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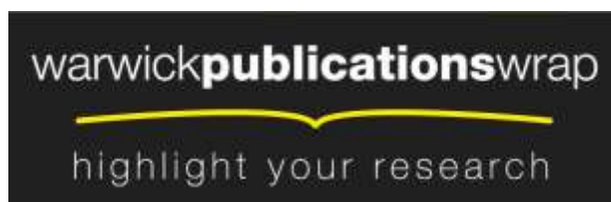
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A local regulatory network around three NAC transcription factors in stress responses and senescence in Arabidopsis leaves

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SUMMARY

A model is presented describing the gene regulatory network surrounding three similar NAC transcription factors that have roles in Arabidopsis leaf senescence and stress responses. *ANAC019*, *ANAC055* and *ANAC072* belong to the same clade of NAC domain genes and have overlapping expression patterns. A combination of promoter DNA/protein interactions identified using yeast 1-hybrid analysis and modelling using gene expression time course data has been applied to predict the regulatory network upstream of these genes. Similarities and divergence in regulation during a variety of stress responses are predicted by different combinations of upstream transcription factors binding and also by the modelling. Mutant analysis with potential upstream genes was used to test and confirm some of the predicted interactions. Gene expression analysis in mutants of *ANAC019* and *ANAC055* at different times during leaf senescence has revealed a distinctly different role for each of these genes. Yeast 1-hybrid analysis is shown to be a valuable tool that can distinguish clades of binding proteins and be used to test and quantify protein binding to predicted promoter motifs.

Keywords: *Arabidopsis thaliana*, *Botrytis cinerea*, NAC transcription factors, gene regulatory network, senescence, stress.

INTRODUCTION

As plants are sessile organisms, constant exposure to biotic and abiotic stresses has driven evolution of ever more complex mechanisms to defend against attack from other organisms and to deal with environmental perturbations. Elicitation of these stress responses requires signal perception, signal transduction and large-scale gene expression changes. Although some gene expression may be specific to each stress there is often a large overlap between stresses, suggesting that common features are likely to be involved (e.g. Kilian *et al.*, 2007).

Genetic studies have highlighted the important role of the plant-specific NAC (NAM/ATAF1, 2/CUC2) family of transcription factors (TFs) in the regulation of stress responses (reviewed in Puranik *et al.*, 2012). NAC proteins share a

conserved N-terminal NAC domain and form one of the largest plant TF families with over 100 members encoded in the Arabidopsis genome (Ooka *et al.*, 2003). NAC TFs have been implicated functionally in a variety of stress-related programs such as the response to drought, high-salinity, bacterial pathogens, fungal pathogens and senescence (Fujita *et al.*, 2004; Tran *et al.*, 2004; Guo and Gan, 2006; Hu *et al.*, 2006; Balazadeh *et al.*, 2010). Many members of the NAC family have overlapping expression patterns and are induced by several stresses, a situation that suggests common roles and regulation in multiple stress responses (Balazadeh *et al.*, 2010; Nakashima *et al.*, 2012).

Three members of the NAC family that have been implicated in overlapping stress responses are ANAC019,

ANAC055 and ANAC072. These TFs form part of a closely related clade of stress-related NAC proteins of the SNAC-A class as defined recently in Nakashima *et al.* (2012) or group III-3 as defined in Jensen *et al.* (2010). There is evidence that these three genes have overlapping functions, but that they may also act individually in different stress responses. Tran *et al.* (2004) have shown that overexpression of any of the three genes results in increased drought tolerance, that all were induced in expression following salt stress, abscisic acid (ABA) or jasmonic acid (JA) treatment and that all three proteins could bind the promoter of the drought-inducible *ERD1* gene. However, ANAC072 (RD26) has been shown to have a role in cold and desiccation stress in addition to a central role in ABA signalling (Fujita *et al.*, 2004), while ANAC019 and ANAC055 are implicated in JA and/or ethylene signalling following pathogen infection (Bu *et al.*, 2008).

The elucidation of the individual roles and regulatory pathways for these NACs is complicated by their overlapping activities, but here we present a dual approach to address this question and to identify common and distinguishing factors that lie upstream and downstream of ANAC019, ANAC055 and ANAC072 in stress-specific gene regulatory networks (GRNs). A key mechanism in the regulation of differential gene expression is through the sequence-specific binding of TFs to DNA motifs present in the promoters of targets. The physical interactions between these *trans*- and *cis*-acting factors and the nature of their expression-controlling activity form the basis of GRNs that tightly coordinate the spatiotemporal expression of gene products that are responsible for stress tolerance at the biochemical and physiological level. High-throughput yeast 1-hybrid (Y1H) assays are becoming an increasingly popular 'gene-centred' approach for delineating GRNs in Arabidopsis (Mitsuda *et al.*, 2010; Brady *et al.*, 2011; Castrillo *et al.*, 2011; Ou *et al.*, 2011). We used Y1H to identify TFs that interact with the promoters of ANAC019, ANAC055 and ANAC072 and, by modelling expression patterns of these potential regulators, excellent candidates for direct upstream regulation in different stress responses were predicted. We complemented this approach with microarray analysis of these potential upstream regulators to test the model predictions and also investigated the consequences of knockout mutants of the NAC genes to reveal genes that lie downstream in the GRN in senescence.

RESULTS

Y1H assays identify TFs that interact with ANAC019, ANAC055 and ANAC072 promoters

The Y1H technique was used to identify TFs capable of interacting physically with the promoters of ANAC019, ANAC055 and ANAC072. Distal effects on transcription in yeast were

minimised using a 'promoter-hiking' (Pruneda-Paz *et al.*, 2009) approach with a series of overlapping promoter sequences of approximately 400 bp and that spanned 800–1200 bp upstream of the predicted transcription start site (TSS). These promoter fragments were used as bait against an Arabidopsis TF library (Castrillo *et al.*, 2011) that was comprised of approximately 1500 Arabidopsis TFs. All interactions were verified pairwise with each fragment–TF pair.

The Y1H assays identified many TFs that were capable of binding to the promoters of ANAC019, ANAC055 and ANAC072 respectively: a total of 71 interactions between TFs and all bait fragments (Table 1). The interacting TFs comprised members of several TF families including bZIP, bHLH, MYB, AP2/ERF and homeodomain and are all expressed above background in green and/or stressed leaf tissues [based on internal gene expression data and/or eFP Browser analysis (Winter *et al.*, 2007)]. *In silico* analysis of each fragment sequence revealed the presence of putative binding sites for all the interacting TFs identified (Figures S1, S2 and S3).

