A library of synthetic transcription activator-like effector-activated promoters for coordinated orthogonal gene expression in plants

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SUMMARY
A library of synthetic promoters containing the binding site of a single designer transcription activator-like effector (dTALE) was constructed. The promoters contain a constant sequence, consisting of an 18-base long dTALE-binding site and a TATA box, flanked by degenerate sequences of 49 bases downstream and 19 bases upstream. Forty-three of these promoters were sequenced and tested in transient assays in Nicotiana benthamiana using a GUS reporter gene. The strength of expression of the promoters ranged from around 5% to almost 100% of the viral 35S promoter activity. We then demonstrated the utility of these promoters for metabolic engineering by transiently expressing three genes for the production of a plant diterpenoid in N. benthamiana. The simplicity of the promoter structure shows great promise for the development of genetic circuits, with wide potential applications in plant synthetic biology and metabolic engineering.

Keywords: synthetic promoters, transcription activator-like effectors, orthogonality, metabolic engineering, transient assays, Nicotiana benthamiana, terpenoid, synthetic biology, plants, technical advance.

INTRODUCTION
One of the goals of synthetic biology is to establish artificial transcriptional networks for the expression of novel signalling or metabolic pathways. While it has been successfully done in unicellular microorganisms (Brophy and Voigt, 2014), the development of such transcriptional networks in higher eukaryotes, in particular in higher plants, presents additional difficulties due to the lack of sufficient available data, multicellularity and chromatin structure. One prerequisite for the construction of such networks is the availability of a range of transcriptional promoters of varying strengths that can be activated by orthogonal transcription factors (TFs). Since such promoters are not available in nature, they have to be designed, synthesized, assembled in functional transcription units and tested. Several approaches have been adopted for promoter engineering, including random mutagenesis, saturation mutagenesis of poorly conserved regions and the engineering of hybrid promoters (Blazeck and Alper, 2013). Expression levels achieved so far have covered a range over one to three orders of magnitude (Blazeck and Alper, 2013), indicating that such sets of promoters are adequate to finely tune the expression of individual genes for engineering of pathways. An additional desirable property of synthetic promoters is their activation by orthogonal TFs. This is particularly relevant for multicellular organisms such as higher plants, for example if the aim is to express a biosynthesis pathway exclusively in one tissue or under specific conditions such as biotic or abiotic stresses. Most promoter libraries described so far in yeast are based on upstream activating sequences (UAS) of endogenous TFs (Blazeck and Alper, 2013). If the TF on which the design of these synthetic promoters is based is expressed constitutively, or only in specific tissues or conditions, this will considerably restrict their potential use. For constitutive TF, this will prevent activation in specific tissues; conversely, for tissue-specific TFs this will restrict the use of...
the promoters to a specific tissue, preventing their general application for other tissues or conditions. A set of promoters whose expression can be induced by an orthogonal TF would circumvent these problems. In this case, the localization and pattern of expression is determined by the orthogonal TF, which itself can be put under the control of a tissue-specific promoter, for example.

The recently discovered transcription activator-like effectors (TALEs) provide the perfect raw material for developing such a tool (Boch et al., 2009; Boch and Bonas, 2010). The modular structure of their DNA-binding site allows the design of truly orthogonal TFs by selecting a DNA sequence that does not occur in the genome, thus minimizing off-target gene activation. Since their discovery, TALEs have been used both for transcription activation and genome editing, the latter by fusing them to nucleases (de Lange et al., 2014). While tools for the construction of custom-designed TALEs are available (Weber et al., 2011), there are, to the best of our knowledge, no described resources for libraries of synthetic promoters that can be activated by custom-designed TALEs. Such a resource would not only allow the coordinated expression of metabolic pathway genes in specific cell types or under given conditions, but would also provide the building blocks to construct artificial transcriptional networks by introducing positive or negative feedback loops, or transcriptional cascades. To fill this gap, and motivated by our interest in plant metabolic engineering, we undertook the design and construction of libraries of synthetic TALE-activated promoters (STAPs), which we describe here. These were assembled in Golden Gate-compatible vectors to facilitate their downstream use (Engler et al., 2014). These STAPs were first evaluated with reporter genes (GUS and GFP) in transient assays in Nicotiana benthamiana, then, as proof of principle, tested in a metabolic engineering set-up for the production of a plant diterpene.

