

1 **An efficient gene excision system in maize**

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16 **Keywords;**

17 *Agrobacterium*; developmentally-regulated promoters; heat-shock promoters; morphogenic

18 genes; marker-free events; rapid maize transformation

19 **Abbreviations;**

20 *Bbm*: Babyboom, *Cre*: CRE recombinase, HSP: Heat-shock promoters, SMG: Selectable marker-

21 free; Pro: promoters, QE: Quality events, UE: Usable event; *Wus2*; Wuschel2

22

23

1 **ABSTRACT**

2 Use of the morphogenic genes *Baby Boom* (*Bbm*) and *Wuschel2* (*Wus2*), along with new ternary
3 constructs, has increased the genotype range and the type of explants that can be used for maize
4 transformation. In addition, altering the ectopic expression pattern for *Bbm/Wus2* has resulted in
5 rapid maize transformation methods that are faster and applicable to a broader range of inbreds.
6 However, expression of *Bbm/Wus2* can compromise the quality of regenerated plants, leading to
7 sterility. We reasoned excising morphogenic genes after transformation but before regeneration
8 would increase production of fertile T0 plants. We developed a method that uses an inducible
9 site-specific recombinase (*Cre*) to excise morphogenic genes. The use of developmentally
10 regulated promoters, such as *Ole*, *Glb1*, *End2* and *Ltp2*, to drive *Cre* enabled excision of
11 morphogenic genes in early embryo development and produced excised events at a rate of 25%-
12 100%. A different strategy utilizing an excision-activated selectable marker produced excised
13 events at a rate of 53.3%-68.4%; however, the transformation frequency was lower (12.9%-
14 49.9%). The use of inducible heat shock promoters (e.g. *Hsp17.7*, *Hsp26*) to express *Cre*, along
15 with improvements in tissue culture conditions and construct design, resulted in high frequencies
16 of T0 transformation (29%-69%), excision (50%-97%), usable quality events (3.6%-14%), and
17 few escapes (non-transgenic; 14%-17%) in three elite maize inbreds. Transgenic events produced
18 by this method are free of morphogenic and marker genes.

1 INTRODUCTION

2 The use of the morphogenic genes *Bbm* and *Wus2* has considerably increased
3 transformation frequencies and reduced genotype dependence in many cereal crops (Lowe et al.,
4 2016;Mookkan et al., 2017;Anand et al., 2018;Lowe et al., 2018). This enabled the development
5 of a rapid transformation method involving direct formation of somatic embryos and T0 plants
6 from immature scutella (Lowe et al., 2018). This approach has facilitated transformation (Lowe
7 et., 2016; Mookkan et al.,2017) and CRISPR/Cas-mediated editing (Chilcoat et al., 2017) in
8 numerous elite maize inbreds, and enabled use of alternate explants, such as embryo slices from
9 mature seeds or leaf segments, for successful maize transformation (Lowe et al., 2016;Lowe et
10 al., 2018). However, ectopic expression of the morphogenic genes often resulted in pleiotropic
11 effects including abnormal shoots/roots and infertile plants (Lowe et al., 2016). The use of
12 promoters that drive high expression levels during the transformation process, but lower
13 expression levels in the vegetative plant, provides one option to ameliorate these problems
14 (Lowe et al.,2018) but the presence of morphogenic genes can still result in some negative effects
15 and is undesirable in commercial products. While fertile T0 plants can be recovered under these
16 conditions, non-visible pleiotropic effects remain a distinct possibility. Similarly, transgenic
17 plants regenerated through *de novo* meristem induction stimulated by morphogenic gene
18 expression also resulted in developmental abnormalities (Maher et al., 2020), and without
19 removal also raise concerns that non-visible pleiotropic effects are possible. Therefore, excising
20 the morphogenic genes is desirable for regenerating healthy plants, for transgene testing and
21 commercial product development. Previously a method using a non-integrating *Wus2* gene
22 expression approach recovered fertile T0 plants free-off morphogenic genes, however this
23 method needed a plant selectable marker gene (SMG) for regenerating events (Hoerster et al.,

1 2020). Here we report an approach that allows excision of both the morphogenic gene and the
2 SMG used in transformation at the same time. As an added benefit this method eliminates any
3 adverse effect from the non-trait genes in commercial products.