Analysis of the biological processes associated with the interacting TFs using gene ontology (GO) classifications revealed significant enrichment for TFs associated with 'response to hormone stimulus' and 'response to ABA'. Expression of all three NAC TFs is induced by ABA (Tran *et al.*, 2004) and thus targeting by TFs that are known to mediate ABA signalling would be expected. Indeed, the key ABA-related TFs ABF3 and ABF4, which recognise the ABA-responsive element (ABRE) (Yoshida *et al.*, 2010), were found to bind to all three promoters (Table 1). ABRE occurs multiple times within each NAC promoter fragment (Figures S1, S2 and S3) and recent motif analysis revealed two ABRE-like motifs in the promoter of ANAC019 that were sufficient to drive expression in response to stress (Zou *et al.*, 2011). The promoter fragments of ANAC019 and ANAC055 also interact with another key mediator of ABA expression, ABI4 (Table 1), an AP2 family TF whose predicted binding motif (Niu *et al.*, 2002) is present in the promoters of these genes (Figures S1 and S2).

Members of a MYB TF family group, MYB2, MYB21, MYB108, MYB112 and MYB116, bound to the promoters of all three NAC genes (Table 1). An example of these interactions is shown in Figure 1(a) and demonstrates the interaction of overlapping fragments of the ANAC055 promoter (ANAC055_F3 and ANAC055_F4) with these five MYB TFs. Consistent with this observation is the presence of a MYB binding motif (TRANSFAC ID: M00218, Solano *et al.*, 1995) in the overlapping region of these two fragments (Figure 1b). The five MYB TFs form part of a phylogenetic clade within the MYB family, based on the similarity of the DNA-binding domain (Stracke *et al.*, 2001). Interestingly MYB21, the least related within this group, shows a weaker interaction (i.e. less growth) with the NAC promoters

Table 1 Positive interactions identified by yeast 1-hybrid (Y1H) screens between named transcription factors (TFs) and the promoter fragments of *ANAC019*, *ANAC055* and *ANAC072*. Fragments are numbered F1–F5 with F1 being closest to the transcription start site (TSS)

AGI	Name	ANAC019					ANAC055				ANAC072			
		F1	F2	F3	F4	F5	F1	F2	F3	F4	F1	F2	F3	F4
AT4G34000	ABF3				•		•		•		•			
AT3G19290	ABF4	•			•		•		•		•		•	•
AT2G40220	ABI4	•					•							
AT1G19350	BES1													•
AT1G59640	BPEP				•									
AT4G25490	CBF1										•			
AT4G25470	CBF2										•			
AT4G25480	CBF3										•			
AT5G51990	CBF4										•			
AT4G36730	GBF1				•		•		•					
AT4G01120	GBF2				•		•		•					
AT2G44910	HB04	•												
AT5G65310	HB05				•									
AT3G61890	HB12				•									
AT5G66700	HB53	•												
AT5G11260	HY5								•					
AT3G17609	HYH								•					
AT5G54680	ILR3				•				•					
AT3G06490	MYB108					•			•	•				•
AT1G48000	MYB112					•	•		•	•				•
AT1G25340	MYB116					•	•		•	•				•
AT2G47190	MYB2					•	•		•	•				•
AT3G27810	MYB21					•	•		•	•				•
AT1G06160	ORA59	•							•		•			
AT5G61270	PIF7	•			•				•	•	•		•	
AT2G22750	bHLH				•				•					
AT4G14410	bHLH104								•					
AT4G34590	bZIP11								•					
AT2G35530	bZIP16								•					
AT1G75390	bZIP44						•		•					
AT1G06850	bZIP52								•					
AT3G44460	bZIP67								•					

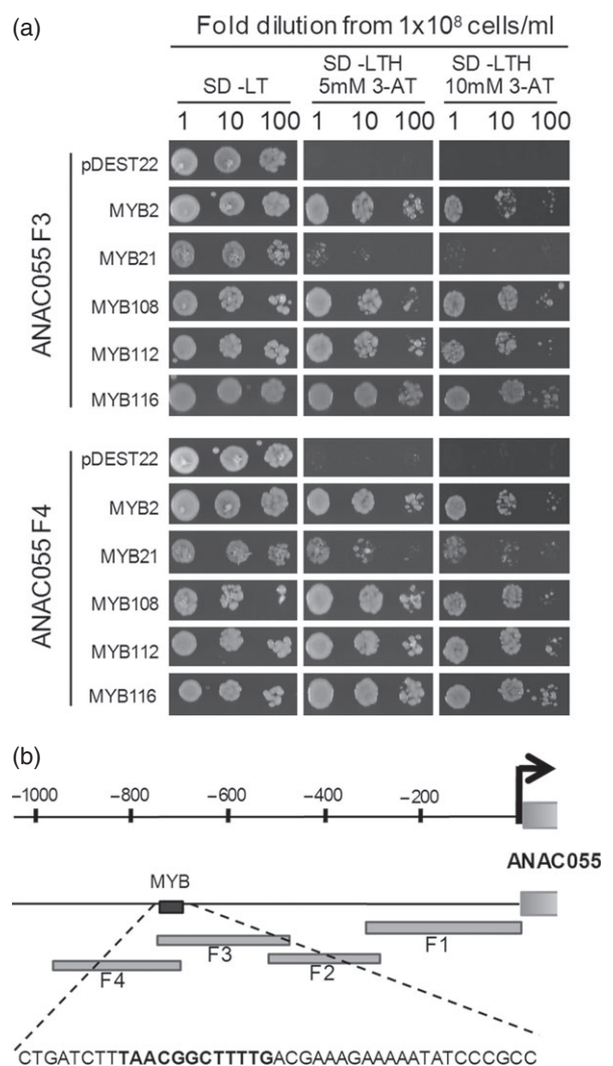


Figure 1. Related MYB transcription factors (TFs) bind to the promoter of *ANAC055*.

(a) Interactions between MYB TFs and *ANAC055* promoter fragments 3 and 4 linked to the *HIS3* reporter gene were transformed into yeast with plasmids carrying an Activation Domain (AD)-MYBTF translational fusion or the control plasmid expressing AD alone. Growth was assessed in the presence of increasing 3-amino-1,2,4-triazol (3AT) concentrations.

(b) The *ANAC055* promoter indicating the putative MYB binding site highlighted in bold (TRANSFAC ID: M00218, Solano *et al.*, 1995) in the overlapping region of fragments 3 and 4.

