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The Wheat GENIE3 Network Provides Biologically-Relevant Information in Polyploid Wheat

Sophie A. Harrington,* Anna E. Backhaus,* Ajit Singh,† Keywan Hassani-Pak,† and Cristobal Uauy*,1

*Department of Crop Genetics, John Innes Centre, Norwich Research Park, Norwich, NR4 7UH, United Kingdom and
†Computational and Analytical Sciences, Rothamsted Research, Harpenden, Hertfordshire, AL5 2JQ, United Kingdom

ABSTRACT Gene regulatory networks are powerful tools which facilitate hypothesis generation and candidate gene discovery. However, the extent to which the network predictions are biologically relevant is often unclear. Recently a GENIE3 network which predicted targets of wheat transcription factors was produced. Here we used an independent RNA-Seq dataset to test the predictions of the wheat GENIE3 network for the senescence-regulating transcription factor NAM-A1 (TraesCS6A02G108300). We re-analyzed the RNA-Seq data against the RefSeqv1.0 genome and identified a set of differentially expressed genes (DEGs) between the wild-type and nam-a1 mutant which recapitulated the known role of NAM-A1 in senescence and nutrient remobilisation. We found that the GENIE3-predicted target genes of NAM-A1 overlap significantly with the DEGs, more than would be expected by chance. Based on high levels of overlap between GENIE3-predicted target genes and the DEGs, we identified candidate senescence regulators. We then explored genome-wide trends in the network related to polyploidy and found that only homeologous transcription factors are likely to share predicted targets in common. However, homeologs which vary in expression levels across tissues are less likely to share predicted targets than those that do not, suggesting that they may be more likely to act in distinct pathways. This work demonstrates that the wheat GENIE3 network can provide biologically-relevant predictions of transcription factor targets, which can be used for candidate gene prediction and for global analyses of transcription factor function. The GENIE3 network has now been integrated into the KnetMiner web application, facilitating its use in future studies.

KEYWORDS Gene networks
GENIE3
Wheat
Senescence
RNA-Seq
Polyploidy

Transcriptional regulation of gene expression is fundamental to all biological processes. Increasingly, this is studied using large-scale datasets obtained from RNA-Sequencing (RNA-Seq) experiments across multiple tissues, genotypes, treatments, and timepoints. As library preparation and sequencing costs fall, more and more RNA-Seq datasets are being published, providing a wealth of transcriptional information. These datasets are then available for integration into large-scale gene regulatory networks covering various biological conditions. In wheat (*Triticum aestivum*), genomic and transcriptomic analysis has historically been hampered by its large, repetitive polyploid genome (Appels et al. 2018). More recently, a high-quality genome and gene annotation has facilitated transcriptomics work in wheat (Appels et al. 2018; Ramirez-González et al. 2018). This has allowed the use of substantial RNA-Seq datasets to build gene regulatory networks and predict transcription factors involved in complex processes such as senescence (Borrill et al. 2019) and grain development (Pfeifer et al. 2014; Chi et al. 2019). However, these studies typically use bespoke RNA-Seq datasets to generate the regulatory networks, rather than exploiting publicly-available data.

Recent efforts to standardize and manually curate public RNA-Seq datasets [8], combined with the publication of large datasets which span wheat development [2], have resulted in unprecedented availability of wheat transcriptome data. These large, curated datasets can now be mined to build new gene regulatory networks covering
many biological processes in wheat. Using 850 RNA-Seq datasets combined from multiple independent studies, gene co-expression networks for root, leaf, spike, and grain tissues, as well as abiotic and biotic stresses, were developed (Ramírez-González et al. 2018). This same study also generated a network of predicting transcription factor - target relationships using the 850 independent RNA-Seq samples from a wide range of developmental, tissue, genotypes and stress conditions (Ramírez-González et al. 2018). This network was created with the Gene Network Inference with Ensemble of trees 3 (GENIE3) algorithm, which uses a Random Forests approach to predict the strength of putative regulatory links between target genes and their putative regulators (i.e., transcription factors) (Huynh-Thu et al. 2010). The program produces a ranked list output of each pairwise comparison ranked from the most confident to the least confident regulatory connection. GENIE3 was able to recapitulate known genetic regulatory networks in Escherichia coli when first tested. Further testing demonstrated that the GENIE3 network was able to predict known transcription factor-gene regulatory connections in a Drosophila melanogaster dataset (Huynh-Thu and Geurts 2018). Intriguingly, in this same study the GENIE3 network was not able to predict gold-standard interactions in a Saccharomyces cerevisiae dataset, performing no better than random. This suggested that the relative performance of the GENIE3 network is data-dependent, highlighting the importance of testing the outputs of a GENIE3 network before utilizing it for de novo predictions. Since its introduction, the GENIE3 algorithm has been used to identify tissue-specific gene regulatory networks in maize (Huang et al. 2018) and key regulatory genes in glaucoma (Chen et al. 2017), as well as to study the drought response in sunflower (Marchand et al. 2014). Notably, previous studies have integrated the GENIE3 network predictions with ChIP-Seq and other proteomic and transcriptomic data and found that the GENIE3 predictions do correspond with independent biological datasets (Walley et al. 2016; Huang et al. 2018).