4 Different strategies have been developed for the removal of helper genes following plant
5 transformation, often focused on removing plant selectable markers. One approach is co-
6 transformation with two constructs, one with the SMG and one with the gene of interest. In a
7 transgenic plant with independent insertions of each of these constructs, the selectable marker
8 can be segregated genetically (Hare and Chua, 2002;Puchta, 2003;Darbani et al., 2007;Ling et
9 al., 2016). Alternatively, SMGs can be removed by excision via homologous recombination
10 (Puchta, 2000;Zubko et al., 2000), elimination by transposition (Maeser and Kahmann, 1991;Gao
11 et al., 2015) or, by the use of recombinases to excise unwanted DNA. Several recombination
12 systems have been used to excise SMGs, including *Cre/lox* from bacteriophage P1 (Hoess et al.,
13 1982;Hoess and Abremski, 1985), *Flp/frt* from *Saccharomyces cerevisiae* (Cox, 1983;Senecoff
14 et al., 1985), R/RS from *Zygosaccharomyces rouxii* (Araki et al., 1985), and *Gin/gix* from
15 bacteriophage (Klippel et al., 1988). Recombinases have been delivered via retransformation
16 (Odell et al., 1990;Dale and Ow, 1991), sexual crosses (Bayley et al., 1992;Kilby et al.,
17 1995;Kerbach et al., 2005), or transient expression (Gleave et al., 1999;Kopertekh et al.,
18 2004;Kopertekh and Schiemann, 2005;Jia et al., 2006). In most of these systems excision takes
19 place after the T0 generation and requires screening multiple plants to find one that has
20 undergone successful excision. A design where the SMG and the recombinase genes are on the
21 same construct between the recombination sites has been referred to as “auto-excision”
22 (Verweire et al., 2007;Moravčiková et al., 2008), and allows generation of SMG-free events. By
23 placing the recombinase under the regulation of an inducible/chemical promoter, an expression

1 system that allowed spatial and temporal control (regulated by external or intrinsic signals) was
2 shown to be faster and less resource-intensive (Chong-Pérez and Angenon, 2013;Yau and
3 Stewart, 2013).

4 We have evaluated three different strategies for auto-excision prior to regeneration to
5 recover stable T0 plants free of morphogenic genes and in some cases the SMG as well: 1) an
6 auto-excision system involving developmentally regulated promoters, 2) an excision-activated
7 marker gene system, and 3) an inducible promoter approach for excising both the morphogenic
8 genes and the SMG. The excision strategies were evaluated to meet key production
9 transformation criteria of 1) high transformation frequency, 2) high quality event (QE, single-
10 copy of T-DNA, backbone and morphogenic gene free) frequency, 3) ability to generate marker-
11 free T0 plants, and 4) applicability to multiple elite maize inbreds. The use of developmentally
12 regulated promoters driving *Cre* enabled auto-excision of morphogenic genes, but resulted in
13 low transformation frequency and QE recovery. These limitations were addressed using heat-
14 shock inducible promoters driving expression of *Cre*, that resulted in higher frequencies of T0
15 transformation, gene-excision and QE recovery.

