Introduction

Phenotypic plasticity in response to adverse conditions determines plant productivity and survival. Abiotic stress results in the largest loss in crop yields worldwide [1] and is a major threat to crop sustainability [2]. Thus, improving abiotic stress resistance is considered to be a main route for sustainable yield growth and will likely become progressively more important as arable land becomes increasingly limited [3] due to (1) the deterioration of water resources [4], (2) predicted increase in the occurrences of climatic extremes [5], (3) the predicted increase in the occurrences of climatic extremes [7].

Plants reduce growth under adverse conditions as a mechanism to avoid potentially lethal stresses [8]. In addition, plants can utilize environmental cues to detect and anticipate imminent, adverse conditions and correspondingly adjust their growth [9]. For example, woody perennials (trees and shrubs) from temperate latitudes stop shoot growth in response to short-day (SD) photoperiods that signal the approaching winter and impending months of dehydration and freezing conditions. The cessation of shoot growth precedes a more permanent growth inhibition known as winter dormancy that can last months, requires development of a specialized organ (e.g., bud), and entails physiological resetting to allow resumption of growth [10].

Gibberellins (GAs) are involved in regulating several aspects of plant growth and development [11–13]. The GA metabolic and signaling pathways have been extensively studied. The GA 2-oxidases (GA2ox) and DELLA domain proteins, like GAI (GA-insensitive) and RGL1 (repressor of ga1-3 like), are important regulators of GA levels and signaling. GA2oxs are enzymes that catalyze the 2-oxidation inactivation of both bioactive GAs and some of their precursors [14–16]. Overexpression of GA2oxs in transgenic plants leads to bioactive GA-deficiency and various levels of dwarfism [15,16]. GA2oxs are encoded by small gene families which regulate specific processes in plants, in part by specific expression patterns [15–18]. DELLA domain proteins are strong repressors of several GA responses and characterized by the conserved DELLA domain which mediates the susceptibility of the protein to proteolytic degradation [19]. Mutant forms of these proteins (gai and gfl1) with truncation of the DELLA domain are resistant to degradation and impart repressive blocks to several GA-mediated responses [20–22].

An accumulating body of evidence suggests that DELLA domain proteins and GA2oxs are involved in plant abiotic stress...
response. For example, activation of DELLA domain proteins appears to be crucial for restraining growth in adverse conditions [23–25]. DELLA proteins are believed to affect both cell expansion [11] and proliferation [26,27]. For instance, DELLA proteins have been shown to inhibit cell proliferation through elevation of cell cycle inhibitors [27], and to promote cell differentiation by reducing inhibitors of the developmental transition from mitosis to endoreduplication that modulate anaphase-promoting complex/cyclosome activity [26]. DELLA proteins not only inhibit growth but also promote plant survival under stressful conditions by limiting the accumulation of reactive oxygen species (ROS), thus delaying cell death [25]. In Arabidopsis (Arabidopsis thaliana), salt stress leads to DELLA protein stabilization and as a consequence, growth inhibition and increased plant survival [23]. In addition, a DELLA protein in Arabidopsis has been shown to bind to the promoter and increase expression of the XERICO gene, which is involved in drought response [28]. DELLA proteins have also been implicated in mediating hormonal cross-talk between GA and abscisic acid (ABA) signaling pathways [23,28]. ABA is a growth inhibiting hormone that regulates one of the two major stress signal transduction pathways in plants [29].

In addition to modulation of GA sensitivity, stressful conditions can directly influence levels of bioactive GAs. For example, cold treated Arabidopsis plants have been shown to have increased expression of three GA 2-oxidase (GA2ox) genes [24], whereas under salinity stress six GA2ox genes were shown to be up-regulated [30]. Furthermore, the cold-inducible CBF1/DREB1b protein in Arabidopsis imparts freezing tolerance, at least in part by activating the expression of GA2ox genes, which in turn leads to reductions in bioactive GAs and suppression of growth [24]. Similarly, in Arabidopsis the DWARF AND DELAYED FLOWERING 1 (DDF1) protein, involved in salt stress response, binds to the promoter and activates the GA2ox7 gene [30]. Though GA2ox7's role in control of seed dormancy has been well substantiated [31,32], their involvement in regulation of winter bud dormancy is based solely on correlative evidence. For instance, in several tree species, SD-induced transition to dormancy is associated with reduction in bioactive GAs [33–35].

Changes in GA catabolism and signaling can have profound effects on tree growth, phenology, morphology, physiological, metabolism, and gene expression [36–39]. In Populus, GA-deficient and GA-insensitive transgensics are semidwarfs of varying degrees of severity [39]. Semidwarfarism has also been associated with other characteristics that could be advantageous under adverse conditions such as increased biomass allocation to roots, reduced stem elongation, and increased water use efficiency [40]. The wide diversity of effects associated with GA modulation suggests that GA metabolism and signaling could be involved in mediating plant adaptive responses to adverse environmental conditions. Here, using a diverse array of evidence, we show that both DELLA and GA2ox encoding genes in hybrid poplar (Populus tremula x Populus alba) constitute a major regulatory circuit mediating growth restraint and physiological adaptation to drought stress and SD photoperiods.