RESULTS

Design and construction of a library of synthetic TALE-activated promoters

The structure of the STAPs is presented in Figure 1(a). They contain an 18-base-long DNA-binding domain, named EBE002, flanked by a 19-base-long degenerate sequence immediately upstream, and a consensus TATA box downstream. The TATA box is followed by a 43-base-long degenerate sequence and the ATG start codon. This design is based on the observation that the DNA-binding sequence of TALEs from bacterial pathogens that activate genes in planta are typically located close to the transcription start site (less than 100 bp) and that insertion of a DNA-binding sequence at position -55 or -40 was sufficient to confer TALE-mediated inducibility (Kay et al., 2009). The library was cloned using the MoClo system based on Golden Gate cloning (Werner et al., 2012). The scheme for the construction of the STAPs library is presented in Figure 1(b). The library was made in a level -1 vector of the MoClo system, yielding pAGT559-L, and transferred in a level 0 vector, yielding pAGT582-L. Level 0 vectors contain parts (typically promoter, cDNA, terminator) which are brought together in level 1 vectors to assemble functional transcription units. Transferring the library from level -1 to level 0 ensures that only fragments containing the correct overhangs are kept, thus limiting the number of false positives and increasing the efficiency of cloning in subsequent steps. Approximately 50 clones from the level 0 library were picked and checked for the presence of a fragment of the expected size. Forty-three clones were thus selected and sequenced (see Data S1 in Supporting Information). All promoter sequences contained the EBE002 DNA-binding site.

Transient assays of the STAPs with a GUS reporter gene

Each STAP was cloned upstream of the GUS reporter gene and the ocs transcription terminator in a level 1 vector, which can also be used for Agrobacterium tumefaciens T-DNA transfer (Figure 2a). The transcriptional

(a) ggag (n)_{19} tccccgcataaagcaacaattataa (n)_{43} aatg

(b) ACAT BpiI + ligase

GGAG Level -1 TATA pAGT559-L

TTTG BsaI + ligase

GGAG Level 0 AATG pAGT582-L

Figure 1. Design and cloning procedure of the synthetic transcription activator-like effector-activated promoters (STAPs). (a) general sequence of the STAPs. The constant designer transcription activator-like effector (dTALE)-binding site (EBE002) is highlighted in blue and the TATA box in green. The atg triplet at the end of the promoter corresponds to the translation start codon. The degenerate sequences upstream and downstream of the constant region are indicated by (n)_{19} and (n)_{43} respectively. (b) The STAPs cloning procedure. The library was synthesized using two degenerate oligonucleotides overlapping the constant region and cloned into a level -1 Golden Gate vector. Subsequent cloning into a level 0 vector allowed elimination of aberrant products (e.g. dimers) and false positives. The level 0 vectors can then be used for assembly into transcription units.

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activity of these promoters was first evaluated in transient assays in *Nicotiana benthamiana* by measuring GUS activity in protein extracts. The designer TALE (dTALE) is under control of the Act2 promoter (Figure 2a), which provides sufficient expression for the activation of transcription. To quantify the expression, leaves were infiltrated in triplicate with the STAP:GUS constructs, with or without the dTALE and with a 35S:GFP construct. The GFP fluorescence was measured and used as a normalization factor. The results (Figure 2b) show a distribution of expression strengths ranging from less than 1% to over 90% of that of the 35S promoter. For some promoters there is significant residual background expression without the TALE, the useful range of expression in this transient assay is estimated to be between 5 and 90% of that of the 35S promoter. For most STAPs, however, no significant expression was detected in the absence of the TALE, indicating that the activity seen with the TALE does not stem from endogenous activators, establishing the orthogonality of the promoters.