16

17 **Excision via developmentally-regulated promoters**

18 The presence of morphogenic genes in transgenic events is undesirable because of unpredictable
19 phenotypes (Lowe et al., 2016). Auto-excision of morphogenic genes occurs early in the
20 transformation process which enables trait evaluation in T0 generation and reduces attrition due
21 to T0 sterility. We evaluated several auto-excision designs, using *Cre* driven by various
22 promoters. These included seven different developmentally regulated (embryo or meristem)
23 promoters, the constitutive maize ubiquitin (*ZM-Ubi*) promoter, and the *Agrobacterium* nopaline

1 synthase (*Nos*) promoter (Table 1). To facilitate excision, the morphogenic genes (*Wus2* and
2 *Bbm*) and the *Cre* gene cassette were flanked with a single pair of directly oriented *loxP* sites
3 (Figure 1 A). The resulting excised events following auto excision is depicted in Figure 1B. We
4 evaluated two different inbreds (HC69 and PH2RT) to identify *pro:Cre* combinations that
5 produced high frequencies of both transformation and excision. Molecular event data is
6 presented in Table 2. All constructs tested produced stable transgenic events with some number
7 of properly excised events. The *Ole_{pro}:Cre* had the highest transformation frequencies (27.2%-
8 37.1%), while the *Glb1_{pro}:Cre* construct produced events with higher QE frequencies (8.6%-
9 18.4%).

10

11 **Excision via marker gene activation**

12 Although we achieved auto-excision with all developmentally regulated promoters tested, even
13 for the best construct the usable events rate was around 2% and 80-90% of events were not
14 excised quality events. To improve efficiency, we designed constructs with SMG that was
15 activated only upon excision of the morphogenic genes. This approach selects directly for
16 excised events and was expected to increase QE frequency. A similar construct design was
17 previously used to optimize tissue culture conditions for recovering high quality maize
18 transgenic events (Chu et al., 2019). A schematic design of the construct is depicted in Figure 2A
19 and the quality excised product in Figure 2B. For these experiments, either the *Glb1* or the *Ole*
20 promoters were used to drive *Cre* expression for evaluation of excision-activated marker gene
21 selection. The data from side-by-side testing of these two promoters using the construct design
22 described in Figure 2 are summarized in Table 3. The construct containing *Glb1_{pro}:Cre* improved
23 T0 transformation and QE frequencies (1.8 and 1.4-fold), compared to the developmentally

1 regulated gene-excision approach. When *Ole_{pro}:Cre* was used, the T0 transformation frequency
2 was similar (>1.1-fold) while the QE frequency increased approximately 1.7-fold. The excision
3 frequency was higher when excision-activated selection was used, with excision frequencies of
4 53.3% (*Ole_{pro}:Cre*) and 68.4% (*Glb_{pro}:Cre*) when compared to the previous approach.
5 Additionally, no null events (escapes) were identified by qPCR analysis.

6 The *Glb_{pro}:Cre* construct design was further evaluated in two additional inbreds, PH84Z
7 and PH85E, alongside HC69 for comparison (Table 4). QEs were recovered in all three inbreds,
8 which were free of the morphogenic genes with no escapes. Excision frequency was similar
9 (55%-61%) across all the inbreds; QE frequencies varied by genotype: 8.7% (HC69), 27.7%
10 (PH85E) and 6.7% (PH84Z) leading to differences in usable quality event frequency (UE,
11 quality events per 100 embryos): 4.3% (HC69), 3.6% (PH85E) and 1.9% (PH84Z).

13 **Excision via stress-inducible promoters**

14 To further improve efficiency, a series of stress-inducible promoters were tested for excision of
15 morphogenic genes. The promoters were selected from a set of genes induced by heat (maize
16 *Hsp17.7* and *Hsp26*) and drought (*ZmRab17*, *SiRAB21*, *BdDRP1*, and *BdDRP12*). The construct
17 design is identical to that described in Figure 1, where stress-inducible promoters drive *Cre*
18 expression as represented by *pro:Cre*. The different steps in the transformation process,
19 selection immature embryo infection, In preliminary screening, embryos derived from HC69
20 were infected with one of the six constructs and, subsequently subjected to one of three different
21 conditions: no heat shock (control), heat shock at 37°C for 1 day, or 42°C for 2h/day for 3
22 consecutive days. The different steps in maize immature embryo transformation process,
23 included embryo infection with *Agrobacterium* strain containing the construct (Figure 3A),

1 selection of transgenic events on media supplemented with selectable marker (Figure 3B), heat-
2 shock treatment step (Figure 3C), regeneration of events on media with selection pressure
3 (Figure 3D) and rooting (Figure 3E), before the events were sent to greenhouse. The auto-
4 excision frequencies under induced and non-induced conditions were determined by qPCR
5 analysis. Somatic embryos on maturation media (18 dpi) with 0.1 mg/L imazapyr were subjected
6 to one of the heat conditions and moved onto a rooting media with 0.1 mg/L imazapyr following
7 heat shock (Figure 3D).

