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Rational design and testing of abiotic stress-inducible synthetic promoters from poplar cis-regulatory elements

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Summary
Abiotic stress resistance traits may be especially crucial for sustainable production of bioenergy tree crops. Here, we show the performance of a set of rationally designed osmotic-related and salt stress-inducible synthetic promoters for use in hybrid poplar. De novo motif-detecting algorithms yielded 30 water-deficit (SD) and 34 salt stress (SS) candidate DNA motifs from relevant poplar transcriptomes. We selected three conserved water-deficit stress motifs (SD18, SD13 and SD9) found in 16 co-expressed gene promoters, and we discovered a well-conserved motif for salt response (SS16). We characterized several native poplar stress-inducible promoters to enable comparisons with our synthetic promoters. Fifteen synthetic promoters were designed using various SD and SS subdomains, in which heptameric repeats of five-to-eight subdomain bases were fused to a common core promoter downstream, which, in turn, drove a green fluorescent protein (GFP) gene for reporter assays. These 15 synthetic promoters were screened by transient expression assays in poplar leaf mesophyll protoplasts and agroinfiltrated Nicotiana benthamiana leaves under osmotic stress conditions. Twelve synthetic promoters were induced in transient expression assays with a GFP readout. Of these, five promoters (SD18-1, SD9-2, SS16-1, SS16-2 and SS16-3) endowed higher inducibility under osmotic stress conditions than native promoters. These five synthetic promoters were stably transformed into Arabidopsis thaliana to study inducibility in whole plants. Herein, SD18-1 and SD9-2 were induced by water-deficit stress, whereas SS16-1, SS16-2 and SS16-3 were induced by salt stress. The synthetic biology design pipeline resulted in five synthetic promoters that outperformed endogenous promoters in transgenic plants.

Keywords: synthetic biology, synthetic promoters, de novo motif analysis, computational design, abiotic stress, cis-regulatory elements, poplar, bioenergy.

Introduction
Abiotic stresses such as water-deficit stress and salinity are major challenges in agriculture. Rising temperatures associated with climate change can exacerbate these challenges by increasing evaporation rates and plant transpiration. Thus, there is great interest and demand for improving abiotic stress tolerance through biotechnology. One approach is to introduce transgenes into crops to endow stress resistance using constitutive promoters, for example the Cauliflower mosaic virus (CaMV) 35S promoter (Wang et al., 2016). However, constitutive expression of stress resistance genes can result in unintended pleiotropic effects on plant growth when stressors are not present (Lee and Pijut, 2018; Li et al., 2012; Okumura et al., 2016; Wei et al., 2017; Yamasaki and Randall, 2016).

Expressing transgenes under the control of stress-specific inducible promoters may be useful to minimize negative pleiotropic effects (Das et al., 2011; Datta et al., 2012; Saint Pierre et al., 2012; Wei et al., 2017), but native plant promoters often result in relatively low and complex patterns of induced gene expression (Halpin, 2005; Hou et al., 2012; Peramuna et al., 2018). Synthetic promoters may be designed to overcome shortcomings of native promoters in genetic engineering (Mehrotra et al., 2011; Liu and Stewart, 2015; Rushton, 2016).

Rational design of synthetic promoters typically starts with appropriate -omics data to identify cis-regulatory elements (CREs). CREs are composed of non-coding DNA that is recognized by transcription factor(s) that specifically regulate downstream genes. Key motifs from these elements can be multimerized upstream of a core promoter sequence (Liu et al., 2014). The simplest architecture consists of a single short DNA motif that is multimerized and fused upstream of core promoter, which results in low transcriptional complexity (Rushton, 2016). Spatial and temporal transgene expression can be made more complex by integrating additional CREs, synthetic motifs and enhancers into the design (Hernandez-Garcia and Finer, 2014; Kassaw et al., 2018). In addition, synthetic promoter CREs may be composed of purely native sequences, those from in silico design, or a mix
(Roberts, 2011; Wu et al., 2018). While synthetic promoters have been quite useful in plant biology (Dey et al., 2015; Hernandez-Garcia and Finer, 2014; Liu and Stewart, 2016), their construction is challenged by our limited knowledge of appropriate CREs and their relationship in controlling gene expression.

**Populus** is an economically important genus and one of the top tree models in research. Poplar is a feedstock for paper, cellulose, wood and fibre and a candidate for next-generation biofuels (Sannigrahi et al., 2009). *Populus* genomes have been sequenced (Tuskan et al., 2006) and are routinely genetically engineered (Han et al., 2000). Because of their experimental tractability, poplar species have been widely used in physiological studies to better understand how deciduous trees respond to changing environments (Jia et al., 2017; Li et al., 2014).

Here, we describe the performance of rationally designed synthetic osmotic stress-inducible promoters in response to treatments including osmotic stress, salinity and water-deficit stress. Published poplar transcriptomic data were mined to discover CREs and motifs that were used to design and construct synthetic promoters. The resulting synthetic promoters were tested for stress induction using poplar leaf protoplasts, transiently transformed *Nicotiana benthamiana* leaves and in whole, stably-engineered *Arabidopsis thaliana* plants using reporter genes.

**Results**

**Co-expressed gene analysis from water-deficit or salt-stressed poplar transcriptomes**

Synthetic promoter design was performed using de novo cis-regulatory DNA motif discovery (Figure 1). Synthetic promoters were built using a unified architecture by multimerizing candidate motifs upstream of the CaMV 35S – 46 core promoter sequence, which was, in turn, fused to GFP.

As a first step to identify core motifs for designing synthetic promoters, we analysed gene expression patterns from published RNA-seq data of different *Populus* species. A subset of water-deficit or salt-responsive genes were identified from six different transcriptomes (four from water-deficit and two from salt stress experiments) (Cosset al., 2014; Filichkin et al., 2018; Jia et al., 2017; Tang et al., 2013; Figure S1). In water-deficit-treated poplar transcriptomes, a total of 210 genes were selected to be mined for motifs (Figure S1). The respective 210 genes were detected in at least three out of four transcriptomic studies, in which there was at least a fourfold increase in gene expression in leaves under stress conditions (Table S1). To identify salt stress-inducible genes, we compiled up-regulated genes from two different root RNA-seq data sets from salt-treated plants (Filichkin et al., 2018; Zheng et al., 2015; Figure S1). Fourteen genes were commonly detected with over fourfold increase in gene expression in leaves under stress conditions (Table S1). To identify salt stress-inducible genes, we compiled up-regulated genes from two different root RNA-seq data sets from salt-treated plants (Filichkin et al., 2018; Zheng et al., 2015; Figure S1). Fourteen genes were commonly detected with over fourfold increase in gene expression in leaves under stress conditions (Table S1). To identify salt stress-inducible genes, we compiled up-regulated genes from two different root RNA-seq data sets from salt-treated plants (Filichkin et al., 2018; Zheng et al., 2015; Figure S1). Fourteen genes were commonly detected with over fourfold increase in gene expression in leaves under stress conditions (Table S1).

**De novo DNA motif selection from promoters of expressed genes**

Promoters from 210 water-stress-related co-expressed and 340 salt stress-related expressed genes were collected from *P. trichocarpa* genome annotation (v 3.0) and analysed for common motifs using MEME Suite (Bailey et al., 2015), MotifSampler (Thys et al., 2002) and Weeder (Pavesi et al., 2004). Thirty and 34 DNA motifs (E < 0.001) were predicted by MEME to be water-deficit- and salt-stress-inducible motifs, respectively (Tables S3 and S4). The same input was used to run MotifSampler and Weeder whose overlapped motifs with MEME output are summarized together in Tables S3 and S4.