**Transient assays of selected STAPs with GFP**

To confirm these results with another reporter gene, a selection of eight STAPs with different expression strengths (STAPs numbers 1, 2, 3, 5, 14, 17, 42 and 44) were cloned in front of the GFP gene and transiently expressed in *N. benthamiana*. The results (Figure 3) are consistent with those of the GUS assay, indicating that the strength of expression for GUS and GFP seems to be independent of the transcribed gene. No GFP signal could be seen in the absence of the TALE for any of the STAPs tested, confirming the high specificity of the synthetic promoters. The strongest promoters (numbers 1, 3 and 5) show a signal intensity approaching that of the 35S promoter, while the weaker promoters (numbers 17 and 42) still produce a detectable signal.

**Using STAPs for diterpene metabolic engineering in *N. benthamiana***

We next sought to validate the utility of these promoters by using them for metabolic engineering of a plant diterpene. As output for our metabolic engineering assay we used cambratrienol synthase (CBTS2a) from tobacco (Wang and Wagner, 2003; Ennajdaoui et al., 2010). This diterpene synthase produces a mix of two stereoisomers (α and β) of the macrocyclic cembratrienol (CBTol) (Figure 4). In plants, the terpenoid precursors isopentenyl diphosphate (IPP) and dimethylallyl diphosphate (DMAPP) are produced by two distinct pathways, the plastidic 2-C-methyl-d-erythritol-4-phosphate (MEP) pathway and the cytosolic mevalonate (MEV) pathway (Vranova et al., 2012). Diterpenes are synthesized in the plastids and therefore predominantly use IPP and DMAPP from the MEP pathway. We have previously shown that co-expression of 1-deoxyxylulose 5-phosphate synthase (DXS), the first step of the MEP pathway, with a geranylgeranyl diphosphate synthase (GGPPS) leads to more than a doubling of diterpene production in *N. benthamiana* transient assays (Brückner and Tissier, 2013). These assays were carried out with the strong cauliflower mosaic virus 35S promoter controlling the expression of all the genes. The DXS2 (tomato), GGPPS2 (tobacco) and the CBTS2a genes were cloned downstream of some of the strongest STAPs characterized above and transiently expressed together with the dTALE. The results show that both with the 35S promoter and the STAPs, the amount of CBTol produced is significantly higher when all genes (CBTS2a, GGPPS2 and DXS2) are co-expressed than with the other combinations (CBTS2a alone, CBTS2a + DXS2 or CBTS2a + GGPPS2).
(Figure 4c), confirming previous results with the 35S promoter (Brückner and Tissier, 2013). However, the amount of CBTol produced with the the STAPs was around threefold lower than with 35S. One explanation for this could be that gene expression is lower. To check this, we measured transcript levels by quantitative RT-PCR from the same samples used for quantification of CBTol. Surprisingly, the transcript levels were higher with the STAPs than with the 35S promoter (1.9 times for CBTS2a, 11.5 for GGPS2 and 1.5 for DXS2) (Figure 5). Thus, the lower CBTol levels cannot be caused by reduced transcript levels of the genes. Plotting the gene transcript levels versus the amount of CBTol produced indicates a negative correlation, regardless of the promoters used (STAPs or 35S) (Figure S1). Thus, this inverse effect of gene expression on metabolite levels does not seem to be specific to STAPs.

These results show that STAPs can be used for metabolic engineering applications and can deliver expression levels which are comparable to those of the strongest known promoters.