Here we have conducted a series of analyses to investigate whether the GENIE3 network provides biologically-relevant information in polyploid wheat. As a first case study, we re-analyzed the RNA-Seq datasets from Pearce et al. (Pearce et al. 2014) which examined gene expression of the NAC transcription factor NAM-A1. This transcription factor is known to affect monocarpic senescence and nutrient remobilisation in polyploid wheat, affecting gene expression even before visual signs of senescence can be observed (Ramírez-González et al. 2018). This network was created with the Gene Network Inference with Ensemble of trees 3 (GENIE3) algorithm, which uses a Random Forests approach to predict the strength of putative regulatory links between target genes and their putative regulators (i.e., transcription factors) (Huynh-Thu et al. 2010). The program produces a ranked list output of each pairwise comparison ranked from the most confident to the least confident regulatory connection. GENIE3 was able to recapitulate known genetic regulatory networks in Escherichia coli when first tested. Further testing demonstrated that the GENIE3 network was able to predict known transcription factor-gene regulatory connections in a Drosophila melanogaster dataset (Huynh-Thu and Geurts 2018). Intriguingly, in this same study the GENIE3 network was not able to predict gold-standard interactions in a Saccharomyces cerevisiae dataset, performing no better than random. This suggested that the relative performance of the GENIE3 network is data-dependent, highlighting the importance of testing the outputs of a GENIE3 network before utilizing it for de novo predictions. Since its introduction, the GENIE3 algorithm has been used to identify tissue-specific gene regulatory networks in maize (Huang et al. 2018) and key regulatory genes in glaucoma (Chen et al. 2017), as well as to study the drought response in sunflower (Marchand et al. 2014). Notably, previous studies have integrated the GENIE3 network predictions with ChIP-Seq and other proteomic and transcriptomic data and found that the GENIE3 predictions do correspond with independent biological datasets (Walley et al. 2016; Huang et al. 2018).

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MATERIALS AND METHODS

GENIE3

The GENIE3 network was previously published in (Ramírez-González et al. 2018) and made available at https://doi.org/10.5447/ipk/2018/7. In brief, it utilized a set of 850 publicly-available RNA-Seq samples in a Random Forests approach to predict targets of wheat 3,384 transcription factors (Huynh-Thu et al. 2010). The top one million connections in the network were used for all analyses in the paper, consistent with previous studies (Huang et al. 2018; Ramírez-González et al. 2018).

RNA-Seq analysis

Mapping: Publicly-available reads from (Pearce et al. 2014) were downloaded from https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE60635. Reads from the wild-type and nam-a1 (ggc-a1) mutant lines from 12 and 22 days after anthesis (DAA) were pseudoaligned with kallisto (v 0.43.1) (Bray et al. 2016) against the v1.1 annotation from the RefSeq genome v1.0 (Appels et al. 2018) using standard settings for single reads (-single -b 30 -l 200 -s 20) (Supplementary Table 1). Only the A and B genomes of RefSeqv1.0 were used for the pseudoalignment, as the raw reads were derived from tetraploid cv. Kronos plants.

Differential expression analysis: Gene expression levels (transcript per million, TPM) were determined using the R package Sleuth (Pimentel et al. 2017) using the default settings for the Wald test (sleuth_wt; v 0.30.0). Sleuth utilizes a general linear model to estimate the biological variance between replicates not due to technical factors, where the bootstraps produced by kallisto are treated as proxies for true technical replicates. We compared the expression of genes between the wild-type and nam-a1 mutant samples at 12 and 22 DAA. We used the cut-off of q < 0.05 (unless otherwise stated) to identify differentially expressed genes, where q is the p-value adjusted for false discovery rate using the Benjamini-Hochberg correction (Benjamini and Hochberg 1995). The list of differentially expressed genes for each timepoint is reported in Supplementary Table 2.

In order to compare our identified differentially expressed genes with those originally identified in (Pearce et al. 2014), we re-analyzed the original set of differentially expressed genes using the same four statistical approaches—DESeq (Anders and Huber 2010), edgeR (Robinson et al. 2009), the Mann-Whitney-Wilcoxon test, and the Student’s T-test—at different p-value cut-offs. We applied the same adjustment as used in (Pearce et al. 2014), where the DESeq and edgeR p-values were adjusted using the Benjamini-Hochberg method (Benjamini and Hochberg 1995) to account for false discovery rate. For consistency with the original paper, we refer to the statistical significance of all four tests as a p-value, though the DESeq and edgeR p-values are comparable to the adjusted q-value generated by Sleuth.

Methods for ID conversion and comparison: The genes and contigs identified as differentially expressed in the original (Pearce et al. 2014) study were converted to RefSeqv1.1 gene models where possible using BLASTn (Altschul et al. 1990). Briefly, the differentially expressed sequences were extracted from the IWGSC CSS genome (IWGSC 2014) and compared with BLASTn (v 2.2.33 -num_alignments 1 -outfmt 6) against the RefSeq v1.1 transcriptome (including both high and low confidence gene models). The BLAST hit with the greatest percent identity to the original CSS sequence was assigned as the equivalent RefSeqv1.1 gene model.

Comparison of the diﬀerentially expressed genes with GENIE3

Calculation of shared ratios: We calculated the level of overlap or shared genes between different transcription factors or datasets as follows:

\[
\frac{|A \cap B|}{|A|}
\]
Where A and B are sets of genes, and |A| < |B|.

For example, given two sets of genes A and B, where A contains 5 genes and B contains 10 genes, if they share two genes in common the shared ratio is 2/5, or 0.4.

This calculation was carried out as implemented in R scripts available at https://github.com/Uauy-Lab/GENIE3_scripts. It was used to create the distribution of shared targets between transcription factors and the differentially expressed genes, as well as between the predicted targets of any two transcription factors.

**Distributions of shared ratios:** Initially, we analyzed the shared ratios between transcription factors in the GENIE3 network and the set of differentially expressed genes obtained from the re-analyzed NAM-A1 RNA-Seq data. The target genes of 1000 randomly selected transcription factors were compared against the differentially expressed genes at 12 and 22 DAA to obtain the distribution of shared ratios (Figure 1B). This calculation was also carried out individually for the targets of NAM-A1 against both timepoints.

