Hairy Root Transformation: A Useful Tool to Explore Gene Function and Expression in Salix spp. Recalcitrant to Transformation

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Willow (Salix spp. L.) species are fast-growing trees and shrubs that have attracted emergent attention for their potential as feedstocks for bioenergy and biofuel production, as well as for pharmaceutical and phytoremediation applications. This economic and environmental potential has propelled the creation of several genetic and genomic resources for Salix spp. Furthermore, the recent availability of an annotated genome for Salix purpurea has pinpointed novel candidate genes underlying economically relevant traits. However, functional studies have been stalled by the lack of rapid and efficient coupled regeneration-transformation systems for Salix purpurea and Salix spp. in general. In this report, we describe a fast and highly efficient hairy root transformation protocol for Salix purpurea. It was effective for different explant sources and Salix purpurea genotypes, with efficiencies between 63.4% and 98.7%, and the screening of the transformed hairy roots was easily carried out using the fluorescent marker DsRed. To test the applicability of this hairy root transformation system for gene functional analysis, we transformed hairy roots with the vector pGWAY-SpDRM2, where the gene SpDRM2 encoding a putative Domain Rearranged Methytransferase (DRM) was placed under the control of the CaMV 35S constitutive promoter. Indeed, the transgenic hairy roots obtained exhibited significantly increased expression of SpDRM2 as compared to controls, demonstrating that this protocol is suitable for the medium/high-throughput functional characterization of candidate genes in S. purpurea and other recalcitrant Salix spp.

Keywords: Salix purpurea, willow, domains rearranged methyltransferase 2 (DRM2), Agrobacterium rhizogenes-mediated transformation, pGWAY-0

INTRODUCTION

Salix spp. L. (willows) are very diverse, comprising more than 400 identified species spread over a wide variety of natural habitats (Sulima et al., 2018). Willows display a high morphological diversity, occurring in the growth forms of trees, shrubs, or subshrubs. Shrub willows (Vetrix sub-genus) are ideal biomass feedstocks for bioenergy and biofuel applications given the ease of vegetative propagation, fast growth in short-rotation coppices (SRC) and high biomass yields (Kuzovkina et al., 2008; Karp et al., 2011). Some Salix spp. can also be used in phytoremediation strategies as they are characterized by physiological adaptations and ecological resilience, rendering them particularly suitable for the clean-up
of environmental contaminants (Kuzovkina and Quigley, 2005; Zalesny and Bauer, 2007). Besides, willow species such as Salix purpurea L. (purple willow) have great potential for the production of natural alternatives to synthetic aspirin, as their bark is a source of salicylic glycosides (SGs) (Sulima et al., 2017). In the last few years, the development of genetic and genomics tools for Salix spp., together with extensive phenotyping efforts, have significantly extended our knowledge on the factors involved in trait determination and phenotypic adaptation in willows (Hanley and Karp, 2014; Sulima et al., 2017; Fabio and Smart, 2018; Sulima et al., 2018; Gouker et al., 2019). Furthermore, the recent availability of the S. purpurea genome (https://phytozome.jgi.doe.gov) coupled to transcriptomic studies (Carlson et al., 2017; Yanitch et al., 2017) allowed this species to become a model for the Salix genus. However, the functional characterization of new candidate genes has been hindered by the lack of rapid and efficient regeneration and transformation protocols for S. purpurea and Salix spp. in general. Salix spp. are recalcitrant to both transformation and in vitro regeneration. There are few reports describing the in vitro regeneration of Salix plants (Grönnroos et al., 1989; Stoehr et al., 1989; Lyrya et al., 2006), but only one reported the regeneration of a significant number of plantlets (Stoehr et al., 1989). Early attempts to regenerate transgenic willow species were also proven ineffective, as no shoots were regenerated from transformed calli (Vahala et al., 1989; Vahala et al., 1993). Yang et al. (2013) developed a coupled regeneration-transformation system for Salix matsudana Koidz using the embryo apical region of mature seeds as initial explant. Shoots were regenerated directly from cotyledonary nodes. The average transformation frequency was low, approximately 7%, and moreover, this method required a laborious screening of transformants, giving the chimeric nature of transgenic plants produced. More recently, Guan et al. (2018) developed an Agrobacterium tumefaciens-mediated genetic transformation system using leaf-based calli of S. mongolia as explants for transformation. Differentiation of adventitious buds and rooting of plantlets was accomplished by adding different ratios of 2,4-dichlorophenoxyacidic acid (2,4-D), 6-benzylaminopurine (BA), and naphthaleneacetic acid (NAA) into Murashige and Skoog (MS) medium. The Agrobacterium-mediated integration of the β-glucuronidase (gus) gene into S. mongolia genome of five transgenic lines was confirmed by Southern blot (Guan et al., 2018), but no data on transformation efficiency were provided.