Among the predicted DNA motifs from water-deficit stress-inducible genes, 9 motifs (motifs 27, 19, 18, 13, 25, 9, 24, 1 and 4) were conserved sequentially on a 110-bp domain in the final localization image of predicted motifs by motif location diagram integrated in MEME Suite. We denote this 110-bp domain as the A domain (Figure 2a,b). The A domain was identified in gene promoters of 16 of the 210 water stress-related genes. Their locations were variably distributed over 2-kb upstream sequence from the start codon among the 16 promoters (Figure 2c; Table S5). All motifs, except motif 25, were predicted simultaneously by both MEME and MotifSampler (Table S3).

To find CREs over the 9 long motifs of interest that were putatively related to water stress, we submitted these 9 long motifs to the database of plant cis-acting regulatory DNA elements (PLACE; Higo et al., 1999). Interestingly, an auxin-responsive motif was identified in motifs 19 and 18 (Table 1). Motif 18 also contained sequences for cold and freezing stress-responsive factors such as MYC and an ICE binding factor (Ohta et al., 2018). A metal-responsive factor-binding sequence was detected in motif 9 (Table 1). Furthermore, these 9 motifs had higher statistical significance among 30 predicted DNA motifs (Table S3). However, motifs 1 and 4 had abundant A and T repeats (Table 1), suggesting that these two motifs may not be stress-specific even though their E-values were significant (Figure 2a). Since these motifs came from water-deficit stress-inducible promoters and were subsequently used to make synthetic promoters, hereafter these elements are referred to as SD (synthetic promoter from drought stress-inducible promoters) motifs.

Thirty-four DNA motifs were predicted by MEME from 340 gene promoter ‘hits’ from salt treatment studies (Tables S2 and S4). These predicted DNA motifs appeared well dispersed throughout candidate endogenous promoters. Most predicted motifs had many A/T or several sequence repeats (Table S4). Nonetheless, we found two GC-rich motif candidates (motifs 10 and 16) that were salt stress-responsive in protoplast transfection assays. Since 4 copies of motif 16 resulted in high GFP induction in transfected protoplast, we selected 20-base motif 16 (SS16, synthetic promoter from salt stress-inducible promoter 16) to make the respective synthetic promoter. SS16 has not been reported in the literature as a CRE (Table 1). The SS16 motif was predicted from information in 34 expressed gene promoters (Figure 3). For our application, we selected two genes, *Potri.002G039300* and *Potri.006G148800*, annotated as basic helix–loop–helix family protein and ethylene-responsive transcription factor genes in *P. trichocarpa* genome for further analysis (Table S5).

**Osmotic stress-related responses of native poplar promoters**

To verify that the 16 water-deficit-inducible promoters containing SD motifs and 34 salt stress-inducible promoters containing SS16 motif were stress-inducible, we performed quantitative reverse transcriptase–PCR (qRT-PCR) assays to measure gene expression
of randomly selected downstream genes in stress-treated poplar. The gene expression was assessed in wilted 717-1B4 leaves by water cessation for 10 days or salt stress by watering 200 mM NaCl for 3 days. Neither stress treatment resulted in mortality nor discernable plant damage (Figure 4a). The expression of four water-deficit-induced genes was up-regulated significantly over 1.2-fold (log2) higher in water-stressed plant leaves as compared to those grown under normal conditions (Figure 4b). Interestingly, all of these genes, except Potri.017G015700, were also highly expressed in leaves from salt-treated plants. Six randomly selected salt stress-responsive genes from 34 genes containing SS16 motif in their promoter were induced in the range of 1.5- to 4.3-fold change (log2) by the salt treatment; the expression of Potri.018G019700 increased 4.3-fold (log2) in salt treatment and was similar in water-deficit stress (Figure 4b). Potri.002G029300 had a 2.1-fold (log2) increase under salt treatment, whereas its expression was reduced under water cessation (Figure 4b). Taken together, the expression of each gene was specifically induced during water-deficit stress and salt stress, respectively. Four genes responded to both water-deficit and salt stress.

Poplar mesophyll protoplast transfection was used throughout the research as a first-tier test of promoter activity in poplar. We fused native promoters of four of the representative genes mentioned above (water stress-induced Potri.017G015700 and Potri.002G039300; salt stress-induced Potri.006G148800 and Potri.006G148800), which were annotated as transcription factors (Figure 4b). All four native promoters drove GFP expression in poplar mesophyll protoplast under 500 mM mannitol or 100 mM NaCl treatments (Figure 4d). These results provided sufficient evidence to further explore the use of the SD and SS motifs for designing functional synthetic sequences in promoters to enable a strong gene induction in response to osmotic stress.

**Synthetic promoter architecture and osmotic stress responses in poplar mesophyll protoplasts**

We developed a vector system that used two fluorescent protein genes for quantitative assessment of specific promoter activity in plant cells. Each stress-inducible synthetic promoter was fused to TurboGFP as a reporter gene. A constitutively expressed 35S::TurboRFP internal control cassette was included in each construct in order to monitor transformation efficiency and relative expression strength (Figures 5a and 6a). A wide range of GFP inducibility was observed in poplar leaf-derived protoplasts among the synthetic promoter variants (Figures 5d and 6d).

Previous results reported that several copies of a length of 4 to 8 DNA sequence could induce reporter gene expression (Liu et al., 2014). Reflecting on this study, we selected repeated formats of 7-8 and 5 bases to design synthetic promoter for water-deficit and salt induction, respectively. Since the tandem version of synthetic promoter enhanced inducible gene expression, we
optimized copy number of a core sequence by comparing GFP expression driven by a range of repeats from trimer to heptamer in transfected poplar protoplast, a logic suggested by Sharon et al. (2012). Increasing of the copy number of a core sequence led to increased GFP expression (Figure S3). Therefore, we employed the promoter format with heptamerized 7–8 base or 5 base for SD or SS motif inducibility test, respectively. Eight different heptamerized promoters of conserved 7–8 bases were synthesized from SD18, SD13 and SD9 motifs whose E-values were higher than other predicted motifs (Figure 2a). Among the discovered sequences in SD18 and SD9 motifs were auxin and calmodulin-binding elements (Table 1). In contrast, SD13 did not harbour any previously reported DNA elements. Based on the motif number designations, we denoted the derived

**Figure 2** Bioinformatics characterization of SD motifs. (a) The LOGO sequence results generated by MEME Suite of 9 predicted SD motifs that were used to construct the domains shown in panel (b). Motifs 1 and 4 contain many T and A repeats; therefore, they were excluded from further analysis. Note that motifs 18, 13 and 9 were selected based on the top scored E-values for subsequent experiments. (b) The ‘A domain’ included conserved nine SD motifs. The motifs and their locations were predicted by the MEME Suite. (c) Location of the conserved A domain on −2 kb promoter sequences of 16 co-expressed genes.

| ID | Motif sequence | Predicted element in PLACE | Sequence | Position (strand) |
|----|----------------|-----------------------------|----------|------------------|
| SD 27 | AAGAGCTTGGGACCAAGAAG | root nodule-specific element | CTCTT | 1(−) |
| 19 | GTGGTCTCAAGTTCGAGCC | auxin-responsive factor | GAGAC | 4(−) |
| 18 | GCTCATATGATGGCCACTGG | MYC, ICE1 binding | CANNIG | 4(−) |
| 25 | GGCTTACATGGTCGTTAACT | auxin-responsive factor | CATATG | 4(+) |
| 9  | GTCGAGGTACGCGCAAGCTG | light induced | GCCAC | 13(+)) |
| 13 | GGGCCCCGTGGGATTA | phyA-induced motif | GGGCC | N.D. |
| 25 | GTCGAGGTACGCGCAAGCTG | copper and oxygen responsive | GTAC | 7(+) |
| 9  | GTCGAGGTACGCGCAAGCTG | calmodulin binding | VCGCB | 9(−) |
| 24 | GCCCGGACACCCACG | N.D. | N.D. | N.D. |
| 1  | TTTCAGTGAGAAAAAAA | T-box | TTWTWTTTWT | 6(−) |
| 4  | TTTCAGTGAGAAAAAAA | T-box | TTWTWTTTWT | 6(+) |
| SS16 | CCGGTCTCAAGTTCGAG | N.D. | N.D. | N.D. |