Results

Poplar DELLA Domain and GA2ox Encoding Genes are Induced by Drought and SDs

We studied expression of four poplar DELLA protein (PtaGAI1, PtaGA2, PtaRGL1-1, and PtaRGL1-2) and seven PtaGA2ox (PtaGA2ox1 to 7) encoding genes (Table 1) in leaves in response to drought and SD photoperiods (Figures 1 and 2). Drought treatment, imposed through water deprivation under greenhouse conditions, increased expression in two of the four DELLA protein encoding genes and four of the seven PtaGA2ox genes (Figure 1). Expression increased weekly, reaching peak levels for most genes at the end of the studied period. The largest increase in expression occurred for PtaGA2ox2 and PtaGA2ox7 which showed seven-fold induction (Figure 1). To study the role of the same genes in growth cessation during SD-induced bud dormancy, we imposed a SD photoperiod (8 h light/16 h dark) under controlled growth chamber conditions (see Materials and methods) and followed changes in expression in the leaves on a weekly basis. Expression of three of the four DELLA protein encoding genes and three of the seven PtaGA2ox genes increased significantly (Figure 2). There was substantial overlap in the expression of genes up-regulated by both drought and SDs (PtaRGL1-1, PtaRGL1-2, PtaGA2ox1, PtaGA2ox3, and PtaGA2ox7). Expression of only two (PtaGAI1 and PtaGA2ox2)

Figure 1. Poplar DELLA domain and GA2ox encoding genes were significantly up-regulated in response to drought stress. Shown are mean±SE of RT-PCR results for three biological reps each consisting of leaf tissue pooled from 2–3 plants for well-watered control (C) and water-withholding (1–3 weeks) treatments. Expression was normalized to Ubq and Cyc. Significant differences between watered and water-withholding treatments were determined by one-way ANOVA followed by Dunnett’s post-hoc test (*, P<0.05).

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Inhibition in Response to Drought and SDs

The induction of *DELLA* and *GA2ox* genes in response to drought and/or SDs suggests that these genes may mediate growth inhibition during both responses. We took advantage of previously, well-characterized, GA-deficient (35S::P*GA2ox*) and GA-insensitive (35S::rgl1 and pGA1:ga1) transgenic poplar with representative, stable, and intermediate/semidwarf phenotypes (see Materials and methods) to study their response to drought stress and SD photoperiods. To ensure that any inherent differences in size between genotypes did not obscure analysis of treatment effects, relative growth rates were used to determine significant differences between WT and transgenic plants. Experiments (drought and SD photoperiod) with transgenic plants were performed in a similar fashion to that of expression analysis and as described in the Materials and methods. Prior to implementing drought and SD photoperiod experiments, weekly relative growth rates were not significantly different between transgenic and WT plants under well-watered conditions and long-day photoperiods (Figure 3). For the drought experiment, to further facilitate valid comparisons between different genotypes, methods recommended by Verslues et al. [41] were employed; whereby, transgenic and WT plants were grown in the same pots so that roots of all genotypes would grow into the same soil and be exposed to similar conditions (see Materials and methods) (Figure S1). Because of the leaves’ importance in controlling water loss and previous results of transgenic genotypes having specific effects on leaf size [39], we measured leaf area and expansion (Figure S1). In support of our previous findings [39], the leaf area of GA-deficient transgenics was significantly different than WT, whereas GA-insensitive transgenics was not. However, expansion rates of newly formed leaves were similar between all transgenics and WT throughout the experiment (Figure S1). After only one week of withholding water, the *gai* and *rgl1* expressing transgenic plants had significantly reduced weekly relative growth rates in height, diameter, and number of nodes compared to WT (Figure 3A). Three weeks post-water deprivation, growth was virtually absent in *gai/rgl1* expressing plants whereas WT, and to a lesser extent *GA2ox* expressing plants, did not completely cease growth until weeks five and six. Interestingly, water deprivation also affected secondary woody growth (stem diameter at the base) in the *gai/rgl1* transgenics, as indicated by their significantly decreased growth rates relative to WT in weeks three through five (Figure 3A).

We also studied transgenics’ growth response to SD photoperiods that induce winter dormancy. The first response to SDs, which precedes and is a prerequisite for dormancy, is cessation of shoot growth. Poplars are highly photoperiod sensitive, and the genotype under investigation, *P. tremula x alba* (717 1B4), cease growth after three to five weeks under SD photoperiod [42]. All transgenics had significantly greater, early reductions in weekly relative growth rate compared to WT plants (as early as one week under SD) (Figure 3B). WT plants had a more gradual reduction in weekly growth and, as in the drought experiment, did not completely cease growth until the fifth week under SD conditions. In contrast to drought, we did not observe any differences between transgenics and WT with respect to reduction of diameter growth under SD conditions. Despite differences in growth cessation, the timing of bud set was not significantly (P>0.05) affected and occurred around week five in both transgenics and WT (data not shown).