**DISCUSSION**

We report here on the design, construction and testing of a library of synthetic promoters activated by a dTALE TF in plants. Based on a remarkably simple architecture consisting of the dTALE DNA-binding site, a TATA box and degenerate sequences flanking these constant regions, these STAPs exhibit properties that make them ideal tools for the development of complex transcriptional networks or for metabolic engineering applications. First, based on the GUS reporter gene, they exhibit a usable range of expression levels in transient assays that varies from 5 to 90% of that of the 35S promoter, which is one of the strongest plant promoters available. However, it is possible that, due to the high load of T-DNAs that are transferred, transient assays in *N. benthamiana* lead to background transcription activity. Whether this is still the case in stable transgenics remains to be seen. Large variations in expression levels of individual genes within a biosynthesis pathway have been observed in plants. For example, genes encoding the cytochrome P450 monooxygenases of the parthenolide pathway in feverfew (*Tanacetum parthenium*) can be expressed at levels several 100-fold higher than the gene encoding germacrene A synthase, which represents the first committed step of the pathway (Liu *et al.*, 2014). Thus, the range of expression levels delivered by STAPs should prove particularly useful for tuning the expression of individual pathway genes in metabolic engineering projects. This can serve to optimize pathway flux but also to avoid excess production of certain enzymes, which may cause metabolic overload of the host cells. Second, in the absence of the dTALE, no or just background levels of
expression were detected, demonstrating the orthogonal nature of the dTALE/STAP system.

We could also validate the utility of STAPs for metabolic engineering in a transient assay. We could confirm that co-expression of GGPPS2 and DXS2 with the diterpene synthase CBTS2a leads to increased levels of CBToI (Figure 4c). However, the levels were lower than with the 35S promoter, which could have been due to lower gene expression levels. Unexpectedly, however, transcript levels measured by quantitative RT-PCR show that STAPs confer significantly higher expression than the 35S promoter, in particular up to more than 10-fold higher for STAP-5 driving the expression of the GGPPS2 gene. Several hypotheses can be proposed to account for this negative correlation between transcript levels and product yield. The high expression of the enzymes could lead to negative feedback on the MEP pathway and thereby reduce flux through this pathway. Another possibility is that CBToI is produced in larger amounts with STAPs but is modified by endogenous N. benthamiana enzymes (e.g. acyl- or glycosyltransferases, glutathione transferases), as has been shown for mono- and sesquiterpenoids (Ting et al., 2013). Here, however, one would have to invoke that the more CBToI that is produced, the more it is modified in proportion. Yet another possibility is that the higher transcript levels are not accompanied by higher translation rates but that on the contrary this leads to reduced translation activity. Testing these different hypotheses will require extensive additional experiments which should be carried out in the future.

Eukaryotic promoters are currently described as containing two major domains, namely the core promoter element, defined as the region where the transcription machinery is recruited and initiates transcription, and the upstream elements, which are bound by TFs that can either activate or repress transcription (Hahn and Young, 2011). It is well established that binding to specific upstream elements by TFs triggers the recruitment of the core transcription machinery and thereby activates transcription (Hahn and Young, 2011). However, hardly anything is known about how the sequence of the core promoter element influences transcriptional activity (Lubliner et al., 2013). In an early study, the analysis of 95 yeast promoters led to the identification of a so-called ‘locator’ signal 30–10 bp upstream of the transcription start site (TSS), which is rich in Ts and poor in As, and is ‘stronger’
in promoters with high activity (Maicas and Friesen, 1990). More recently, a survey of 859 yeast core promoters identified T-richness upstream of the TSS as a strong predictor of promoter activity, regardless of whether the promoters are constitutive or inducible (Lubliner et al., 2013). As far as we could tell from the available literature, nothing is known about the situation in plants. Because of their short sequence and the presence of an identical DNA-binding site, STAPs constitute an appropriate material for studying the influence of the base composition of the core promoter on transcription activity. However, the identification of significant correlations between expression level and sequence content in our dataset would require a larger number of sequences.