N.D., not detected.
synthetic promoters as SD18-1, SD18-2, SD18-3, SD13-1, SD13-2, SD9-1, SD9-2 and SD9-3 (Figure 5b,c). Protoplast transfection frequency ranged from 52.9% to 59.3%, which was calculated from scoring constitutively expressed RFP fluorescence in protoplasts (Figure 5d and Figure S4; black bars). GFP was induced in the range of 2.0%–46.6% of protoplasts, depending on the synthetic promoter; promoters exhibited various levels of induction (Figure 5d and Figure S4a; white bars). SD18-1, SD13-1, SD9-2, and SD9-3 exhibited the highest induction in the mannitol treatment (Figure 5e), whereas lower induction was observed in SD18-3 and SD9-1-transfected cells. SD18-2 and SD13-2 induction was extremely low in the mannitol treatment. Transfection frequency was not different among promoter constructs tested (Figure S4a). Image analysis was used to gauge GFP intensity and protoplast counts, which appeared to positively correspond to one another, with the exception of SD13-2 (Figure 5f).

For the purpose of discovering the minimal submotifs responsible for salt stress induction, we produced a permutable library using the SS16 motif on Potri.002G039300’s promoter. The library contained heptamerized 5-base-long fragments (referred to as SS16-1 through SS16-7) (Figure 6a–c). Transfection frequency ranged from 52.3%–58.8% among the synthetic promoter constructs (Figure 6d and Figure S4b). Three synthetic promoters, SS16-1, SS16-2 and SS16-3, were highly induced under salt treatments, whereas SS16-5 displayed lower relative induction (Figure 6d,e). SS16-4, SS16-6 and SS16-7 had a moderate level of salt induction (Figure 6d,e). As above, GFP intensity appeared to positively correspond to GFP protoplast frequency count (Figure 6f). Synthetic promoters SS16-1, SS16-2 and SS16-3 were selected for further experimentation in other plant expression systems, which included agroinfiltrated Nicotiana benthamiana leaves and stably transformed Arabidopsis.

In summary, 12 of the 15 synthetic promoters we designed were stress-inducible in transfected poplar leaf protoplasts (>50% of GFP/RFP-positive protoplast count), 9 of which were deemed to have relatively high induction: six SD- and three SS-based synthetic promoters.

Native and synthetic promoter inducibility under osmotic stress in transient agroinfiltrated Nicotiana benthamiana leaves

From the protoplast screens, six SD synthetic promoters and three SS synthetic promoters were selected for leaf agroinfiltration assays (Figures 5 and 6). In addition, one native water-deficit-inducible promoter of Potri.017G015700 and one native salt-inducible promoter of Potri.002G039300 (Figure 4d) were included in agroinfiltration assays to compare the induction driven by native and synthetic promoters. Native and synthetic promoter constructs shared the same vector architecture
Preliminary research (data not shown) was conducted to determine appropriate water-deficit and salt stress treatments for *N. benthamiana* experiments. Treatments included 200 mM NaCl, 100 mM mannitol, and watering cessation for testing promoter activities (Figure 7b). These treatments were not lethal to *N. benthamiana* plants although leaf wilting was observed in water-deficit conditions (Figure 7c). *N. benthamiana* leaves were undamaged in salt or mannitol treatment.

The 1 kb native water stress-inducible (*Potri.017G015700*) promoter drove higher GFP expression than mock controls in the mannitol and water-deficit treatments, respectively (Figure 7d), but was not induced in the salt treatment. Several synthetic promoters were highly induced under mannitol treatment, as measured by the increase in GFP fluorescence: SD18-1, SD18-3, SD9-2 and SD9-3 (Figure 7d). After 4 days from the last irrigation, we observed increased GFP fluorescence in leaves transformed with the following synthetic promoters: SD18-1, SD18-3, SD13-1, SD9-2 and SD9-3 (Figure 7d). Synthetic promoters SD18-1, SD18-3, SD13-1 and SD 9-2 all controlled significantly higher induction than the native *Potri.017G015700* promoter in the water-deficit stress treatments (Figure 7d).

There was a variable response of the synthetic promoters designed to be salt stress-inducible treatments. All three synthetic SS motif-containing promoters (SS16-1, SS16-2 and SS16-3) were induced in the salt treatments (Figure 7d). SS16-2 and SS16-3 were significantly induced by salt and mannitol (Figure 7d), while the native poplar promoter *Potri.002G039300* was not induced in any treatments in the agroinfiltration experiment.
Stably transformed Arabidopsis as a synthetic promoter chassis

Five synthetic promoter constructs (SD18-1, SD9-2, SS16-1, SS16-2 and SS16-3) were selected for stable transformation in Arabidopsis on the basis of the agroinfiltration experiments. T2 transgenic lines appeared to have no morphological differences in stress treatments (100 mM NaCl in tissue culture media over 10-d water cessation) (Figure 8a). Each of the SS synthetic promoters was significantly induced under salt treatment with up to 279.8 and 345.0% increase in GFP gene expression driven by SS16-1 and SS16-3 promoters, whereas GFP was not induced in SD18-1 and SD9-2.
Since synthetic promoters SD18-1 and SD9-2 responded to water-deficit stress treatments in transiently transformed N. benthamiana leaves, transgenic plants expressing GFP under the control of SD18-1 and SD9-2 were tested under water-deficit stress. Watering of 4-week-old Arabidopsis plants grown in soil was stopped for 10 days, until the rosette leaves displayed signs of wilting (Figure 8a). GFP fluorescence increased in both SD18-1 and SD9-2 plants relative to controls, which was commensurate with increases in GFP transcript abundance in the same plants (Figure 8c, d). Water-use efficiency (WUE), which was measured to validate the water-deficit condition, indeed increased after 10-d water cessation (Figure 8e).

(Figure 8b). Since synthetic promoters SD18-1 and SD9-2 responded to water-deficit stress treatments in transiently transformed N. benthamiana leaves, transgenic plants expressing GFP under the control of SD18-1 and SD9-2 were tested under water-deficit stress. Watering of 4-week-old Arabidopsis plants grown in soil was stopped for 10 days, until the rosette leaves displayed signs of wilting (Figure 8a). GFP fluorescence increased in both SD18-1 and SD9-2 plants relative to controls, which was commensurate with increases in GFP transcript abundance in the same plants (Figure 8c, d). Water-use efficiency (WUE), which was measured to validate the water-deficit condition, indeed increased after 10-d water cessation (Figure 8e).