A. rhizogenes-mediated hairy root transformation systems are particularly useful for species recalcitrant to transformation by A. tumefaciens, since much higher transformation efficiencies are obtained, and shorter transformation periods are attainable in comparison to A. tumefaciens-mediated transformation systems. Hairy root transformation systems have been previously established in woody species such as Prunus spp. (Bosselut et al., 2011), Populus spp. (Yoshida et al., 2015), Eucalyptus grandis (Plasencia et al., 2016), Camelia sinensis (Alagarsamy et al., 2018), and Dryas spp. (Billault-Penneteau et al., 2019). In poplar, hairy roots were used to study the role of the transcription factor MYB182 on the regulation of proanthocyanidin and anthocyanin biosynthesis (Yoshida et al., 2015). In Eucalyptus, hairy roots were shown to be a suitable system for the functional characterization of genes involved in lignin biosynthesis, such as the Eucalyptus cinnamoyl-CoA reductase1 (EgCCR1) (Plasencia et al., 2016). To our best knowledge, in Salix spp., the induction of hairy roots has only been described in Salix alba L. (Hauth and Beiderbeck, 1992) in a report dating back to the early nineties. However, the aim of this study was solely to improve root biomass by the production of A. rhizogenes-induced hairy roots, as the normal roots cultures of Salix alba presented slow growth rates. Given the nature of this study, no confirmation of putatively transformed hairy root lines was done. Therefore, there is still a need to develop a hairy root transformation protocol that would allow the rapid characterization of gene function in Salix spp.

Here, we propose a reproducible, rapid and highly efficient A. rhizogenes-mediated hairy root transformation system for S. purpurea (Figure 1; details in Supplementary Material S1). In this method, the transformed hairy roots are detectable by fluorescent markers, allowing an easy and fast selection of transgenic roots. Our results suggest that this transformation system can potentially be applied to different genotypes, enabling gene functional studies in selected S. purpurea genotypes.

Hairy Roots: A Highly Efficient Strategy for Purple Willow Transformation

Different hairy root transformation protocols were tested to check the effect of explant age, plant culture media, and genotype on the transformation efficiency. We initially tested four transformation protocols combining one genotype at two developmental stages and two culture media (Supplementary Table S2.1—Experiment A). DiRed fluorescence was easily detected in neo-formed calli at wounded sites and in emerging roots in average 7 days after inoculation (dpi), i.e. (Figure 1). The transformation efficiencies obtained 21 dpi were higher for two-week-old S. purpurea plantlets grown in MS30 (reaching 83.33%) with the other three tested conditions presenting lower efficiencies (between 63.41% and 67.86%). To validate the results of the first experiment and to check if the developed protocol could be applicable to different S. purpurea genotypes, a second transformation experiment (Supplementary Table S2.1—Experiment B) was performed comparing two non-related genotypes, ELB3/6 and ELB2/5, using two-week-old in vitro clonal lines of S. purpurea grown in MS30, as this was shown to be the best condition to maximize the transformation efficiency. In this second experiment, the transformation efficiencies were slightly higher in both genotypes (above 86%), showing that the developed protocol is reproducible, rapid, and highly efficient (Supplementary Table S2.1—Experiment B). Moreover, we observed that isolated transformed hairy roots were able to continue to grow in solid media, and composite plants could be grown in hydroponic culture, allowing the maintenance and multiplication of composite plants and hairy roots for further experiments.

