THE DEVELOPMENT AND EVALUATION OF TRANSGENIC SORGHUM LINES FOR CONFERRED PHENOTYPES

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THE DEVELOPMENT AND EVALUATION OF TRANSGENIC SORGHUM LINES FOR CONFERRED PHENOTYPES

BY

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A THESIS SUBMITTED IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR DEGREE OF MASTER OF SCIENCE IN

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UNIVERSITY OF RHODE ISLAND

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OF

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2020
ABSTRACT

Agrobacterium-mediated transformation of sorghum (Sorghum bicolor) requires minimally 9-months from wild type immature embryo explants to T0 seeds. More efficient methods for sorghum transformation are necessary to conduct routine transgenesis for genome editing purposes in this important crop. With this in mind, there were two primary objectives to this thesis. The first was to evaluate and possibly improve upon methods for characterizing putative transformants once produced through Agrobacterium-mediated transformation of sorghum. The second was to evaluate and possibly improve upon transformation efficiencies in sorghum using an available Cas9 construct that would provide, long-term, a platform in sorghum for gene editing purposes. The first objective was addressed by evaluating a previously generated transgenic sorghum line designed to improve overall grain yield through the introduction of a maize silkless gene (skl) construct. The maize silkless gene was first used in maize by Hayward et al., 2016 to produce transgenics that conferred feminization on maize male flower by down regulating the jasmonic acid synthesis pathway. The same construct SK1ΔSVL:Citrine:SVL used in Hayward et al., 2016 was used to transform sorghum based on the understanding of high homology between the sorghum and maize genomes. Through a series of analyses, the presence and expression of SK1ΔSVL:Citrine:SVL was confirmed, but the predicted phenotype of flower feminization and improved yields was not observed in T1 transgenic sorghum lines. The second objective was designed to introduce into sorghum a vector containing Cas9 to test the stable expression of Cas9 for genome editing in transgenic lines. The first step towards this objective was to generate and characterize the
required transgenic lines and appropriate controls. The pNG111-ZmUbi::TaCas9_PvUbi::1GFP construct was used to integrate TaCas9 into the wild type BTx430 sorghum genome and molecularly characterize these events. The pNG108PvUbi::1GFP construct served as a negative control for pNG111 since it is lacking the TaCas9 cassette. Both constructs contain constitutively expressed mGFP which is detected as a visible reporter, and the bar gene served as a selectable marker conferring resistance to the herbicide bialaphos. In addition, transgenic lines for both constructs were molecularly characterized by PCR, Southern blot analysis, and the ‘paint assay’ to detect the functional expression of the bar gene. Functional analysis of the stably integrated TaCas9 will be conducted in future studies. By analyzing the newly developed transgenic sorghum lines with constitutively expressed TaCas9, we hope to contribute to the development of a new platform for genome editing in sorghum.
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LITERATURE REVIEW

I. Plant Transformation

The development of genome editing applications through the use of plant transformation technology in cereal crops is critical for future crop improvement. This is a required technology to develop and analyze genome modified plants. Plant transformation is used to introduce valuable transgenes into the plant genome, followed by the recovery of regenerated, fertile plants with stably integrated transgenes that confer trait enhancement or improvement. Improvements on existing transformation technologies would ideally allow for genotype independent and almost tissue culture free strategies. Improvements in current plant transformation technology are both required and necessary to functionalize genomic analyses through genome editing. This is essential to functionally link genes to biological processes that would allow for the modification of existing metabolic pathways.

Crop improvement strategies based on biotechnology strategies will require concurrent advances in plant genomics including advanced genomics for gene annotation and functional analysis and genome editing through plant transformation (Kausch et al. 2019; Council for Agricultural Science and Technology (CAST), 2018; Altpeter et al. 2016). In fact this very topic has been the subject of intensive review over the past several years (Songstad et al. 2017). Robust genomics and computational biology tools provide ways to identify target sequences for genome editing. Further, advanced genome editing approaches then allow for desired modifications in a target species genome, such as, single base pair change, insertions and deletions (Council for
Agricultural Science and Technology (CAST), 2018; Songstad et al. 2017). The application of these technologies is highly dependent on the capability to recover fertile genome modified plants. Plant transformation for any cereal species or variety will make possible direct gene analysis, targeted trait modification and provide a new basis for the application of the principles of synthetic biology.

Standard plant transformation protocols can be applied to knock-out (down) gene expression, make specific adjustments in protein structure and function, and observe over-expression and ectopic characteristics as an enabling technology in basic plant biology. However, the limitations of current standard plant transformation protocols have created a daunting bottleneck for functional genomic analyses and genome editing (Altpeter et al. 2016). Ideally, plant transformation should not be limiting with regard to plant species, genotype or explant source. An additional preference is that the technology be untethered, as much as possible, to reliance on tissue culture. The improved technology should be able to efficiently modify any genomic sequence in any variety and efficiently produce stably heritable events.

With increasing global human population growth and food consumption, many research efforts have focused on biotechnology based crop improvement to overcome the previous issues for sustainable agriculture (Jiao et al. 2018). Plant transformation is a major challenge and bottleneck for creating the transgenics and the recovery of genome edited events for functional genomics. The role of plant transformation is central to introducing specific DNA components, or genome editing of specific gene targets to recover stable, heritable genetic events. The gene modified plants will express the stable and heritable events of interest which can then allow for phenotypic
characterization for their conferred traits. However, the regeneration ability still exists as a significant bottleneck for most plant species to allow for efficient plant transformation (Altpeter et al. 2016).

Even if the related approaches have been well developed to transfer DNA into a single plant cell that can be regenerated into a whole, intact transgenic plant, plant transformation protocols still contain limiting parameters, as described previously, including genotype dependence and low or no regeneration efficiency in several important crop species. Moreover, some plant tissues are difficult to obtain as usable plant transformation explants (e.g., immature embryos or immature inflorescences). Therefore, these bottlenecks prevent the accessibility and transfer of technologies from academic laboratories in the public sector.

Plant transformation technology has been a well-recognized challenge for decades (Altpeter et al. 2016) and the technology has been incrementally improved upon over a long period on time. However, the current state-of-the-art is still ineffective for many crops, because current plant transformation requires an appropriate genotype, experienced labor, improvements for low frequencies, more efficient editing tool and huge screening efforts for inconsistencies of gene expression. Crop functional genomic research suffers from specific plant species that are recalcitrant to plant transformation (Altpeter et al. 2016; Kausch et al. 2019). Therefore, efficient Agrobacterium-mediated transformation is typically applied to a small range of plant species. One of the major challenges is the long tissue culture periods that are used to cultivate modified cells and tissues to transgenic plants. This extended tissue culture period has resulted in many obstacles for generating transgenic
and genome edited plants. Even though the tissue culture takes a long time to produce the transformed events, the frequency is low for engineered tissues to regenerate stably in many plants and cultivars.

There are two DNA delivery methods which are predominately used to edit organisms in most academic and industry laboratories: *Agrobacterium*-mediated transformation and particle bombardment-mediated gene delivery. The current transformation protocols for most plants currently require extensive tissue culture, but it is important to simplify and optimize the protocols for all crops to allow for wider application.

The complete systems for developing plant transformation technologies must be comprehensive, since it has involved the understanding of molecular biology, plant tissue culture, plant physiology and plant developmental biology (Kausch et al. 2019). However, the difficulties inherent in plant transformation are often underappreciated even though the technology provides the fundamental platform to develop transgenic organisms. The transgenic plants created provide the basis for analysis of conferred trait genetic value, basic plant biology, improve the metabolic pathways in plants, and produce stable resistance to environmental or biotic stress. For example, transgenic plants can express genes which confer resistance to herbicides (Devos et al. 2008) and viral or microbial pathogens (Ferreira et al. 2002). It is almost impossible to efficiently produce innovative improvement on plant traits, plant gene discovery, and functional genomics without plant transformation technologies.

*Agrobacterium* infection and gene transfer to plant cells has been well studied for the past three decades (Altpeter et al. 2016). A general schematic for
*Agrobacterium* infection and gene transfer to plants is shown (Figure 1). When the plant cells are injured, especially the dicot, the response is secretion of phenolic compounds, including acetosyringone. *Agrobacterium* responds to these signals through the VirA/VirG two-component sensing system which will stimulate the expression of the virulence (*vir*) genes (Altpeter et al. 2016). In this set of *vir* genes, *virD1* and *virD2* will combine together and form specific nucleases to cut the T-DNA region between the borders in the Ti plasmid in *Agrobacterium*. Recently designed binary vectors are capable of delivering the T-DNA containing any gene construct of interest including genome editing components. When the T-DNA region is excised out from the Ti plasmid, the VirD2 proteins will bind to the single-strand T-DNA and direct transfer of the T-DNA into the infected plant cell via the type IV secretion system. In the plant cell, the single DNA binding protein VirE2 covers and protects the T-DNA/VirD2 strand (Altpeter et al. 2016). The complex, formed by T-DNA/VirD2 strand with VirE2 and other plant proteins, can target the nucleus. Once the T-DNA is transferred into the nucleus, proteins will be dissociated from the T-DNA strand. The transferred T-DNA is then replicated to form double-stranded DNA which is in a non-integrated form (transient transformation). After integration into the plant genome, the T-DNA forms a stably transgenic cell which can be selected through the use of selectable markers, such as *bar*, *hra*, *als*, *hpt* and others (Kausch et al. 2019). The next step in the process is to regenerate these cells to fertile transgenic plants capable of use for breeding purpose (Altpeter et al. 2016).
Figure 1. Schematic of *Agrobacterium*-mediated transformation (Altpeter et al. 2016).