**Discussion**

Synthetic biology will certainly revolutionize agriculture (Wurtzel et al., 2019). One of its important applications is in advanced biotechnology to improve crops and cropping systems (Goolod et al., 2018). The most mature technologies in plant synthetic biology include gene editing, and synthetic promoters and transcription factors (Liu and Stewart, 2015; Liu and Stewart, 2016; Liu et al., 2013). Synthetic promoters have been employed to widely diversify the repertoire of tools for tuning gene expression (Hernandez-Garcia and Finer, 2014; Kassaw et al., 2018). One of the significant challenges to design synthetic

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Figure 7 Transient expression analysis by leaf agroinfiltration of SD and SS synthetic promoter constructs in Nicotiana benthamiana leaves. (a) Binary vectors contained native promoters fused to GFP. Native promoters were represented by ca. 1 kb of the native promoter region of the water stress-inducible Potri.017G015700 or the salt stress-inducible Potri.002G039300. Vectors also contained a 35S::RFP reporter cassette that served as an internal positive control for constitutive expression to use for normalization of GFP fluorescence. (b) The experimental setup of stress treatments and fluorescence measurements on agroinfiltrated leaves. (c) A representation of plants from the agroinfiltration experiment. Mock and 100 mM mannitol or 200 mM NaCl-treated plants were not damaged by 3-day watering while halting watering resulted in wilt after 4 days. (d) GFP fluorescence in agroinfiltrated leaf under mock treatment, 200 mM NaCl, 100 mM mannitol and 4 days water cessation. GFP fluorescence and RFP fluorescence were measured at emission wavelength of 502 nm and 574 nm under a fixed excitation wavelength of 482 nm and 553 nm for GFP and RFP, respectively. The mock treatment (control) consisted of watering every 2 days. Bars represent count per second (CPS) value of normalized GFP fluorescence under stress versus and mock control. Values are the mean ± SD of three independent transformations (n = 3); asterisks denote significant differences via t-tests: * P < 0.05, ** P < 0.01.
The identification of functional CREs (Roccaro et al., 2013). One end goal is to understand the regulation of plant gene expression, especially transcription regulation, which will enable parts (e.g. promoters and transcription factors) to be manufactured and installed in plant genomes to render high predictive functionality.

The combination of high-throughput gene expression profiles and bioinformatic tools is a powerful approach to design synthetic promoters (Koschmann et al., 2012). Synthetic promoter design for predictable transcript inducibility largely rests on the interaction of induction conditions and critical CREs in native promoters. One route to discovery of CREs is via -omics data (e.g. Liu et al., 2014), which is now quite amenable in poplar. Poplar has a reference genome and deep transcriptomic data for abiotic stress traits. Indeed, co-expressed gene data from microarray data have been used to construct inducible synthetic promoters (Koschmann et al., 2012; Liu et al., 2014), but only recently has sufficient RNA-seq transcriptomic data been available for poplar, which we used exclusively in our study (Figures 2 and 3 and Figure S1).

Figure 8  Stress induction analysis of SD and SS synthetic promoters in stable transgenic Arabidopsis. Stable transgenic Arabidopsis plants were generated via the floral dip method with binary vectors harbouring SD and SS synthetic promoters used in both protoplast transformation and agroinfiltration. (a) T2 SD18-1 transgenic Arabidopsis plants grown on MS plates supplemented with 100 mM NaCl for 3 days, and in pots after watering was stopped for 10 days. No lethal effects were observed after stress treatment (bar = 1 cm). (b) GFP gene expression in transgenic Arabidopsis harbouring SD18-1, SD9-2, SS16-1, SS16-2 and SS16-3 synthetic promoter constructs. Three T2 Arabidopsis seedlings (10-day-old) for each independent transgenic line were grown on each of three MS plates with 100 mM NaCl for 3 days. GFP gene expression in mock or treated samples by transgenic construct and event was calculated relative to the AtActin2 internal reference gene. The numerals in red font above each statistically significant bar denote reporter gene induction as a percentage over the control. (c) Relative GFP gene expression under the control of either SD18-1 or SD9-2 synthetic promoters in three independent T2 transgenic Arabidopsis lines under a 10-day water stress (no watering). Transgenic control plants were watered every 2 days in the mock treatment (control). (d) GFP fluorescence in transgenic Arabidopsis harbouring either SD18-1 or SD9-2 promoter under water-deficit stress. GFP fluorescence and RFP fluorescence were measured as described in Figure 7 at 10 days after water cessation, at which time GFP fluorescence between water stress and mock treatments was calculated. GFP values in mock and treated sample were used for GFP normalization. (e) WUE of SD18-1 and SD9-2 transgenic Arabidopsis 10 days after watering cessation. WUE was determined by calculating the ratio of carbon uptake (a) and the rate of gas exchange transpiration (e) measured as described in Methods section. In panels b, c and d, the bars represent means ± SD of 9 plants grown among three plates (n = 9). Each transgenic construct was represented by three independent transgenic T2 lines. qRT-PCR data for each plant (single leaf) were based on three technical replicates. In panel e, one representative plant from each independent T2 line was measured to assess water stress (n = 3) for each construct. Statistical significance is shown via t-tests between treated water cessation and mock (*P < 0.05, **P < 0.01).
plant species. The subsequent validation, through agroinfiltration assays on leaves of *N. benthamiana* and stable transgenic *Arabidopsis*, demonstrated orthologous responsiveness across eudicot species that are not closely related (Figures 5–8).

A persistent challenge in synthetic promoter design has been false-positive hits in de novo DNA motif discovery algorithms (Hu et al., 2005; Lihu and Holban, 2015; Tompa et al., 2005), and early studies achieved up to only 13% in sensitivity and 35% in precision (Hu et al., 2005). Although high-quality sequence data are increasingly becoming available, motif discovery, synthetic promoter design and testing are still a hindrance to increase promoter precision (Lihu and Holban, 2015). One attempt to improve motif discovery has been explored through ensemble approaches, which combine different de novo motif-finding tools, offering improved prediction over any single algorithm (Harbison et al., 2004; Hu et al., 2006; Liu et al., 2014; MacIsaac and Fraenkel, 2006). In the present study, we applied an ensemble approach to coordinate three motif-finding tools coupled with high-quality genome sequencing and transcriptomic data; 15 synthetic promoter candidates were built. Of these, 12 synthetic promoters appeared to be osmotic stress-inducible in poplar protoplasts, 9 of which were confirmed to be inducible in agroinfiltration studies (Figure 7). We conclude that improved -omics data enabled more efficient and reliable CRE discovery, which led to better inducible synthetic promoter design. Nonetheless, we speculate that CRE selection is suboptimal by the current protocol that uses position weight matrix alignment of long DNA sequence based on co-expressed gene clustering. Thus, improved bioinformatics algorithms may be needed to enable more predictive CRE selection. In addition, high-throughput ChIP-seq experiments using known stress-responsive transcription factors may also be coupled with bioinformatics approaches to identify CREs.

Developing versatile synthetic promoters by constructing repeats of an appropriate short-length motif(s) appears to be a powerful strategy with regard to abiotic stress induction. SD18-1, SD18-3, SD9-2 and SD9-3 were highly inducible under mannitold and water-deficit stress treatments, whereas these promoters were weakly inducible to salt stress. On the other hand, the GFP expression driven by SS16-1, SS16-2 and SS16-3, under salt stress was higher than the expression induced by mannitol and water-deficit treatments. Our results were consistent with two previous studies showing a common response of synthetic promoters, consisting of DRE/C-repeats (GCGCAG), to dehydration, high salinity, and cold stress in transgenic *Arabidopsis* (Hou et al., 2012; Kim et al., 2002). Therefore, repeated short-length motifs in synthetic promoters appear to be a good strategy to endow promoter induction to abiotic stress. In contrast, native promoters tested in the present study were specifically induced either by water-deficit or high salinity (Figure 7d), suggesting that different CREs may need to be synchronized with the designed synthetic promoters in order to acquire individual stress specificity. Furthermore, the present study was limited in scope. We focused on synthetic promoter activities in leaves only, while acknowledging the potential relevance in other tissues. Future research should combine research on additional tissues with synthetic promoters comprising tissue-specific CREs for enhancing induced gene expression in space and time. Orthogonal regulatory systems expressing synthetic transcription factors to regulate synthetic promoters are a good strategy to further control gene expression in plants. A new plant orthogonal regulatory system was recently reported that leveraged a yeast transcriptional regulator to induce synthetic activators or repressors (Belcher et al., 2020). Such a system or a new synthetic Neurospora-based ‘Q-system’ (Persad et al., 2020) could be coupled with designed synthetic promoters to tune signal specificity and sensitivity in plants.

