic for stress-related gene sets. For instance, they contained many genes that are annotated with the GO term ”response to stress” (see Figure 3). This term is significantly enriched in the heat stress data set (p-value: 1.3*10^{-2}), while the other five treatment sets contain it with considerable, but not significantly enriched frequencies (p-values ≥ 5*10^{-2}). In addition, the heat stress and genotoxic stress data sets showed the expected enrichment of genes that are associated with heat response and DNA repair processes, respectively (GO:0009408, p-value: 4.9*10^{-3} and GO:0006281, p-value: 6.1*10^{-5}).

In summary, the above large-scale DEG study identified a comprehensive set of candidate PKF and PUF genes that are involved in generic and specific stress response pathways. These results suggest the existence of one or more abiotic stress response regulons in Arabidopsis similar to the environmental stress regulon (ESR) described in yeast (Gasch et al., 2000; Gasch, 2002). Furthermore, the generated data sets represent an important resource for other scientists, who are interested in addressing more specific questions relevant to abiotic stress research by querying the generated DEG information in alternative ways (see Supplement S7 & online database).

**Plant Unknown-eome and Gene Expression Databases**

To provide efficient access to the extensive data sets of this study, we have developed two publicly available online portals: the Plant Unknown-eome Database (POND, http://bioweb.ucr.edu/scripts/unknownsDisplay.pl) and the Plant Gene Expression Database (PED, http://bioweb.ucr.edu/PED). The POND interface provides query and download options for the latest PUF sets from Arabidopsis. Their predictions are based on the three search methods used for this study: (1) BLASTP searches against the PKFs from Swiss-Prot, (2) HMM searches against the Pfam domain database and (3) retrieval of the ‘unknown’ annotations from the Gene Ontology system (MF).

The PED integrates our diverse co-expression data with a variety of online tools for user-friendly DEG analysis, cluster visualization and data mining (Figure 4). The aim of this service is not to duplicate or compete with the excellent web resources that are already available for array-based
expression data from plants, such as GEO, Genevestigator, BAR, AtGenExpress, ATC, PageMan, CSB.DB and MetNet (Barrett et al., 2006; Grennan, 2006; Zimmermann et al., 2004, 2005; Toufighi et al., 2005; Schmid et al., 2005; Jen et al., 2006; Usadel et al., 2006; Steinhauser et al., 2004b; Yang et al., 2005). Instead PED complements the available resources by providing a subset of the publicly available Affymetrix expression data from Arabidopsis in pre-analyzed form using various statistical methods for DEG identification combined with expression cluster information for co-regulation analysis. To provide high-confidence data, the database is restricted to data sets with two or more replicates. The following text provides a brief overview of the most interesting features of the database.

All expression data in PED were normalized with the RMA and MAS5 algorithms (Irizarry et al., 2003; Qin et al., 2006). The incorporation of the expression values from both normalization methods increases the utility spectrum of the provided data sets. The quantile-based RMA method generates more accurate expression measures for weakly expressed genes, whereas the MAS5 scaling approach is more appropriate for comparisons between expression studies (Lim et al., 2007). The option to identify DEGs by statistical modeling is a very unique feature of this online service. For this, PED provides the results of experiment design-based expression changes from several statistical methods, such as LIMMA (Smyth, 2004, 2005). The corresponding experiment analysis strategies are available for online viewing and download. A combinatorial query page allows searching for DEGs by specific treatments and filtering by various quantitative values to obtain candidate gene lists with strategies that resemble typical microarray analysis routines. Furthermore, the expression intensity and DEG data in PED are fully integrated with a comprehensive set of gene co-expression data from correlation and cluster analyses. To identify for a gene of interest its most positively or negatively co-regulated neighbors, the interface contains a correlation tool that provides for every gene on the arrays the Pearson and Spearman correlation profiles against all other genes. Information on discrete expression clusters is combined with the correlation data. It contains the four separate HTC cluster data sets that were generated by this study using as distance measures the two correlation coefficients in their signed and absolute forms (see previous section). An expression profile plotting tool is available for evaluating the quality of expression clusters or visualizing the expression patterns for custom gene sets across all samples in the database. This utility offers convenient options for inspecting the vast number of expression clusters of this study efficiently. Extensive download options for imports into local spreadsheet programs are available on all query levels for intensity, DEG, correlation and cluster data.