Transcript levels were measured by quantitative RT-PCR for three of the promoters (STAPs 1, 3 and 5; Figure 5). The STAP-1 and STAP-3 promoters showed transcript levels that were similar to those of the 35S promoter, a result which is consistent with the GUS expression assays. Surprisingly the STAP-5 promoter showed transcript levels around 10 times higher than the 35S promoter, although GUS expression levels were similar to those with the 35S promoter. The base content between the TSS and the translation initiation codon [i.e. the 5'-untranslated (UTR) region] is known to have a dramatic impact on translation efficiency leading to differences of up to 100-fold in translation efficiency (Rojas-Duran and Gilbert, 2012). It is therefore quite possible that in the case of STAP-5 the high transcript levels do not result in correspondingly high protein expression due to an unfavourable 5'-UTR sequence. One way to circumvent this issue would be to introduce a 5'-UTR sequence known to enhance translation efficiency, such as the Ω-translation enhancer of the tobacco mosaic virus (Sleat et al., 1987).

Our design for the STAPs was based on reports that identified the presence of AvrBs3-binding sites within 100 bp of the transcription start site of genes activated by AvrBs3, a TAL-effector from the plant pathogenic bacterium Xanthomonas campestris (Kay et al., 2007, 2009; Romer et al., 2007). Furthermore, Kay and collaborators showed that a TATA box immediately downstream of the TALE-binding site leads to greatly increased expression (Kay et al., 2009). Although our design gave satisfactory results, further variations could provide additional regulatory options. For example, adding multiple TALE-binding sites could lead to increased expression, as shown in mammalian cells (Perez-Pinera et al., 2013). However, since the strongest STAPs are almost on a par with the viral 35S promoter this may be of limited interest. More relevant would be the insertion of TALE-binding sites for positive or negative feedback regulation. Positive feedback can simply be achieved by adding a DNA-binding site for another

Figure 5. Transcription activator-like effector (TALE)- and 35S-driven expression of the CBTS2a, DXS2 and GGPPS2 genes in agro-infiltrated Nicotiana benthamiana leaves.

Transcript levels were quantified by RT-PCR in three selected biological replicates from samples used in Figure 4 (rep1-3) from agro-infiltrated leaves co-expressing p19, CBTS2a, DXS2 and GGPPS2 driven by synthetic transcription activator-like effector-activated promoters (TALE) or the 35S promoter (35S). Mean expression values of three technical replicates ± SD and the corresponding CBTol amounts for each sample are given.
dTALE in the promoter driving the first dTALE. Negative feedback can be achieved in the same way but by replacing the activation domain of the second TALE by a repressor domain. Transforming dTALEs into transcriptional repressors has been successfully carried out in various eukaryotic organisms including plants (Blount et al., 2012; Mahfouz et al., 2012). Thus, our STAPs provide the starting material for designing complex regulatory circuits that could include transcriptional cascades, Boolean operators, as well as positive and negative feedback loops.

Given the ever-increasing number of genome sequences becoming available, including from higher plants, it will be possible to design TALE DNA-binding sequences that do not, or only minimally, interfere with endogenous gene expression, and conversely that are not bound by endogenous TFs. Thus, the dTALE/STAPs system can be considered a truly orthogonal system that neither depends nor interferes with endogenous transcription activation networks. The relevance of this system for metabolic engineering is obvious. Our transient expression assay for diterpene production in N. benthamiana demonstrated that several genes can be co-expressed by the dTALE/STAPs system at levels that are comparable to the strong CaMV 35S promoter. While using the same promoter to co-express multiple genes in a transient assay does work properly, in stable transgenic plants this can lead to undesirable consequences such as gene silencing (Peremarti et al., 2010). Therefore, a particularly relevant application of these promoters will be in stable transgenic plants, where coordinated tissue-specific expression of pathway genes can be tested without requiring the identification of multiple endogenous promoters conferring the same expression profile and with different expression levels. Hence, by simply changing the promoter driving the expression of the activating dTALE and keeping the rest of the construct identical, it will be possible with minimum cloning effort to test different tissues in parallel for their performance in producing a compound of interest. For example, one could compare tissues such as the seed endosperm, the epidermis, glandular trichomes or root hairs, for which specific promoters are available (Potenza et al., 2004; Tissier, 2012).