(Figure 1a). Members of this MYB subgroup are implicated in regulating the same stress responses as the NAC TFs. *MYB2* is involved in the ABA-mediated regulation of salt and drought-responsive genes (Abe *et al.*, 2003), whilst *MYB108* is induced in response to ABA, JA and ethylene and is involved in regulating the response to infection by *B. cinerea* and other pathogens (Mengiste *et al.*, 2003; Mandaokar and Browse, 2009). The observation that only these related members of the MYB family demonstrate positive interactions with the three NAC promoters

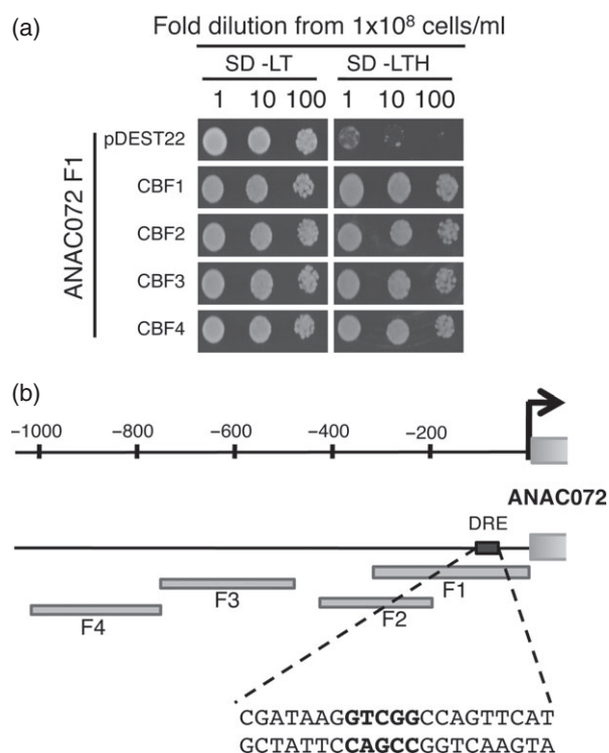


Figure 2. The C-repeat/dehydration-responsive element (DRE) binding factor genes (CBF1–CBF4) bind to the promoter of *ANAC072*.

(a) Interactions between CBF1–CBF4 and fragment 1 of the *ANAC072* promoter (*ANAC072_F1*) were confirmed by transforming yeast that carried the *HIS3* reporter gene under the control of *ANAC072_F1* with plasmids that expressed the AD-CBF fusion proteins or a control plasmid that expressed AD alone and by assessing growth on media that lacked histidine.

(b) The DRE motif (consensus (A/G)CCGAC, bold text) in the *ANAC072* promoter on the reverse strand reading 5' to 3'.

suggests that this binding occurs in a selective, sequence-specific manner.

In contrast to the common interactions of ABA-related and MYB family TFs with all three NAC promoters, binding of four highly related proteins, CBF1 (C-repeat/dehydration responsive element (DRE) Binding Factor 1) CBF2, CBF3 and CBF4, was detected to the *ANAC072* promoter only (Figure 2a). The CBFs bind via the DRE, which has the consensus (A/G)CCGAC (Sakuma *et al.*, 2002) and is present in the interacting promoter fragment (Figure 2b). To confirm that CBF1–CBF4 are interacting via the DRE motif, mutational analysis was performed and the protein/DNA interactions were quantified. Firstly the DRE motif was completely replaced with the sequence TTTTT (Figure 3a – M1). This replacement resulted in the abolition of the interaction between the *ANAC072* promoter fragment 1 and all four of the CBFs (Figure 3c). Hao *et al.* (2002) showed that the core DRE was CCGAC by testing the effect on CBF1 binding of mutating the nucleotides at either end of this motif. As this study identified only the boundaries of the DRE motif, mutation of the central nucleotide (CCGAC) was

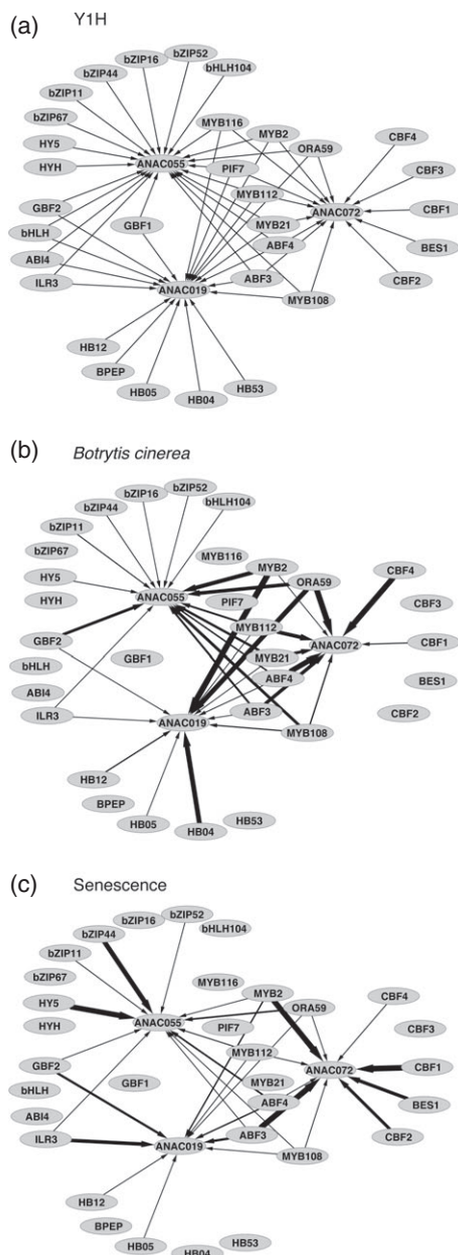


Figure 4. Hierarchical causal structure identification (hCSI) modelling predicts stress-specific sub-networks using different stress timecourse datasets based upon the core yeast 1-hybrid (Y1H) network (a). Networks for (b) *B. cinerea* infection and (c) developmental senescence are shown. True Y1H connections predicted by CSI are shown as solid black lines in which the thickness of the line represents the likelihood of that interaction, and the arrows show direction.

contrast, CBF3 is predicted to have the most influence in cold, osmotic and salt stress (Figure S4).

The hCSI analysis indicates that, of the five MYB proteins shown to bind the three NAC promoters in Y1H, MYB116 appears to have no significant role in developmental senescence or the stress responses modelled here

(Figures 4 and S4). Interestingly the predicted regulatory role of MYB2 is different between senescence and *B. cinerea*. Although predicted to influence all three NACs, the models suggest that MYB2 is highly likely to regulate *ANAC019* and *ANAC055* in *B. cinerea* infection (Figure 4b), whilst in developmental senescence its main role is predicted to be in the regulation of *ANAC072* (Figure 4c). MYB108 is predicted to influence the expression of all three NACs in all stresses modelled, particularly *ANAC055* in *B. cinerea* infection (Figure 4b) and salt stress (Figure S4a). These predictions were tested experimentally as described in the next section.