The molecular mechanisms involved *Agrobacterium*-mediated DNA transfer (left) transfers the T-complex into plant cells (center), which can then be regenerated to produce transgenic plants from the single totipotent cell.

*Agrobacterium tumefaciens* has been the subject of research for decades [reviewed by: (Altpeter et al. 2016; Kausch et al. 2019)]. Early work on *Agrobacterium* dates to the early 1940s with investigations on its plant pathogenicity as the causative agent for crown gall disease. In 1977, *Agrobacterium* DNA transfer to plant cells was reported via *Agrobacterium* Ti plasmid (Altpeter et al. 2016). The Ti plasmid contains the T-DNA which is imported into the plant to nucleus, but designed and engineered DNA constructs could not at that time be integrated into Ti plasmid. The genes involved with pathogenesis were removed (Fraley et al. 1983) to create a disarmed Ti plasmid which could be subsequently engineered (Fraley et al. 1983). Then, the binary vector in *Agrobacterium*-mediated transformation was developed after the expression of bacterial genome was reported in plant cell (Kausch et al. 2019). The developed binary vector was used to transfer foreign DNA into plant cells, which was able to stably genetically modify the plant genome (De Block et al. 1984). In the
binary vector system, antibiotic resistance genes were used to select the transformed organism from non-transformed groups and various promoters were explored to drive the expression of the inserted gene(s) (Bevan et al. 1985).

The early plant transformation systems required *Agrobacterium*-mediated gene delivery, which contain the tissue culture system for the DNA delivery, transformant selection and plant regeneration. However, the initial plant transformation systems were not able to be used with monocotyledonous plant. Even though extensive research efforts to develop plant transformation systems for monocots (Kausch et al. 2019), stable gene delivery through *Agrobacterium* and recovery of fertile transgenic plants was not accomplished. In a very innovative approach to overcome this obstacle to the gene transfer barriers, John Sanford and Ted Klein invented ‘the Gene Gun’ (microprojectile bombardment) in 1987 (Kausch et al. 2019) as an alternative to the *Agrobacterium*-mediated method (Kausch et al. 2019). In 1988, the first transgenic monocot plant was produced by microprojectile bombardment (Kausch et al. 2019). This direct DNA delivery method also eliminated the difficult and ineffective protoplast systems for maize transformation, an agronomically important monocot species. This technology was then applied successfully for the transformation of many cereals, but it also had several drawbacks, including: 1. gene silencing by multicopy gene insertion; 2. non-essential DNA insertions, such as the gene construct plasmid backbone; and 3. truncated or rearranged transgene cassette integration. The biolistic technology brought a novel approach to plant transformation, and produced significant research on basic plant biology and gene regulation. Eventually and fortunately, the *Agrobacterium*-mediated transformation methodology was improved upon and
extended to successfully recover transgenic monocot plants. This was accomplished in 1995 by researchers at Japan Tobacco by modification of the virulence genes to create super-virulent *Agrobacterium* strains which were capable of infecting and transferring DNA to monocot species. These super virulent strains have in turn, been modified and improved over the years to result in highly effective gene transfer methods now applicable to most monocots (Kausch et al. 2019).

In conclusion, plant transformation is a critical platform for the development of genome editing and transgenic genome engineering in plants. Furthermore, the *Agrobacterium*-mediated and microprojectile bombardment transformation systems remain as the most the reliable approaches for plant transformation for most plants. The main processes for plant transformation can be summarized in three critical steps: 1. the specific DNA delivery and integration into recipient cells; 2. selection of the successful stable integrant; and, 3. the regeneration of the transgenic plant from a single transformed totipotent cell to a fertile plant. The specific transformation protocols are variable depending on the unique features of monocots and dicots, which will be an important foundation for further development of functional genomics in both monocot and dicot plants.

II. Genome Editing

Extensive and time consuming conventional breeding programs produce the bulk of the varieties used for most of our current commercial crop plant production, the remainder being only a few undomesticated wild plants being used for crop production. All of the human food supply is produced from domesticated organisms
that are either naturally occurring mutants or the result of selection for desirable traits that confer environmental and yield advantages (Meyer et al. 2012). After early stages of plant domestication and selection of mutants through conventional breeding, mutational techniques were developed to promote higher frequencies of mutations and increased the efficiency of the selection stages for new varietal development. These induced mutation techniques include chemical mutagenesis such as EMS, physical mutagenesis, such as various radiation techniques, and the insertional mutagenesis such as transposon or gene tagging. Induced mutagenesis has successfully produced over 3200 officially released new varieties (FAO/IAEA 2014). Nonetheless, more efficient and precise techniques are still needed for increasing production levels.

New gene editing tools have been developed and used to efficiently edit genomic sequence in different species [reviewed in: (Weeks et al. 2016); and, (Songstad et al. 2017)]. For application of most gene editing tools, their editing function is dependent on how to identify and target specifically the desired sequence in the target genome. In order to accurately recognize a target sequence, site-directed nucleases (SDNs) would be used to recognize the target sequence. The SDNs directed system would make double stranded breaks (DSBs); then, DSBs would be repaired by endogenous non-homologous end joining (NHEJ) or homology-directed recombination (HDR). Typically, NHEJ would make some small sequence deletions or insertions which may result in genetic change at the target site. There have been several gene editing approaches that have been developed using SDNs, such as, meganucleases (Gao et al. 2010), Zinc Finger Nucleases (ZFNs) (Shukla et al. 2009), Transcription Activation-Like Effector Nucleases (TALENs) (Clasen et al. 2015) and
Clustered Regularly Interspaced Short Palindromic Repeats (CRISPRs) (Jinek et al. 2012). Currently, the CRISPR system is preferred and has been widely applied in academic research laboratories and industry. The CRISPR system has been developed from naturally occurring defense mechanism in *Streptococcus pyogenes* (Jinek et al. 2012; Cong et al. 2013; Mali et al. 2013). Compared to ZFN and TALENs, both of which use a protein to recognize the target DNA sequence, the CRISPR system is more precise and easier to apply in genomic research because RNA was used to recognize the target DNA sequence (Sander and Joung 2014; Schiml and Puchta 2016). The CRISPR system has been successfully applied in a wide range of plant species for genomic modifications (Jinek et al. 2012; Cong et al. 2013; Mali et al. 2013). Advanced genome editing provide a significant opportunity for modern plant breeding, which provides multiple directions for phenotypic improvements in plants (Bortesi and Fischer 2015; Jiang et al. 2013).