Following this, the shared ratio was calculated for 1000 randomly selected pairs of transcription factors from the GENIE3 network (see Figure 3A). The same analysis was carried out for individual transcription factor super-families, based on the family assignments from https://opendata.earlham.ac.uk/wheat/under_license/toronto/Ramirez-Gonzalez_et_al_2018-06025-Transcriptome-Landscape/data/data_tables/transcription_factors_to_use_high_confidence.csv (Ramírez-González et al. 2018). All pairs selected were unique, and where a transcription factor family was not large enough to contain 1000 unique pairs, the maximum number of unique pairs was sampled (e.g., in the family CCAAT_HAP3, N = 3 and thus the number of unique pairs sampled was 6). This calculation was also carried out for all homeolog pairs, where triads were classified as in https://opendata.earlham.ac.uk/wheat/under_license/toronto/Ramirez-Gonzalez_et_al_2018-06025-Transcriptome-Landscape/data/TablesForExploration/Triads.rds (Ramírez-González et al. 2018). The subset used for this analysis only included syntenic 1:1:1 triads (see Ramírez-González et al. 2018 for definition), resulting in a total of 708 triads and 2,124 individual genes.

**Movement ratios:** The shared ratios of homeologous pairs were also distinguished by movement classifications, as defined previously (Ramírez-González et al. 2018). In brief, triads were classified into three categories (“Dynamic”, “Middle 80”, and “Stable”) based on variation in their homeolog expression bias across tissues. Dynamic triads show more variation in relative homeolog expression levels across tissues than stable triads. The assignment of triads to each category is found here: https://opendata.earlham.ac.uk/wheat/under_license/toronto/Ramirez-Gonzalez_et_al_2018-06025-Transcriptome-Landscape/data/Triad_Subsets_Movement/. Triads in the “HC_CS_no_stress_movement_top_10pc.txt” file were defined as Dynamic, the “HC_CS_no_stress_movement_middle_80pc.txt” as Mid 80, and the “HC_CS_no_stress_movement_low_10pc.txt” as Stable. This analysis was implemented in “Gene3_Statistics_SharedRatios_Homoeologs_MovementCategories_Figure4_SuppFig3.Rmd” at https://github.com/Uauy-Lab/GENIE3_scripts.

**Developmental expression datasets:** Public datasets were used for the expression analysis in Figure 2. The developmental time course was first published in (Ramírez-González et al. 2018), from the spring wheat cv. Azhurnaya. This dataset was included in the generation of the GENIE3 network. The senescence-specific time course was first published in Borrill et al. (2019), from the spring wheat cultivar Bobwhite, and was not included in the GENIE3 network.

**Gene Ontology (GO) term analysis**

GO-term enrichment analysis was carried out as previously described in (Borrill et al. 2019), using the GOSeq (v 1.3.4.1) package in R (Young et al. 2010).

**Data visualization, manipulation, and statistical analyses**

Graphs were made in R, principally using the ggplot2 (v 3.1.1)(Wickham 2016) and ggpubr (v 0.2)(Kassambara 2019) packages as well as the “heatmap” function of the NMF package (v. 0.21.0)(Gaujoux and Seoighe 2010). Networks in Figures 3B and C were visualized using Cytoscape (v 3.7.1) (Shannon et al. 2003). Data manipulation was also carried out in R, using the packages dplyr (v 0.8.0.1)(Wickham et al. 2019) and tidyr (v 0.8.3)(Wickham and Henry 2018) in scripts as linked throughout the methods. Statistical analyses were carried out in R, as detailed in the results section. The sign test was carried out using the R package BSDA (v 1.2.0) (Arnholt and Evans 2017).

**Data availability**

All scripts used for analysis in this paper are available on Github at https://github.com/Uauy-Lab/GENIE3_scripts. Links to the public datasets used in the analysis are included within the Materials and Methods, where appropriate, or are linked in the scripts on Github. Datasets from the Ramírez-González et al. 2018 paper are available at https://opendata.earlham.ac.uk/wheat/under_license/toronto/Ramirez-Gonzalez_et_al_2018-06025-Transcriptome-Landscape/. The original GENIE3 network is deposited at https://doi.org/10.5447/ipk/2018/7. Supplementary Table 1 contains Kallisto mapping statistics. Supplementary Table 2 contains the list of de novo differentially-expressed genes. Supplementary Table 3 contains the list of candidate senescence regulators, as shown in Figure 2. Supplementary File 1 contains Supplementary Figures 1-4. Supplementary File 2 contains the enriched GO terms for the GENIE3 targets of all transcription factor families and of the candidate senescence regulators (see Supplementary Table 3). Further information on the contents of this file is available in the README.txt file within Supplementary File 2. Supplemental material available at figshare: https://doi.org/10.25387/g3.11953728.

**RESULTS**

**RNA-Seq analysis**

In 2014, Pearce et al. analyzed the differences in gene expression between wild-type (WT) Kronos, a tetraploid wheat cultivar, and a NAM-A1 loss-of-function mutant (nam-a1 or gpc-a1) which contained a premature stop codon (W114*) (Pearce et al. 2014). Their analysis was carried out before the release of a complete wheat genome or gene annotation, relying upon a bespoke gene annotation. Here, we reanalysed the RNA-Seq datasets for the wild-type and nam-a1 single mutant lines at 12 and 22 DAA using the most recent wheat genome annotation and an updated alignment method (Appels et al. 2018). Reads were pseudoaligned to the A and B genomes of the RefSeqv1.1 transcriptome using kallisto (Bray et al. 2016), a software which has been shown to differentiate well between homeologs during alignment and is thus appropriate for use with polyploid wheat (Borrill et al. 2016; Ramírez-González et al. 2018). Each sample contained on average 35 million reads, with the exception of one sample with 85 million reads, of
which on average 78% were pseudoaligned to the transcriptome (Supplementary Table 1).