While the backend of the database is based on PostgreSQL and the web interface is implemented in Java, the framework of data analysis and online tools is largely designed around R and BioConductor utilities (R Development Core Team, 2006; Gentleman et al., 2005). The latter design feature will allow us to routinely add to PED’s online services in the future additional useful tools from the wide spectrum of statistical data analysis packages that are provided by the R open source community.
Conclusion
We present here one of the most comprehensive gene co-regulation studies that are currently available for Arabidopsis. Our study is unique by focusing on the analysis on PUF genes and their systematic association with functional annotations of PKF genes. By applying a combination of genome-wide cluster and DEG analysis methods, we identified many interesting groups of potentially co-regulated genes from a wide range of biological processes and stress response pathways. This approach allowed us to assign 1,541 PUF genes to relative specific and functionally informative GO terms. These gene associations provide a valuable resource for guiding future functional characterization experiments of PUF and PKF genes. In addition, the developed large-scale expression data analysis methods and the associated database represent important components of a future open-source framework for other scientists who are interested in performing similar studies, or utilizing public gene expression resources more efficiently. Finally, users of the provided data sets should keep two limitations in mind. First, the generated associations are hypotheses and not final proofs of gene functions. Second, even the most careful statistical approaches for large-scale data can only reduce, but not fully eliminate errors in the decision making processes associated with the interpretation of microarray data.
Material and Methods

Sequence Similarity and Domain Searches
Sequence similarity searches of the Arabidopsis proteome against the SwissProt database were performed with the BLASTP program (Altschul et al., 1997) using an E-value of $1 \times 10^{-6}$ as cutoff and the default settings for the remaining parameters. The Arabidopsis protein sequences were obtained from the TAIR site (version 7 release, ftp://ftp.arabidopsis.org/home/tair/Sequences) and the SwissProt sequences (Wu et al., 2006) were downloaded from the ExPASy site (release 54.4, ftp://ftp.expasy.org/databases/uniprot). To query only the functionally characterized protein space, all entries annotated as sequences of unknown function were removed from the SwissProt data set.

To identify protein domains of known function in the above Arabidopsis proteins, domain searches against the hidden Markov models of the Pfam database (Bateman et al., 2004) were performed with the HMMPFAM program (Eddy, 1996) using an E value of $1 \times 10^{-2}$ as cutoff. The global models of the Pfam release 22 were used for these searches (ftp://ftp.sanger.ac.uk/pub/databases/Pfam/). Matches against domains of unknown function (DUF) were ignored in the post-processing of the search results in order to identify only candidate sequences with domains of known functions.

GO Analysis
The Arabidopsis gene-to-GO mappings from TAIR/TIGR were used for all GO analysis steps of this study. They were downloaded from the Gene Ontology site (10-12-2007 release, http://geneontology.org). Direct assignments to the root node of each ontology were considered as unknown function annotations. These root assignments, in combination with the evidence code ND (No biological Data available), are the new official GO terms for sequences of unknown function. The former terms, molecular function unknown (GO:0005554), biological process unknown (GO:0000004) and cellular component unknown (GO:0008372), were discontinued by the consortium on 10-17-2006. In the subsequent GO term enrichment analysis steps, the new unknown annotations to the root were considered as artificial terminal annotations. This was necessary, because the root node is connected with all other genes in the GO network, which makes it impossible to obtain for the new unknown annotations meaningful enrichment data with most GO analysis approaches. This modification does not affect the results for any of the other GO nodes.