Application of Hairy Roots Transformation of S. purpurea for Gene Functional Analysis

To confirm that this transformation approach was suitable for functional gene characterization and gene function hypothesis-testing, we transformed roots with SpDRM2 (SapurV1A.0571s0130), a gene encoding a putative DNA methyltransferase with rearranged catalytic domains (For...
FIGURE 1 | Workflow of the hairy root transformation of Salix purpurea in vitro plantlets by A4RS harboring a DsRed-based binary vector. Emerging DsRed fluorescent roots are already detected 7 days post inoculation (dpi). (A) 6-week-old S. purpurea in vitro plantlets are sub-cultured to MS medium with full strength macroelements and 30 g L⁻¹ sucrose (MS30). (B) 14-day-old S. purpurea in vitro plantlets were infected by stabbing the stem with a needle swabbed with Agrobacterium rhizogenes. (C) Infected plants were co-cultivated with A. rhizogenes for 14 days on MS30 supplemented with acetosyringone under dim light. (D) Plants were transferred to MS30 medium supplemented with Augmentin. (E) Generated hairy roots were examined at 21 dpi under a stereo fluorescence microscope. (F) Co-transformed roots were excised and collected for further analysis. (G) In vitro culture of hairy roots and composite plants. Figure adapted from Plasencia et al. (2016), introducing the protocol specificities of hairy roots transformation for S. purpurea.
details on cloning of SpDRM2 see Supplementary Data S1, Supplementary Table S2.2, and Supplementary Figure S1). This gene is an ortholog of DRM2 of Arabidopsis thaliana (AT5G14620), where it was shown to encode an enzyme involved in de novo DNA methylation and gene silencing (Cao and Jacobsen, 2002). Transgenic hairy roots potentially containing the construct pGWAY-SpDRM2 were screened using DsRed and transformation efficiencies were comparable to those obtained with pGWAY-0 (98.75% for 82 analyzed plants). Gene expression analysis by RT-qPCR confirmed the overexpression of SpDRM2 in pools of hairy roots transformed with pGWAY-SpDRM2, collected from three different composite plants (Figure 2) (Supplementary Data S1, and Supplementary Table S2.3). Comparing wild-type roots to roots transformed with pGWAY-0 (empty vector), no significant difference was observed in the transcript level of SpDRM2, indicating that the endogenous gene expression was not affected by the transformation process. In contrast, the expression level was significantly increased by 2.9-fold to 6.9-fold in SpDRM2-overexpressing lines as compared to wild-type (For details on cloning of SpDRM2 and RT-qPCR analysis of transformed hairy roots check S1- Detailed Material and methods, Supplementary Table S2.2 and Supplementary Table S2.3). Based on the results obtained, we expect that this transformation method will be a valuable tool for the medium-/high-throughput functional characterization of candidate genes in S. purpurea.