To identify genes differentially expressed between the WT and nam-a1 mutant at the two developmental timepoints (12 and 22 DAA) we used sleuth, a program designed for analysis of RNA-Seq experiments for which transcript abundances have been quantified with kallisto (Pimentel et al. 2017). Using an adjusted p-value cut-off of \(q < 0.05\), we identified 866 differentially expressed genes (DEGs) between WT and the nam-a1 mutant at 12 DAA and 130 DEGs at 22 DAA (Supplementary Table 2). This set of DEGs will be referred to as the de novo DEGs throughout the paper. We carried out gene ontology (GO) term enrichment analysis on the two sets of DEGs and found that DEGs at both timepoints are highly enriched for terms related to metal ion transport, including zinc, manganese, and copper (\(P < 0.001\), adjusted for false discovery rate; Supplementary File 2). This correlates closely with the findings from the original analysis, which found that GO terms related to transporter function were highly regulated by NAM-A1 (Pearce et al. 2014). This also supports previous physiological studies of the NAM genes which found them to be important in nutrient remobilisation and transport (Uauy et al. 2006; Waters et al. 2009).

We used BLASTn to compare the differentially expressed sequences from the Pearlce dataset to the RefSeqv11.1 transcriptome annotation which was used for the de novo analysis (Appels et al. 2018). We successfully assigned 453 out of 517 DEG identified by Pearce et al. to RefSeqv11.1 gene models (442/504 DEG at 12 DAA and 11/13 DEG 22 DAA). Based on this common nomenclature, we then directly compared the de novo DEGs identified by sleuth with the DEGs identified originally (Pearce et al. 2014). At 12 DAA, 177 of the 442 DEGs (40%) were present in the de novo differential expression set, whereas at 22 DAA, 7 of the 11 DEGs (64%) were found in the de novo set.

At both developmental time points, our reanalysis identified more transcripts as significantly differential expressed compared to the original study (Pearce et al. 2014). This is possibly due in part to the study being carried out before the advent of a complete wheat genome annotation, relying instead on mapping of transcriptome assemblies to incomplete chromosome contigs. We have shown previously that earlier gene annotations could be incomplete or mis-annotated (Brinton et al. 2018). However, this discrepancy in numbers of differentially expressed genes is most likely a result of the more liberal significance cut-off used in the current study, as compared to the stringent combination of four statistical tests which were used to reduce false positive discoveries in the original study (Pearce et al. 2014). Specifically, the earlier analysis combined pairwise comparisons of samples using the programs DESeq (Anders and Huber 2010) and edgeR (Robinson et al. 2009), both adjusted for false discovery rate using the Benjamini-Hochberg method (Benjamini and Hochberg 1995), in combination with the Mann-Whitney-Wilcoxon test and the Student’s t-test. For a gene to be considered differentially expressed, the \(p\)-value was required to be \(< 0.01\) for all tests except the Mann-Whitney-Wilcoxon, which utilized a cut-off of \(P < 0.05\). To test whether the more stringent cut-off was driving the reduced number of identified DE genes in the original analysis, we reanalysed the original data from Pearce et al. (2014) using a cut-off of \(P < 0.05\) for all four tests. Focusing on 12 DAA, with the highest proportion of DE genes, 837 genes were found to be differentially expressed under this cut-off, consistent with the 866 DEGs identified in our analysis. Conversely, if we increased the stringency of our re-analysis using a cut-off of \(q < 0.01\) in Sleuth, we recover only 509 DEGs, which is similar to the identification of 504 DEGs at 12 DAA by Pearce et al. (2014).

To determine the impact of the cut-off value on the calling of DEGs, we ranked the de novo DEGs by \(q\)-value and recorded the position of the 177 shared DEGs at 12 DAA. We found that the majority of shared DEGs (53%) ranked in the top quarter of the list of de novo DEGs (Supplementary Figure 1A). However, 9% of the common DEGs were found in the bottom quarter of the de novo DEGs. We then looked at the distribution of \(q\)-values across the universe of genes included in the RNA-Seq experiment. For both timepoints, there is a peak in the density of \(q\)-values around 0, representing the expected enrichment of differentially expressed genes (Supplementary Figure 1B). We found that by selecting a cut-off of 0.05, we could capture most of the peak present in the 12 DAA dataset. In contrast, selecting a cut-off of 0.01 would be too conservative, excluding much of the enriched peak. Together, this suggests that the cut-off value of \(q < 0.05\) is appropriate to maximize the identification of relevant DEGs.

The GENIE3 network predictions overlap with known DEGs

To investigate whether the GENIE3 network provides biologically relevant information, we compared the GENIE3 predicted targets for NAM-A1 against the list of differentially expressed genes identified between the wild-type and nam-a1 mutants from the de novo RNA-Seq analysis. As the RNA-Seq experiment was carried out in tetraploid wheat, we only considered target genes on the A or B genome. We focused on the 12 DAA and 22 DAA timepoints, which captured the onset and intermediate stages of senescence, respectively. At 12 DAA, we found that of the 79 genes predicted to be targets of NAM-A1 in the GENIE3 network, 12 were shared with the set of de novo DEGs (15.2%; Figure 1A). However, at 22 DAA only 4 of the 79 GENIE3 predicted targets were shared with the DEGs (5.1%; Figure 1A). The decrease in overlap between 12 and 22 DAA is consistent with NAM-A1 primarily acting early in senescence (Uauy et al. 2006; Pearce et al. 2014).