The hypergeometric distribution was used to test gene sets for the over-representation of GO terms. To perform this test, we developed a set of modular functions using the R language for statistical computing for their implementation (R Development Core Team, 2006). The corresponding GOHyperGAll script computes for a given sample population of genes the enrichment test for all nodes in the GO network, and returns raw and adjusted p-values. As an adjustment method for multiple testing, it uses the Bonferroni method according to Boyle et al. (2004). GOHyperGAll is based on the GOSTats package (Falcon and Gentleman, 2007) from the BioConductor project (Gentleman et al., 2005), and it provides similar utilities as the hyperGTest function included in this package. The main differences of our method are that it simplifies the usage of custom gene-to-GO mappings, and it contains various utilities for efficiently analyzing large numbers of gene sets from cluster analyses in batch mode. All functions of the GOHyperGAll script are available in Supplement S9.

Microarray Analysis
A total of 1,310 Affymetrix raw data Cel files were downloaded from the AtGenExpress and GEO sites (Schmid et al., 2005; Barrett et al., 2006; Kilian et al., 2007). All of them are derived from the Affymetrix ATH1 gene GeneChip microarray for Arabidopsis, and the corresponding samples contained at least two replicate samples. A summary of the utilized experiment sets is provided in
Table III, while a detailed description of the analyzed data with their experimental design parameters is provided in Supplement S2. The required probe set-to-locus mappings for the ATH1 chip were obtained from TAIR (ftp://ftp.arabidopsis.org/home/tair/Microarrays/Affymetrix, version 2-5-2007). All ambiguous probe sets on this chip were treated in the gene enumeration steps of this study in the following manner: controls and probe sets matching no or several loci in the Arabidopsis genome were ignored in the downstream analysis steps. In addition, redundant probe sets that represent the same locus several times were counted only once.

The normalization of the raw data Cel files was performed in R using the MAS5 and RMA algorithms, that are implemented in the affy package form the BioConductor project (Irizarry et al., 2003, 2006; Qin et al., 2006). To allow in the DEG analysis comparisons between the different samples of an experiment set, the RMA normalization was performed in batches for entire experiments sets (Table III). This batch normalization is only required for the quantile-based RMA approach, but not for the MAS5 scaling approach. The present call information of the non-parametric Wilcoxon signed rank test was computed with the affy package to estimate the amount of unexpressed genes (Liu et al., 2002; McClintick and Edenberg, 2006). The obtained expression values from both normalization methods were uploaded to the PED database.

For the DEG analysis, the replicates and the most appropriate sample comparisons were determined manually for each experiment set. The generated analysis strategies were recorded in experiment definition tables (Supplement S2). These tables were used to control the downstream DEG analysis steps in an automated manner by providing all information on replicates and sample comparisons to the statistical test methods. The actual analysis of DEGs was performed with the LIMMA package from Smyth (2004, 2005). The Benjamini & Hochberg method was selected to adjust p-values for multiple testing and to determine FDRs (Benjamini and Hochberg, 1995). As confidence threshold we used an adjusted p-value of \( \leq 0.01 \) in combination with a minimum fold-change filter of 2. All DEG analyses were performed on both the MAS5 and RMA normalized data sets. While both DEG analysis results were uploaded to the PED database, only the RMA set is discussed in this study, because the RMA algorithm provides more accurate measurements on weaker expressed genes (Qin et al., 2006).

**Cluster Analysis**

The correlation and cluster analysis steps were performed in R on the MAS5 normalized expression data set. For this, the mean values from replicated biological measurements were combined in one large expression matrix. The RMA data were not used for cluster analysis, because they are less reliable for correlation studies than MAS5 data (Lim et al., 2007). The Pearson and Spearman correlation coefficients were calculated with the cor function in R. The obtained correlation coefficients were transformed into a correlation-based distance matrix after subtracting their values from 1. Four separate distance matrices were calculated for the Pearson and Spearman correlation coefficients in their signed and absolute forms. The matrices were passed on to the hclust function (Murtagh, 1985; R Development Core Team, 2006) that performs agglomerative hierarchical clustering. Complete linkage was used as cluster joining method.