Recently, the discovery of the bacterial immune system based on the CRISPR/Cas9 system has opened novel opportunities for genome engineering and synthetic biology (Fineran and Dy, 2014; Hsu et al., 2014; van der Oost et al., 2014). The CRISPR/Cas9 enzyme is a nuclease whose sequence specificity is determined by a guide RNA (gRNA) containing the target sequence (van der Oost et al., 2014). As such, CRISPR/Cas9 is perfectly suited for genome editing, with the possibility of simultaneously targeting several loci by co-expressing or even concatenating multiple gRNAs, which can be easily cloned due to their small size (Hsu et al., 2014). In this regard, CRISPR/Cas9 genome editing is simpler to implement than TALE-based editing, because for each locus to be targeted, two new dTALEs need to be designed and expressed (de Lange et al., 2014). The CRISPR/Cas system was also modified to generate TFs (CRISPR/Cas-TF), whose specificity is determined by gRNAs (Nissim et al., 2014). Typically, gRNAs are expressed from RNA pol III promoters which are constitutive, and therefore provide no flexibility for tissue- or condition-specific expression. One solution to this problem is to express the gRNAs by RNA pol II promoters and edit the RNA with the Csy4 enzyme to ensure processing of the gRNA in the nucleus (Qi et al., 2012), but the functionality of Csy4 remains to be shown in plants. In addition to this added complexity, the fact that the specificity of the CRISPR/Cas-TF is determined at the RNA level, and not at the protein level as with the TALEs, constitutes a limitation for the design of more complex regulatory networks with activating or repressing steps. Because all CRISPR/Cas proteins will recognize the gRNAs and their target regardless of whether they are modified to be transcriptional activators or repressors, co-expression of both types of CRISPR/Cas-TFs in a single cell would result in unpredictable expression of the target genes. Also, a single CRISPR/Cas-TF protein controlling the whole network cannot act simultaneously as a repressor and activator on different targets. Thus, to introduce a negative regulatory step in a CRISPR/Cas regulatory network, Nissim et al. (2014) had recourse to a miRNA. In contrast, distinct dTALEs with either repressor or activating properties can be co-expressed, thus allowing much more flexibility and possibilities for the design of regulatory networks (de Lange et al., 2014).

These STAPs can be used for metabolic engineering of complex biosynthetic pathways requiring the simultaneous and tunable expression of multiple genes. This is particularly relevant in plants, where specific tissues or conditions can then be tested and compared for their performance in producing specific classes of compounds. Furthermore, these STAPs provide the basis for the design and construction of complex regulatory networks that contain transcriptional cascades and positive and negative feedback loops. Thus, their potential reaches beyond the field of metabolic engineering.

**EXPERIMENTAL PROCEDURES**

**Construction of a STAPs library**

To generate a library of promoter sequences as described in Figure 1, two degenerate primers, Talpro7 and Talpro8 (see sequences below), containing the TALE DNA-binding site (effector binding element 2 or EBE-002, described in Weber et al. (2011) as the binding site for dTALE-2) and degenerate sequences on either side of the DNA-binding site were made and annealed at a concentration of 0.2 μM each in a 50 μl reaction mix with the KOD polymerase (Merck Millipore, https://www.merckmillipore.com). Primer extension was carried out by first denaturing for 5 min at 94°C, then annealing at 60°C for 30 sec and extending at 72°C for 1 minute. A primer extension reaction was done for each primer to generate libraries of de novo promoter sequences. Sequences of the primers are given below:

| Primer       | Sequence                  |
|--------------|---------------------------|
| Talpro7      | GGTGATATGTATGTATGTATATG   |
| Talpro8      | GTAGATATGTATGTATGTATG     |