We then compared the lists of DEGs at 12 and 22 DAA against the targets of all 3,384 transcription factors included in the GENIE3 dataset (Figure 1B). The median number of shared targets between the DEGs and predicted targets of a given transcription factor was 0, with a maximum of 33.3%. Comparing the overlap between random transcription factors and the RNA-Seq dataset, we found a significantly higher level of overlap between the GENIE3-predicted targets of NAM-A1 and genes differentially expressed in the nam-a1 mutant at both timepoints (\(P < 2.2e-16\), Sign Test; Figure 1B). This result suggests that the GENIE3 network has value in directing focus toward targets with independent experimental support.

We also investigated whether the predicted targets of NAM-A1 shared enriched GO terms with the differentially expressed genes in the nam-a1 mutant. We identified 32 GO terms which were significantly over-represented among the predicted targets of NAM-A1, of which 11 were shared with the 137 GO terms enriched in de novo DEGs at 12 DAA. Of these 11 GO terms, all but one had 9 or more of its associated genes included in the list of predicted targets. These shared terms included known NAM-A1-related processes such as iron-ion binding (GO:0005506), oxidoreductase activity (GO:0016491, GO:00055114) and transmembrane transport (GO:0055085). This again supports the assertion that the GENIE3 network is predicting biologically relevant putative targets of NAM-A1.

Identification of senescence associated transcription factors

We hypothesized that transcription factors which also share predicted targets with the de novo DEGs may have roles in senescence.
We therefore identified transcription factors that had a higher percentage of shared targets with the de novo DEGs than NAM-A1 itself (Figure 2C, Supplementary Table 3). In total, we identified 20 such transcription factors, 0.6% of all transcription factors in the network. Five transcription factors were identified through comparison to the 12 DAA DEGs, 11 with the 22 DAA DEGs, and a further four which had a higher shared ratio at both time points. We obtained expression data for all 20 of the transcription factors across a developmental time-course and a senescence-specific time-course (Ramírez-González et al. 2018; Borrill et al. 2019). A diverse range of transcription factor families were represented, including four NACs, from the same family as NAM-A1 (Supplementary Table 3). Only one pair of homeologs was identified, from the C2C2-CO-like family. Using the developmental time-course, we calculated the fold-change in gene expression between non-senescent tissues and senescing leaf tissues as previously described (Borrill et al. 2019). We found that 14 of the 20 genes showed an increase in expression in the senescing tissues, four of which were enriched more than twofold (Figure 2A). Based on a published analysis of the senescence-specific time-course, six of the 20 transcription factors were identified as differentially regulated during flag leaf senescence in wheat (Borrill et al. 2019). Analysis of the GENIE3 predicted targets for these 20 genes identified nine transcription factors which shared at least one target gene with NAM-A1 (Figure 2B).

To investigate the potential functions of these 20 transcription factors further, we identified GO terms which were enriched in the GENIE3-predicted targets of these transcription factors (Supplementary Table 3; Supplementary File 2). At 12 DAA, the targets of three of the nine transcription factors were enriched for transporter-related GO terms, while others were enriched for senescence-related GO terms such as catabolism, phosphatase activity, and chlorophyll biosynthesis. Of the transcription factors identified at 22 DAA, GO terms related to transporters, senescence, circadian rhythms, and stress responses were enriched in the GENIE3-predicted targets. By integrating the information from the GENIE3 network with independent senescence-related expression data, we were able to identify a robust set of candidate senescence-associated transcription factors to prioritize for functional characterization.

**Non-homeologous transcription factors share few targets in the GENIE3 network**

After establishing that the GENIE3 network can provide biologically-relevant predictions, we then turned to using the network to interrogate genome-wide patterns in transcription factor targets. We first investigated the extent to which transcription factors share the same targets. To do this, we carried out pairwise comparisons between randomly selected transcription factors and calculated the overlap between their targets. Following 1,000 iterations, we found that any two random transcription factors typically have no targets in common, while the vast majority of comparisons shared 0 targets (Figure 3A, B; Table 1). However, there was a long tail to the right of the graph, highlighting that in some cases, certain non-homeologous transcription factors do share a substantial portion of targets.

The set of transcription factors in the GENIE3 network was then split into separate transcription factor super-families, as previously annotated (Appels et al. 2018). The same analysis was performed within each transcription factor family, carrying out pairwise comparisons between transcription factor targets. We found that for the majority of transcription factor families, the distributions of shared targets were very similar to that found for the full set, as illustrated by
the representative NAC and C2C2-Dof families (Figure 3A; Table 1; Supplementary Figure 2). Not all transcription factor families were large enough to support 1,000 unique pairwise comparisons, and in these cases the distribution clearly deviates from the whole (e.g., in CCAAT_HAP3, N = 3).

Roughly 70% of wheat homeolog triads (composed of A, B, and D genome copies) show balanced expression patterns, that is, similar relative abundance of transcripts from the three homeologs across tissues (Ramírez-González et al. 2018). We therefore hypothesized that homeologs would be more likely to share predicted targets than randomly selected transcription factors, even within the same family. To test this, we randomly selected syntenic triads from the GENIE3 dataset, and calculated the percentage of shared genes for each of the three pairwise comparisons (A-B, A-D, B-D). This was carried out for the 708 syntenic triads included in the network, a total of 2,124 pairwise comparisons, and showed that homeologs are indeed more likely to share targets with each other than randomly selected triads are (Figure 3A, C; Table 1).

Dynamic triads share fewer targets than stable triads

Wheat genome triads can be classified into different categories based on how the expression levels of the homeologs varies across tissues (Ramírez-González et al. 2018). So-called ‘dynamic’ triads represent the 10% of triads which show the highest variation in relative expression levels of the different homeologs across tissues, while ‘stable’ triads represent the 10% of triads with the lowest variation. Dynamic triads are under more relaxed selection pressure and we hypothesized that they may represent the initial steps toward neo or sub-functionalization of wheat homeologs (Ramírez-González et al. 2018). This hypothesis would suggest that homeologs in dynamic triads are more likely to have distinct functions, and thus may be less likely to share predicted targets. To test this, we compared the level of overlap between targets of homeologous transcription factors in dynamic and stable triads, as well as the ‘Mid 80’ intermediate set. We found that the dynamic triads do indeed have significantly less overlap in targets than the stable triads (P < 0.05, Wilcoxon test; Figure 4A; Table 1), supporting the neo/sub-functionalization hypothesis.