In order to obtain from hierarchical dendrograms discrete clusters, we developed a new hierarchical threshold clustering (HTC) method for this project. This method identifies sub-clusters in dendrograms based on a minimum tolerable similarity cutoff between all cluster members. This is achieved by applying an all-against-all similarity test for the clusters from all possible sub-trees. At the same time, unique cluster memberships are maintained and all items in the processed dendrogram are assigned to clusters with one or more members. The corresponding HTC R script is available in Supplement S10. As cutoff we used for this cluster selection procedure a correlation coefficient of \( \geq 0.6 \). This cutoff was chosen because it resulted in the highest enrichment of functionally related genes compared to alternative cutoffs settings (Supplement S4). As a result of this method, the members of every
identified cluster shared with all other members of the same cluster correlation coefficients between 0.6 and 1.0.
Acknowledgments

We thank the community projects - R, BioConductor, TAIR, TIGR, GEO and AtGenExpress - for providing the excellent software and data resources that were used by this project. This work was supported by grants from the National Science Foundation (2010 Program #0420033, #0420152, and IGERT Program DGE 0504249) and the National Institutes of Health (INBRE Program P20 RR-016464). TG acknowledges support from the Bioinformatics Core Facility, the Center for Plant Cell Biology (CEPCEB) and the Institute for Integrative Genome Biology (IIGB) at UC Riverside.
References


URL http://www.hubmed.org/display.cgi?uids=17028152


URL http://www.hubmed.org/display.cgi?uids=12582260


Figure Legends

**Figure 1: Relative Amount of Detectable Genes.**
The relative amount of present calls is plotted for all genes (ALL), the PKF set and the PUF set using the five frequency intervals (bins): 0, 1-25, 26-50, 51-75 and 76-100% present calls. All experiment sets of this study were used for generating this plot. The complete present call data set for the individual experiment categories is available in Supplement Table S3.

**Figure 2: Cluster Distributions.**
The numbers of clusters (a) and genes (b) are plotted for the cluster size intervals (bins) that are given along the abscissa. Each set of four bars, from left to right, contains the data for the clustering results using PCC, absolute PCC, SCC and absolute SCC values as distance measures, respectively.

**Figure 3: Generic and Specific Stress DEGs.**
The number of PUF and PKF encoding genes are plotted that were identified as generic and specific stress DEGs. The values above the bars provide the corresponding numbers of genes that are currently annotated with the GO term "response to stress" (GO:0006950 in BP ontology). The different stress types are given along the abscissa. Genes responding to the majority of the 10 abiotic stresses were considered as generic stress DEGs (Generic), while those responding predominantly to a specific type of stress were classified as specific stress DEGs. The following filters were used for assigning genes to the two stress categories. (1) Generic stress-response genes are those that showed in at least 80% of all stress treatments one or more significant changes. (2) Whereas, specific stress-response genes are those that showed in ≥25% of all comparisons of a given stress significant changes, and exhibited there ≥4 times as many changes than in the other nine stresses. For both filters, the observed expression changes were only counted when they meet our confidence criteria of a FDR ≤0.01 and a fold change ≥2. The specific stress data for the four treatment sets - light, oxidative, drought and wounding - are not plotted here, because their data sets did not contain any DEGs that meet our specific stress criteria.

**Figure 4: Plant Gene Expression Database (PED).**
The outline illustrates important utilities of the database (URL: [http://bioweb.ucr.edu/PED](http://bioweb.ucr.edu/PED)).
Tables

Table I: Functional Classification by Gene Ontologies.
The numbers of protein coding loci from Arabidopsis are given for custom categories of evidence codes of the three gene ontologies: MF, Molecular Function, BP, Biological Process and CC, Cellular Component. A description of the evidence codes is available on the Gene Ontology project site (http://www.geneontology.org GO.evidence.shtml). The number of loci with annotations in Any of the three ontologies are given in the last two rows. The percentage values are calculated relative to the total number of protein coding genes represented in the three ontologies. The evidence codes are grouped into the following custom categories of functional assignments: Empirical data (IC, IDA, IGI, IMP, IPI, TAS), Large-Scale experiments (IEP, RCA, NAS, NR), Sequence similarity or feature predictions (IEA, ISS) and PUFs lacking functional data (ND). The column Missing accounts for genes that lack annotations within the listed ontologies.