**Physiological Changes in Response to Drought Stress**

Drought stress has a profound effect on a number of physiological parameters in plants, and measurement of these parameters is useful in determining their stress response and resistance [43,44]. To test if the *GA2ox* and *DELLA* expressing transgenic plants differ with respect to their physiological responses to drought, we measured several important parameters before and during drought stress response. We quantified pigment concentrations, photosynthetic rate, transpiration, and stomatal conductance, which are frequently used to measure the degree of stress imposed on plants (Table 2). In general, GA-insensitive and GA-deficiency had similar effects on measured responses (Table 2). Under well-watered conditions, all transgenics had significantly higher photosynthetic rates compared to WT (Table 2). Increases in photosynthetic rate corresponded to significantly higher total chlorophyll and carotenoids for all transgenic types (Table 2). Typically, drought stress reduces photosynthesis and causes
In support of our quantitative pigment measurements, signs of senescence after water deprivation were more visible in WT leaves compared to transgenics (Figure 5). Percent of senescing leaves was not significantly different, but an apparent trend of more advanced senescence compared to WT but only those expressing gai were significantly different (Table 2).

To assess the overall stress resistance of the different genotypes, we measured wilting and electrolyte leakage (EL). The DELLA expressing transgenics showed significantly lower levels of wilting compared to WT (Figure 4A). The GA2ox transgenics also showed a slight decrease in water use efficiency as compared to WT but only those expressing gai were significantly different (Table 2).

Delayed Senescence in GA-deficient and GA-insensitive Poplars

In support of our quantitative pigment measurements, signs of senescence after water deprivation were more visible in WT leaves compared to transgenics (Figure 5). Percent of senescing leaves was not significantly different, but an apparent trend of more advanced senescence in WT as compared to transgenics was evident (Figure 5).

Leaf senescence preceding winter dormancy is typically initiated by SD photoperiods but is enhanced by low temperatures (<10°C). To investigate senescence in GA-deficient and GA-insensitive plants, we subjected transgenic and WT plants to six weeks of SDs at 21°C followed by three weeks at 4°C and measured the extent of leaf senescence. In a manner similar to their response to drought, all transgenic plants showed delayed senescence when compared to WT (Figure 6).

Increased Axillary Shoot Outgrowth in GA-deficient and GA-insensitive Poplars

As with bud set, there were no significant differences in the timing of the first bud to flush after winter dormancy between transgenics and WT (Figure 7). However, in WT plants apical buds were always the first to initiate growth, whereas in transgenics axillary lateral buds typically flushed first (Figure 7).

In GA2ox overexpressing transgenics, this was followed by a significant increase in lateral branch outgrowth as compared to WT plants (Figure 8A and B). In contrast to GA2ox transgenics, the flushed axillary buds in the DELLA expressing plants never elongated and remained in a leaf rosette stage. Because of the significant increase in lateral branches in GA2ox transgenics, we measured total height and branch growth three months after bud flush. Although GA2ox transgenics were significantly shorter, their branch length was significantly greater compared to WT (Figure 8C and D).

Differential Expression of a Large Number of Genes in GA-modified Transgenics

Because the GA-modified transgenics showed accelerated response to drought and SDs, we hypothesized that the mechanisms associated with these responses are constitutively elevated even under control conditions (well-watered and long-day photoperiods). We therefore used whole-genome poplar microarray to compare transcriptomes of transgenic and WT leaves from plants grown under a control environment. We found 2,890 differentially expressed genes (ANOVA, P<0.01) (Table S1). Gene ontology (GO) analysis was used to gain insight into the global patterns of gene expression. Consistent with our results showing increased resistance to drought stress, we found 'response to stress' (GO:0006950) in the top 10 most significantly-enriched biological categories in the transgenic plants (Table S2). Among the genes associated with stress response were orthologs of CBF1 and CBF3 that encode AP2/ERF type transcription factors, and previously, CBF1-mediated cold stress response was shown to involve reductions in bioactive GA through increased GAI expression, which promoted DELLA protein accumulation [24].

Table 1. Names, models, and primer sequences for genes used in expression analysis.

| Name           | Model                  | Forward                   | Reverse                   |
|----------------|------------------------|---------------------------|---------------------------|
| PtaGA2ox1      | Potri.001G378400       | TGAGATTTCTGAAATGATGGCT    | GCCTATTTACGGCAATCTGGAG    |
| PtaGA2ox2      | Potri.002G191900       | TGATGGTAAAGGCGATGAG     | TCAAGATTTGGAAGTGGGAG     |
| PtaGA2ox3      | Potri.004G065000       | GCTGATCCCTGTAAACCAAGAA   | TCAATAGCCCAAGCTGTAACGTCAC |
| PtaGA2ox4      | Potri.008G101600       | ATGTTAATGGTACCTCCAAAC   | TACTCGAATGTGTGGGCA       |
| PtaGA2ox5      | Potri.010G149700       | ATCTGATTTGAGCAGAGATTG   | GTTACGCTTTGGTGGCTCAC     |
| PtaGA2ox6      | Potri.011G095600       | AATGATTATTTGGTGGACAC    | TAAACCTATTTCTGAAGTATTTGGCAAG |
| PtaGA2ox7      | Potri.014G117300       | TATTCCATGAGGCTCAGAGAGC   | GATTTTGCTGGAACACATGTG    |
| PtaKL1_1       | Potri.004G089800       | TCAACGGTTGAAGATGATCAAG   | CACCTCTGGAACACTCTCC      |
| PtaKL1_2       | Potri.017G125200       | GGTGTTGCTGGGAAATCTT     | TCAAACTCTCCAAACTGACC     |
| PtaGA11        | Potri.008G131700       | CACCGAGTCTGGGCTGTT      | CAAATTAGCGCTCCTAAACTCACT |
| PtaGA12        | Potri.010G110700       | TTATACCCCTGAAAAATACCAG   | TACTGATTGCTAGCTGTCGCT    |
| PtaCYC         | Potri.004G168800       | GCCTAATTTTGCCGATGAGA   | AGCTCACCTCCATCACAACAC    |
| PtaUBQ         | Potri.002G224700       | AGAGGTGTGAGAGAGAGAGAG   | CGAGCACGATCATCAAACAGAG    |

Gene models are according to poplar genome v.3.