We next examined whether the targets of a transcription factor may hold signatures of the evolutionary origin of that transcription factor. We hypothesized that a transcription factor is more likely to have targets that reside on the same genome as the transcription factor itself; e.g., GENIE3-predicted targets for an A genome transcription factor will be enriched for A genome targets compared to B and D genome targets. We found no significant association between

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**Figure 2** Candidate senescence regulators. Twenty transcription factors were identified which had a higher shared ratio between the GENIE3-predicted targets and the RNA-Seq DEGs than NAM-A1 itself. (A) Their expression pattern is shown across a developmental time course (Ramírez-González et al. 2018) and a senescence-specific time course (Borrill et al. 2019). The TPM reported is averaged across three samples of the same tissue and timepoint from each dataset. Genes upregulated in senescence are highlighted with a light blue box, and those with a greater than twofold increase are highlighted with a purple box (left side). Genes present in the list of 341 candidate transcription factors based on the senescence time course in Borrill et al. 2019 are indicated with a green asterisk. (B) The number of targets shared between the transcription factors and NAM-A1. (C) The shared ratio for each gene against the 12 DAA (gray) and 22 DAA (black) DEGs. Note that genes which had a higher shared ratio at both 12 and 22 DAA are shown with split bars.

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We next examined whether the targets of a transcription factor may hold signatures of the evolutionary origin of that transcription factor. We hypothesized that a transcription factor is more likely to have targets that reside on the same genome as the transcription factor itself; e.g., GENIE3-predicted targets for an A genome transcription factor will be enriched for A genome targets compared to B and D genome targets. We found no significant association between
the genome of origin of the transcription factor and the genomes of its
targets (Figure 4B). Likewise, we found no significant association
between transcription factor genome and target genome when triads
were assigned into their respective movement categories. For exam-
ple, an A genome transcription factor from a Dynamic triad had
similar numbers of GENIE3-predicted targets from each of the three
genomes. (Supplementary Figure 3).

GO-term enrichment of predicted targets suggests
novel biological functions
We next examined if we could use the GENIE3-predicted targets to
gain a more general overview of functional trends within transcrip-
tion factor families. Using the GO terms described in (Ramírez-
González et al. 2018), we carried out a GO-term enrichment on the
targets of individual transcription factor families (Supplementary File
2). To test the validity of this method, we first checked the predicted
GO terms of the Heat Stress Factor (HSF) transcription factor family.
Based on their known role in responding to heat and light stress
(Kotak et al. 2007), we expected to see GO terms highly enriched for
heat and light stress responses. This was the case, suggesting that this
approach can provide accurate insights into the large-scale functions
of transcription factor families.

We then identified enriched GO terms for the remaining 56 tran-
scription factor families. Some families were enriched for few or no
GO terms, and we found that this was typically the case for families
with few members. We restricted our analysis, therefore, to the
34 transcription factor families with at least 30 members. In many
cases, enriched GO terms supported known functions of transcrip-
tion factor families. For example, the MADS_II family was enriched
for, among other terms, floral organ identity (GO:0010093) which
corresponds to their known role in flower patterning in wheat and
other species (Ng and Yanofsky 2001). The mTERF family was
strongly enriched for chloroplast-related terms, corroborating their
known role in organelle function (Kleine 2012; Quesada 2016) while
the WRKY family was highly enriched for stress-response GO terms
(Rushton et al. 2010; Phukan et al. 2016; Jiang et al. 2017).

Beyond the expected enriched GO terms, we also identified some
cases where highly significantly enriched GO terms may point toward
a previously unknown function within that transcription factor
family. For example, one of the more highly enriched GO terms
for the C2C2-Dof family is positive gravitropism (GO:0009958). This,
combined with other GO terms related to amyloplasts and auxin
responses, suggests that members of the C2C2-Dof family may play a
role in regulating the gravitropic growth of roots (Su et al. 2017).
We also see that the TUB family, of Tubby-like transcription factors, is
highly enriched in cell-cycle related GO terms. This includes
specific terms involved in microtubule movement and spindle
formation (GO:0007018 and GO:0051225) as well as in regulation
of cell cycle progression and transition (GO:0010389, GO:0051726
and GO:0000911). Plant TUB proteins contain an F-box domain,
suggesting they may function in tandem with other F-box-con-
taining proteins, such as SCF E3 ubiquitin ligases, which also
regulate the cell cycle in plants (Gusti et al. 2009).

The GENIE3 network is accessible through KnetMiner
We have made the GENIE3 network available in the KnetMiner
discovery platform for wheat (https://knetminer.org/) (Hassani-Pak
et al. 2016; Hassani-Pak et al. 2020). The top one million TF-target
predictions were integrated onto the wheat genome-scale knowl-
edge graph (Hassani-Pak et al. 2016) as directed relations between
genesis (A regulates B). The interaction weight for each predicted
relationship was also included in the network, where larger weights

![Figure 3](https://example.com/figure3.png)

**Figure 3** Non-homeologous transcription factors share few targets in the GENIE3 network. (A) Comparison of the shared targets of 1000 random transcription factor pairs found that the majority of transcription factors share few to no targets in common. This was also found to be the case within the majority of transcription factor families, showing here NAC and C2C2_Dof transcription factors. However, pairs of homeologous transcription factors shared many more targets in common, with a mean shared ratio of 39.7%. (B) An example of shared overlap between two homeologous transcription factors, the NAC transcription factors NAM-A1 (TraesCS6A01G108300) and NAM-D1 (TraesCS6D01G096300) is shown, with the two homeologs sharing 62% of possible targets. (C) The two randomly chosen transcription factors, in this case TraesCS1D01G436500 (Sigma 70-like family) and TraesCS4B01G383400 (HSF family), share no targets.
correspond to more strongly supported relationships. The data are accessible in conjunction with other information types (e.g., protein-protein interactions, literature co-occurrence, ontologies, homology, etc.) and can be searched through the KnetMiner web-app or web-services.