**CRISPR/Cas9**

Clustered regularly interspaced short palindromic repeats (CRISPR) along with CRISPR-associated proteins represent components of microbial defense mechanisms found in most Archaea and many Eubacteria. This is a mechanism to defend against viral and plasmid cellular invaders (Thurtle-Schmidt and Lo 2018). The CRISPR component contains many non-contiguous repeats and spacers, spacers are foreign genome elements (Figure 2A; blue and green elements) between repeats (Figure 2A). During the evolutionary adaptation of this mechanism, Archaea and Eubacteria achieve a cellular memory of the invading virus or plasmid (some pieces of foreign
DNA), and these pieces of foreign DNA are integrated into the CRISPR genomic locus. When CRISPR incorporates a foreign sequence, the system allows for the production of a target RNA, termed crRNA that can direct the Cas9 protein to bind to foreign target DNA (invaders) and cleave the sequence (Figure 2). Because crRNA recognizes some nucleotide sequences of the invader, it can direct Cas9 proteins to bind specifically to the DNA of invaders instead of itself (Thurtle-Schmidt and Lo 2018). The CRISPR systems contain 3 types of mechanism, type I and III are found in both bacteria and archaea, type II is found only in bacteria. Type I contains the Cas3 gene that encodes a large protein with divergent helicase and DNase activities. Type III contains polymerase and RAMP (repeat-associated mysterious proteins) modules. The RAMP superfamily does not present an autonomous functional unit but it can catalyze the processing of the long spacer-repeat-containing transcript into the mature crRNA (Thurtle-Schmidt and Lo 2018). The type II mechanism of the CRISPR/Cas9 system is best adapted to use for editing because it requires just one Cas9 protein and two RNA components. Before generating of crRNA, the CRISPR sequence is transcribed and combined with the foreign sequence to form pre-crRNA (Figure 2). The upstream portion of the CRISPR sequence is also transcribed. This is termed the trans-activating CRISPR RNA (tracrRNA). The tracrRNA can be complementary to the repeat region in the CRISPR sequence. It will bind with pre-crRNA to form a double-stranded RNA. Rnase III will recognize and cleave this double-strand RNA to form crRNA: tracrRNA (contain one spacer) complex. When this complex combines with Cas9 protein, the Cas9 protein can be activated to cleave a targeted DNA sequence (Thurtle-Schmidt and Lo 2018).
To function properly the Cas9 protein needs to recognize the protospacer-adjacent motif (PAM) first. Cas9 contains two domains to cleave the targeted DNA sequence (Figure 2B). One is the HNH domain that is complementary to crRNA; another is the RuvC-like domain that is not complementary to crRNA. However, designed single guide RNA can bind the crRNA and tracrRNA complex. This modified CRISPR/Cas9 system is much simpler to design, compared to either TALENs or ZFNs, for gene editing purposes. It is easier to construct DNA vectors than having to design the protein binding domains of TALENs and ZFNs. The induced sgRNA and Cas9 protein can make multiple double-strands break simultaneously, resulting in enhanced mutagenesis and gene editing. However, the modified CRISPR/Cas9 system also has been limited by the restriction of target sequence, large protein size and off-target mutation (Thurtle-Schmidt and Lo 2018). The PAM site is the main limitation for selection of a target sequence and different bacterial species have different PAM sites for the Cas9 protein. Scientists have created a Cas9 variant, namely SpCas9, that can recognize a different PAM site (Nakade et al. 2017). Another use of SpCas9 is an RNA-targeting Cas9 (Rcas9) system, which requires a PAM-presenting oligonucleotide hybridizing with a target single-strand RNA to act as PAM motif (Nakade et al. 2017).
Figure 2. (A) Schematic of CRISPR/Cas9 in vivo (Biolabs 2014) and (B) Function domain of Cas9 protein (A) shows 3 general working processes of CRISPR/Cas9: the foreign DNA sequences (same with target sequence) insert into CRISPR loci, then, CRISPR loci transcribes and binds with tracrRNA, this combined complex will bind to target site and direct Cas9 protein cleaving the sequence. (B) indicates the specific functional domain of Cas9 protein: RuvC domain and HNH domain.
III. Sorghum Transformation

The fact that cereal crops are important to global agriculture, food security, world economy, and international stability is well documented and widely understood (Belide et al. 2017). Sorghum (*Sorghum bicolor*) represents the third largest cereal crop in the world and the fifth largest in the US (Belide et al. 2017; Jiao et al. 2018). The grain is widely used for food, animal fodder and biofuels (Jiao et al. 2018). Application of modern biotechnology approaches for genetic improvement of sorghum is important to expand agricultural uses for this crop and address basic biological questions. Biotechnology approaches include advanced genomics, transgenics, and genome editing leading to improvement in traits such as enhanced yields, insect and pest resistance, abiotic stress tolerance, and increased nutrition. Basic biological questions, including genetic control of plant development, water use efficiency and photosynthesis can also be evaluated. In order to accomplish these goals, a reliable and robust plant transformation protocol is a fundamental requirement.

The protocols for sorghum transformation are well established (Figure 3). While protocols for sorghum transformation are well established, they are still not routine and subject to low efficiencies. The beginning of sorghum transformation requires large scale sorghum cultivation of wild type plants for donor material to provide enough immature embryos for continuous sorghum transformation. Also, the need for a constant supply of immature embryos requires intensive and expensive labor. However, while most research suggest that immature embryos are ideal explants over others for plant transformation (Belide et al. 2017; Wu et al. 2014), other explants, such as leaf material are currently being explored. The need for healthy and vigorous
immature embryos as explants has significance for transformation efficiency since they influence the embryogenic callus induction (Zhao et al. 2000). The transgene can be transferred into isolated immature embryos via Agrobacterium-mediated transformation or particle bombardment. The particle bombardment has several drawbacks (as previously described); therefore, Agrobacterium-mediated transformation has become the main approach for the sorghum transformation for routine trait manipulation. The transformed cells in sorghum immature embryos are selected from non-transformants and regenerated to complete fertile sorghum plant. However, some negative tissue culture factors, including the accumulation of phenolic compounds and continuous sub-culture, will gradually decrease the poor regeneration efficiency (Belide et al. 2017). The entire sorghum transformation protocol requires 9 to 12 month from the wild type sorghum embryos to the T0 gene modified sorghum fruit. Therefore, routine and robust sorghum transformation protocols are essential for investigating fundamental questions.

There are some challenges for sorghum transformation. The stable and continuous sorghum plant supply is required to provide appropriate explant for transformation experiments. Tissue culture is an essential step in sorghum transformation and also the need for experienced labor and corresponding facilities. The efficiency of transformation always suffers from genotype dependence, long tissue culture time and callus culture intermediates. These challenges interfere with the necessary enhancement of efficiency for sorghum transformation. In comparison, advances in genomics and gene editing approaches have shown tremendous progress. Recently, a breakthrough in monocot transformation provided an opportunity for the
improvement of sorghum transformation. BABY BOOM (BBM) AND WUSCHEL (WUS2) was confirmed as transcription factors genes, which are involved in somatic embryogenesis (Mookkan et al. 2018; Mookkan et al. 2017; Nelson-Vasilchik et al. 2018). The expression of BBM and WUS2 produce morphogenic regulators that can induce efficient somatic embryogenesis. Their use in transformation constructs could potentially improve the efficiency of transformation for sorghum and some other recalcitrant plant species (Lowe et al. 2018; Lowe et al. 2016; Nelson-Vasilchik et al. 2018). Necessary improvements for sorghum transformation efficiency can allow for improvements in strategies for sorghum genomic modifications of agriculture importance.

Figure 3. Representative timeline for standard sorghum transformation (Altpeter et al. 2016). Sorghum immature embryo explants (12 days post pollination, are used for plant transformation; the entire procedure starting from the bottom left corner to bottom right corner requires 9 to 12 months, and each part of the procedure is shown with the corresponding time.
CHAPTER 1: SEX DETERMINATION IN SORGHUM
Chapter 1: Sex Determination in Sorghum

Introduction:

Cereal crops that feed the world include rice, maize, sorghum, wheat, barley, oats and other minor cereal crops such as fonio and teff. Morphologically and genetically the cereal crops share significant homology, therefore, discovery of a gene in one of them has potential application as an orthologue in another. The *silkless 1* (*sk1*) gene of maize is known to play a key role in sex determination (Hayward et al. 2016). It is, therefore, plausible that a gene from maize could function in sorghum and carry out a similar function. There are many examples where across broad species orthologous genes have functioned well (Hayward et al. 2016), as well as a few where they do not.

Maize and sorghum are both hermaphroditic plants with similar flower morphologies, but not identical. Maize flowers are unisexual while sorghum flowers are bisexual. Having unisexual flowers is highly advantageous in hybrid crop production and the *sk1* gene may be the key distinguishing feature giving rise to maize and sorghum flower structure differences (Hayward et al. 2016).

The *sk1* gene has been characterized in the sex determination pathway in maize (Hayward et al. 2016). The sex determination pathway in maize and sorghum is complicated because several genes and phytohormones are involved. In particular, in maize, *silkless 1* (*sk1*), *TASSELSEED 1* (*TS1*), and *TASSELSEED 2* (*TS2*) are important in sex determination and influence each other (Li and Liu 2017). The *TS2* generates cell death signals for the pistil and *TS1* controls the expression of *TS2*. 
Moreover, both of them contribute to biosynthesis of jasmonic acid (JA) that is involved in pistil elimination and plays a key role in the developing stamen. The skl gene product is a protector for pistil formation because it prevents pistil elimination mediated by JA (Li and Liu 2017; Hayward et al. 2016). Therefore, it was hypothesized that enhanced expression of the maize skl in transgenic sorghum may influence sex determination for sorghum in a similar developmental pathway to what is observed in maize.