Synthetic promoter strength and specificity may be affected by combining multiple CRE sequences and altering their copy number and spacing (Liu and Stewart, 2016). To date, the basic gene construct design has largely included synthetic sequences upstream of a core promoter sequence such as the 46 35S promoter. Recently, a minimal synthetic promoter was designed for constitutive response by various arrangement of CREs at different positions to TATA box area (Cai et al., 2020). In our study, no downstream gene expression was observed in synthetic promoters including three copies of single CREs (Figure S3). However, 7–8 copies of the same short CRE induced GFP induction in our study. Therefore, optimizing copy number of CRE may be a pivotal factor to design useful synthetic promoters.

The present study used repeats of short motif without spaces between each motif. While our synthetic promoters were responsive to osmotic stress, we made no attempt to optimize spacing of repeats. Such optimization could be desirable. A correlation between the copy number of cis-elements and productivity of synthetic promoters was observed in various plant species such as *Arabidopsis* (Sahoo et al., 2014), tobacco (Sawant et al., 2005) and rice (Wu et al., 1998), a pattern that was observed in our study. Abscisic acid and salicylic acid responsiveness of two copies of ACGT was dependent on the spacing between the two copies, indicating that spacing is an important factor for synthetic promoter design (Mehrotra and Mehrotra, 2010). In addition to the spacing between multiple copies of motifs, the spacing between CREs and the core promoter is also important (Rushton, 2016); many unknowns remain about optimal spacing and repeats of CREs (Ali and Kim, 2019; Rushton, 2016).

In the past, well-characterized CREs have been used to construct synthetic promoters inducible by abiotic stress. These cis-elements include ABRE, DRE/C-repeat and coupling factors derived from the cis-regulatory region of wheat *HVA21*, *Arabidopsis* COR, *RD29A* and *ERD1* constructed by classical promoter deletion assays (Shen et al., 1996; Simpson et al., 2003; Thomashow, 1999; Yamaguchi-Shinozaki and Shinozaki, 1994). In one recent study, a 6 × ABRE CRE repeat was used to make a synthetic abscisic acid-inducible promoter, which was tested in Arabidopsis (Wu et al., 2018). Although the present synthetic promoters were designed using motifs that are novel, from a prior characterization standpoint, their inducibility was similar to that of synthetic promoters using known promoter motifs, such as ABRE and DRE. Therefore, our present workflow is enabling to discover novel sequences with functional cis-regulations to use in synthetic promoters.

Even though trees are subjected to various abiotic stresses throughout their lifetimes, there has been few studies that focus on abiotic stress-inducible in poplar. Recently, five *Populus trichocarpa* C-repeat binding factor (CBF)/DREB1 family proteins were found in 2263 annotated genes, but that study did not specifically address abiotic stress (Li et al., 2017). The promoter of *Populus euphratica* ascorbate peroxidase 2 (PeAPX2), a mediator of freezing tolerance through scavenging reactive oxygen species, is recognized by a cysteine-2/histidine-2 type zinc finger transcription factor (PeSTZ1) (He et al., 2019). That study points the way to a route for cold-inducible synthetic promoter construction. Of interest is that our synthetic promoters SD18-1, SD18-3 and...
SD 9-2 included auxin responsiveness (CATATG), ICE1-binding (CANNNTG) sequences, and the calmodulin-binding sequence (ACGCCGA), respectively. Therefore, other types of stress such as cold and freezing can be sensed by reconstructing expanded sequences into the present synthetic promoter construct architecture.

In our study, these synthetic promoters had relatively high induction under osmotic stress (Table 1, Figures 5–8). The molecular/physiological function of SD and SS motifs as abiotic stress-responsive CREs in poplar will be further characterized in upcoming field experiments.

**Experimental procedures**

**Poplar plants**

Female *P. tremula* × *P. alba* interspecific hybrid (717-1B4) aspen clones were kindly donated by Dr. Steven Strauss (Oregon State University). Plants were grown on ProMix BK25 (Premier Tech, Quakertown, PA) in 4-L pots in a greenhouse. Plants were watered once every 2 days. Poplar micropropagation via apical shoots from pot-grown 2-month-old plants was conducted whereby apical shoots were sterilized by shaking them in 50% commercial bleach for 5 min followed by washing in 70% ethanol for another 5 min. Subsequently, cut shoots were washed five times with autoclaved water. Sterilized shoots were grown on rooting media (1 × MS salt with Gamborg’s vitamin mixture (Caisson Labs, Smithfield, UT), 20 mM MES (Sigma, St. Louis, MO), 3% sucrose (Sigma), 0.5% activated charcoal (Sigma), 0.15% Gelrite (Sigma) and adjusted pH to 5.7) in Magenta GA7 boxes (Bioworld, Dublin, OH). Regenerated plants were propagated every 2 months.

**Nicotiana and Arabidopsis plants**

*Nicotiana benthamiana* and *Arabidopsis thaliana* (Col) were grown on ProMix BK25 soil in 1-L pots after stratification at 4°C for 4 days in the sterile water without light. Plants were watered three times per week depending on conditions. All plants were grown under 16 h/8 h for light/dark cycles at constant 22°C with 150 μmol/m²s light intensity.

**Bioinformatic analysis for selection of stress-responsive motif**

All promoter sequences for bioinformatic analysis were profiled from BioMart (Smedley et al., 2015) integrated in Phytozome v12.1.6 (www.phytozome.org). Two kilobase upstream sequences flanking the ATG start codon were collected from *P. trichocarpa* v 3.0 genome annotation (Tuskan et al., 2006). We subjected these promoter sequences to command line application of MEME Suite (v5.0.5), MotifSampler (v3.2.2) and Weeder (v 2.0). DNA motifs up to 20 bases in length were predicted to design reliable core sequences for downstream experiments. To find cis-elements over DNA motifs, we submitted them to the database of plant cis-acting regulatory DNA elements (PLACE; https://www.dna.affrc.go.jp/PLACE/) with default analysis parameters.

**Construction of fusion plasmids with fluorescent reporter genes**

To screen inducible native or synthetic promoters against osmotic stress, we generated two different backbone genes harbouring GFP constructs with or without constitutive RFP expression gene constructs (Figures 4c, 5a, 6a and 7a). For native promoters used in protoplast transformation (Figure 4c), the backbone gene construct including TMV 5′ UTR (Ω) leader and TurboGFP was constructed with Gateway cloning sites via the Golden Gate modular cloning technique using the MoClo system (Engler et al., 2014; Figure 4c).

Through a BLAST search with the coding sequence of *P. trichocarpa* annotation of osmotic stress-responsive genes to INRA 717-1B4 genome assembly (Mader et al., 2016: http://urgi.versailles.inra.fr), about 1-kb upstream promoter sequences from ATG were collected from the aspen genome. The native promoter was amplified from 717-1B4 genomic DNA extracted following Kang’s method (Kang and Yang, 2004) using Phusion DNA polymerase with gene-specific primers for Gateway cloning (Thermo Fisher Scientific, Waltham, MA, Table S7). The PCR was performed as described below; 1 cycle of 95°C for 2 min, 30 cycles repeating 95°C for 30 s, 57°C for 30 s and 72°C for 1 min. The insert was destined into the backbone construct by LR clonase (Invitrogen, Carlsbad, CA).