KnetMiner can be searched using keywords, genes or genomic regions to identify connections between genes and hidden links to complex traits. Searching for “NAM-A1” returns two hits, TRAESC-S6A02G108300 and TRAESC6D02G096300, which correspond to NAM-A1 and its D-genome homeolog NAM-D1, respectively. Using the KnetMiner network, we can visualize the relationships between NAM-A1 and NAM-D1, seeing, for example, that they target each other in the GENIE3 network (Figure 5). Associated traits are also included in the network, linking NAM-A1 and NAM-D1 to “Grain Protein Content” (Figure 5). Links to orthologous genes in other species are also included in the network, such as the Arabidopsis thaliana ortholog ANAC018 (Figure 5).

**DISCUSSION**

Here we have shown that a GENIE3 network provides biologically-informative predictions of targets for transcription factors in polyploid wheat. We have used the network in conjunction with independent RNA-Seq datasets to identify a set of candidate senescence regulators. We have also shown the value of the network to analyze genome-wide patterns of homeologous transcription factors and transcription factor families.

**Re-analysis of an RNA-Seq dataset identifies a high-quality set of differentially expressed genes**

In our analysis of the GENIE3 network, we used a previously published RNA-Seq dataset to test its ability to predict the targets of a well-characterized transcription factor, NAM-A1 (Uauy et al. 2006). In doing so, we needed to re-align the raw RNA-Seq reads to the most current transcriptome, RefSeqv1.1 (Appels et al. 2018). After carrying out de novo pseudoalignment and differential expression analysis, we obtained a larger set of differentially expressed genes between the wild-type and mutant nam-a1 lines than the original study (Pearce et al. 2014). In part, this is likely due to the less strict cut-off used in our study (q < 0.05). In the original study (Pearce et al. 2014), a combination of four statistical tests were used to reduce the number false positive discoveries. However, as our intention was not to reduce the rate of false positives, but rather that of false-negatives or incorrectly removed genes, we used a less stringent cut-off, based solely on adjusting the original q-values for false discovery rate. We found that the DEGs identified in Pearce et al. (2014) were found throughout the de novo list of DEGs, suggesting that the cut-off chosen was not overly generous (Supplementary Figure 1). We also closely recapitulated their findings in that our de novo DEGs were highly enriched for metal ion transport-related GO terms, indicating

![Figure 4](image-url) **Figure 4** The effect of relative homeolog expression variation on shared ratios. (A) There is a significant reduction in shared targets between homeologs in dynamic triads compared to stable triads (Wilcoxon test, P = 0.021). There is a near-significant decrease between the Mid 80 subset and the Dynamic triads (P = 0.051), and a slight non-significant decrease from Stable to Mid 80 triads (P = 0.21). (B) We found no evidence that transcription factors from a specific genome were more or less likely to have targets from the same genome (Two-sample Kolmogorov Smirnov test, FDR adjusted).
that the de novo DEGs are consistent with NAM-A1 function. These enrichment results also corroborate our understanding of NAM-A1 as a critical regulator of nutrient remobilisation during senescence.

The GENIE3 network provides biologically relevant transcription factor - target relationships

Gene networks are increasingly used in plant genetics research as a way to establish relationships across large gene sets and for hypothesis-generation. Initial assessment of the biological relevance of these gene networks often rely on enrichment analyses using GO terms. These methods are useful in identifying trends within large gene sets, as we found when carrying out GO term enrichment of transcription factor families. The targets of families with known functions, such as the Heat Stress Factors, were enriched for the expected GO terms (Kotak et al. 2007). However, these enrichment analyses are limited by the information that is used to develop the GO terms themselves. Very few GO term annotations are supported by experimental evidence even in model species such as Arabidopsis thaliana (Rhee and Mutwil 2014). GO terms associated to the RefSeqv1.0 transcriptome were based primarily on automated annotation and orthologs in other plant species, with over 96% of the GO terms assigned to genes based on inference from sequence orthology (ISO) (Appels et al. 2018). As a result, while enrichment analyses with GO terms can provide useful information particularly with large-scale datasets, they must be combined with external data to validate their predictions.

As a result, while the GO term enrichment analyses suggested that the network produced biologically relevant results, more rigorous testing of the network required the use of experimentally-derived data. By using independent RNA-Seq datasets, which were not used in the creation of the GENIE3 network, we were able to show that the predictions made by the in silico network hold up under comparison to in vivo datasets. It is important to note, however, that the overlap between the predicted targets of NAM-A1 and the differentially expressed genes from the RNA-Seq data are far from complete. At 12 DAA, the timepoint where NAM-A1 is thought to first start exerting its effect, only 12 genes are shared out of 79 predicted genes and 866 differentially expressed genes. While this gives a shared ratio of approximately 15%, significantly higher than that expected by chance, the GENIE3 network and the differentially expressed genes do not contain identical targets. However, based on the differences in the datasets used, it is likely that a large portion of this discrepancy is due to the fact that the GENIE3 network was derived from 850 distinct RNA-Seq samples, spanning different tissues, ages, stresses, and varieties, while the RNA-Seq dataset came from single timepoints taken from flag leaf tissue (Pearce et al. 2014; Appels et al. 2018). While NAM-A1 is expressed in the flag leaf during senescence, it is also expressed in the peduncle during senescence (Borrill et al. 2016; Ramirez-González et al. 2018), and at lower levels in various leaf, stem, and even spike tissues during development (Supplementary Figure 4). It is possible that many, if not most, of the predicted targets from the GENIE3 network may be regulated or influenced in some way by NAM-A1, but not in the flag leaf and at the precise 12 DAA and 22 DAA timepoints captured in the independent RNA-Seq data.