Maize is monoeocious with flowers that are initially bisexual. The maize sex determination system results in inflorescences with imperfect florets, the tassel and the ear, through organ arrest. Genetic analysis has shown that the expression of the skl is required to protect pistils in ear spikelets from tasselseed-mediated elimination. Recent studies by (Hayward et al. 2016) in maize showed that plants transformed with a skl transgene (SK1ΔSVL:Citrine:SVL) driven by a constitutive cauliflower mosaic virus (CaMV) 35S promoter displayed a pistillate phenotype, where the tassel inflorescence was completely feminized. The SVL domain is a putative peroxisomal targeting sequence. The reporter, citrine, was demonstrated to localize to peroxisomes (Hayward et al. 2016). These results indicate that skl expression is sufficient and necessary to block the tasselseed-mediated elimination of pistils in both ear and tassel spikelets, resulting in a completely feminized plant. This implies a mechanism of skl protection by prevention of jasmonic acid mediated pistil elimination (Hayward et al. 2016). Many related grasses, such as sorghum, develop two types of flowers on a panicle: one of these, known as the sessile spikelet (SS) is fertile and develops seeds; the other type called pedicellate spikelets (PS) do not make seeds. Single-copy
orthologs of sk1 have been identified in sorghum, although the SS florets are perfect because they are fertile and produce seed (bisexual). It is hypothesized that constitutive overexpression of the maize sk1 in sorghum could result in seed production in PS. It is also known that the msdl gene in sorghum (Jiao et al. 2018) participates in the jasmonic acid pathway and mutants result in rescued pedicellate flowers.

Using the same vector as in the maize experiments (35S:SK1ΔSVL:Citrine:SVL), 26 T₀ independent events were generated via Agrobacterium-mediated transformation of sorghum and selection for resistance to the herbicide phosphinotricin conferred by the selectable marker, bar, in the transformation vector (Dellaporta, personal communication 2017). The results from the current study involve analysis of T₁ plants produced from the T₀ events. Some of these plants showed resistance to the herbicide, and the presence of the transgene was confirmed by PCR and Southern blots compared with the segregating wild-type plants. Analysis using confocal microscopy was performed for citrine fluorescence, and seed morphology in SS and PS was examined in mature inflorescences.

Methods

Agrobacterium-mediated transformation

Agrobacterium-mediated transformation of sorghum cv BTx430 was performed following the method described in chapter 2 of this thesis (see also, Nelson-Vasilchik et al. 2018; Do et al. 2018) using the same construct described by Hayward et al. (2016) (pYU2996 SK1ΔSVL:Citrine:SVL; see Figure 4). The resulting
transgenic plants were then eventually transferred to Plant Cons, and finally to soil (Metro-mix). The transformation experiments were performed by Kimberly Nelson-Vasilchik and the resulting transgenic plants were made available for this study.

Paint Assay

The T₀ plants were swabbed with 3% bialaphos (referred to as the ‘paint assay’) to evaluate the presence and active expression of the bar gene. The ‘paint assay’ is non-destructive and allows for accurate diagnosis of resistant and sensitive plants.

Wild type BTx430 plants were used as controls. Transgenic T₁ individuals were also confirmed for phenotypic resistance on the swabbed region indicating the presence and active expression of the bar gene.

Genomic DNA extraction

Genomic DNA was extracted from T₁ plants (Chen and Dellaporta 1994). Leaf tissue was collected from T₁ plants, then ground with a mortar and pestle to a fine powder in liquid nitrogen. The plant tissue powder was then incubated with lysis buffer UEB3 (mixed by Urea, Tris, EDTA, Na₂SO₄, N-laurysarcosine and PVP) to break down the cellular membranes. Those samples were extracted with phenol:chloroform and centrifuged to precipitate most of the extracted plant material. The resulting suspension solution was centrifuged with isopropanol to precipitate DNA pellet. The DNA pellet would be dissolved by TE, Acetate and Qiagen RNase, Qiagen RNase was used to remove the RNA from DNA. The dissolved DNA solution was centrifuged with phenol:chloroform to precipitate rest waste from previous steps, the upper suspension from centrifuged tube contain most DNA materials and was centrifuged with isopropanol to form the DNA pellet. The collected DNA pellets were
dissolved by TE, then purified with 95% ethanol and 2.5M ammonium acetate. Purified DNA samples were rinsed by 70% ethanol and prepared for PCR and Southern blots.

PCR

The polymerase chain reaction (PCR) was used to determine the presence of *bar* and *citrine* cassettes in T1 plants. The PCR reactions were performed with the KAPABIOSYSTEMS KAPA3G Plant PCR kit. Primer information for the two genes is given in the Appendices (Table 2 a, b). Because the Tm for two gene primers is 60.0°C and the product size is approximately 500 bp each for the *bar* and *citrine* cassettes, the chosen annealing temperature was 55°C with a 30 second elongation time for 35 cycles in the thermocycler program.

Southern blot

Southern blots were used to determine approximate insert copy number. The Southern blots were performed with the Roche DIG Southern blot kit and the *bar* primer was used to produce the DIG-labeled probe for the hybridization step. The Southern blot protocol was performed according to the instructions in the Roche DIG Southern blot kit. The extracted genomic DNA was digested by restriction enzyme HindIII-HF (New England BioLabs). There no HindIII digestion sites in the complete *bar* gene sequence. Therefore, a positive band on blot membrane represents a complete, intact *bar* gene copy. Therefore, the number of bands on a given blot sample indicates the gene copy number for the gene. The digestion procedure was set up in 37°C water bath for 16 hours. The digested genomic DNA would be separated to different sequence size by gel electrophoresis. Then, the separated and digested
genomic DNA would be transferred from gel to blot membrane. The blot membrane would be hybridized with bar probe in DIG easy hybridization solution at 65°C for 16 hours. The hybridized blot membrane needed to be washed to remove undesired probe by using stringency buffer, and let bar probe connecting to antibody in block solution with Anti-DIG-AP. Then, the blot membrane would be covered with chemiluminescent CSPD solution and exposed to Lumi-Film. The hybridization result on the blot membrane would be revealed on the Lumi-Film.

Confocal microscopy

Citrine was imaged using water immersion confocal microscopy according to established protocols (Hayward et al. 2016). This work was done at the Leduc Bioimaging Facility in Brown University. Because citrine is fused with skl in the pYU2996 SK1ΔSVL:Citrine:SVL construct used in this study (Figure 4), the expression of Citrine also indicates the co-expression of SK1.

Figure 4. The skl transgene (pYU2996 SK1ΔSVL:Citrine:SVL) construct (Hayward et al. 2016). From T-DNA right border (RB) at left and 5’ to 3’, the constitutively expressed double 35S promoter (green) from cauliflower mosaic virus (CaMV) is used to drive expression of the bar CDS (coding sequence) selectable marker (light blue) fused to the TR7 terminator (brown). From the T-DNA left border (LB) at right and 5’ to 3’, the constitutively expressed double 35S promoter (green) is again used to drive expression of the silkless 1 coding sequence SK1CDS (dark blue) fused directly to the coding sequence for citrine (yellow) and the SK1 SVL domain for peroxisomal targeting (pale green) and the 35S termination signal Ter (grey).
Results

Transformed T₀ sorghum plants were produced using Agrobacterium harboring the pYU2996 SK1ΔSVL:Citrine:SVL construct (Figure 4) for transformation in the sorghum variety BTx430. The T₀ plants were selfed to produce a segregating population of T₁ seeds which were used for further characterization. Seeds were harvested at maturity and stored at 25°C in darkness.

The T₁ seeds harvested from pYU2996 Event #1, plant #1, were used to grow 25 segregating plants. Three plants (#10, #11, #21) of the 25 plants showed sensitivity to bialaphos by the ‘paint assay’ indicating segregation of the bar gene in the T₁ population. The remaining plants showed resistance to bialaphos in the ‘paint assay’.

Thirteen plants were chosen at random for further characterization (Figures 5, 6, 7 and 8). The lines #1, #2, #3, #4, #6, #7, #9, #11, and #13 showed resistance to bialaphos in the ‘paint assay’ (Figure 5), confirming the presence and active expression of the bar gene. The lines #5, #8, #10 and #12 were sensitive to bialaphos indicating segregation of the transgene in the T₁ generation.

| Sample # | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 |
|----------|---|---|---|---|---|---|---|---|---|----|----|----|----|
| ‘Paint Assay’ for bialaphos resistance | + | + | + | + | − | + | + | − | + | − | + | − | + |

Figure 5. ‘Paint Assay’ for pYU2996 transformants. ‘Paint assay’ results for sensitivity(−) or resistance(+) to the herbicide bialaphos indicating the absence or presence, respectively, on segregating T₁ plants shown above. Lines #5, 8, 10 and 12 are sensitive to bialaphos and negative for the bar gene by PCR (see Figure 6) of 13 plants. The other lines #1, #2, #3, #4, #6, #7, #9, #11, and #13 show resistance to bialaphos, indicating expression of the bar gene, which is confirmed by PCR results in Figure 6.

Molecular characterizations were performed using PCR for bar and citrine (Figures 6 and 7, respectively) analysis. PCR analysis was conducted for all T₁ plants.
in this study using bar and citrine primers (Figure 6, 7. Also see Table 2 a, b; Appendices). PCR analysis for the bar gene should result in an expected size for the product of 513 bp. Plants #5, #8, #10, and #12 are negative for the presence of the bar sequence. Plants #1, #2, #3, #4, #6, #7, #9, #11, and #13 have positive bands with the expected molecular weight of 513 bp. The pattern of PCR results is for all samples consistent with the bialaphos ‘paint assay’ results shown in Figure 5. The PCR analysis of the citrine gene cassette has an expected size of 421 bp. The pattern of results is consistent with the bialaphos ‘paint assay’ results shown in Figure 5 and the PCR results for the bar gene shown in Figure 6. Plants #5, #8, #10, and #12 are negative for the presence of the citrine sequence. Plants #1, #2, #3, #4, #6, #7, #9, #11, and #13 all have positive bands.