<table>
<thead>
<tr>
<th></th>
<th>Empirical</th>
<th>Large-Scale</th>
<th>Sequence</th>
<th>PUFs</th>
<th>Missing</th>
</tr>
</thead>
<tbody>
<tr>
<td>MF</td>
<td>1,918</td>
<td>9,061</td>
<td>4,677</td>
<td>8,665</td>
<td>2,228</td>
</tr>
<tr>
<td>%</td>
<td>7</td>
<td>34</td>
<td>18</td>
<td>33</td>
<td>8</td>
</tr>
<tr>
<td>BP</td>
<td>3,731</td>
<td>4,777</td>
<td>4,462</td>
<td>10,194</td>
<td>3,385</td>
</tr>
<tr>
<td>%</td>
<td>14</td>
<td>18</td>
<td>17</td>
<td>38</td>
<td>13</td>
</tr>
<tr>
<td>CC</td>
<td>3,333</td>
<td>1,661</td>
<td>8,527</td>
<td>8,426</td>
<td>4,602</td>
</tr>
<tr>
<td>%</td>
<td>13</td>
<td>6</td>
<td>32</td>
<td>32</td>
<td>17</td>
</tr>
<tr>
<td>Any</td>
<td>5,837</td>
<td>11,005</td>
<td>12,851</td>
<td>14,071</td>
<td>0</td>
</tr>
<tr>
<td>%</td>
<td>22</td>
<td>42</td>
<td>48</td>
<td>53</td>
<td>0</td>
</tr>
</tbody>
</table>

Table II: PUF Identification by Different Methods.
The table provides a matrix representation of the number of PUFs determined by the three different identification methods: BLASTP searches against the SWP database, HMMpfam searches against Pfam and the GOMF approach from Table I. The amount of PUFs common between pairwise comparisons of methods are provided in the corresponding row and column intersects of the matrix. The numbers of PUFs identified by All three methods or by at least one of them (Any) are given in the last two rows, respectively. The percentage values are calculated relative to the total number of protein coding genes. The complete gene lists for the PUF sets are available in Supplement S1.

<table>
<thead>
<tr>
<th>Method</th>
<th>SWP</th>
<th>Pfam</th>
<th>GOMF</th>
</tr>
</thead>
<tbody>
<tr>
<td>SWP</td>
<td>8,681 (32%)</td>
<td>6,260 (23%)</td>
<td>5,456 (20%)</td>
</tr>
<tr>
<td>Pfam</td>
<td>8,272 (31%)</td>
<td>5,788 (21%)</td>
<td></td>
</tr>
<tr>
<td>GOMF</td>
<td>8,665 (32%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All</td>
<td>4,667 (17%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any</td>
<td>12,781 (47%)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table III: Analyzed Gene Expression Arrays.
The table provides an overview of the different categories of GeneChip microarray experiments (1st column) that were analyzed in this study. The following numeric columns contain the number of raw data (Cel) files, the amount of the corresponding biosamples (Samples), the number of performed comparisons in the DEG analysis (Comp) and the number of experiment sets (ExpSet) the raw data are derived from. A more detailed list of this data is available in Supplement S2.