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Likely because GA is highly integrated into the regulatory network of other hormones [45], we found that at least one aspect of each of the five major hormones (e.g., auxin, cytokinin, brassinosteroid, ABA, ethylene) was significantly affected in the GA-modified transgenics (Table S2). GO categories associated with ethylene were prevalent in a number of categories and enrichment significance (Table S2). The trends in expression of these ethylene-associated genes are indicative of increased production and signaling through the ethylene signal transduction pathway. For example, genes encoding biosynthetic enzymes, such as acetyl-CoA carboxylase and acetyl-CoA synthetase, were up-regulated while the catabolizing ETHYLENE OVERPRODUCER 1 was down-regulated. Furthermore, downstream AP2/ERF type transcription factor genes were also up-regulated.

GA metabolism affects both cell expansion and proliferation [11]. The growth restraining effects of DELLA proteins has been shown to be, at least in part, due to regulation of cell proliferation [26,27]. Previously, Achard et al. [27] showed that DELLA proteins restrain cell production by causing up-regulation of cell cycle inhibitors. We found the Kip-related protein (KRP3), a negative regulator of cell division, to be up-regulated in both GA-modified transgenics (Table S1), which is suggestive of a common mechanism for growth restraint through inhibition of cell proliferation.

**GA-deficiency and GA-insensitivity Share Common Transcriptome Responses with Dormancy and Drought**

Because our studies suggest that GA-insensitive and GA-deficient plants have faster and more robust responses to dormancy-inducing conditions and drought, we speculated that alterations in GA metabolism and signaling in transgenics may result in transcriptome level changes that are shared with plants that are responding to stress (i.e., SD photoperiod and drought). Therefore, we compared the transcriptome of GA-deficient and GA-insensitive poplars to previously published transcriptomes of WT poplars during dormancy induction and drought response [46,47]. Unfortunately, the data for dormancy induction was based on apices; microarray data for expression changes in poplar leaves was not available. Nevertheless, transcriptome comparisons could be useful in identifying differentially expressed genes that are not tissue-confounded. Indeed, nearly a quarter of all differentially expressed genes in the GA-modified transgenics were common
with the genes found to be regulated during both dormancy induction and drought (Figure 9). In addition, the transgenics' transcriptomes shared additional overlaps with either dormancy induction or drought. Among the most significantly enriched GO terms in the common transcriptome were processes associated with chloroplast biogenesis and function (Table S3).

Discussion

Survival and productivity of long-lived perennial plants in temperate zones are dependent on robust responses to prolonged and seasonal cycles of unfavorable conditions. Here we report whole-genome microarray, expression, physiological, and transgenic evidence suggesting that GA catabolism and repressive signaling mediate shoot growth inhibition in response to both drought and SD-induced bud dormancy. Both water deprivation and SDs elicited up-regulation of a suite of poplar GA2ox and DELLAs' protein encoding genes (Figures 1 and 2). We also found that the two environmental factors elicited similar responses in the same set of genes with the only exception being PtaGAI1 and PtaGA2ox2, which were up-regulated by one but not the other factor. Particularly instructive are the trends in expression of the GA2ox genes, for which we have performed a significant amount of previous work [17,18]. For example, we found that GA2ox3 was up-regulated by both drought and SD photoperiods (Figures 1 and 2). PtaGA2ox3 is predominantly expressed in the apex, which contains the shoot apical meristems (SAM); furthermore, it was the only gene for which we failed to regenerate RNAi-suppressed

Table 2. Physiological changes in GA-deficient and GA-insensitive poplars under well-watered and drought conditions.