In both PCR tests, the positive control plasmid has a strong band and non-template control H2O does not have any band. These results show that the PCR assay is valid and without any contamination or artifacts. Also, comparison between the PCR results for the bar gene (Figure 6) and citrine gene (Figure 7) show that plants #1, #2, #3, #4, #6, #7, #9, #11, and #13 have positive bands, demonstrating the presence of both the bar and citrine genes. Plants #5, #8, #10, #12 were PCR negative for both bar and citrine genes and probably the result of segregation in the T1 generation.
Figure 6. PCR result for the presence of the bar gene cassette. The expected product is 513bp. The pattern of results is same with the bialaphos ‘paint assay’ results shown in Figure 5. Plants #5, #8, #10, and #12 are negative for the presence of the bar sequence. Plants #1, #2, #3, #4, #6, #7, #9, #11, and #13 have positive bands. The DNA ladder serves as a PCR product size ruler. The plasmid skl construct as positive PCR control (+C). The H₂O lane is the non-template control to determine the presence of contamination in PCR reaction reagents.

Figure 7. PCR result for the presence of the citrine gene cassette. The expected product is 421bp. The pattern of result is same with the bialaphos ‘paint assay’ results shown in Figure 5 and the PCR results for the bar gene shown in Figure 6. Plants #5, #8, #10, and #12 are negative for the presence of the citrine sequence. Plants #1, #2, #3, #4, #6, #7, #9, #11, and #13 have positive bands. The DNA ladder serves as a PCR product size ruler. The plasmid skl construct as positive PCR control (+C). The H₂O lane is the non-template control to determine the presence of contamination in PCR reaction reagents.

Southern blots and expression of the citrine CDS further confirm the integration of the T-DNA cassettes (Figures 8 and 10, respectively). Southern blots were conducted to re-confirm the presence of the transgene and also to determine the transgene copy number in T₁ plants. The bar primer (See Table 2 a, b; Appendices) was used to produce probe, so the bar sequence on the membrane would be targeted during the hybridization to determine integration of the transgene. As previously shown by the ‘paint assay’ results and the PCR analyses, plants #5, #8, #10, and #12 are negative for the presence of the bar sequence and plants #1, #2, #3, #4, #6, #7, #9,
#11, and #13 have positive bands. All positive samples show the single bar gene copy insert in the blot result. Therefore, the T₁ transgenic plants have a single SK1ΔSVL:Citrine:SVL construct copy. This result rules out the possibility of anomalous expression of the transgene which is often observed in transgenic plants with multiple gene copies. The black exposed dots in this blot are unexpected background but do not weaken interpretation of the positive results.

Figure 8. Southern blots for determination of bar gene copy number. The molecular weight ladder (DIG) is from Roche DIG Southern blot kit. In this blot, the bar primer was used to produce the probe. The targeted band in each sample line is the bar sequence, and the number of bands for each sample is the corresponding copy number.

A Zeiss water immersion confocal microscope was used to determine the expression of citrine in T₁ plants (Figures 9 and 10). E. coli expressing citrine and non-transformed E. coli were used as controls for confocal microscope imaging shown in Figure 9.
Figure 9. Confocal images of *E. coli* expressing citrine (A) and *E. coli* negative control (B)
A: Citrine is expressed in *E. coli* serving as a convenient positive control for imaging citrine in the confocal microscope. B: The *E. coli* without expressing Citrine served as negative control, it does not have any yellow fluorescence.

Figure 10. Confocal images of sorghum expressing citrine (A) and sorghum negative control (B) Confocal images of PCR positive (A) and PCR negative plants (B) from a segregating population in etiolated leaves show citrine positive structures consistent with the size of peroxisomes. Note that the negative control however does show some background autofluorescence.

Figure 9A shows *E. coli* expressing citrine and Figure 9B shows the non-transformed *E. coli* negative control. These results confirm that the confocal microscope is capable of detecting citrine expression. Young leaves were collected
from etiolated plants that were PCR positive for *citrine* and negative control plants. The *citrine* PCR positive leaves show *citrine* positive structures consistent with the size of peroxisomes (Figure 10A). The PCR negative samples also show some yellow fluorescence in background (Figure 10B), which may be autofluorescence. However, clearly these samples show stark differences. These putative results therefore need further investigation, but indicate that the *citrine* cassette is being expressed in plants positive for the presence of the transgene.

The presence of the transgene was confirmed in all PCR positive plants. However, there were no phenotypic differences between PCR positive and negative plants when their inflorescences were compared (Figure 11). The PS were aborted in PCR positive inflorescences (Figure 11A-D), and the SS had mature fruits. This phenotypic result of PCR positive (Figure 11A-D) is almost the same compared with PCR negative plants (Figure 11E-H). When the whole size of the PCR positive inflorescence (Figure 12A) was compared to the PCR negative (Figure 12B) equivalent, there was no obvious phenotypic differences between their morphological architecture. The positive plants do not exhibit a protected phenotype for the pedicellate flower, and both have approximately the same number of seeds. These results indicate that the same construct used in maize (Hayward et al. 2016) does not confer a phenotype in sorghum.
Figure 11. Dissecting light scope images of pYU2996 SK1ΔSVL:Citrine:SVL PCR positive inflorescence (Top row) and negative control (Bottom row). Developmental floral morphology of PCR positive plants (A-D) compared with PCR negative plants (E-H) show no phenotypic differences in the development of the pedicellate (PS) or sessile (SS) flowers, and are equivalent to those of non-transgenic plants.
Figure 12. The inflorescences of pYU2996SK1ΔSVL:Citrine:SVL positive plants (A) and pYU2996SK1ΔSVL:Citrine:SVL negative plants (B). The inflorescences of the positive plants (A) compared with their negative segregating controls (B) show no phenotypic differences in morphological architecture. The positive plants do not exhibit a protected phenotype for the pedicellate flower, and both have approximately the same number of seeds.

**Discussion**

Phenotypic analysis (paint assay) and genotypic analysis confirmed the presence of the Hayward et al. (2016) construct in T1 transgenic sorghum plants. After the pYU2996SK1ΔSVL:Citrine:SVL constructs integrated into the sorghum genome it conferred bialaphos resistance from the constitutively expressed *bar* gene. Therefore, the plants showing sensitive results by the ‘paint assay’ indicates that these plants are not expressing the *bar* gene and are likely non-transgenic, and lacking the T-DNA
insert after T₁ segregation. The resistant plants indicate that they are transgenic sorghum lines with at least one functional bar gene. The PCR bar and citrine positive T₁ plants further confirmed the ‘paint assay’ results. The single copy number of skl construct in T₁ plants was confirmed by southern blot analysis. The expression of citrine was detected via confocal microscope, and the PCR positive T₁ plants do express strong fluorescence. Since the skl is fused with citrine (Figure 4), the expression of citrine in the PCR positive T₁ plants also indirectly indicated the expression of skl. All of these tests indicated the presence and expression of the pYU2996SK1ΔSVL:Citrine:SVL construct. The skl construct was hypothesized to protect the pedicellate flower via promoting the expression of skl in sorghum. The overexpression of the maize skl construct did protect the female primordia in maize, but it had no apparent impact on the protection of the pedicellate flower in sorghum. The PS were still aborted in PCR positive plants, and were phenotypically identical to the PCR negative plants. There are several possible explanations for the observed results. First, maize is monocious; its floral development is different with sorghum. The maize genome is highly homologous with sorghum, but the maize skl gene product may not function in the sorghum jasmonic acid pathway. In addition the skl orthologue in sorghum is only 72% homologous to the maize skl gene at the protein level. However, similar experiments with constructs from other plants used in transgenic experiments, even from distantly related plants, have been successful (Kausch and Altpeter, personal communication) in conferring the expected phenotype. Therefore, constitutive overexpression of maize skl may not be able to protect the pedicellate flower in sorghum transformed with the pYU2996SK1ΔSVL:Citrine:SVL
construct. Secondly, even though the corresponding genotypic and phenotypic analysis had confirmed the presence and expression of the maize sk1 construct in transformed sorghum, the expression level of sk1 could be insufficient to protect the pedicellate flower. Third, the sorghum variety used in this project was cv BTx430, and the expected phenotype may require a different sorghum line having the appropriate genetic background.