The GENIE3 network can be used for hypothesis generation with large gene sets

The fact that any two random transcription factors share, on average, zero targets in common in the GENIE3 network highlights that the network is not overwhelmed by spurious connections between transcription factors and biologically irrelevant targets. The network is also not so conservative that transcription factors with similar functions share no targets in common, as is made clear by the more normally-distributed homeologs (Figure 3A). Nevertheless, as most transcription factors do not share targets, this suggests that the original cut-off chosen for the network (1,000,000 connections) was appropriately stringent to avoid noisy, low-confidence connections. More recently, the GENIE3 network has been integrated into the RTP-STAR pipeline for gene regulatory network inference which trimmed the network based on the proportion of transcription factors present in the initial dataset (Clark et al. 2019). This has been
demonstrated to maximize the precision of the network (Shibata et al. 2018), though with the caveat that this analysis was focused on identifying direct downstream targets of two specific transcription factors. In preliminary investigations, we found that trimming the GENIE3 network by edge weight did lead to a slight increase in the shared ratio between the predicted targets and the de novo DEGs, however this was mostly due to the loss of predicted targets of NAM-A1 (data not shown). Therefore, where the aim is a global analysis of gene regulation, a less stringent cut-off such as that used here might be recommended to minimize type II errors.

The presence of overlapping targets between transcription factors suggests that the transcription factors may be acting in similar pathways. However, it is important to recognize the limitations of the network, in particular that predicted targets of a transcription factor are not necessarily true targets. Therefore, integrating the GENIE3 network with other sources of regulatory information, such as RNA-Seq datasets as demonstrated here, can provide cumulative evidence toward new hypotheses and candidate genes. By combining the DEGs obtained from a nam-a1 mutant line and the GENIE3 network, we have produced a set of candidate transcription factors which may act in the same or similar pathways as NAM-A1. We have then compared these candidates with an independent senescence time-course (Borrill et al. 2019), highlighting six candidate genes which were identified through both methods and are good candidates for further exploration. Moving forward, information generated from networks such as the GENIE3 will need to be validated functionally and in planta. Recently, NAM-A2, a previously uncharacterized member of the NAM family of transcription factors, was predicted to be involved in senescence based on expression and network data (Borrill et al. 2019). The function of this transcription factor was then validated using independent mutant TILLING lines (Krasileva et al. 2017) demonstrating the ability of the networks to predict biologically-relevant candidates.

**Homeolog expression variation in dynamic triads may be indicative of functional divergence**

Previous work showed that triads of homeologs can display variable patterns of genome dominance across different tissues (Ramírez-González et al. 2018). Triads with the most variable relative homeolog expression patterns, ‘Dynamic’ triads, were also found to have higher Ka/Ks values, suggesting they were under reduced selection pressure. It was proposed that the variation in relative expression patterns across tissues arises as a result of this relaxed selection pressure, facilitating both neo- and sub-functionalization following polyploidy. We found that dynamic triads were less likely to share GENIE3 targets in common than stable triads, supporting the hypothesis that dynamic triads are in the process of diverging functionally (Figure 4B). However, we found no correlation between the genome of origin of the transcription factor and the target genomes in dynamic triads (Supplementary Figure 3).

At what stage, then, did the targets of homeologs in dynamic triads begin to diverge? These results may suggest that the variation in expression seen between the homeologs arose following polyploidisation, as there is no bias toward the genome of origin. However, we do not know enough about the behavior of transcription factors following polyploidisation to draw clear conclusions. For example, we do not know to what extent transcription factors gain the ability to regulate homeologous genes on other genomes after hybridization. The application of methods such as ChIP-Seq (Park 2009), DAP-Seq (Bartlett et al. 2017), and large-scale yeast two-hybrid interaction screens (Brückner et al. 2009) to transcription factors in diploid and polyploid wheat will provide experimental data on homeologous transcription factor interactions and binding preferences. Until these datasets become available it is premature to draw conclusions on the evolutionary origins of transcription factor homeolog functional divergence. Nevertheless, genome-wide analyses of datasets such as the GENIE3 network and the expression datasets on expVIP have pointed to the dynamic triads as a source of genetic functional variation (Borrill et al. 2016; Ramírez-González et al. 2018).

**CONCLUSIONS**

Using publicly available datasets, we have shown that the wheat GENIE3 network provides biologically-relevant information that can be used to guide hypothesis generation in wheat. The utility of the network lies particularly in enrichment analyses of larger gene sets and in integration with other datasets, such as independent RNA-Seq experiments, for candidate gene selection. New germplasm resources in wheat, such as the in silico TILLING resource (Krasileva et al. 2017), can be rapidly leveraged for functional characterization of candidate genes in planta. Transgenic approaches such as CRISPR (Rey et al. 2018) and virus-induced gene silencing (VIGS) (Lee et al. 2012) can now be used in wheat to validate gene function. The GENIE3 network can be accessed through the KnetMiner web application and using R scripts available from https://github.com/Uauy-Lab/GENIE3_scripts. We predict that gene regulatory networks such as GENIE3 will play an increasingly important role in wheat genetics as more transcriptomic datasets become publicly available.

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