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30 sec. The product was then purified on a Qiaquick column (Qiagen, http://www.qiagen.com/) and inserted into vector pAGM1311 (universal level –1 cloning vector) (Engler et al., 2014) using a standard Golden Gate cloning procedure with BsaI and T4 DNA ligase (Werner et al., 2012). An aliquot (100 μl) of the transformed bacteria was spread over two Petri dishes for evaluation of the complexity of the library, and the remainder of the transformation inoculated in liquid culture for library preparation. Then DNA from the whole library was prepared and further subcloned into the level 0 vector pICH41295 (Engler et al., 2014) using BpiI and T4 DNA ligase. The strategy of cloning the library in two steps (level –1 and 0) results in a higher amount of promoters with a correct structure compatible with Golden Gate cloning. In particular, the two successive cloning steps that use BsaI and then BpiI for cloning result in the elimination of promoters that may have contained BpiI or BsaI sites in internal random sequences. The resulting level 0 constructs (pAGT582-n) were then picked individually, checked for the presence of single promoter sequences after digestion with BsaI, sequenced and used for further cloning. The sequences of the 43 promoters are provided in Data S1. Talpro7: 5′-ttgctc aacat GGAG (n)19 tccccgcatagctgaacatc. Talpro8: 5′-ttt ggtctc aacat CATT (n)43 ttatata ggtctcagctatgcgggg (the capital letters indicate sequences that will become part of the BsaI cleavage sites in the final level 0 modules).

**Reporter constructs**

The STAPs coming from the pAGT582-n vectors were cloned either in front of a GUS reporter gene (plasmid pCH75111) (Engler et al., 2014) and the ocs terminator (pICH41432) (Engler et al., 2014) to give pAGT615-n vectors, or in front of the GFP reporter gene (pICH4153) (Engler et al., 2014) and the ocs terminator to give the pAGT917-n vectors. The constructs were made in the T-DNA vector pICH75044 (Engler et al., 2014). All clonings were made using standard Golden Gate cloning procedure with the appropriate restriction enzymes (Engler et al., 2008).

**GUS transient assays**

All T-DNA vectors were transformed into A. tumefaciens strain GV3101 (Koncz and Schell, 1986). Nicotiana benthamiana plants were grown in a greenhouse maintained at a constant temperature of 22°C with 82% humidity. For the GUS assays, the constructs were co-infiltrated with the TALE containing the DNA-binding domain specific for the EBE002 sequence (construct pICH74043; Weber et al., 2011) and a 35S:GFP construct as internal reference (construct pAGM4731; Engler et al., 2014) to give pAGT615-n vectors, or in front of the GFP reporter gene (pICH4153) (Engler et al., 2014) and the ocs terminator to give the pAGT917-n vectors. The constructs were made in the T-DNA vector pICH75044 (Engler et al., 2014). All clonings were made using standard Golden Gate cloning procedure with the appropriate restriction enzymes (Engler et al., 2008).