| Condition | Genotype | Total Chlorophyll (mg g\(^{-1}\)) | Chlorophyll a/b ratio* | Carotenoid (mg g\(^{-1}\)) | Photosynthesis Rate (μmol m\(^{-2}\) s\(^{-1}\)) | Transpiration Rate (mmol m\(^{-2}\) s\(^{-1}\)) | Stomatal Conductance (mol m\(^{-2}\) s\(^{-1}\)) | Water Use Efficiency (mol mmol\(^{-1}\)) |
|-----------|----------|---------------------------------|------------------------|-----------------|------------------------|------------------------|------------------------|------------------------|
| Watered   | WT       | 1.28 ± 0.05c                    | 1.96 ± 0.12a           | 0.24 ± 0.01c    | 12.32 ± 0.22b          | 6.00 ± 0.29a           | 0.32 ± 0.07ab          | 2.08 ± 0.07b           |
|           | 35S::GA2ox | 2.37 ± 0.09a                    | 1.68 ± 0.16ab         | 0.36 ± 0.01a    | 15.65 ± 0.56a          | 6.18 ± 0.29a           | 0.39 ± 0.07ab          | 2.51 ± 0.06a           |
|           | pGA::Gal | 1.67 ± 0.05b                    | 1.38 ± 0.10b          | 0.26 ± 0.01b    | 14.07 ± 0.17a          | 5.92 ± 0.36a           | 0.35 ± 0.07ab          | 2.54 ± 0.14a           |
| Drought   | WT       | 1.55 ± 0.03b                    | 1.94 ± 0.11a          | 0.28 ± 0.01b    | 6.56 ± 0.64c           | 1.73 ± 0.18b           | 0.07 ± 0.01b           | 3.93 ± 0.17a           |
|           | 35S::GA2ox | 1.99 ± 0.07a                    | 1.96 ± 0.12a          | 0.34 ± 0.01a    | 10.3 ± 0.45ab          | 2.97 ± 0.3a            | 0.17 ± 0.02a           | 3.72 ± 0.20ab          |
|           | 35S::rgl1 | 1.98 ± 0.05a                    | 1.94 ± 0.17a          | 0.32 ± 0.01a    | 8.44 ± 0.65b           | 3.18 ± 0.48a           | 0.20 ± 0.04a           | 3.12 ± 0.28ab          |
|           | pGAI::GAI | 2.12 ± 0.06a                    | 1.36 ± 0.06b          | 0.33 ± 0.01a    | 9.05 ± 0.39ab          | 3.01 ± 0.39a           | 0.18 ± 0.04a           | 3.36 ± 0.22b           |

Means ± SE are based on at least three biological replications. Well-watered and drought conditions were analyzed separately. Statistically significant differences were determined by one-way ANOVA followed by Tukey's post-hoc test (P < 0.05). Letters following mean ± SE indicate statistical significance, where treatment (i.e., Genotype) means with the same letter are not significantly different.

* Drought measurements 5 weeks after withholding water.

** Drought measurements 3 weeks after withholding water.
transgenic plants, strongly suggesting that this gene is involved in organization of SAM [18]. Because growth, and particularly primary growth which originates in SAM, is most affected by drought and SDs, we believe that the up-regulation of PtaGA2ox3 is likely an important mechanism for controlling growth through modulation of SAM activity. Another gene that was up-regulated by both drought and SDs was PtaGA2ox7. We have previously found that PtaGA2ox7 and its paralog PtaGA2ox2 are predominantly expressed in roots under optimal conditions and positively regulate lateral root proliferation [18,37]. The fact that they are up-regulated by drought in leaves suggests they are involved in a mechanism for coordination of shoot-root ratio in relation to resource availability. Decreased aerial growth with increased lateral root growth would produce a plant that demands less of limited resources while still actively exploring the soil environment.

The commonalities in the transcription responses to drought and SDs are interesting because the signaling mechanisms induced by these two types of environmental cues are quite different. One involves a response to osmotic stress [48], while the other encompasses the light signaling pathway [49]. This suggests that GA2ox and DELLA protein genes are downstream of different signal transduction pathways that converge on the same types of genes. One possible convergence point could be through cross-talk with ABA biosynthesis or response. For example, DELLA proteins have been shown to regulate XERICO, which encodes a ubiquitin-ligase that positively regulates ABA biosynthetic encoding genes [28]. Indeed, our microarray work showed that a poplar ortholog of XERICO was highly up-regulated in the GA-modified transgenic plants. Alternatively the DELLA proteins themselves could be a convergence hub, as an increasing body of evidence suggests that they are focal points of multiple signaling pathways in plants [50]. Finally, ethylene can also play a role in the cross-talk, as our microarray results indicate a strong up-regulation of genes involved in ethylene biosynthesis and response.

Using previously characterized GA-deficient and GA-insensitive transgenics, with increased expression of GA2ox and DELLA protein encoding genes and representative semidwarf phenotypes [36,39,40], we show that GA-modified transgenics have significant changes in growth and physiological responses under both drought and SDs. Overall, overexpression of GA2ox and DELLA proteins in poplar caused hypersensitive growth inhibition under both
drought and SDs. Relative growth rates were not significantly different between transgenic and WT plants under well-watered conditions and long-day photoperiods. In comparison, transgenics generally responded to water withholding and SDs by reducing relative growth on average one to three weeks earlier than WT (Figure 3). Nevertheless, traits were affected differently under drought and/or SD conditions. Decrease in GA concentration and sensitivity in transgenic plants caused inhibition of primary growth (e.g., height growth and number of internodes) under drought and/or SDs. In contrast, the transgenic GA modulation had an impact on secondary growth (e.g., stem diameter growth) only under drought but not SDs. In addition, some genotype-specific response differences were found among the transgenics under the two conditions. Most notably, under drought conditions only GA-insensitive (and not GA-deficient) transgenics had significantly reduced relative growth as compared to WT plants (Figure 3A).

Figure 7. Bud flush in transgenic and WT Populus. Arrows point to the first bud to flush. Bars indicate mean±SE of at least three biological replications. Relative position of the first bud to flush was expressed as percent of buds from the top of the plant, 0% being the apex of the plant and 100% being the very bottom bud (closest to the soil). Significant differences between transgenic and WT treatments were determined by one-way ANOVA followed by Dunnett’s post-hoc test (*, P<0.05).