In the agroinfiltration test, native promoters fused with TurboGFP as described above were linked with TurboRFP driven by 2 × CaMV 35S promoter (Figure 7a). For synthetic promoter construct generation, a binary backbone plasmid constructing two different gene constructs was generated by the Golden Gate cloning technique (Figures 5a, 6a and 7a): i) a gene construct for synthetic promoter cloning site including BsaI restriction site, CaMV 35S minimal promoter (−46 bp) and TMV 5′ UTR (Ω) leader, fused with TurboGFP, and ii) a gene construct including 2 × CaMV 35S short promoter driving TurboRFP expression (Engler et al., 2014).

Synthetic promoter fragments were generated by reannealing two complementary single-stranded oligonucleotides consisting of heptamer repeats of a DNA motif plus a 5′ overhangs of Bsal digestion sites on the backbone plasmid (Integrated DNA Technologies, Coralville, IA, Table S6). A pair of complementary strands was combined together and incubated at 95°C for 2 min followed by 65°C for 5 min, and then was cooled down gradually to 4°C using a thermal cycler. The annealed double strands were cloned into a Bsal-digested final destination vector. All gene construct DNA sequences and ligation conditions were verified by Sanger sequencing analysis (UT Genomics Core, Knoxville, TN).

**Poplar mesophyll protoplast transformation assay**

Poplar mesophyll protoplasts were isolated from leaves of 1-month-old 717-1B4 in vitro culture as described above. Cellulose enzyme treatment and recovery of extracted protoplasts were performed as described in Guo’s protocol (Guo et al., 2012). Ten micrograms of plasmid was transfected to 2 × 10⁴ protoplasts. Since WI solution contains 0.5 M mannitol, osmotic condition was induced simultaneously with incubation for 2 days at room temperature. To find optimal NaCl concentration, we tested protoplast cell viability in 50 mM, 100 mM and 150 mM NaCl complemented with WI solution. Based on this result, the transformed protoplasts were incubated for 2 days at room temperature in 100 mM NaCl. The GFP and RFP fluorescent images were taken by EVOS M7000 (Thermo Fisher Scientific) and analysed by ImageJ software (Schneider et al., 2012). The fluorescent ratio was determined by counting fluorescent-detected protoplast out of viable protoplasts in a bright field image. The relative fluorescence ratio was determined by calculating the number of GFP-expressing protoplasts over the number of RFP-expressing protoplasts.

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Transient agroinfiltration assay in *N. benthamiana* leaves

The same plasmids for protoplast assay were transfected into *Agrobacterium* strain EHA105 by the liquid nitrogen transformation method (Höfgen and Willmitzer, 1998). For infiltration, *Agrobacterium* was cultured in LB medium containing 50 mg/L rifampicin and 50 mg/L kanamycin overnight shaking at 28°C. Resuspended cell culture was infiltrated into the abaxial side of the third and fourth leaves by pushing the syringe as previously described (Ma et al., 2012).

The GFP fluorescence and RFP fluorescence were measured by Fluorolog®-3 (HORIBA, Kyoto, Japan) on the abaxial side of the agroinfiltrated leaf at the third day and the fourth day after salt or mannitol treatment or water cessation, respectively (Figure 7b). The fluorescence was measured at the emission wavelength of 502 and 574 nm under fixed excitation at 482 and 502 nm for TurboGFP and TurboRFP, respectively. GFP fluorescence increase was determined by calculating increase ratio of GFP fluorescence in stress-treated leaves over that in mock. GFP intensity was normalized by RFP fluorescence ratio between stress-treated and mock-treated cells.

Stable transgenic Arabidopsis generation

To generate stable transgenic *Arabidopsis*, we used the floral dip technique on 1-month-old *Arabidopsis* plants (Clough and Bent, 1998). The same EHA105 *Agrobacterium* lines described above were transformed into *Arabidopsis*. Transgenic *Arabidopsis* plants were screened on 1/2 × MS plates including Gamborg’s vitamins (Caisson), 1% sucrose and 0.8% Phytoagar (Sigma) complementing 50 μg/mL kanamycin. For genotyping, genomic DNA was extracted from leaves of 3-week-old T2 generation *Arabidopsis* seedlings among multiple independent transgenic lines per construct (Kang and Yang, 2004). Two-hundred and fifty nanograms of genomic DNA was subjected to DreamTaq Green PCR Master Mix (Thermo Scientific) including 200 nM of specific forward and reverse primers of GFP (Table S7). PCR was performed as 1 cycle of 95°C for 5 min, 30 cycles repeating a chain of 95°C for 30 sec, 60°C for 30 s, and 72°C for 30 s, and final extension at 72°C for 10 min. We selected three independent transgenic lines from each synthetic promoter (Figure S5).

Stress treatment for the osmotic inducibility experiment

To optimize osmotic stress conditions in present studies, 717-1B4, *N. benthamiana* and *Arabidopsis* were first exposed various concentrations of mannitol or NaCl, or halting watering up to 10 days. We selected the chemical concentration or duration of water stress to detect enough reporter gene expression without severe damage in stress-treated plants.

To verify endogenous poplar gene expression under water stress and high salinity (Figure 4b), we transplanted 717-1B4 after 2 months of regeneration in Magenta GA7 boxes to 4-L pots of potting mix. These plants were grown another 1 month with watering every 2 days. For drought condition, we ceased watering until leaves began to wilt (around 10 days after water cessation). The salt treatment consisted of watering plants with 200 mM NaCl solution every 3 days. Control poplar was watered with tap water.

In the agroinfiltration assay in *N. benthamiana*, the infiltrated plants were stressed by watering either with 200 mM NaCl or with 100 mM mannitol (Figure 7a). We ceased watering for 4 days after agroinfiltration until the leaf began to wilt.

To quantify GFP gene expression in transgenic *Arabidopsis* under osmotic stress (Figure 8), 10-day-old T2 seedlings from three different lines per each synthetic promoter were screened first on 1/2 × MS plates with 50 μg/ml kanamycin. The screened *Arabidopsis* plants were transferred to 1/2 × MS plates with 100 mM NaCl for 3 days. For water cessation treatment, screened T2 plants were transplanted and grown in pots watered every 2 days. After 4 weeks, watering was stopped for 10 days.

To determine water-deficit stress on leaf surface, we measured CO2 uptake (A) and transpiration (E) by LI-6800 portable photosynthesis system (LI-COR Biosciences, Lincoln, NE). WUE (A/E) was used as a marker to assess drought stress (Yang et al., 2019).

Mock plants transformed with the same plasmids by the same process were watered every 2 days with deionized water.

qRT-PCR

The qRT-PCR process is summarized in Figure S6. Total RNA was extracted via Plant RNA Extraction reagent following the manufacturer’s manual (Invitrogen, Carlsbad, CA). One microgram of purified total RNA was used to perform the reverse transcription reaction with RevertAid First Strand cDNA Synthesis Kit (Thermo Fisher Scientific, Waltham, MA). The five-time diluted single-strand cDNA was subjected into 20 μl of total volume with 1 × PowerUp SYBR Green Master Mix including gene-specific primers as listed in Table S7 (Applied Biosystems, Foster City, CA). qPCR profile was 50°C for 2 min, 95°C for 10 min and 40 cycles of 95°C for 15 s and 60°C for 30 s. The relative gene expression change was determined by the 2^{-ΔΔCt} equation (Livak and Schmittgen, 2001).

Statistical analysis

Technical and biological repeat numbers are described in individual figure legends. Statistical analyses of all measurements were performed by Student’s t-test integrated in R (R core team, 2014).

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Author contributions

YY and JHL equally contributed to designing and performing the experiments, analysing the data and writing the manuscript. MRP and YS assisted with the experiments and plant care. WL and SCL consulted to find DNA motif and construct working plasmid. AA, EB and CNS conceived of the study and its design and coordination, and assisted with interpretation of results and revisions to the manuscript. All authors read and approved the final manuscript.