8 All promoters except *Hsp26* were leaky under non-induced conditions, resulting in gene-
9 excision rates from 3.4% (*Rab17_{pro}*) to 36% (*BdRab21_{pro}*) compared to zero in the *Cre*-minus
10 construct. For a subset of the promoters (*Hsp1.7*, *Hsp26*, *Drp1* and *Drp12*), higher excision
11 frequencies ranging from 43% to 100%, were observed in the 42°C, 2h/day for 3 days heat
12 treatment. Longer exposure of the somatic embryos at 37°C adversely effected T0 event
13 recovery, compared to a short pulse of heat shock at 42°C (2hr/day for 3 days). Based on the
14 recovery of excised T0 events with *Hsp26_{pro}* construct at 42°C treatment compared to 37°C
15 treatment, this promoter appeared to be induced only at higher temperatures.

16 Additional experiments were performed to further evaluate gene excision and optimize
17 heat shock conditions using three of the inducible promoters (*Hsp17.7*, *Hsp26* and *Drp12*). HC69
18 embryos infected with the three constructs were subjected to heat shock treatment at the
19 maturation stage (Figure 3C). One of three different treatments were applied 1) no heat shock
20 (control), 2) 42°C for 2h and 3) 42°C, 2h on 3 consecutive days to determine frequencies of
21 excision and UE recovery (Table 6). Consistent with the previous observation, *Hsp17.7_{pro}*
22 driving *Cre* expression under both heat treatments resulted in higher excision rates (62.5%-
23 69.2%) resulting in higher UE rates (10 to 18) compared to *Hsp26_{pro}* and *Drp12_{pro}*. Based on the

1 data we identified *Hsp17.7_{pro}* as the preferred promoter for auto-excision with heat shock of
2 42°C for 2h.

3

4 **Optimization of heat-shock conditions to improve auto-excision**

5 Further experiments were designed with *Hsp17.7_{pro}* and *Hsp26_{pro}* to optimize excision
6 conditions. After three weeks of selection, somatic embryos at the maturation stage (Figure 3)
7 were subjected to one of three different heat conditions 1) 42°C, 2h/day for 2 d, 2) 42°C for 24h,
8 or 3) 45°C for 2h/day to determine frequencies of excision and UE. Across the treatments,
9 transformation frequencies ranged from 35%-54.9%, except in the 42°C for 24h treatment of
10 embryos with *Hsp17.7_{pro}* driving *Cre* expression, which was lower (Table 7). The heat
11 treatments increased excision rates, which varied with the conditions applied. Of the two *Hsp*
12 promoters tested, *Hsp17.7_{pro}* resulted in events with higher excision frequency (75% at 42°C for
13 24h and 76.6% at 45°C for 2h) compared to *Hsp26_{pro}* (66.7% and 61.9%). The treatment, 45°C
14 for 2h worked best for both *Hsp* promoters.

15

16 **Concurrent elimination of morphogenic and plant selectable marker genes**

17 Next, we developed a strategy that simultaneously excised both the morphogenic genes and the
18 SMG. Two different SMGs, *HRA* and *NPTII* were tested. The construct design was slightly
19 changed to enable excision of the SMG by including it as part of the excised DNA (morphogenic
20 genes and *Cre*) flanked with a single pair of directly oriented *loxP* sites (Figure 4A) and the
21 resulting excised events are free of SMG (Figure 4B). The binary construct designs with
22 different selectable marker, morphogenic gene and a reporter gene *Zs-GREEN* is illustrated in
23 Figure 4A. Following transformation and selection (either 0.1 mg/L imazapyr for the *HRA* gene