Expression of *ANAC019*, *ANAC055* and *ANAC072* is perturbed in *myb2* and *myb108* mutants in different conditions

As shown above, a related group of five MYB TFs can bind to the promoters of *ANAC019*, *ANAC055* and *ANAC072* in Y1H, with a predicted stress-dependent differential contribution to the control of the expression of the three NACs. Following *B. cinerea* infection, MYB108 was predicted to regulate the expression of all three NAC genes, but with a more significant influence on *ANAC055* (Figure 4b); and MYB2 was predicted to influence the expression of *ANAC019* and *ANAC055* and, to a lesser extent, *ANAC072* (Figure 4b). The hCSI algorithm does not give the sign (activation or inhibition) for each interaction. However, through comparison of MYB and NAC expression profiles in *B. cinerea* infection and developmental senescence (Figure S5; Windram *et al.*, 2012 and Breeze *et al.*, 2011 respectively) it is predicted that the action of the MYBs would be to activate NAC expression. To test these predictions the expression of the three NAC genes was determined in leaves of *myb108* and *myb2* T-DNA insertion mutants at two time points following infection with *B. cinerea* (Figure 5a). None of the NACs shows a significant expression change in *myb108* compared with wild type (WT) at 26 h post infection (hpi) however, at 30 hpi all genes show reduced levels of upregulation as was predicted in our model, with *ANAC055* showing the greatest reduction in expression. In the *myb2* mutant, as with *myb108*, there is no differential expression of the three NAC genes at the earlier time point (24 hpi), but by 30 hpi the expression of *ANAC019* is statistically significant lower whereas *ANAC072* expression is not altered significantly. However, the predicted effect on *ANAC055* expression is not observed.

To test the predictions of the senescence model, the expression of the three NAC genes was determined in leaves of the *myb108* and *myb2* mutants subjected to dark-induced senescence (DIS) at two time points termed DIS1 and DIS2 (8 and 9 days after darkness respectively). In the senescence hCSI model, MYB108 is predicted to have an effect on the expression of all three NACs.

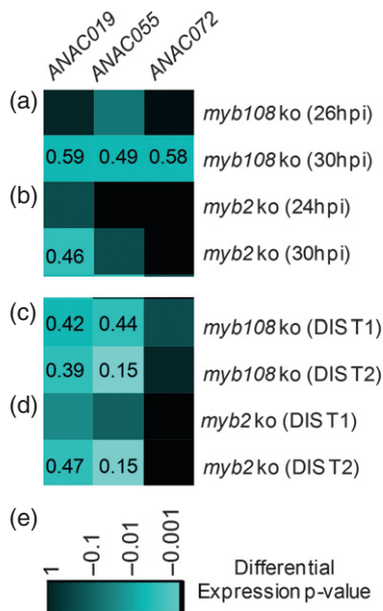


Figure 5. Expression levels of *ANAC019*, *ANAC055* and *ANAC072* in *myb2* and *myb108* mutant backgrounds under different stress conditions. Heat-map representation (*P*-values) of expression changes of *ANAC019*, *ANAC055* and *ANAC072* shows downregulation in response to infection by *B. cinerea* (hours post infection; hpi) (a, b) and dark-induced senescence (DIS) at two time points (T1, T2) (c, d). Numbers shown on heat map are ratios of gene expression in mutant compared with wild-type (WT). (e) Colour scale for *P*-value significance.

However, in the *myb108* mutant, at both time points only *ANAC019* and *ANAC055* show a significant reduced level of expression (Figure 5c). The expression of *ANAC072* is not significantly altered in the mutant. In contrast, MYB2 is predicted to significantly affect the expression of *ANAC072*, whilst having less influence on the expression of *ANAC019* and *ANAC055*. In the *myb2* mutant expression of both *ANAC019* and *ANAC055* is significantly reduced at DIS2 but *ANAC072* is unaffected (Figure 5d). The discrepancy between the predicted and experimental results with respect to expression of *ANAC072* indicates that additional components are involved in the regulation of *ANAC072* during senescence. The data points in the senescence time course used for the CSI modelling are taken at 2-day intervals. In contrast, at 2 hourly intervals the data points in the *B. cinerea* time course are more highly resolved thus providing a more detailed picture of the immediate consequences of temporal changes in gene expression. Additionally, the CSI modelling used data from a developmental senescence experiment. In contrast, analysis of the *myb2* and *myb108* mutants was performed following DIS.

The use of expression data to predict the most likely interactions from the protein DNA-binding studies generated testable stress-specific models. In this example, we tested the predicted interaction of two of the many upstream interactors in two different stresses. Most of the

predictions tested were confirmed, a finding that indicated that many of the additional interactions shown in Figures 4 and S4 are also likely to be true.

Microarray analysis identifies genes and pathways regulated by *ANAC019* and *ANAC055* during senescence

To identify downstream components in the NAC GRN, microarray analysis of T-DNA mutants *anac019* and *anac055* was performed at five time points (TP1 to TP5) during developmental leaf senescence. Firstly genes that were differentially expressed over the whole time course were identified using a Gaussian process two-sample test (GP2S) (Stegle *et al.*, 2010). Compared with WT, 2785 and 7457 genes demonstrated altered expression during the time course in the *anac019* and *anac055* mutants respectively. To identify genes that were differentially expressed from WT at each time point, rather than overall, a *t*-test analysis was performed with the data at each time point (Tables S1 and S2). This double selection considerably refined the number of differentially expressed genes for GO term analysis.

Gene ontology (GO) term analysis for groups of up- or downregulated genes revealed that there was little overlap in pathways affected by mutations in the two NAC genes, indicating that they have clearly different roles in the senescence response (Tables 2 and S1). Expression of both NAC genes was significantly downregulated at every time point in the relevant T-DNA mutant, but was not affected by the other mutation and indicated that neither gene is dependent on the other for expression (Figure 6a). Several photosynthesis genes, including *LHCA4* (Figure S6a), *LHCB6*, *PSBX* and *PSAK*, showed increased expression in *anac019* compared with WT – indicating a delay in aspects of the senescence process. Conversely, certain chloroplast-related genes, such as *SBPase*, are downregulated in *anac055* – possibly indicating a more advanced decay and accelerated senescence in this mutant (Figure S6a).