In conclusion, the desired transgenic lines containing a heritable and functional version of maize sk1 were successfully produced and analyzed. The maize sk1 may not be able to protect pedicellate flowers in sorghum based on the above described possibilities. However, all of the T₁ plants were cultivated from single transgenic event #1. The possibility exists that the inserted sk1 construct may be not complete and identical to the original construct because of some occasional insertions or deletions within the sk1 construct. This possibility could explain the inability of the inserted construct to produce the expected phenotype. In the follow-up work, similar analyses as applied in this current study may need to be done on T₁ plants from different transgenic events. Also, western blots could be used to further confirm definitively the expression of sk1. Future studies should also focus on the use of genome editing to create knockouts not only of sk1 but other candidate genes such as the sorghum msd1 gene.
CHAPTER 2: THE DEVELOPMENT OF TRANSGENIC SORGHUM WITH CAS9
Chapter 2: The Development of Transgenic Sorghum with Cas9

Introduction

Sorghum (*Sorghum bicolor*) represents the third largest cereal crop in the world and the fifth largest in the US (Belide et al. 2017; Jiao et al. 2018). The sorghum grain is widely used for food, animal fodder and biofuels (Jiao et al. 2018). Any agronomic improvements for sorghum will contribute toward enhancing the value of this important crop. Modern approaches to enhance traditional breeding and selection strategies, including genetic transformation strategies and associated genome editing approaches, are necessary and important tools to contribute toward these improvements. Recently, a new gene editing tool has been developed, which is termed Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) along with CRISPR-associated proteins. The modified CRISPR/Cas9 system is much simpler to design than previous genome editing approaches such as Transcription Activator-Like Endonucleases (TALENs) or Zinc Finger Nucleases (ZFNs). It is easier to construct CRISPR vectors than was previously possible because CRISPR relies on precise nucleotide base pairs in contrast to the less precise protein binding domains of TALENs and ZFNs. The induced sgRNA and Cas9 protein can make multiple double-stranded breaks simultaneously, resulting in enhanced mutagenesis and gene editing. The CRISPR/Cas9 system has been applied in several plant genome editing projects to date (Bortesi and Fischer 2015; Kausch et al. 2019). For example, a nicotine-free and nontransgenic tobacco has been developed via CRISPR/Cas9 editing (Schachtsiek and Stehle 2019).
In the project described here, transgenic sorghum lines were developed to constitutively express \textit{bar}, \textit{gfp} and \textit{TaCas9}. This stable transgenic line would provide a valuable tool for future genome editing projects in sorghum since only the guide RNAs would need to be introduced. The development of transgenic sorghum lines with constitutively expressing \textit{TaCas9} would be an important tool for producing other improvements of sorghum through targeted genetic modification. This project focuses on the generation and molecular characterization of the transgenic events for the \textit{TaCas9} containing construct. The long term goal of this project is to verify the efficiency of stable Cas9 expression for generating edited events in future transgenic sorghum lines.

Two gene constructs are involved in this project and made by the Voytas lab at the University of Minnesota: pNG111-ZmUbi::TaCas9_PvUbi::1GFP; and pNG108PvUbi::1GFP [See Appendices Figure 1 and 2]. The genes in the vector for \textit{TaCas9} in the T-DNA region are driven by the maize ubiquitin promoter to provide for constitutive expression [Figure 13 and 14]. The T-DNA in pNG111-ZmUbi::TaCas9_PvUbi::1GFP is described (Figure 13). Toward the 3’ end of the cassette insert, the \textit{bar} gene is driven by the constitutive ubiquitin promoter, PvUbi2, from \textit{Panicum virgatum} (Pv; switchgrass). This promoter is ligated along with the 5’ untranslated region (UTR) and the PvUbi2 intron1 to the \textit{bar} gene and the CMV 35S termination sequence as the selectable marker. This construct should constitutively express bialaphos resistance. Toward the 5’ end of the cassette, the \textit{gfp} gene is driven by the PvUbi1 promoter which is fused to the 5’ untranslated region and the PvUbi1 intron1. The \textit{mGFP} coding sequence (CDS) and the Psrbc S E9 termination sequence
serve as a constitutively expressed visible marker. The TaCas9 cassette is inserted between the bar and gfp cassettes and consists of the maize (Zm) ubiquitin promoter fused to the 5’ UTR and the ZmUbi1 intron1 ligated to the TaCas9 CDS gene and the heat shock protein (HSP) termination sequence. This cassette to be designed to constitutively drive TaCas9 expression. This entire, intact construct should constitutively express bar, gfp, and TaCas9. The pNG111 vector has the TaCas9 region, which is required to evaluate the efficiency of stably expressed TaCas9. The pNG108 vector serves as a control, lacking the TaCas9 construct. Both vectors contain both bar and gfp also driven by a constitutive promoter, namely the Ubiquitin promoter from switchgrass (Panicum virgatum). The selectable marker in both constructs, pNG111 and pNG108, is the bar gene conferring resistance to the herbicide bialaphos. The expression of gfp is used as a visible marker to detect the presence of the cassette. The pNG108 has the bar and gfp region without the TaCas9 sequence, and serves as negative control for pNG111. The same cassettes for pNG111 is described (Figure 13). Both bar and gfp expression were used in pNG108. Once the stable transgenic sorghum lines are developed and characterized containing both constructs, the designed guide RNAs will then be delivered into plants via particle bombardment and tested in future experiments. The efficiency of editing can be evaluated and quantified based on the degree of gene modification success by CRISPR/Cas9.
Figure 13. T-DNA in pNG111-ZmUbi::TaCas9_PvUbi::1GFP. To the left of the cassette insert, the bar gene is driven by the constitutive ubiquitin promoter from Panicum virgatum (Pv; switchgrass) PvUbi2 (pale blue arrow) which is fused to the 5’ untranslated region (UTR in dark yellow), and the PvUbi2 intron1 (pale green) ligated to the bar gene (BAR in maroon) and the CMV 35S Termination sequence (in purple) as the selectable marker. To the right, the gfp gene is driven by the PvUbi1 promoter (deep blue arrow), which is fused to the 5’ untranslated region (UTR in dark yellow), and the PvUbi1 intron1 (orange) ligated to the mGFP coding sequence (CDS) gene (mGFP CDS in green ) and the Psrbc S E9 Termination sequence (in brown) as a visible marker. The TaCas9 cassette is inserted between the bar and gfp cassettes and consists of the maize ubiquitin promoter (ZmUbi in grey) which is fused to the 5’ untranslated region (UTR in dark yellow), and the ZmUbi1 intron1 (pale yellow) ligated to the TaCas9 CDS gene (in pale orange) and the heat shock protein (HSP) Termination sequence (in brown) to code for the which encodes the Cas9 protein. This construct should constitutively express bar, gfp, and TaCas9.

Figure 14. T-DNA in pNG108PvUbi::1GFP. This construct will serve as a negative control for previous pNG111, it does not have the TaCas9 cassette. The same cassettes for pNG111 as described in Figure 13 for both bar and gfp expression were used in pNG108.

Methods

Agrobacterium-mediated transformation

Standard Agrobacterium-mediated transformation of sorghum cv BTx430 was conducted using immature embryos 12-14 days post-pollination as explants. The standard protocols for sorghum transformation were followed (Nelson-Vasilchik et al.)
The specific media specifications are shown in the Appendices (Table 1). The plasmids for pNG111 and pNG108 were independently isolated from their *E. coli* cloning vectors and transferred to the *Agrobacterium* strain AGL1 for sorghum transformation. Transformed AGL1 colonies were selected on YEP agar plate with antibiotic kanamycin and rifampicin, and grown following standard protocols (Nelson-Vasilchik et al. 2018). The isolated colonies were used to grow overnight cultures which were inoculated into infection media for transformation. These cultures were used to inoculate wild type sorghum cv BTx430 immature embryos which are oriented abaxial side up. Incubation with *Agrobacterium* harboring either pNG111 or pNG108 was for 3 days at 28°C in the dark. These embryos were then transferred to resting medium for 14 days, lacking the herbicide bialaphos as the selective agent for the presence of the *bar* gene. Prior to selection, this medium promotes development of somatic embryos, an essential central criterion for successful transformation in sorghum (Kausch et al. 2019). The transferred embryos were selected from non-transgenic cells on bialaphos selection medium. The resistant callus would subsequently be transferred to media to promote somatic embryo germination and then shoot growth and eventually transferred to rooting medium. These plants were then finally transferred to Plant Cons, and finally to a soil medium (Metro-mix).

**Paint Assay**

A ‘paint assay’ was performed on the T₀ plants as described in Chapter 1 of this thesis to evaluate the presence and expression of the *bar* gene. The ‘paint assay’ is non-destructive and allows accurate identification of resistant and sensitive plants. Wild type BTx430 plants were used as controls. Transgenic T₀ individuals would
show phenotypic resistance on the swabbed region indicating bar gene expression. The regenerated plants were grown to maturity in the greenhouse and selfed in order to recover T₁ seed.