1 or 150mg/L G418 for the *NPTII* gene), the somatic embryos were heat-shock treated at 45°C for
2 2h. Transformation data are presented in Table 8. Both *HRA* and *NPTII* constructs produced T0
3 plants free of morphogenic genes and SMG in the three inbreds tested. With the *HRA* construct,
4 lower frequencies of QEs and UEs were observed and 2-fold more null events were produced
5 compared to the *NPTII* construct. The excision frequency was comparable in both *HRA* and
6 *NPTII* constructs. Irrespective of the differences, both selectable markers produced high
7 frequencies of single copy, backbone-free events which are free of the morphogenic and marker
8 genes.

10 **Progeny analysis**

11 To study the inheritance and segregation of the morphogenic and SMG-free events, we screened
12 single-copy T0 plants free of morphogenic gene and SMG produced from the *NPTII* construct.
13 Thirteen T0 QE plants, six plants from HC69 and seven plants from PHR84Z, were selected for
14 progeny analysis. These plants were selected and self-pollinated in the greenhouse to enable
15 segregation analysis. Plants from all 13 events produced seeds, 100 to 200 seeds per plant. T1
16 plants were evaluated for zygosity using qPCR to evaluate copy number of *Cre* and *NPTII* genes
17 (excised DNA). Twelve of the 13 events showed the expected Mendelian inheritance of a single
18 copy T-DNA integration (1:2:1; chi-square p-value>0.05) in the T1 generation (Table 9).

20 **DISCUSSION**

21 In maize, direct induction of somatic embryos capable of rapidly germinating from immature
22 embryos (without a callus phase) has been demonstrated using the auxin-inducible promoter
23 *Axig1* driving *Wus2* expression in combination with *Bbm* driven by a maize *PLTP* promoter

1 (Lowe et al., 2018). Continued expression of morphogenic genes results in abnormal phenotypes
2 (Lowe et al., 2016). Therefore, removing morphogenic genes is imperative for accurate construct
3 evaluation and product development and, therefore, a prerequisite for broad application of the
4 technology. Morphogenic gene excision was accomplished using a drought-inducible *Rab17*
5 promoter driving *Cre* recombinase expression (Vilardell et al., 1991). Although this approach
6 was used for successful excision, the requirement for a desiccation step significantly reduced
7 stable event recovery and excision frequency (Lowe et al., 2016).

8 In order to develop a more efficient system promoters of seven developmentally
9 regulated genes, the *Knotted-1 (Kn1)* (Bolduc et al., 2012), *Leafy cotyledon1 (Lec1)* (Pelletier et
10 al., 2017), barley *Lipid transfer protein2 (Ltp2)* (Kalla et al., 1994), an early embryo response
11 gene (*End2*) (Casper et al., 2005), *Globulin1 (Glb1)* (Belanger and Kriz, 1991), and *Olesin (Ole)*
12 (Anand et al., 2017b) were evaluated for their ability to express *Cre* and excise morphogenic
13 genes. *Glb1*, *Ole*, and *End2* promoters unlike inducible promoters did not need either physical or
14 chemical induction for auto-excision. While morphogenic gene removal was observed using
15 developmentally regulated promoters, this generally resulted in lower QE frequencies. A
16 possible explanation is that premature expression caused by early unintended low-level
17 expression from the developmentally regulated promoters resulted in low levels of *Cre*
18 expression.