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Figure 8. Decreased apical dominance in the GA-deficient (35S::PcGA2ox) plants. (A) WT are on the left and 35S::PcGA2ox plants are on the right. Inset shows plants prior to entering dormancy for comparison purposes. In (B) to (D), bars represent mean±SE of eight ramets/line and eight WT. Statistically significant differences between WT and transgenic plants were determined by t-test (*P<0.05, and **P<0.01).

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Figure 9. Significant overlap between GA-deficient and GA-insensitive (GA), dormancy, and/or drought transcriptomes. See text for details.

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(discussed below); whereas, under SD conditions all transgenic types had significantly reduced relative growth (Figure 3B).

In addition to growth cessation, reduction of bioactive GA levels and GA signaling in poplar transgenics significantly modified and/or delayed the onset of several other physiological changes that are typically associated with stress response. For instance, stressful conditions typically cause a reduction in photosynthesis and promote leaf senescence, which in turn negatively affects productivity [43]. Thus, delayed leaf senescence and the ability to sustain high photosynthetic capacity in adverse conditions are highly desirable traits [51]. Under drought stress, GA-deficient and GA-insensitive transgenics sustained higher rates of photosynthesis (Table 2). Although we did not measure the photosynthetic rate during SD-induced dormancy, we did observe a delay in leaf senescence in transgenic plants (Figure 6), which is suggestive of a similar response. In support of our findings, GAs have been implicated in the repression of light-regulated genes [52]. Decreases in GA levels via up-regulation of GA2ox genes is associated with increases in the expression of light-regulated genes [52], photosynthesis [53], chlorophyll content, and light harvesting chlorophyll proteins [54]. It has also been shown that DELLA proteins are directly involved in photomorphogenesis [55]. In GA-deficient and GA-insensitive transgenics, sustained increases in photosynthesis coupled to reduced shoot growth demands could facilitate shunting of resources from primary to secondary/storage metabolism and/or investment in root growth. Previously, we have shown experimental evidence supporting these hypotheses. For example, the same transgenics also accumulate increased levels of secondary metabolites [36] and show increased levels of root production [36,37,40].

In addition, carotenoids are antioxidants that can increase stress resistance via detoxification of ROS [56], and are precursors to the stress-responsive hormone ABA [57]. In Arabidopsis, stress induced DELLA protein accumulation and increased activity of ROS-detoxification enzymes led to reductions in ROS levels [25]. Our microarray data also showed enrichment of GO categories associated with stress response (Table S2, GO:0006950) in the GA-deficient and GA-insensitive transgenics. A number of genes encoding enzymes associated with detoxification of ROS were found in this category, including peroxidases, glutathione-S-transferase, superoxide dismutase, glyoxylate reductase, mono-dehydroascorbate reductase (Table S1 and 2). Furthermore, genes associated with ABA biosynthesis, such as nine-cis-epoxycarotenoid dioxygenase 3, were also highly up-regulated. In summary, DELLA proteins and GA2oxs likely not only repress growth but enhance plant resistance to stress via activation of ROS-detoxification enzymes, increased carotenoid production and ABA biosynthesis.

To better understand the effects of overexpressing GA2ox and DELLA protein encoding genes on drought response, we must also consider the implications of GA-deficient and GA-insensitive transgenics’ semidwarf phenotype. Soil drying experiments can be difficult to interpret because the plants dictate the severity of stress associated with increases in the expression of light-regulated genes [25]. Moreover, plants tend to increase root growth, which would be advantageous to water uptake and further promote drought avoidance. Indeed, GA-insensitivity has been previously found to be associated with increased ratios of root mass to leaf area which have been suggested could help sustain higher transpiration rates and decreases in water use efficiency [40]. In summary, the GA-modifications caused complex responses that likely contribute to both drought avoidance and tolerance mechanisms.

Similar to recent findings by Mauriat et al. [58], we show that the GA2ox expressing transgenics have increased branch production which is indicative of decreased apical dominance (Figure 8). However, in contrast to this previous finding, in the present study, the increased branching phenotype developed only after winter dormancy. This difference could be attributed to the different promoters, genetic backgrounds or growing conditions used in this study (note sylleptic branching is very sensitive to environmental quality). It appears that the observed reduction of apical dominance caused by GA2ox up-regulation is a result of decreased auxin concentration and transport [58]. We also show that up-regulation of DELLA proteins leads to a similar phenotype, suggesting that the effect of GA on apical dominance, through modulation of auxin biosynthesis and transport, is downstream of and mediated through a DELLA signaling hub. However, in the DELLA transgenics, branch outgrowth after dormancy was very short-lived and resulted in a very short, stubby branch. In contrast, GA2ox transgenic plants produced very well developed branches from almost all axillary buds. This suggests that DELLA proteins have differential effects on apical dominance before and after dormancy release. Furthermore, this would imply that apical dominance before and after dormancy may be regulated via different branches of the GA signaling pathway or by a completely different mechanism involving different hormones and regulators. These two phenotypic differences in the GA2ox and DELLA
transgenics had a significant and dramatic impact on crown shape and size in the field (e.g., wide, ball-shape in GA2ox; narrow compact in the DELLA transgenics) [39].