The GO terms ‘response to stress’, ‘response to stimulus’ and ‘defence response’ are over-represented in genes upregulated in the *anac019* mutant. The most consistent differentially expressed stress response gene is *SAT32*, a salt responsive gene, that is upregulated at all five time points (Figure S6b). Overexpression of this gene in *Arabidopsis* increases tolerance to salt (Park *et al.*, 2009). Other upregulated stress-related genes include *PHYTOALEXIN DEFICIENT 3 (PAD3)*, *PR4*, *PDF1.2*, *ACCELERATED CELL DEATH 6 (ACD6)* and *RECEPTOR LIKE PROTEIN 46 (AtRLP46)*, all of which are involved in the defence response and could improve stress tolerance (Table S1 and Figure S6b). In contrast, the *anac055* mutant demonstrates reduced or delayed expression of several abiotic stress response genes, including *LEA14*, *SAT32*, *RHA2A*, *EDS5* and *ERD5* (Table S2 and Figure S6c).

Table 2 Enriched gene ontology (GO) terms in groups of genes showing higher or lower expression in the NAC gene knockout mutants compared with wild-type (WT) at different times during senescence

Days after sowing	<i>anac019</i> mutant		<i>anac055</i> mutant	
	Up-regulated	Down-regulated	Up-regulated	Down-regulated
23	Response to stress JA biosynthesis Response to JA		Response to SA Response to stimulus	
29			Response to stimulus Cell wall	Response to chitin JA stimulus
31	Indole derivative biosynthesis Defence response Photosynthesis	Flavonoid biosynthesis SA signalling	Response to stimulus Transporter activity	Response to stress JA stimulus
33	Chromatin assembly Response to biotic stimulus Photosystem	Flavonoid biosynthesis SA signalling	Response to chitin Defence response	Chloroplast Response to stress
35	JA biosynthesis Response to stimulus Response to JA		Response to chitin Response to stress	Response to stimulus Response to stress

JA, jasmonic acid; SA, salicylic acid.

Gene ontology (GO) term analysis indicates that these two closely related NACs may have reciprocal roles in the regulation of JA and salicylic acid (SA) signalling. ANAC055 may be required for normal JA signalling, while ANAC019 could enhance SA and repress JA signalling. Enriched GO terms for JA biosynthesis and signalling at TP1 and 5 in the *anac019* mutant are illustrated by increased expression of JA biosynthesis genes, including *LOX2* and *ALLENE OXIDE SYNTHASE (AOS)* (Figure S6b), and JA response genes including *PR4* and *PDF1.2* (Table S1). In contrast, reduced expression of JA signalling genes such as *JAZ7* and *JAZ10* is observed in the *anac055* mutant (Figures 6b and S6c and Table S2). SA signalling appears to be the dominant pathway in the *anac055* mutant with GO term enrichment for this response associated with upregulated genes at TP1. This is illustrated by early enhanced expression of genes such as *CELL WALL-ASSOCIATED KINASE (WAK1)*, usually expressed in response to SA (Figure S6c) (He *et al.*, 1999), and *EDS1*, which is involved in SA-mediated signalling in plant defence (Feys *et al.*, 2001). This change may result in the apparent inhibition of the JA pathway in the *anac055* mutant. Downregulation of the SA pathway in the *anac019* mutant is illustrated by decreased expression of *AHBP-1B* (Figure S6b), a transcriptional repressor implicated in SA signalling (Fan and Dong, 2002).

ANAC019 may act to enhance expression of the flavonoid biosynthesis pathway but repress the activity of the camalexin pathway (possibly as a consequence of the repression of JA signalling). The flavonoid biosynthesis pathway is significantly downregulated in the *anac019* mutant, including genes such as *DFR*, *LDOX*, *F3H*, *TT4* and *TT5*, which show reduced expression at TP3 and TP4 in the mutant when compared with WT (Table S1 and Figure 6c).

In addition, two potential regulatory genes *MYB90* and *TT8*, both of which are TFs implicated in regulation of flavonoid biosynthesis (Borevitz *et al.*, 2000; Baudry *et al.*, 2004), are also downregulated, indicating that these TFs could be a primary target for ANAC019. Example expression patterns (for *DFR* and *TT8*) in Figure 6(c) show that the rapid induction of expression of these genes in the WT is blocked in the mutant, but that expression recovers to WT levels by TP5, indicating that the lack of this TF can be compensated for later in senescence.

Genes involved in the synthesis of the antifungal phytoalexin camalexin are associated with the GO term 'indole biosynthesis', over-represented in upregulated genes in the *anac019* mutant. These genes include *ANTHRANILATE SYNTHASE ALPHA SUBUNIT 1 (ASA1)*, *PHOSPHORIBOSYL ANTHRANILATE TRANSFERASE 1*, and *INDOLE-3-GLYCEROL PHOSPHATE SYNTHASE (IGPS)*, which all function in the biosynthesis pathway from chorismate to tryptophan and in addition to *PAD3* and *CYTOCHROME P450 MONOOXYGENASE 79B2 (CYP79B2)*, which are required for the production of camalexin from tryptophan (Schuhegger *et al.*, 2006). The presence of ANAC019 causes a delay in the early expression of camalexin synthesis genes (see *PAD3* example in Figure S6b); by TP5 the levels of expression are the same in both mutant and WT.

In the *anac055* mutant there is a striking group of genes with the GO annotation 'response to chitin' that show lower expression than WT at TP2, but that exhibit higher expression at TP4 and TP5 (illustrated by *WRKY53* expression in Figure 6b). This group contains several TFs, including *WRKY33*, *WRKY53*, *WRKY11* and *ERF5*, all of which are enhanced in expression in response to chitin and in defence responses (Libault *et al.*, 2007). These genes show a peak in expression at TP2 that is considerably delayed in