19 Developing a method for regenerating events that are free of morphogenic genes using an
20 excision-activated marker gene system may increase excision frequency and QE recovery is
21 described (Chu et al., 2019). In a similar manner, developmentally regulated promoters *Glb1*
22 and *Ole* that are active during late embryo development (Kriz et al., 1990; Anand et al., 2017b),
23 were used to drive *Cre* expression for auto-excision. This strategy resulted in the reconstitution

1 of the *HRA* marker gene, which conferred herbicide resistance (Chu et al., 2019) and would grow
2 in the presence of selective agent. As anticipated, the strategy resulted in improved frequencies
3 of T0 transformation and QE that resulted in approximately a 2-fold increase in UE production.
4 Despite excision of the morphogenic genes and activation of selectable marker, a large
5 proportion of T0 events were multi-copy and non-excised. One possible explanation is the
6 dosage effect of the *HRA* gene on rapid maize transformation, leading to enrichment of events
7 with stable insertions of more than one copy of the transgene. The other possibility is the
8 restricted activation of the developmental promoters leading to partial/incomplete excision,
9 which does not work in rapid maize transformation for enriching quality events.

10 To achieve controlled expression of recombinases genes for excision, inducible
11 promoters have been an attractive choice. These promoters predominantly fall into two
12 categories; 1) heat shock- or stress-inducible promoters (Kilby et al., 1995;Cuellar et al.,
13 2006;Zhang et al., 2006;Du et al., 2019) and, 2) chemical inducible promoters (Gatz, 1996;Zuo
14 and Chua, 2000). Expressing the recombinase under the control of promoters requiring inducers
15 (heat, osmotic, or chemical) has allowed tighter control of gene expression, while minimizing the
16 negative effect of ectopic gene expression. Among the stress-inducible promoters tested,
17 *Hsp17.7_{pro}* and *Hsp26_{pro}* produced the best results for auto-excision based on a higher frequency
18 of T0 transformation, gene excision and UE rate. In maize, the regulation of *Hsp* promoters in
19 response to stresses has been described (Pegoraro et al., 2011), including accumulation of *Hsp*
20 proteins under temperatures over 32-33°C (Ristic et al., 1991;Vierling, 1991) and enhanced
21 *Hsp70* synthesis under drought and/or heat (Hu et al., 2010). The heat-inducible auto-excision
22 system was previously described using a construct design that involves *Hsp70_{pro}* driving the *Cre*
23 recombinase for elimination of the SMG (egfp) while a second marker gene, expressing the

1 anthocyanin pigmentation (Rsc) gene, was used for event sorting (Du et al., 2019). While
2 successful, the strategy has limited practical application requiring tracking of transgenes in the
3 T1 generation and subsequent segregation, which is resource- and time-intensive.

4 Taking a methodological approach, a system was developed to obtain morphogenic gene-
5 free events at high frequencies (66%-77% of the total events generated). The overall strategy was
6 to develop an efficient auto-excision system for rapid maize transformation, with the objective of
7 eliminating both morphogenic and marker genes, that is highly efficient to meet the needs of
8 high throughput maize transformation. The method we developed resulted in the elimination of
9 morphogenic and marker genes at the maturation stage of transformation at high frequencies
10 (ranging from 60%-97%) in multiple elite inbreds. This was achieved by optimizing tissue
11 culture conditions, optimization of heat shock treatment and identifying a versatile SMG. The
12 stably transformed plants were normal, produced seeds and showed stable transmission of the
13 integrated T-DNA to the next generation.

14

15 **MATERIALS AND METHODS**

16 **Plant Material**

17 Pioneer temperate maize inbreds (R03, PH2RT, PH85E and PH84Z) were used in this study. All
18 plants used for source immature embryos were grown in the greenhouse. One of the inbred lines
19 (R03) is nonproprietary and publicly available. The other three inbred lines described here are
20 proprietary (PH2RT, PH85E and PH84Z). In order to protect Corteva Agriscience proprietary
21 germplasm, such germplasm will not be made available except at the discretion of Corteva
22 Agriscience and then only in accordance with all applicable governmental regulations.