Genomic DNA extraction

DNA was isolated from T₀ plants for the molecular analysis and transgene presence confirmation (Chen and Dellaporta 1994). Leaf tissue was collected from T₀ plants. Purified DNA samples were prepared for PCR and Southern blots.

PCR

The polymerase chain reaction (PCR) was used to determine the presence of bar, gfp, and TaCas9 cassettes in T₀ plants using protocols described in Chapter 1 of this thesis. The primer information for the three genes in pNG111 is shown in the Appendices (Table 2; a, c, d). The primers for the bar and gfp genes in pNG108 are the same as those used for pNG111. Because the Tm for all three primers is approximately 60.0° C and their product sizes are shorter in the range of approximately 500 bp, a 55° C degree annealing temperature was used with a 30 second elongation time for 35 cycles for the thermocycler program.

Southern blot

Southern blot analysis on T₀ plants was performed as described in chapter 1 of this thesis. The bar primers were used to produce the DIG- labeled probe for the hybridization step.

GFP Microscopy
GFP expression was evaluated on a Zeiss Discovery v20 microscope with the magnification 10-409 and mGFP 470 filters to detect the GFP expression in the T₀ plants. Wild type sorghum served as negative control.

**Results**

Multiple T₀ transgenic events for the two constructs, pNG111 and pNG108, were produced and successfully grown to maturity to yield T₁ seed. As Table 1 shows, pNG111 produced 6 independent events with a total 32 plants, and pNG108 produced 1 event with a total 7 plants. Transformation experiments with pNG108 that are still in progress to generate additional independent events.

The process of stable plant transformation for sorghum requires significant experience especially with the tissue culture steps. As shown (Figure 15), colonies of resistant calli were recovered and often showed recalcitrance during the plant regeneration steps (see Table 1). All experiments with pNG111 and pNG108 were conducted using bialaphos as a selection agent for the *bar* gene selectable marker (see Table 1 in Appendices). Molecular analysis (described below) revealed that there were no ‘escapes’ in these experiments. Escapes would be putative transformants which do not test positive for the *bar* gene. Developing transgenic calli grew well under selection but often showed a decline on regeneration medium. The transformed pNG108 calli were developing well-formed somatic embryos on regeneration medium (Figure 15 A).
Variety | Construct | # of Events | Total transgenic plants | # of Embryos | Frequency
---|---|---|---|---|---
BTx 430 | pNG111 | 6 | 32 | 734 | 0.82%
BTx 430 | pNG108 | 1 (in process) | 7 | 499 | 0.2%

Table 1. Sorghum transformation results with pNG111 and pNG108. The pNG111 lines have been developed and T₁ plants have been analyzed. The pNG108 lines are still in process.

Transformed calli with pNG111 showed similar signs of decline on regeneration medium (Figure 15 A and C). This is typical of sorghum transformation and does not appear to be construct-specific.

The ‘paint assay’ confirmed the resistance to bialaphos in all T₀ plants. All of the plants that are bialaphos resistant reflect the expression of the bar gene in transgenic lines with no sensitive escapes (Figure 16). In general, all samples were confirmed to be transgenic, by the ‘paint assay’, PCR and Southern blots.

| Sample ID | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
|---|---|---|---|---|---|---|---|---|---|---|
| Bialaphos-Resistant | + | + | + | + | + | + | + | + | + | + |

Figure 16. ‘Paint assay’ for pNG111&pNG108. ‘paint assay’ results on representative T₀ plants transformed with the pNG11) vector and swabbed with 3% bialaphos.
PCR analysis was conducted on all T0 plants for the *bar*, *gfp* and *TaCas9* genes (Figure 17-19). The non-template controls for all three analyses (+C lane in Figures 17-19) show clean results indicating that all positive bands are valid without any contamination.

PCR analysis for the *bar* gene cassette in T0 transgenic sorghum transformed with pNG111 confirmed the presence of the *bar* gene (Figure 17) using *bar* primers (see Appendices, Table 2). The expected PCR product size for *bar* is 513 bp. All of the tested samples were positive.

![Bar gel](image)

**Figure 17.** PCR analysis for the *bar* gene cassette in T0 transgenic sorghum transformed with pNG111 confirmed using *bar* primers (see Appendices). The expected PCR product size for *bar* is 513bp. DNA ladder serves as PCR product size ruler. +C is plasmid serves as positive control. H2O is non-template control to determine the presence of contamination in PCR reaction reagents.

These same plants were also tested for the presence of *gfp* and *TaCas9* (Figure 18 and 19, respectively). The expected PCR product size for *gfp* is 234bp. The expected PCR product size for *TaCas9* is 363bp. These results show that all of the pNG111 T0 plants exhibit the presence of *bar*, *gfp* and *TaCas9* by PCR (Figure 17, 18, 19). For the pNG108 event, PCR validates the presence of both the *bar* and *gfp* genes. The pattern of those PCR results are consistent with the ‘paint assay’ results and show the presence of the *bar* and *gfp* genes and negative for *TaCas9*.
The PCR results clearly show the presence of the introduced vector for all T₀ plants for the *bar*, *gfp* and *TaCas9* genes (Figure 17-19). Southern blot analyses were conducted on these same plants to determine transgene copy number (Figure 20). The *bar* primer was used to produce the probe to hybridize to the digested genomic DNA on the membrane. Samples #1, #2, #3, #4, #5, #6, #9 and #10, all indicate single gene insertion events when probed for the *bar* gene. Plant #7 and #8 indicate multiple insertion copies, containing at least 8 *bar* gene copies. There is no apparent phenotypic consequences in these plants in comparison to the others or wild-type plants. Background noise, appearing as black spots are associated with these blots but do not alter or interfere with the analysis.
Transgenic plants which had been previously shown to have the bar, gfp, and TaCas9 genes from pNG111 were used to observe GFP in T0 and T1 plants. The presence of the gfp cassette had been confirmed by PCR (Figure 18). The gfp gene is driven by the PvUbi1 promoter which is fused to the 5’ untranslated region (UTR) and the PvUbi1 intron1 ligated to the mGFP coding sequence (CDS) gene and the Psrbc S E9 Termination sequence and serves as a constitutively expressed visible marker. The expression of the gfp gene was detected using a Zeiss Discovery v20 microscope.
Root tips collected from T₀ plants were used to test for GFP expression (Figure 21A). T₁ immature embryos growing in the panicles of selfed T₀ plant also indicate a positive result for the GFP expression in segregating plants (Figure 21B). Root tips collected from wild type sorghum served negative control (Figure 21C.), show no fluorescence from GFP or autofluorescence.

Figure 21. GFP expression analysis. A. Root tip collected from a transgenic T₀ plant which had been previously shown to have the bar, gfp, and TaCas9 genes from pNG111 observed using a Zeiss Discovery v20 B. T₁ immature embryos were harvested from mature T₀ transgenic plants with the pNG108 vector showing positive GFP fluorescence. C. Root tip from a non-transformed wild type BTx430 sorghum plant shows no GFP fluorescence or autofluorescence and serves as negative control.

**Discussion**

The significance of cereal crops to global agriculture, the economy, food security and international stability is well documented and widely understood. With the dramatic increase of the human population over the previous three decades, many consequences have been observed, including; climate change resulting in droughts, floods and fires, loss of habitat and a decrease in available arable land, decrease in water availability, resulting in a threat to global food security. In addition there has been a rise in the consumption of many natural resources including energy, resulting for a need to increase research on renewable bioenergy (Belide et al. 2017). Sorghum is a significant crop globally for food feed and bioenergy. In addition, the functional
development of a genome-level knowledge base linking genes to phenotypes through
the use of transgenics in sorghum is critical to understanding fundamental
physiological functions important to crop improvement. Therefore, the capability to
create, test and cultivate transgenics has enabled some of the most innovative and
important scientific discoveries and agricultural achievements over the last three
decades.

Thus, sorghum transformation for crop improvement is central to future
agricultural enhancement. The goal here was to produce a transgenic sorghum line that
would be used to test whether stably expressed TaCas9 would be useful for future
genome editing functions using CRISPR sgRNAs in subsequent transformations. To
address this problem, transgenic sorghum lines were produced using a vector that
contained cassettes to express the bar, gfp and TaCas9 genes (pNG111) and a second
vector (pNG108) containing only the bar and gfp genes as a negative control for the
TaCas9.

Stable transgenic sorghum lines were developed using Agrobacterium-
mediated transformation using the pNG111 and pNG108 vectors and bialaphos
selection. The plants were grown to maturity under greenhouse conditions and selfed
to produce T1 seeds. The frequency of sorghum transformation with these two
constructs is quite low and there are still experiments in process with the pNG108
vector to increase the number of independent events. This inefficiency in production
of the desired outcome is explained primarily by the long term protocols for
transformation (9-12 months to T1 seed). The regeneration frequency for the two
constructs are less than 1% (see Table 1). This is not an unusual situation in sorghum
transformation biology and is probably not construct-specific. Sorghum is referred to as a recalcitrant plant for plant transformation. The transformation protocol for sorghum exhibits low efficiency yet highly reliability. If enough effort is put into these experiments adequate numbers of transgenics will be recovered. The low frequency of successful transformation indicates that the protocol of plant transformation still needs improvement.