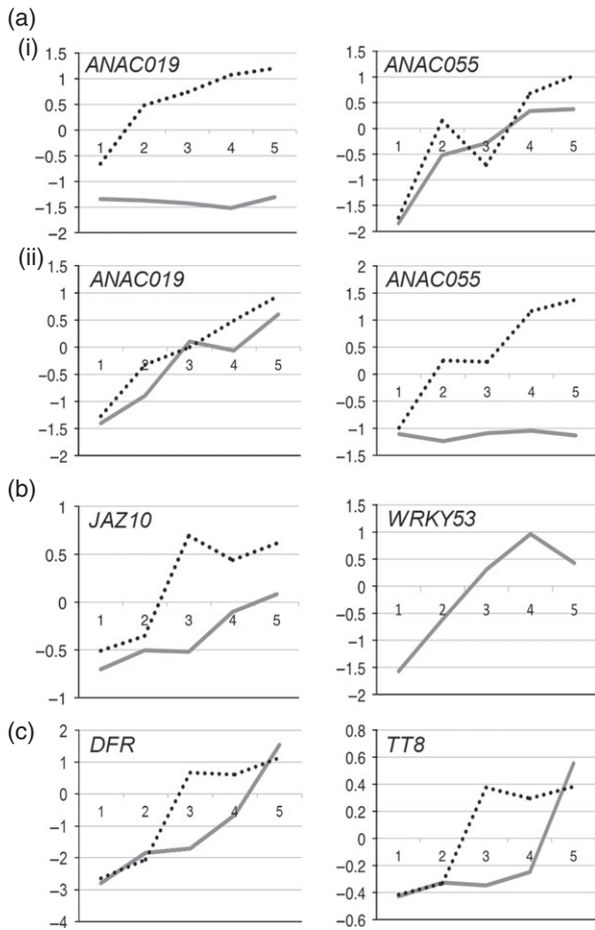


Figure 6. Expression patterns of selected differentially expressed genes in the NAC gene mutants. (a) Expression of the target NAC genes *ANAC019* and *ANAC055* in each knockout (KO) mutant: (i) in *anac019* KO; and (ii) in *anac055* KO. (b) Expression of chitin response (e.g. *WRKY53*) and jasmonic acid (JA) signalling genes (e.g. *JAZ10*) showing altered expression in the *anac055* KO mutant. (c) Expression of genes involved in flavonoid biosynthesis (e.g. *DFR* and *TT8*) showing delay in expression in the *anac019* mutant. The solid line shows expression of the gene indicated in the mutant, a dashed line shows the wild-type (WT) expression.

the mutant implying a requirement for *ANAC055* for this rapid increase in expression. At subsequent time points, expression in the WT decreases while that in the mutant increases. *ANAC055* may play a role in the WT induction of these genes, with other genes able to bring about the induction, although less efficiently, in the absence of *ANAC055*.

In summary, the microarray analysis of the two NAC mutants indicates that they have very different roles to play in regulating gene expression during developmental leaf senescence. Different downstream pathways are affected, for example the flavonoid pathway depends on *ANAC019*, while the pathogen response pathway induced by chitin appears to be dependent on *ANAC055*. Also, the

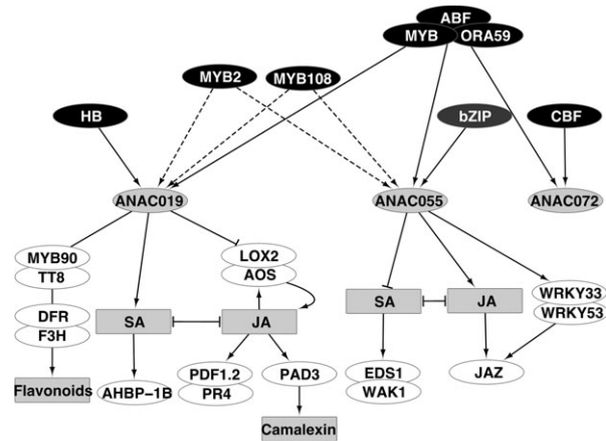


Figure 7. Schematic showing predicted gene regulatory network around *ANAC019*, *ANAC055* and *ANAC072*. Upstream genes were predicted by yeast 1-hybrid (Y1H) assays in combination with context-specific network modelling and mutant analysis. Dashed lines indicate interactions validated in senescence. Downstream components were predicted from microarray analysis of mutants *anac019* and *anac055* during developmental senescence.

genes appear to have opposite roles in regulation of the JA and SA antagonistic interaction.

DISCUSSION

The results in this paper predict a regulatory network around three stress-related NAC genes, *ANAC019*, *ANAC055* and *ANAC072* (Figure 7). Potential upstream regulatory genes were identified by a combination of Y1H, stress-specific modelling and mutant analysis; downstream genes were predicted from microarray data.

This study exploited the use of Y1H to identify interactions between promoter DNA and specific TF proteins. This method is an excellent technique to show protein/DNA binding but is prone to false-negative results as many TFs may not bind in isolation or may require binding at a fixed distance from the start of transcription. For instance, this study did not detect the interaction between the MYC2 protein and the three NAC promoters recently described by Zheng *et al.* (2012). MYC2 may not be made or processed properly in yeast, or may require other factors to enable binding. It has been demonstrated in yeast that transcriptional activation diminishes with increasing distance of the binding element from the TSS (Dobi and Winston, 2007), thus many true interactions may be missed. However, we identified many different interacting TFs, including members of a phylogenetic clade of the MYB TF family, which bind to all three promoters including a conserved MYB recognition site in two overlapping fragments of *ANAC055* (Figure 1a). This result indicates that the chosen fragment length of approximately 400 bp does not necessarily preclude the identification of TFs binding either end of the promoter fragment.

The binding of members of this clade of MYBs shows some degree of specificity; in other experiments other MYBs within our TF library have bound different promoters in the Y1H assay. Current predicted binding motifs are oversimplified but knowledge of such sequences, in combination with Y1H, enables binding specificity to be investigated, as our mutation analysis of the DRE motif has demonstrated. This combination should allow the intricacies of sequence-specific binding to be investigated thus revealing specific TF protein binding motifs beyond the simple gene family motifs we currently employ.

Many TFs occur in large families sharing a similar DNA-binding domain, including NACs, bZIPs and homeodomain TFs (Riechmann *et al.*, 2000). Promoter evolution has been suggested to drive functional differences between members of several stress-related TF families. For example, the CBF TF family, comprised of CBF1, 2, 3 and 4, are important for regulating responses to drought and cold stress. CBF1, 2 and 3 are induced by low temperatures but not dehydration or ABA (Gilmour *et al.*, 1998; Liu *et al.*, 1998; Medina *et al.*, 1999) while CBF4 is induced by dehydration and ABA but not cold (Haake *et al.*, 2002). All members have high similarity at the protein level, yet the CBF4 promoter differs considerably from those of the other CBF genes (Haake *et al.*, 2002). The differential expression of members of TF families such as the CBFs is crucial and it is demonstrated in this paper that although all four members of this family have the ability to bind to the promoter of *ANAC072* in Y1H, modelling indicates that their contribution to the regulation differs amongst this family in a context-dependent manner.

Phylogenetic analysis of the promoters of *ANAC019* and *ANAC055* indicates that they are extremely similar at the promoter level (Ooka *et al.*, 2003; Tran *et al.*, 2004) and this study demonstrates a large overlap in the binding TFs. However, there are also clear differences (Figures 4a and 7). The *ANAC019* promoter is bound by a group of homeodomain TFs that show no interaction with the *ANAC055* promoter, which is instead bound by a selection of bZIP proteins. Such observations suggest that promoter evolution has refined the regulation of these two NAC genes thus adding to the complexity of GRN in which they act.