<table>
<thead>
<tr>
<th>Category</th>
<th>Cel</th>
<th>Samples</th>
<th>Comp</th>
<th>ExpSet</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abiotic Stress</td>
<td>524</td>
<td>254</td>
<td>129</td>
<td>10</td>
</tr>
<tr>
<td>Biotic Stress</td>
<td>200</td>
<td>72</td>
<td>55</td>
<td>6</td>
</tr>
<tr>
<td>Chemical Treatment</td>
<td>99</td>
<td>46</td>
<td>35</td>
<td>9</td>
</tr>
<tr>
<td>Tissue &amp; Development</td>
<td>237</td>
<td>79</td>
<td>40</td>
<td>1</td>
</tr>
<tr>
<td>Genotype</td>
<td>86</td>
<td>29</td>
<td>28</td>
<td>4</td>
</tr>
<tr>
<td>Hormone Treatment</td>
<td>164</td>
<td>80</td>
<td>46</td>
<td>11</td>
</tr>
<tr>
<td>Sum</td>
<td>1310</td>
<td>560</td>
<td>333</td>
<td>41</td>
</tr>
</tbody>
</table>

Table IV: Overview of GO Term Enrichment Analysis.
The amount of clusters and genes are provided for different cluster prioritization filters that were applied to the GO term enrichment data of Supplement S6. The values in parentheses represent the corresponding number of clusters containing PUF genes and the number of PUF genes in these clusters, respectively. The first row contains the counts for the unfiltered data set that considered only clusters with ≥5 members. The subsequent rows refer to the counts after applying the following two-component filter with four different stringency settings. (1) To select clusters with enriched GO terms, the clusters had to contain one or more over-represented GO terms in at least one of the three ontologies based on the Bonferroni corrected p-values of the enrichment analysis. The four different p-value cutoffs used for this filter are given in the first column. (2) In addition, ≥20% of the cluster members needed to be associated with the selected GO term in order to favor functionally homogeneous clusters.

<table>
<thead>
<tr>
<th>Filter</th>
<th>Clusters</th>
<th>Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>916 (794)</td>
<td>11,077 (2,884)</td>
</tr>
<tr>
<td>0.05</td>
<td>519 (429)</td>
<td>6,262 (1,541)</td>
</tr>
<tr>
<td>0.01</td>
<td>373 (301)</td>
<td>4,893 (1,126)</td>
</tr>
<tr>
<td>0.001</td>
<td>212 (170)</td>
<td>3,315 (744)</td>
</tr>
<tr>
<td>1e-06</td>
<td>66 (53)</td>
<td>1,279 (277)</td>
</tr>
</tbody>
</table>

Table V

<table>
<thead>
<tr>
<th>CLID</th>
<th>CLSZ</th>
<th>PUF</th>
<th>Sample</th>
<th>P-value</th>
<th>Ont</th>
<th>GO Term</th>
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</thead>
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<tr>
<td>Reproduction</td>
<td>115</td>
<td>20</td>
<td>5</td>
<td>2</td>
<td>3.00E-06 BP</td>
<td>GO:0010344: seed oil body biogenesis</td>
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<td>-----</td>
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<td>------------</td>
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</tr>
<tr>
<td></td>
<td>115</td>
<td>20</td>
<td>5</td>
<td>11</td>
<td>0.014 CC</td>
<td>GO:0016020: membrane</td>
</tr>
<tr>
<td></td>
<td>115</td>
<td>20</td>
<td>5</td>
<td>4</td>
<td>4.30E-07 MF</td>
<td>GO:0045735: nutrient reservoir activity</td>
</tr>
<tr>
<td>Carbohydrate metabolism</td>
<td>95</td>
<td>21</td>
<td>4</td>
<td>4</td>
<td>5.90E-06 BP</td>
<td>GO:0006073: glucan metabolic process</td>
</tr>
<tr>
<td></td>
<td>95</td>
<td>21</td>
<td>4</td>
<td>8</td>
<td>1.80E-10 CC</td>
<td>GO:0005618: cell wall</td>
</tr>
<tr>
<td></td>
<td>95</td>
<td>21</td>
<td>4</td>
<td>4</td>
<td>1.90E-07 MF</td>
<td>GO:0005199: structural constituent of cell wall</td>
</tr>
<tr>
<td></td>
<td>131</td>
<td>18</td>
<td>1</td>
<td>6</td>
<td>1.10E-10 BP</td>
<td>GO:0006007: glucose catabolic process</td>
</tr>
<tr>
<td></td>
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<td>GO:0005739: mitochondrion</td>
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<tr>
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<td>GO:0004738: pyruvate dehydrogenase activity</td>
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<tr>
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<td>248</td>
<td>11</td>
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<td>3</td>
<td>9.30E-07 BP</td>
<td>GO:0005982: starch metabolic process</td>
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<td>2</td>
<td>9</td>
<td>1.70E-07 CC</td>
<td>GO:0009507: chloroplast</td>
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<td></td>
<td>248</td>
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<td>2</td>
<td>5</td>
<td>0.003 MF</td>
<td>GO:0016740: transferase activity</td>
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<tr>
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<td>300</td>
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<td>3</td>
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GO:0006412: translation
GO:0003735: structural constituent of ribosome
GO:0005840: ribosome