23

1 **Donor material and tissue culture**

2 Seeds were germinated and grown in a greenhouse at temperature set-points of 25.5/20.0°C
3 (day/night), and 16-h daylight. After 21 d, seedlings were transplanted into 5.9 L pots containing
4 a soil-less substrate composed of 38% Canadian sphagnum peat, 51% composted bark, 8%
5 perlite, and 3% vermiculite by volume and adjusted with lime to a pH of 6.0. Maize ears from
6 the Pioneer inbred lines HC69, PH2RT, PH84Z and PH85E were collected from the greenhouse
7 (Johnston, Iowa) at 10 to 11 d after pollination, when the immature embryos were 1.5-2.0 mm in
8 length. Ears were sterilized with 20% Clorox (final sodium hypochlorite concentration of
9 1.65%) for 15 min and rinsed three times with sterile distilled water.

10 **Culture media used for transformations and plant regeneration**

11 Briefly, maize immature embryos (1.5-2 mm) were harvested and used for *Agrobacterium*-
12 mediated transformation, using the media, selection and regeneration methods described
13 previously (Lowe et al., 2018;Chu et al., 2019;Hoerster et al., 2020). All media recipes are
14 described by (Lowe et al., 2018;Chu et al., 2019;Hoerster et al., 2020). For selection, 0.1 mg/L
15 imazapyr was supplemented to somatic embryo formation medium or 150 mg/L G418 was
16 substituted for imazapyr.

17

18 ***Agrobacterium*-mediated transformation**

19 Constructs used in these experiments are illustrated in Figures 1, 2, and 4 and the individual
20 expression components such as promoters, structural genes and terminators are listed in Table S
21 1. The materials reported in this article contain selectable markers (*HRA* and *NPTII*) and reporter
22 genes (*ZS-Green* and *Zs-Yellow*) are owned by third parties. Authors may not be able to provide
23 materials including third party genetic elements to the requestor because of certain third-party

1 contractual restrictions placed on the author's institution. In such cases, the requester will be
2 required to obtain such materials directly from the third party. The authors and authors'
3 institution do not make any express or implied permission(s) to the requester to make, use, sell,
4 offer for sale, or import third party proprietary materials.

5
6 All transformations were done using the thymidine auxotrophic *Agrobacterium*
7 *tumefaciens* strain LBA4404 THY- containing pVIR9 (Anand et al., 2018) at OD₅₅₀ of 0.5. The
8 conditions for *Agrobacterium* suspension culture preparation following embryo isolation and
9 infection has been previously described (Lowe et al., 2018;Hoerster et al., 2020). Two selectable
10 markers were used in experiments: *HRA* (Green et al., 2009), a sulfonyleurea herbicide resistance
11 marker, driven by the sorghum *Als* promoter for selection with 0.1 mg/L imazapyr in culture
12 medium, or the *Ubi_{pro}::NPTII* gene for selection with 150 mg/L G418 in culture medium.

14 **Excision conditions**

15 For the developmentally regulated *pro::Cre* testing, no optimization was required. These
16 experiments were performed on two inbreds, HC69 and PHR2HT. The initial heat shock
17 treatment for excision involved three different conditions: no heat shock (control), heat shock at
18 37°C for 1 day, or 42°C for 2h/day for 3 consecutive days, were tested. We further optimized the
19 heat shock condition testing three additional heat treatments 1) 42°C, 2h/day for 2 d, 2) 42°C for
20 24h, or 3) 45°C for 2h/day to identify a treatment that is best and simple for implementation.

22 **Molecular analyses**

1 All molecular analysis and transgene copy number determination methods were previously
2 described (Wu et al., 2014;Lowe et al., 2016;Hoerster et al., 2020). qPCR data was used to
3 confirm recombinase-mediated excision based on the absence the transgenes flanked by the *loxP*
4 sites, determine the copy number of structural genes outside the excision DNA, and to screen for
5 the presence of *Agrobacterium* binary construct backbone integration. Genomic DNA samples
6 were extracted from a single piece (200 ng) of fresh leaf tissue from each plant (Truett et al.,
7 2000). Non-transgenic maize inbred lines were used as the negative controls. Quantification
8 was based on detection of amplified gene sequences using gene-specific forward and reverse
9 primers, along with the corresponding gene-specific FAMTM or Vic[®]-based MGB fluorogenic
10 probes (Applied Biosystems). The 2- $\Delta\Delta$ CT method (Livak and Schmittgen, 2001) was used to
11 estimate copy number. Events which are single copy for all the transgenes and excised was used
12 to calculate the excision frequency. The events which are excised with a single copy (SC) of all
13 the transgenes without vector backbone integration were defined as a quality event (QE). The
14 usable event (UE) frequency was calculated as transformation frequency times QE frequency.
15 Data collected from different experiments were analyzed separately by analysis of variance
16 (ANOVA), with mean separation by LSD (P=0.05) using JMP Pro 12.2.0 Statistical Discovery
17 software package (SAS Institute Inc., Cary, NC).