A further level of complexity is seen when we consider the context in which the interactions are observed *in vivo*. The use of modelling algorithms allows prediction of true interactions by considering them in the context of stress-specific expression data. Such analysis indicates that although several members of the same TF family have the ability to bind to the promoters in question, they may not actually bind under all conditions *in vivo*, as is predicted here with the MYB and CBF TFs. This analysis also demonstrated the importance of highly

resolved time series expression data, with the highly resolved *B. cinerea* dataset providing more accurate predictions than the senescence dataset. In some cases there may be functional redundancy between TFs, which would prevent testing of the model using knockout mutants but should allow the prediction of likely functional homologues. Additionally, it is important to consider that TFs that are required for activation of a gene do not necessarily need to be differentially expressed and would not be predicted using the hCSI algorithm as it relies on differential expression patterns.

Expression analysis of the mutants *anac019* and *anac055* during developmental senescence indicated involvement of these genes in different signalling pathways. Gene expression in the *anac019* mutant indicates that ANAC019 may be an activator of senescence with a role in activating flavonoid and anthocyanin biosynthesis. Conversely, early downregulation of chloroplast-related genes in the *anac055* mutant hints at accelerated senescence and this TF appears to be involved in the response to chitin. These genes also appear to have opposing roles in regulating the antagonistic JA and SA pathways (Figure 7). Furthermore the observation that certain genes, including *WRKY33* and *WRKY53*, showed an apparent delay in expression in the *anac055* mutant illustrates the importance of measuring the dynamic effects of a mutation.

Previous studies have described similar roles for *ANAC019*, *ANAC055* and *ANAC072* when they were constitutively and ectopically expressed. In this paper we describe the use of a combination of experimental and theoretical tools to create a network model around the three genes to identify upstream regulatory genes and downstream pathways. This analysis has illustrated common features in upstream regulators, but also a distinct set of specific interactions that may modulate the expression of each gene depending on the stress experienced. Also, analysis of pathways predicted to be downstream of *ANAC019* and *ANAC055* has shown that the two genes have very different roles, at least in the process of developmental senescence.

EXPERIMENTAL PROCEDURES

Y1H library screen

The TF library (REGIA + REGULATORS; RR Library) (Castrillo *et al.*, 2011) is a kind gift from the authors and comprises approximately 1500 TFs fused to an N-terminal GAL4 activation domain in pDEST22 (Invitrogen, <http://www.invitrogen.com>). Yeast strain AH109 (*MATa* – Clontech, <http://www.clontech.com>) was transformed with the individual TF clones as detailed by the manufacturer and 24 clones pooled per well in a 96-well plate, in two arrangements.

Gateway Conversion (Invitrogen) was performed on the pHIS-LEU2 vector described in Çevik *et al.* (2012) to generate pHIS-LEU2GW. Overlapping promoter fragments of approximately

400 bp were amplified in a two-step polymerase chain reaction (PCR) from *Arabidopsis* (Col-0) genomic DNA using KOD DNA polymerase (Merck, <http://www.merckmillipore.com>) for *ANAC019*, *ANAC055* and *ANAC072*. Fragments were amplified with sequence-specific oligonucleotides containing half attB Gateway recombination sites (Table S3). Second round PCR was performed with generic attB oligonucleotides (Table S3). Promoter fragments were cloned into the pDonrZeo vector (Invitrogen) using BP clonase II (Invitrogen) and then recombined into pHISLEU2GW using LR clonase II (Invitrogen). Yeast strain Y187 (*MAT α*) (Clontech) was transformed with the pHISLEU2GW-promoter clones to generate bait strains.

The pooled library and bait strains were grown in SD-Trp or SD-Leu media respectively. 3 μ l of each promoter strain was spotted onto YPDA (yeast, peptone, dextrose, adenine) plates and overlaid with 3 μ l of TF library pools. After incubation for 24 h at 30°C, diploid cells were replica plated onto selective plates [SD-Leu-Trp and SD-Leu-Trp-His \pm 1–100 mM 3-amino-1,2,4-triazol (3AT)]. Following overnight incubation, plates were replica-cleaned, then incubated for 4 days. Growth was scored and positive colonies patched onto selective plates and grown overnight at 30°C. Colony PCR was performed by adding a colony to 20 mM NaOH, boiling for 10 min, then these were used as a template in a PCR reaction using pD22 oligonucleotides (Table S3). Products were sequenced to identify the TF showing a positive interaction.

To verify the Y1H results, Y187 was transformed with all promoter constructs and then with pDEST22 or the appropriate TF clone. Cultures were grown in SD-Leu-Trp, diluted to 10⁸ cells/ml, 3 μ l spots of serial 10-fold dilutions plated onto selective plates (SD-Leu-Trp and SD-Leu-Trp-His \pm 3AT) and grown at 30°C for 3 days before scoring.

Prediction of transcription factor binding sites

Position specific scoring matrices (PSSMs) that model DNA-binding specificities for TFs isolated from the Y1H screen were retrieved from the TRANSFAC (Matys *et al.*, 2006) and PLACE (Higo *et al.*, 1999) databases. PSSMs for a similar TF were used when absent from the databases. The matrix similarity score (Kel *et al.*, 2003) was computed at each position and converted to *P*-values based on a score distribution of that PSSM on random sequence. Motif instances that achieved a score <0.001 were judged to be candidate binding sites.

Promoter mutations and quantification of Y1H interactions

Promoter mutations were generated by inverse PCR on entry clones containing the relevant promoter sequences using oligonucleotides shown in Table S4. Entry clones were recombined with the pHISLEU2GW plasmid using LR clonase II. Serial five-fold dilutions of Y187 strains containing the promoter mutant clones and relevant TF were plated as described above. Three independent isolates of each promoter-TF pair were plated in triplicate onto selective plates (SD-Leu-Trp and SD-Leu-Trp-His \pm 3AT) and grown at 30°C for 3 days before scoring. Using a consistent sized circle, the integrated density function in ImageJ was used to measure the growth of each yeast spot, normalized by subtracting the integrated density of an adjacent equal sized area of empty agar.

GO analysis

Gene ontology (GO) annotation analysis was performed using BiNGO 2.3 (Maere *et al.*, 2005). Over-represented categories were identified using a hypergeometric test with a significance thresh-

old of 0.05 after Benjamini–Hochberg false discovery rate (FDR) correction (Benjamini and Hochberg, 1995) with the whole annotated genome as the reference set except for the analysis of interacting TFs in the Y1H experiment in which all TFs were used as the reference set.