Conflicts of interest

The authors declare no conflicts of interest.

References

Ali, S. and Kim, W.-C. (2019) A fruitful decade of using synthetic promoters in the improvement of transgenic plants. Front Plant Sci. 10, 1433.
Kang, T.J. and Yang, M.S. (2004) Rapid and reliable extraction of genomic DNA from various wild-type and transgenic plants. BMC Biotechnol. 4, 20.

Kassaw, T.K., Donayre-Torres, A.J., Antunes, M.S., Morey, K.J. and Medford, J.I. (2018) Engineering synthetic regulatory circuits in plants. Plant Sci 273, 13–22.

Kim, H.J., Kim, Y.K., Park, I.Y. and Kim, J. (2002) Light signalling mediated by phytochrome plays an important role in cold-induced gene expression through the C-repeat/dehydration responsive element (C/DRE) in Arabidopsis thaliana. Plant J. 29, 693–704.

Koschmann, J., Machers, F., Becker, M., Niemeyer, J., Schulze, J., Bulow, L., Stahl, D.J. and et al. (2012) Integration of bioinformatics and synthetic promoters leads to the discovery of novel elicitor-responsive cis-regulatory sequences in Arabidopsis. Plant Physiol. 160, 178–191.

Lee, J.H. and Pijut, P.M. (2018) Optimization of Agrobacterium-mediated genetic transformation of Fraxinus nigra and development of black ash for possible emerald ash borer resistance. Plant Cell Tiss Org. 134, 217–229.

Lehmann, M., Kanofsky, K., Hanko, E.K., Ahrendt, S., Wehrs, M., Machens, F. and Hehl, R. (2016b) Functional dissection of a strong and specific microbe-associated molecular pattern-responsive synthetic promoter. Plant Biotechnol. J. 14, 61–71.

Li, J., Sima, W., Qiang, Y., Wang, T., Zief, K., Luo, Z., Liu, L. et al. (2012) Tomato SIDREB gene restricts leaf expansion and internode elongation by downregulating key genes for gibberellin biosynthesis. J. Exp. Bot. 63, 6407–6420.

Li, Y., Song, Y., Xu, B., Xie, J., Zhang, D. and Cooke, J. (2017) Poplar CBF1 functions specifically in an integrated cold regulatory network. Tree Physiol. 37, 98–115.

Li, Xiong, Yang, Yunqiang, Sun, Xudong, Lin, Huaming, Chen, Jinhui, Ren, Jian, Hu, Xiangyang and et al. (2014) Comparative physiological and proteomic analyses of poplar (Populus yunnanensis) plantlets exposed to high temperature and drought. PLoS One 9, e4107605.

Lihu, A. and Holban, S. (2015) A review of ensemble methods for de novo motif discovery in ChIP-Seq data. Brief Bioinform. 16, 964–973.

Liu, W., Mazarei, M., Peng, Y., Fethe, M.H., Rudis, M.R., Lin, J., Millwood, R.J. et al. (2014) Computational discovery of soybean promoter cis-regulatory elements for the construction of soybean cyst nematode-inducible synthetic promoters. Plant Biotechnol. J. 12, 1015–1026.

Liu, W. and Stewart, C.N. Jr. (2015) Plant synthetic biology. Trends Plant Sci. 20, 309–317.

Liu, W. and Stewart, C.N. Jr. (2016) Plant synthetic promoters and transcription factors. Curr. Opin. Biotechnol. 37, 36–44.

Liu, W., Yuan, J.S. and Stewart, C.N. Jr. (2013) Advanced genetic tools for plant biotechnology. Nat. Rev. Genet. 14, 781–793.

Livak, K.J. and Schmittgen, T.D. (2001) Analysis of relative gene expression data using real-time quantitative PCR and the 2(-delta delta C(T)) method. Methods. 25, 402–408.

Ma, L., Lukaisik, E., Gawehns, F. and Takken, F.L.W. (2012) The use of agroinfiltration for transient expression of plant resistance and fungal effector proteins in Nicotiana benthamiana leaves. In Plant Fungal Pathogens Methods in Molecular Biology (Methods and Protocols), 835. (Bolton, M. and Thomina, B., eds), pp. 74–76. New York: Humana Press.

Maclsaac, K.D. and Frankeel, E. (2006) Practical strategies for discovering regulatory DNA sequence motifs. PLoS Comput. Biol. 2, e36.

Mader, M., Le Paslier, M.C., Bounon, R., Berard, A., Favre Rampart, P., Fladung, M., Leple, J.C. and et al. (2016) Whole-genome draft assembly of Populus tremula x Populus alba clone INRA 717–184. Silvae Genetica. 65, 79–79.

Mehrotra, R., Gupta, G., Sethi, R., Bhalothia, P., Kumar, N. and Mehrotra, S. (2011) Designer promoter: an artwork of cis-engineering. Plant Mol. Biol. 75, 527–536.

Mehrotra, R. and Mehrotra, S. (2010) Promoter activation by AGCT in response to salicylic and abscisic acids is differentially regulated by the spacing between two copies of the motif. J. Plant Physiol. 167, 1214–1218.

Ohta, Masanu, Sato, Aiko, Renhu, Na, Yamamoto, Tsuyoshi, Oka, Nodoka, Zhu, Jian-Kang, Tada, Yasuomi et al. (2018) MYC-type transcription factors, MYC67 and MYC70, interact with ICE1 and negatively regulate cold tolerance in Arabidopsis. Sci. Rep. 8, 11622.
Okumura, A., Shimada, A., Yamazaki, S., Horino, T., Iwata, Y., Koizumi, N., Nishihara, M. and et al. (2016) CaMV-35S promoter sequence-specific DNA methylation in lettuce. Plant Cell Rep. 35, 43–51.

Pavesi, G., Meregghetti, P., Mauri, G. and Pesole, G. (2004) Weeder Web: discovery of transcription factor binding sites in a set of sequences from co-regulated genes. Nucleic Acids Res. 32, W199–203.

Peramuna, A., Bae, H., Rasmussen, E.K., Dueholm, B., Waibel, T., Critchley, J.H., Brzézek, K. et al. (2018) Evaluation of synthetic promoters in Physcomitrella patens. Biochem. Biophys. Res. Commun. 500, 418–422.

Persad, R., Reuter, D.N., Dice, L.T., Nguyen, M.-A., Rigoulot, S.B., Layton, J.S., Schmidt, M.I. et al. (2020) The Q-system as a synthetic transcriptional regulator in plants. Front Plant Sci. 11, 245.

R Core Team (2014) R: A Language and Environment for Statistical Computing. Vienna, Austria: R Foundation for Statistical Computing. http://www.R-project.org/

Roberts, M.L. (2011) The use of functional genomics in synthetic promoter design. In: Compt. Biol. Appl. Bioinform. Intech (Lopes, H.S. ed), pp 375–396.Available from: http://www.intechopen.com/books/computational-biology-and-applied-bioinformatics/the-use-of-functional-genomics-in-synthetic-promoter-design

Roccavo, M., Ahmadinejad, N., Colby, T. and Somsisch, I.E. (2013) Identification of functional cis-regulatory elements by sequential enrichment from a randomized synthetic DNA library. BMC Plant Biol. 13, 164.

Rushton, P.J. (2016) What have we learned about synthetic promoter construction? In Plant Synthetic Promoters Methods in Molecular Biology, 1482. (Hehl, R., ed), pp. 1–13. New York: Humana Press.

Sahoo, D.K., Sarkar, S., Raha, S., Maiti, I.B. and Dey, N. (2014) Comparative analysis of synthetic DNA promoters for high-level gene expression in plants. Planta, 240, 855–875.