**Diterpene metabolic engineering constructs**

The coding sequences of the SIDX2x (pAGT345), NtGPPPS2 (pAGT189) and CBTS2a (pAGT238) genes were those described in Bruckner and Tissier (2013). They were cloned behind the promoters from pAGT582-3, pAGT582-5 and pAGT582-1, respectively. The terminators used were Sl vATPase1 (pICH71431), Ocs (pICH41432), and CBTS (pAGT168), respectively (Bruckner and Tissier, 2013; Engler et al., 2014). These fragments were assembled into the level 1 vectors pICH7751, pICH7742 and pICH7802, respectively. The vectors were transformed into A. tumefaciens strain GV3101 and used for N. benthamiana leaf infiltration as described above. The corresponding constructs with the 35S promoter were as described in Bruckner and Tissier (2013). Metabolite extraction and diterpene GC-MS measurements were performed as described in Bruckner and Tissier (2013) with the following modifications. The 1 ml hexane extracts from the six leaf discs per sample were not evaporated and 160 μl of these extracts were combined with 40 μl of internal standard (IS) (50 μm sciarol stock solution in hexane), mixed, centrifuged and transferred to a GC-vial. Samples were measured on a QP2010 Ultra GC-MS system (Shimadzu, http://www.shimadzu.com/) in the following conditions. One microlitre was injected splitless at 250°C with a 1-min sampling time in a Rxi-5Sil MS 30 m, 0.25-mm diameter column (Restek, http://www.restek.com/). Elution was done with a constant He flow of 0.9 ml min⁻¹. The oven temperature was raised from 50°C to 300°C (7°C min⁻¹), then to 320°C (50°C min⁻¹) and held for 2 min. The interface and the ion source temperatures were 300°C and 250°C, respectively, and the ionization voltage was set at 70 V.

**Quantitative gene expression analysis in infiltrated leaf tissue**

Total RNA was extracted from infiltrated leaf discs from which diterpenes were extracted by a prior hexane wash. Leaf discs were homogenized to a fine powder using a Mixer Mill MM-400 (Retsch, http://www.retsch.com/) and RNA was extracted with the Spectrum™ Plant Total RNA Kit (Sigma-Aldrich, http://www.sigma-aldrich.com/) according to the manufacturer’s specifications. Any genomic contamination was removed before cDNA synthesis using the DNA-free Kit (Ambion, http://www.lifetecnologies.com/uk/en/home/brands/ambion.html) according to the manufacturer’s instructions. The first strand of cDNA was synthesized from 1 μg
total RNA using a Maxima H Minus First Strand cDNA Synthesis Kit (Thermo Scientific, http://www.thermoscientific.com/) and a mix of oligo(dT)18 and random hexamer primers. The cDNA was diluted 10-fold and 1 μl was used in a 10-μl reaction mix containing 1 μl (2 pmol μl⁻¹) of both forward and reverse primers and 2 μl my-Budget 5 × EvaGreen QPCR Mix II (Bio&Sell, http://www.bio-sell.de/). Quantitative real-time PCR was performed in 96-well plates using the CFX Connect real-time PCR detection system (Bio-Rad, http://www.bio-rad.com/). The thermal profile of the reaction was as follows: an initial denaturation at 95°C for 15 min, followed by 40 cycles at 95°C for 15 sec and 58°C for 30 sec with fluorescence acquisition after each cycle. To verify primer specificity and dissociation curve was finally generated by increasing the temperature from 65 to 95°C. Quantitative (q)PCR primers for the amplification of CBTS2a, DXS2 and GGPPS2 genes were designed using the Primer3 program (Undergrasser et al., 2012). The N. benthamiana eukaryotic elongation factor 1a was selected as the endogenous reference for normalization of transcript levels and PCR amplified using primers as described in Liu et al. (2012). To check the specificity of the CBTS2a, DXS2 and GGPPS2 primers, qPCR was also performed on N. benthamiana cDNA and no amplification could be detected in any of the control samples (Figure 5). The sequences of the oligonucleotides used for real-time PCR are provided in Data S2. All the cDNA samples were amplified in three technical replicates from the same RNA preparation and the mean value was considered. Expression of transgenes in infiltrated leaf tissue was calculated using the 2(−ΔΔCt) method (Livak and Schmittgen, 2001).

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CONFLICT OF INTEREST

The authors declare that they have a conflict of interest. A patent application was filed. The promoters will be made available free of charge for non-commercial research purposes.

SUPPORTING INFORMATION

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

Figure S1. Correlation between gene expression and cembratrien-ol (CBTol) levels.

Data S1. Sequences of the 43 synthetic transcription activator-like effector-activated promoters (STAPs).

Data S2. Sequences of the primers used for quantitative gene expression analysis.

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