Causal structure identification

The Gaussian process two-sample (GP2S) approach was used to determine differential expression of each gene in the cold, osmotic and salt stress datasets from Kilian *et al.* (2007). GP2S was implemented as described in Windram *et al.* (2012), except that a log-likelihood ratio of >8 was chosen as the threshold for indicating differential expression. For the *B. cinerea* and senescence time series differential expression was from our previous studies (Breeze *et al.*, 2011 and Windram *et al.*, 2012 respectively). The hCSI approach (Klemm, 2008; Penfold and Wild, 2011; Penfold *et al.*, 2012) was used to infer a separate network topology for the three NAC genes using data from each of five datasets described above, using the Y1H network as a constraining hypernetwork. Initial hyperparameters and prior distributions over the hyperparameters for the Gaussian process priors were set as in Penfold *et al.* (2012). The maximum number of TFs that could bind simultaneously within the algorithm was limited to five if the total number of putative regulators was <15 and 4 otherwise, due to the combinatorial scaling. Five Markov chain Monte Carlo chains were run in parallel, each generating 50 000 samples network structures with the first 10 000 sampled discarded to allow equilibration of the algorithm. The remaining 200 000 samples were thinned by a factor of 5 and used to calculate the marginal probability for each pairwise connection in the Y1H network.

Plant material and stress treatments

The *myb2*, *myb108* and *anac055* lines were T-DNA insertion lines Salk_045455, Salk_024059 and Salk_011069 respectively (obtained from the Nottingham Arabidopsis Seed Centre). The *anac019* dSpm insertion mutant was identified with gene-specific primers in a pool of SLAT line DNA (Tissier *et al.*, 1999). Arabidopsis plants were grown mostly as described by Windram *et al.* (2012). For the developmental senescence timecourse, *anac019* and *anac055* mutants and their WT controls, Col-5 and Col-0, were grown as described by Breeze *et al.* (2011); leaf 7 was tagged with cotton 18 days after sowing (DAS) and harvested from five randomly selected plants, 8 h into the light period, at 23, 29, 31, 33 or 35 DAS (full senescence).

Botrytis cinerea pepper strain spores (Denby *et al.*, 2004) were prepared and Arabidopsis leaves treated as described in Windram *et al.* (2012). Col-0, *myb2* and *myb108* leaves were inoculated with several 10 μ l droplets of *B. cinerea* spores. Replicate samples for the comparison between *myb108* or *myb2* and Col-0 were harvested at 26 and 30 hpi or 24 and 30 hpi respectively.

For the dark induces senescence screen, nine 3-week old Col-0, *myb2* and *myb108* rosettes, were cut and transferred to water-saturated filter paper and stored at 20°C in complete darkness. Plates were photographed daily and RGB colour values calculated for leaf 5 of each rosette using the Color Histogram function in ImageJ. RGB intensities were normalized using a white-background reference point and average red–green ratios provided a quantitative measure of leaf yellowing. A red–green ratio of around 0.8 indicates the initiation of senescence. When the average ratio of Col-0 samples was >0.8, leaf 5 for Col-0, *myb2* and *myb108* lines was harvested (four biological replicates). The same sampling procedure was then performed on consecutive days to sample as senescence progressed.

Microarray analysis

Total RNA was extracted from four leaves per line, labelled and hybridized to CATMA v4 arrays (Allemeersch *et al.*, 2005; <http://www.catma.org>) as described (Breeze *et al.*, 2011). For analysis of Col-0, *myb2* and *myb108* samples four replicates were pooled and labelled twice with each dye giving four technical replicates. Comparisons were made pairwise between WT and mutant under each condition. For analysis of the Col-0 and *anac055*, and Col-5 and *anac019*, biological replicates were labelled separately, twice with each dye, and comparisons made within and between time points in a 'loop design' (Kerr and Churchill, 2001). Analysis of expression differences between Col-0 and *myb2* and Col-0 and *myb108* under each condition was performed using the R Bioconductor package limmaGUI (Wettenhall and Smyth, 2004) applying Print-Tip lowess transformation and quantile-normalization. The data were fitted to a linear model using a least squares method, *P*-values adjusted to control the false discovery rate (Benjamini and Hochberg, 1995). Analysis of the *anac019* and *anac055* time course experiment was performed using a local adaptation of the MAANOVA package as described (Breeze *et al.*, 2011). The GP2S approach was used (as described in Windram *et al.*, 2012) to identify differentially expressed genes (log-likelihood ratio of >5) in the *anac019* and *anac055* mutants compared with WT. A *t*-test analysis was then performed to identify genes that were differentially expressed at each time point. Genes expressed at a higher or lower level than WT (ratio >1.7 or <0.6 respectively, *P*-value <0.1) were identified (Tables S1 and S2).

Data repository

The microarray data used in this paper have been deposited in NCBI's Gene Expression Omnibus (Edgar *et al.*, 2002) and have been given a GEO Series accession number GSE46318. These data will be released on publication.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article.

Figure S1. Identification of putative binding locations for transcription factors that interact with fragments of the *ANAC019* promoter in Y1H assays.

Figure S2. Identification of putative binding locations for transcription factors that interact with fragments of the *ANAC055* promoter in Y1H assays.

Figure S3. Identification of putative binding locations for transcription factors that interact with fragments of the *ANAC072* promoter in Y1H assays.

Figure S4. Hierarchical CSI modeling was used to identify a treatment-specific subnetwork for various stress datasets (developmental senescence, *Botrytis cinerea* infection, salt, osmotic and cold stresses) based upon the core Y1H network.

mental senescence, *Botrytis cinerea* infection, salt, osmotic and cold stresses) based upon the core Y1H network.

Figure S5. Expression of *MYB2* and *MYB108* is positively correlated with that of *ANAC019*, *ANAC055* and *ANAC072* during *B. cinerea* infection and developmental senescence.

Figure S6. Gene expression patterns of selected genes representing GO terms identified as being enriched at one or more time-points during the experiment.

Table S1. Genes differentially expressed in the *anac019* mutant compared to Col-5 WT control over the whole time course were identified using GP2S sampling (Stegle *et al.*, 2010) and genes with a GP2S score >5 were retained.

Table S2. Genes differentially expressed in the *anac055* mutant compared to Col-0 WT control over the whole time course were identified using GP2S sampling (Stegle *et al.*, 2010) and genes with a GP2S score >5 were retained.

Table S3. Oligonucleotides used to generate promoter fragment clones for Y1H screens.

Table S4. Oligonucleotides used to generate mutations in the DRE motif.

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