GO:0006412: translation
GO:0005576: extracellular region
GO:0045735: nutrient reservoir activity

GO:0019915: sequestering of lipid
GO:0005576: extracellular region
GO:0045735: nutrient reservoir activity

GO:0019374: galactolipid metabolic process
GO:0031967: organelle envelope
GO:0042578: phosphoric ester hydrolase activity

GO:0019915: sequestering of lipid
GO:0005576: extracellular region
GO:0045735: nutrient reservoir activity

GO:0045036: protein targeting to chloroplast
GO:0009536: plastid
GO:0003674: molecular function (PUF term)

GO:0045036: protein targeting to chloroplast
GO:0009941: chloroplast envelope
GO:0022804: transmembrane transporter activity

GO:0006810: transport
GO:0005794: Golgi apparatus
GO:0005215: transporter activity

GO:0006606: protein import into nucleus
GO:0005635: nuclear envelope
GO:0008565: protein transporter activity

GO:0006820: anion transport
GO:0005741: mitochondrial outer membrane
GO:0008308: voltage-gated ion channel activity

GO:0015979: photosynthesis
GO:0044436: thylakoid part
GO:0010242: oxygen evolving activity

GO:0015979: photosynthesis
GO:0004045: aminoacyl-tRNA hydrolase activity
GO:0004176: ATP-dependent peptidase activity

GO:0015979: photosynthesis
GO:0004950: tetrapyrrole binding
GO:0046906: tetrapyrrole binding

GO:0006091: generation of metabolites and energy
GO:0005739: mitochondrion
GO:0004449: isocitrate dehydrogenase activity

GO:0006902: photosynthetic electr. transport chain
555 5 1 3 3.30E-06 MF GO:0009055: electron carrier activity
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Cell organization and biogenesis
77 24 5 6 4.20E-15 BP GO:0009834: secondary cell wall biogenesis
77 24 5 3 0.011 CC GO:0031225: anchored to membrane
77 24 5 6 1.70E-05 MF GO:0016757: transferring glycosyl groups
108 18 7 2 0.00084 BP GO:0009831: cellulose and pectin modification
108 18 7 18 1.30E-09 CC GO:0016020: membrane
349 9 0 7 1.00E-15 BP GO:0009664: structural constituent of cell wall
349 9 0 6 1.70E-05 MF GO:0005199: structural constituent of cell wall
953 5 1 2 9.70E-07 BP GO:0010020: chloroplast fission
953 5 1 2 0.029 CC GO:0009507: chloroplast
953 5 1 4 0.044 MF GO:0005488: binding