**DISCUSSION**

Current A. tumefaciens-mediated transformation procedures of willows (Vahala et al., 1989; Yang et al., 2013; Guan et al., 2018) still limit functional studies in Salix spp. As they are laborious, time-consuming, genotype-dependent, and hindered by low transformation efficiencies. Furthermore, no stable transformation protocol is currently available for S. viminalis and S. purpurea, two relevant willow species for the production of biofuel and pharmaceutical compounds, respectively. Thus, the use of A. rhizogenes to induce transformed hairy roots represents a novel approach in Salix, as these species are recalcitrant to regeneration and transformation with A. tumefaciens. Previous reports suggest that hairy roots can be induced on a wide range of woody species, using *in vitro* (Alpizar et al., 2006; Bosselut et al., 2011; Yoshida et al., 2015; Plasencia et al., 2016) and *in planta* strategies (Alagarsamy et al., 2018), and from different explant types, e.g. seedlings, roots, stems, and leaves. In the present study, A. rhizogenes strain A4RS harboring the vector PGWAY-0 (Plasencia-Casavedell, 2015; Plasencia et al., 2016) was proven highly effective at inducing transgenic hairy roots in S. purpurea in explants with different ages and different genotypes. Transgenic hairy roots were easily detectable using the fluorescent marker DsRed, with emerging transformed roots appearing in average 7 days after A. rhizogenes inoculation. The fluorescent marker DsRed facilitates the identification of transgenic roots (which can be selected in the absence or in addition to antibiotic selection), allowing the non-destructive and precocious identification of transgenic roots (Limpens et al., 2004; Chabaud et al., 2006; Meng et al., 2019). High efficiency of hairy root transformation (ranging from 63% to 98%) was achieved for all tested conditions. The hairy root transformation efficiencies obtained in this report are similar to those reported for other woody species (Alpizar et al., 2006; Bosselut et al., 2011; Plasencia et al., 2016, Alagarsamy et al., 2018), legumes (Estrada-Navarrete et al., 2007; Aggarwal et al., 2018), and tomato (Ho-Plágaro et al., 2018). However, most available protocols use seedlings as explant for hairy root induction while in this report we used *in vitro*-cultured plantlets of *S. purpurea* as explants. The use of *in vitro*-cultured clonal lines has the advantage of not requiring the sterilization and germination of seeds prior to transformation, which can be limiting steps in the protocol in case of seed contamination, poor seed germination, and limited seedling growth (Alagarsamy et al., 2018). Contrastingly, *in vitro*-cultured clonal lines only require simple maintenance procedures and can be an almost continuously available source of explants for hairy root induction experiments. Besides, the use of clonal lines allows testing the expression of target genes in a plant material with identical genetic background.

Using the developed protocol (Figure 1, and Supplementary Material S1) we were able to achieve very high hairy root co-transformation efficiencies for both tested *S. purpurea* genotypes, suggesting that this hairy root transformation protocol can potentially be transferred to other *S. purpurea* genotypes. Nonetheless, more experiments should be done to test the susceptibility to *A. rhizogenes* in different *S. purpurea* genotypes and other willows species. A main limitation of this protocol is that it does not allow the transformation of above ground plant tissues other than hairy roots. Still, this can also represent an advantage when investigating root-shoot interactions in composite plant systems, to study, for example, the transport.
of small regulatory and signaling molecules (e.g. sRNAs, peptides and metabolites) between roots and aerial components of the plant. Transgenic hairy roots and composite plant systems can be used to study resistance against different biotic (Mellor et al., 2012; Xue et al., 2017) and abiotic stresses (Kajikawa et al., 2010; An et al., 2017; Du et al., 2018; Li et al., 2019), mycorrhizal associations, and root symbioses (Clemow et al., 2011; Indrasumunar et al., 2015; Billault-Penmeteau et al., 2019). Furthermore, the use of the CRISPR/Cas9 technology with hairy root transformation has emerged as an efficient tool for plant genome editing and gene functional studies (Ron et al., 2014; Wang et al., 2016; Du et al., 2018), thus opening a whole new range of applications for Salix spp., such as the manipulation of target biosynthetic pathways through multiplexed genome editing. Indeed, as each single hairy root represents a single transformation event and can continue to grow autonomously, the system can be particularly useful as a medium-high-throughput tool for functional analysis and biotechnological applications. The overexpression of SpDRM2 in hairy roots transformed with the pGWAY-SpDRM2 construct demonstrated the potential of S. purpurea hairy root transformation as a homologous, versatile, and efficient system that will enable the rapid validation of candidate genes and gene mining in S. purpurea and other recalcitrant willow species.

DATA AVAILABILITY STATEMENT

The datasets generated for this study are available on request to the corresponding author.

AUTHOR CONTRIBUTIONS

JG-P and JP conceived the study and designed the experiments. CG, AP, and JP performed the experiments and analysed the results. CG, AP, JG-P, and JP wrote the manuscript. All authors read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2019.01427/full#supplementary-material

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