The transformation procedure used for sorghum in the research presented in this thesis requires using cv BTx430, because of collaborators restrictions, and selection with the bar gene for bialaphos resistance. The research presented here show that the bar gene does confer resistance to bialaphos during the selection phase of the transformation process without any escapes. Also, this research show the resistance to bialaphos in the ‘paint assay’ is valuable for confirming the presence of bar gene expression. These results were confirmed by PCR for the bar gene with 100% fidelity.

During this research the pNG108 and pNG111 constructs were successfully introduced into sorghum and mature fertile plants were recovered. The PCR results for pNG111 clearly show the presence of the introduced vector on all T0 plants for the bar, gfp and TaCas9 genes, with the one exception for event #1 using pNG111. While all samples tested positive for the bar gene, event #1 for pNG111 was also PCR positive for TaCas9 genes, but this event tested negative for gfp. The most reasonable explanation for this anomaly is that the gfp gene was truncated during transformation. As shown (Figure 13), the mGFP CDR is located near the 5’ end of the T-DNA and close to T-DNA left border. This position has been shown be susceptible to deletion during T-DNA integration (Che et al. 2018). The Southern blot analysis showed that
the majority of events are single gene copy integrations, while two show multiple copy insertions. This is significant because previous studies have shown expression and inheritance complications in plants with multiple copy insertions (Belide et al. 2017). For this reason only events with single gene copy insertions will be used in future phenotypic analysis. The results for pNG108 show all T₀ plants for event #1 for the *bar, and gfp* genes as expected. Transformation experiments using pNG108 are also still in process to increase the number of events.

The expression of TaCas9 will need to be confirmed by Western blot analysis in future evaluations of these lines. After the molecular confirmation of presence and expression of *TaCas9* gene cassette, guide RNAs will be designed by our collaborators in the Voytas Lab at the University of Minnesota and delivered into T₁ immature embryos via particle bombardment. The transferred guide RNAs would be used to quantitatively determine the efficiency of CRISPR/Cas9 in sorghum.

These lines will be used in future experiments to characterize the efficiency of genome editing in the presence of stably expressed TaCas9. For example, one experiment would be to use the pNG111 line to determine the frequency of conversion of the *gfp* sequence to *hfp*. This conversion requires two single amino acid changes (Glaser et al. 2016) and could be visualized using confocal microscopy and quantitated (see Figure Legends in Supplemental Information). Despite tremendous improvements in plant transformation in recent years (Lowe et al. 2018, 2019) sorghum transformation remains a major bottleneck and is still far from routine (Altpeter et al. 2016; Kausch et al. 2019). The procedure for sorghum transformation is labor and material expensive and requires significant laboratory expertise. Recently a program
has been established to focus specifically on transformation technology improvement across a wide range of species (Gordon-Kamm, personal communication) with the overall goal to “bring transformation to the masses”. This program seeks to develop protocols that will allow any researcher in capable laboratories to conduct routine plant transformation for research purposes.
## APPENDICES:

### Table 1. Media Specifications Used for Sorghum Transformation

| Sorghum media components (per liter) | Infection | Co-cultivation | Resting | Callus proliferation | R-I | R-II |
|--------------------------------------|-----------|----------------|---------|----------------------|-----|------|
| MS salts                             | 2.15 g    | 2.15 g         | 4.3 g   | 4.3 g                | 4.3 g | 2.15 g |
| MES                                  | 0.5 g     | 0.5 g          | 0.5 g   | 0.5 g                | 0.5 g |
| L-Proline                            | 0.7 g     |                |         |                      | 0.7 g |
| Glucose                              | 36 g      | 10 g           |         |                      |      |
| Sucrose                              | 68.5 g    | 20 g           | 30 g    | 30 g                 | 60 g | 30 g |
| 2,4-D, 1mg/ml                        | 1.5 ml    | 2 ml           | 2 ml    | 1.5 ml               |      |
| Agar                                 | 8 g       |                |         |                      | 8 g  |
| Phytigel                              |           |                |         |                      |      |
| pH(HCl/KOH)                           | 5.2       | 5.8            | 5.8     | 5.8                  | 5.6  | 5.6  |
| Bs(vitamin), 100x                     | 10 ml     | 10 ml          | 10 ml   | 10 ml                |      |
| Acetosyringone, 100 mM                | 1 ml      | 1 ml           |         |                      |      |
| Ascorbic acid                        |           |                |         |                      |      |
| Casamino acids                       | 1 g       |                |         |                      |      |
| Asparagine                           | 0.15 g    |                |         |                      |      |
| Coconut water                        | 100 ml    |                |         |                      |      |
| Timentin                             | 150 mg    | 150 mg         | 150 mg  | 150 mg               |      |
| Zeatin, 1mg/ml                       |           |                |         |                      | 0.5 ml|
| IAA, 1mg/ml                          |           |                |         |                      | 1 ml  |
| ABA, 0.025 mg/ml                     |           |                |         |                      | 1 ml  |
| TDZ, 0.5 mg/ml                       |           |                |         |                      | 0.2 ml|
| IBA, 1mg/ml                          |           |                |         |                      |      |
| NAA, 1 g/ml                          |           |                |         |                      |      |
| MS vitamin, 1000x                    |           |                |         |                      | 1 ml  |


Table 1. The sorghum medium information.

| PVPP (1% final) | 10 g | 10 g | 10 g | 10 g | 5 g |
|-----------------|------|------|------|------|-----|
| Sterilization procedure | Filter | A/C 20 min | A/C 20 min | A/C 20 min | A/C 20 min |

aABA, abscisic acid; 2,4-D, 2,4-dichlorophenoxyacetic acid; IAA, indole-3-acetic acid; IBA, indole-3-butyric acid (auxin); MES, morpholine-4-ethanesulfonic acid; MS salts, Murashige and Skoog basal salt mixture; MS vitamin, Murashige and Skoog basal medium with vitamins; NAA, 1-naphthaleneacetic acid; PVPP, polyvinylpolypyrrolidone; TDZ, thidiazuron.

R: Regeneration; A/C: Autoclave.

Table 2. PCR Primer Specifications Used to Analyze the bar, citrine, gfp, and TaCas9 genes

| Oligo | Length | tm | GC% | Sequence |
|-------|--------|----|----|----------|
| Forward | 20 | 60.03 | 55.00 | GGATCTACCATGAGCCCAGA |
| Reverse | 20 | 60.00 | 55.00 | GAAGTCCAGCTGCCAGAAAC |
| Product Size: 513 |

Table 2 (a). bar primer

| Oligo | Length | tm | GC% | Sequence |
|-------|--------|----|----|----------|
| Forward | 20 | 60.04 | 50.00 | ACGTAAACGGCCACAAGTTC |
| Reverse | 20 | 60.41 | 50.00 | ATGCCGTTCTTCTGCTTGTC |
| Product Size: 421 |

Table 2 (b). citrine primer

| Oligo | Length | tm | GC% | Sequence |
|-------|--------|----|----|----------|
| Forward | 20 | 60.05 | 50.00 | TCAAGGAGGACGGAAACATC |
| Reverse | 20 | 59.97 | 50.00 | AAAGGCGAGATTGTGTGGAC |
| Product Size: 234 |

Table 2 (c). gfp primer

| Oligo | Length | tm | GC% | Sequence |
|-------|--------|----|----|----------|
| Forward | 20 | 60.01 | 55.00 | AGACCGTGGAAGGTTGAC |
| Reverse | 20 | 60.00 | 55.00 | ACCTGGTGAGGACCTTGT |
| Product Size: 421 |

Table 2 (d). TaCas9 primer
Figure 1. Vector map for pNG111

This vector should constitutively express \textit{TaCas9} and \textit{gfp}. This means that we can analyze for \textit{gfp} and assume that the \textit{TaCas9} cassette is also present because they are linked. In addition presence of both \textit{TaCas9} and \textit{gfp} can be verified molecularly in small plants during regeneration. GFP expression can be analyzed through the development cycle of the transgenic lines from callus to plants. In addition, we can design guide RNAs which will edit the \textit{gfp} gene to convert it to \textit{bfp}. The result should appear as blue foci against a GFP background. The frequency of edits can be quantitatively determined. This information will be extremely useful to predict editing frequencies in studies where a visual marker is not involved.
Figure 2. Vector map for pNG108

pNG108- PvUbi1::GFP
This vector should provide a negative control exhibiting constitutive expression of gfp, but without the presence of TaCas9. Therefore when the same guide RNAs (used in pNG111) are delivered, no BFP expression should be observed.
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