Conclusions

In summary, we found that GA catabolism and repressive signaling are part of a highly interactive mechanism for sensing and responding to immediate (drought) and imminent (SD signaling approaching winter) conditions involving ceasing or reducing growth as well as increasing physiologically acclimative responses to stress. These findings suggest that regulation of GA metabolism and response may be a focal point for evolution of various adaptive strategies in response to short-term and prolonged unfavorable conditions. It also suggests novel venues to engineer and breed for improved stress resistance in crop plants.

Materials and Methods

Plant Material

Genetic background for all transgenic plants was the INRA 717-1B clone (Populus tremula x P. alba). For consistency, all experiments involving WT plants were performed using this same genotype. Generation of 35S::PcGA2ox, pGAI::gai, and 35S::gaiI transgenics was previously described [17,36]. The Arabidopsis (Arabidopsis thaliana) DELLA genes gai and gaiI have complete truncations of the DELLA domains which confers a gain-of-function dominant mutation with constitutive repression of GA signaling. The gai gene was under the control of the native Arabidopsis promoter (pGAI) while gaiI was under the control of the cauliflower mosaic virus 35S promoter (35S). The GA2ox1 gene was from Phaseolus coccineus (PcGA2ox) and was also under the control of the 35S promoter. Because constructs were previously well-characterized with respect to transgene presence, expression, GA content and stability of phenotype over many years [37,39], for each transgenic type one representative line with multiple ramets was selected for use in subsequent studies. The differences in the levels of expression of these transgenes in independent transformation events elicits a gradient of phenotypic responses ranging from severe dwarfism to nearly wild-type like [39]. To avoid confounding effects in severely affected plants, we selected lines with intermediate (semi-dwarf) phenotypes.

Experimental Design

The drought experiment consisted of four genotypes in a completely randomized block design with eight replications. Each block represented a rectangular pot (height = 32 cm, length = 50 cm, and width = 35 cm) assigned four plants: one from each of three transgenic genotypes (35S::PcGA2ox, pGAI::gai, and 35S::gaiI) and a WT (untransformed control). Plants were randomized and spaced in a rectangular pattern (25 × 20 cm apart from each other) within pots. The confiding nature of the pots, allowed plants to be grown in a proximity where their roots were exposed to similar conditions to minimize effects of any differences in water usage between genotypes [41]. The use of relatively large, deep pots facilitated a slow, gradual soil drying process (Figure S2) that provided plants sufficient time for lengthy responses. Plants used in the experiment were propagated and grown in vitro for one month prior to their transfer to a greenhouse. In the greenhouse, plants were placed in a mixture of peat, top soil, perlite, and vermiculite (4:1:1:1, v/v) and grown for approximately two months prior to experiment implementation. The drought treatment consisted of three weeks of well-watered conditions (daily watering regime), followed by a drought stress imposed by completely withholding water for five weeks. A similar experimental regime was used on control plants used in expression analysis (Figure 1) with the exception that plants were grown in a completely randomized design (without blocks), in separate pots, and in a growth chamber (Conviron, Pembina, ND, USA).

SD photoperiod experiments were performed in a completely randomized design with eight replications. Propagation and acclimation of plants to the greenhouse were performed as described above for the drought experiment. After two months of greenhouse growth, plants were transferred to a growth chamber (Conviron, Pembina, ND, USA), with three weeks under LDs (16 h light/8 h dark, at 21°C) followed by six weeks of SDs (8 h light/16 h dark, at 21°C). Plants were then transferred to a cold room (4°C) for 11 weeks to fulfill the chilling requirement needed for resumption of growth. Finally, plants were transferred to LD conditions and bud flush was monitored on a daily basis.

Biometric and Physiological Measurements

Phenotypic measurements were made on a weekly basis. To avoid problems with soil level changes, all height measurements were made from permanent markers at the base of the plant stem. Net photosynthesis rate, transpiration rate, and stomatal conductance were measured with the portable photosynthesis system LI-6400 (LI-COR, Lincoln, NE, USA), using an air temperature of 25°C and photosynthetically active radiation of 1500 μmol m⁻² s⁻¹. Instantaneous water use efficiency was calculated as net photosynthesis rate divided by transpiration rate. Gas exchange measurements were made between 8:00 and 10:00 am weekly and at leaf plastochoxin index (LPI) = 10±1. After five weeks of withholding water, additional measurements were made including percent leaf senescence, percent leaf wilt, and electrolyte leakage (EL). EL was measured with a conductivity meter (Model 32, Yellow Springs Instrument Inc., Yellow Springs, OH, USA) using procedures modified from Ren et al. [59] and Verslues et al. [41]. Leaves (LPI = 10±1) were thoroughly washed in distilled water and three 1 cm² discs obtained from leaves were placed in tubes containing 15 ml of distilled water. Tubes were gently agitated for 4 h and initial conductivity was measured. Samples were then autoclaved for 15 min and conductivity was again measured. EL measurements were repeated three times per plant, and values were averaged and expressed as percentage of initial conductivity. Chlorophyll and carotenoid content was measured on non-stressed (well-watered) plants subjected to drought stress (after withholding water for five weeks). Pigments were extracted in N,N-dimethylformamide according to Porra et al. [60]. Pigments were quantified with a spectrophotometer using equations from Porra et al. [60] for chlorophyll a and b, and from Wellburn et al. [61] for carotenoid. Two 1 cm² discs were measured for each plant from LPI = 8±1. Methods similar to those in [62] were used to quantify the relative intensity of green in senescing leaves. Digital images of leaf surfaces were processed using ImageJ version 1.43 (http://rsbweb.nih.gov/ij/) and the Threshold Colour plugin (http://www.dentistry.bham.ac.uk/landinig/software/software.html).