Secondary metabolism
12 73 13 3 0.0076 BP GO:0046148: pigment biosynthetic process
12 73 13 34 1.30E-18 CC GO:0009536: plastid
12 73 13 3 0.00059 MF GO:0003746: translation elongation factor activity
143 17 2 4 8.70E-08 BP GO:0046148: pigment biosynthetic process
143 17 2 8 1.40E-05 CC GO:0009536: plastid
143 17 2 5 0.0023 MF GO:0016491: oxidoreductase activity
347 10 2 3 1.20E-07 BP GO:0009686: gibberellin biosynthetic process
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Response to stimulus
68 22 3 7 5.40E-07 BP GO:0006952: defense response
68 22 3 11 0.02 CC GO:0016020: membrane
68 22 3 5 1.10E-06 MF GO:0004888: transmembrane receptor activity
85 23 9 10 6.70E-18 BP GO:0009408: response to heat
85 23 9 10 0.21 CC GO:0005575: cellular component (PUF term)
85 23 9 2 0.043 MF GO:0005516: calmodulin binding
90 22 5 6 1.60E-09 BP GO:0009408: response to heat
90 22 5 5 0.04 CC GO:0005634: nucleus
90 22 5 3 0.00052 MF GO:0051082: unfolded protein binding
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346 10 3 9 6.20E-08 CC GO:0009536: plastid
346 10 3 3 0.57 MF GO:0003674: molecular function (PUF term)
356 9 9 8 1.10E-15 BP GO:0009733: response to auxin stimulus
356 9 9 3 0.021 CC GO:0043231: intracellular membrane-bound organelle
356 9 9 7 7.70E-05 MF GO:0003674: molecular function (PUF term)
480 8 1 3 3.80E-05 BP GO:0006979: response to oxidative stress
480 8 1 7 1.10E-08 CC GO:0005739: mitochondrion
480 8 1 2 7.20E-05 MF GO:0046933: hydrogen ion transporting ATP synthase
586 7 1 3 4.20E-05 BP GO:0009737: response to abscisic acid stimulus
586 7 1 2 0.00013 CC GO:0008287: serine/threonine phosphatase complex
Table V: GO Term Enrichment Data for Prioritized Clusters.
The GO annotations for the most conservative cluster prioritization filter from Table IV are provided. The three filtering criteria for selecting the presented clusters are described in the previous legend. Based on space and readability considerations, only the highest ranking GO term within each ontology.
is included here. As a result of our prioritization criteria, every cluster listed has at least one GO term assigned that meets both, the enrichment (p-value ≤ 10^{-6}) and uniformity (≥ 20%) criteria. If an ontology did not contain a GO term passing these filters then the candidate with the lowest p-value was chosen. GO slim terms are used as table subtitles to organize the clusters based on a general biological process classification schema. The different columns provide the identifiers of each cluster (CLID), the number of genes (CLSZ), the number of PUF genes, the number of genes matching a given GO term (Sample), the Bonferroni corrected p-value of the hypergeometric distribution test (P-value), the ontology type (Ont) and the corresponding GO Term, respectively. The complete list of enriched GO terms and the associated gene identifiers for these clusters are available in Supplement S6.

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**Table VI: Abiotic Stress Treatments.**
The table provides an overview of the different types of abiotic stress experiment sets (Stress) that were used in the DEG analysis of this study. The numeric columns contain the number of the analyzed GeneChip microarrays (Stress), the number of the corresponding biosamples (Samples) and the number of the performed comparisons (Comp). A more detailed list of this data is available in Supplement S2.

**Supplemental Data**
S1: PUF sets (Excel table)
S2: GeneChip R microarray experiments and analysis strategies (Excel table)
S3: PMA data (Excel table)
S4: HTC cutoff selection (PDF)
S5: Cluster data (Excel table)
S6: GO analysis of clusters (Excel table)

S7: DEG analysis (Excel table)

S8: GO analysis of DEGs (Excel table)

S9: R script for GO term enrichment analysis (text file)

S10: R script for HTC clustering (text file)
### Summary Table

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Complete correlation profile against all genes to identify positively and negatively co-expressed neighbors

### Profile Viewing Tool

[Graph showing gene expression profiles over time]