Saint Pierre, C., Crossa, J.L., Bonnett, D., Yamaguchi-Shinozaki, K. and Reynolds, M.P. (2012) Phenotyping transgenic wheat for drought resistance. J. Exp. Bot. 63, 1799–1808.

Sannigrahi, P., Ragauskas, A.J. and Tuskan, G.A. (2009) Poplar as a feedstock for biofuels: A review of compositional characteristics. Biofuels Biopro. Biofer. 4, 209–226.

Sawant, Samir V., Kiran, Kanti, Mehrrotta, Rajesh, Chaturvedi, Chandra Prakash, Ansan, Sauriya A., Singh, Pratibha, Lodhi, Niraj and et al. (2005) A variety of synergistic and antagonistic interactions mediated by cis-acting DNA motifs regulate gene expression in plant cells and modulate stability of the transcription complex formed on a basal promoter. J. Exp. Bot. 56, 2345–2353.

Schaumberg, K.A., Antunes, M.S., Kassaw, T.K., Xu, W., Zalewski, C.S., Medford, J.I. and Prasad, A. (2016) Quantitative characterization of genetic parts and circuits for plant synthetic biology. Nat. Methods 13, 94–100.

Schneider, C.A., Rasband, W.S. and Eliceiri, K.W. (2012) NIH Image to ImageJ: 25 years of image analysis. Nat. Methods 9, 671–675.

Sharon, E., Kalma, Y., Sharp, A., Raveh-Sadka, T., Levo, M., Zeevi, D., Keren, L. et al. (2012) Inferring gene regulatory logic from high-throughput measurements of thousands of systematically designed promoters. Nat. Biotechnol. 30, 521–530.

Sheen, J. (2001) Signal transduction in maize and Arabidopsis mesophyll protoplasts. Plant Signal 127, 1466–1475.

Shen, Q., Zhang, P. and Ho, T.H. (1996) Modular nature of abscisic acid (ABA) response complexes: composite promoter units that are necessary and sufficient for ABA induction of gene expression in barley. Plant Cell 8, 1107–1119.

Simpson, S., Nakashima, K., Narusaka, Y., Seki, M., Shinozaki, K. and Yamaguchi-Shinozaki, K. (2003) Two different novel cis-acting elements of erd1, a cPa4 homologous Arabidopsis gene function in induction by dehydration stress and dark-induced senescence. Plant J. 33, 259–270.

Smedley, Damian, Haidar, Syed, Dunific, Steffen, Pandini, Luca, Provero, Paolo, Allen, James, Arnaiz, Olivier et al. (2015) The BioMart community portal: an innovative alternative to large, centralized data repositories. Nucleic Acids Res. 43, W589–598.

Tang, S., Liang, H., Yan, D., Zhao, Y., Han, X., Carlson, J.E., Xia, X. and et al. (2013) Populus euphratica: the transcriptomic response to drought stress. Plant Mol. Biol. 83, 539–557.

Thijs, G., Marchal, K., Lescot, M., Rombouts, S., De Moor, B., Rouze, P. and Moreau, Y. (2002) A Gibbs sampling method to detect overrepresented motifs in the upstream regions of coexpressed genes. J. Comput. Biol. 9, 447–464.

Thomashow, M.F. (1999) Plant cold acclimation: freezing tolerance genes and regulatory mechanisms. Annu. Rev. Plant Physiol. Plant Mol. Biol. 50, 571–599.

Tompa, Martin, Li, Nan, Bailey, Timothy L., Church, George M., De Moor, Bart, Eskin, Eleazar, Favorov, Alexander V. et al. (2005) Assessing computational tools for the discovery of transcription factor binding sites. Nat. Biotechnol. 23, 137–144.

Tuskan, G.A., Difazio, S., Jansson, S., Bohlmann, J., Grigoriev, I., Hellsten, U., Putnam, N. et al. (2006) The genome of black cottonwood, Populus trichocarpa (Torr. & Gray). Science, 313, 1596–1604.

Wang, H., Wang, Shao, H. and Tang, X. (2016) Recent advances in utilizing transcription factors to improve plant abiotic stress tolerance by transgenic technology. Front. Plant Sci. 7, 67.

Wei, T., Deng, K., Zhang, Q., Gao, Y., Liu, Y., Yang, M., Zhang, L. et al. (2017) Modulating ADRB1C expression improves drought tolerance in Salvia miltiorrhiza. Front. Plant Sci. 8, 52.

Wu, R., Duan, L., Pruneda-Paz, J.L., Oh, D.H., Pound, M., Kay, S. and Dinneny, J.R. (2018) The 6xABRE synthetic promoter enables the spatiotemporal analysis of ABA-mediated transcriptional regulation. Plant Physiol. 177, 1650–1665.

Wu, C.Y., Suzuki, A., Washida, H. and Takaiwa, F. (1998) The GCN4 motif in a rice gliadin gene is essential for endosperm-specific gene expression and is activated by Opaque-2 in transgenic rice plants. Plant J. 14, 673–683.

Wurtzel, E.T., Vickers, C.E., Hanson, A.D., Millar, A.H., Cooper, M., Voss-Fels, K.P., Nikel, P.I. et al. (2019) Revolutionizing agriculture with synthetic biology. Nat. Plants, 5, 1207–1210.

Yamaguchi-Shinozaki, K. and Shinozaki, K. (1994) A novel cis-acting element in an Arabidopsis gene is involved in responsiveness to drought, low-temperature, or high-salt stress. Plant Physiol. 100, 858–865.

Yamaguchi-Shinozaki, K. and Shinozaki, K. (1994) A novel cis-acting element in an Arabidopsis gene is involved in responsiveness to drought, low-temperature, or high-salt stress. Plant Physiol. 100, 858–865.

Yamasaki, Y. and Randall, S.K. (2016) Functionality of soybean CBF/DREB1 transcription factors. Plant Sci. 246, 80–90.

Yang, H., Shukla, M., Mao, X., Kang, S. and Du, T. (2019) Interactive regimes of reduced irrigation and salt stress depressed tomato water use efficiency at leaf and plant scales by affecting leaf physiology and stem sap flow. Front. Plant Sci. 10, 160.

Zheng, L., Meng, Y., Ma, J., Zhao, X., Cheng, T., Ji, J., Chang, E. et al. (2015) Transcriptomic analysis reveals importance of ROS and phytohormones in response to short-term salinity stress in Populus tomentosa. Front. Plant Sci. 6, 678.

Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1 The subset of published poplar RNA-seq results of independent experiments used for the basis of promoter discovery in this study.

Figure S2 Cloned regions of four selected native promoters showing domains of interest.

Figure S3 Effect of copy number repeats of core sequences used in synthetic promoter constructs.

Figure S4 The ratio of GFP- and RFP-positive transfected protoplasts in each respective experimental population.

Figure S5 PCR analysis of T2 transgenic Arabidopsis.

Figure S6 Scheme of qRT-PCR process.

Table S1 Selected up-regulated genes from four different RNA-seq datasets: from leaves taken from drought-treated poplar.

Table S2 Gene lists from two different RNA-seq datasets from roots of salt-stressed poplar.

Table S3 The predicted motifs in water-stressed RNA-seq promoter data (5D motifs) by de novo motif-detecting software.
Table S4 The predicted motifs in salt-stressed RNA-seq promoter data (SS motifs) by de novo motif-detecting software.

Table S5 Selected expressed genes from six different RNA-seq datasets and included SD in A domain or SS motif on their promoters.

Table S6 DNA sequence for generating synthetic promoter fragment to construct binary plasmid using the reannealing method.

Table S7 List of qRT-PCR and genomic DNA PCR primers.