Expression Analysis

All samples used in expression analysis were collected at the same time weekly, immediately frozen in liquid nitrogen, and stored at −80°C for further use. Samples were analyzed in three biological replications, each consisting of tissues pooled from two to three plants. Total RNA was isolated using the RNeasy Plant Mini Kit (Qiagen, Valencia, CA, USA) with an on-column DNA digestion using RNase-Free DNase (Qiagen), cDNA was synthesized from 2 μg of DNaseI-treated total RNA using Superscript III.
reverse transcriptase (Invitrogen, Carlsbad, CA, USA) with an oligo-dT primer. One µl of RT reaction was used for RT-PCR. Primers used in the expression analyses are shown in Table 1.

Statistical Analysis
Statistical analysis was performed using SAS 9.1 (SAS Institute Inc., Cary, NC, USA). In transgenic experiments, treatments consisted of the four genotypes under investigation. For both experiments, analysis of variance (ANOVA) was used to test for overall treatment effects (α = 0.05), and separate ANOVAs were done for each week of the experiment. When significant differences were present, Tukey’s Honestly Significant Difference (HSD) was used to make comparisons among WT and transgenics, or Dunnett’s test (P<0.05) was used when we were interested in making comparison to WT only. For analyses of growth-related parameters, the response variable is the weekly relative growth rate, calculated as: \[ Y = \frac{(X_{n+1} - X_0)}{X_0}, \] where \( Y \) is weekly relative growth rate, \( X \) is the measured growth parameter, and \( n \) is the week.

Microarray Hybridization and Data Analyses
We used a total of three individual genotypes as follows: WT, 35S::PcGA2ox and 35S::gai1. Leaves from two independent biological replicates per genotype were used, each pooled from twenty clonally propagated plants. RNA was isolated as previously described using the Qiagen RNeasy Plant Kit [17]. Prior to labeling, RNA quality was assessed by Agilent Bioanalyzer (Agilent Technologies, Santa Clara, USA) and three µg of total RNA was used to prepare biotinylated complementary RNA (cRNA). The labeling, hybridization, and imaging procedures were performed according to Affymetrix protocols at the Center for Genomics Research and Biocomputing, Oregon State University (USA), using the Affymetrix Poplar GeneChip (Affymetrix Santa Clara, CA, USA).

Collection and analysis of data were compliant with MIAME standards (Brazma et al., 2001). Data were analyzed using T-MeV [63,64]. Raw data was first normalized using RNA algorithm [65]. One-way ANOVA (P<0.01) was used to isolate differentially expressed genes. Gene ontology (GO) analyses for significant enrichments of various categories were performed using GOEAST [66] and the corresponding AGI loci using default parameters. Data for transcriptome changes during drought and dormancy were derived from previously published studies [16,47]. Because of the differences in the microarray platforms and genome versions used in these studies, we used the AGI of the closest Arabidopsis ortholog to compare the three data sets. We have previously validated the results from the array analysis using RT-PCR for 12 genes (six up-regulated and six down-regulated) for the roots from the same plants [37]. The microarray data have been deposited in the Gene Expression Omnibus (GEO) database as accession number GSE38390.

Supporting Information
Figure S1 Representative phenotypes and leaf measurements of GA-modified transgenic poplar. (A) Transgenic (35S::PcGA2ox; top left, pGAI::gai1; bottom left, and 35S::gai1; bottom right) and wild-type (WT; top right) plants grown in pots (25×20 cm apart from each other) for three weeks under well-watered conditions, prior to being subjected to five weeks under water-withholding conditions. (B) Bars indicate mean±SE of total area (cm²) of at least eight mature leaves per a genotype. (C) Leaf expansion (cm²) was measured weekly under well-watered (week 0) and water-withholding conditions (weeks 1 to 5) on eight ramets/line and eight WT plants. Measurements in G represent the total area of expansion of the first unfurled leaf [leaf plastochon index 1] after one week. Leaf measurements were made from digital images in ImageJ version 1.43 [http://rsbweb.nih.gov/ij/]. Significant differences between transgenic and WT plants were determined by one-way ANOVA followed by Dunnett’s post-hoc test (*, P<0.05). Scale bar = 25 cm (A).

Figure S2 Estimation of volumetric soil moisture content (g cm⁻³) in pots under well-water (week 0) and water-withholding conditions (weeks 1 to 5). A soil auger was used to sample whole vertical profiles of pots. Samples were oven dried at 100°C for at least 24 hrs. Values are means±SE of at least three samples taken from at least two pots.

Table S1 Differential expressed genes in GA-modified transgenics compared to WT poplar. (XLSX)

Table S2 Gene ontology analyses of genes differentially regulated in the GA-modified transgenics. (XLSX)

Table S3 Gene ontology analyses of genes regulated by the GA modifications, drought and winter dormancy. (XLSX)

Author Contributions
Conceived and designed the experiments: CZ VBB. Performed the experiments: CZ VBB. Analyzed the data: CZ VBB. Contributed reagents/materials/analysis tools: CZ VBB. Wrote the paper: CZ VBB.

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