18

19 **CONCLUSION**

20 Despite the recent progress in developing a rapid maize transformation, the presence of
21 morphogenic genes in the transgenic event have shown to result in pleiotropic phenotypes and is
22 not recommended for transgene testing or commercial product development. The first generation
23 of rapid maize transformation method was designed to improve the transformation rates and to

1 extend transformation capabilities to many genotypes. Subsequently, we demonstrated a viable
2 second-generation alternative, using a mixture of an *Agrobacterium* strains, one with non-
3 integrating *Wus2* gene and the other with a combination of structural genes to regenerate
4 transgenic plants free of morphogenic genes. Even though this simplifies vector construction,
5 however, the process still relies on SMG for recovery of stable transgenic events. This study
6 demonstrated a viable third alternative, relying on inducible promoters for auto-excision of both
7 the morphogenic genes and the SMG in the early stages of maize transformation. The stable
8 transformed plants recovered by this method are free of the morphogenic genes and marker
9 genes, a desirable quality for transgene evaluation and in commercial products.

10

11 **AUTHOR CONTRIBUTION STATEMENT**

12 A.A., E.W., L.K., W.G-K., T.J., and N.D.A conceived the research idea, A.A., E.W., L.K., and
13 W.G-K. designed constructs and research, and N.W., MA., HG. and R.L conducted maize
14 transformation and optimization; E.W. and A.A, performed data analysis; A.A., W.G-K . T.J.,
15 and N.D.C. wrote the manuscript.

16

17 **CONFLICT OF INTEREST**

18 NW, MA, HG, RL, EW, LK, W.G-K and AA are inventors on pending applications on this work
19 and a related work are current employees of Corteva Agriscience who owns the pending patent
20 applications. TJ and NDC are current employees of Corteva Agriscience.

21

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44

45

1 Table 1. List of the promoters, their source, and their expression pattern in plants.

Promoters	Source	Expression	Reference
<i>Kn1</i>	Maize	Apical Meristem	Gen bank AY312169
<i>Lec1</i>	Maize	Early Embryo	(Shane, 2007)
<i>End2</i>	Maize	Early Embryo	(Casper et al., 2005)
<i>Ltp2</i>	Maize	Early Embryo	(Kalla et al., 1994)
<i>Glb1</i>	Maize	Late Embryo	(Liu et al., 1998)
<i>Ole</i>	Maize	Late Embryo	(Anand et al., 2017b)
<i>Rab17</i>	Maize	Late Embryo/Stress	(Busk et al., 1997)
<i>Nos</i>	<i>Agrobacterium tumefaciens</i>	Constitutive	(An, 1986)
<i>Ubi_{pro}</i>	Maize	Constitutive	(Christensen et al., 1992)
<i>Hsp17.7</i>	Maize	Heat shock inducible	(Anand et al., 2017a)
<i>Hsp26</i>	Maize	Heat shock inducible	(Anand et al., 2017a)
<i>Rab21</i>	<i>Seteria itallica</i>	Drought inducible	Previously unpublished Corteva Agriscience sequence Si026926m
<i>Drp12</i>	<i>Brachypodium distachyon</i>	Drought inducible	Previously unpublished Corteva Agriscience sequence Bradi3g43870.1
<i>Drp1</i>	<i>Brachypodium distachyon</i>	Drought inducible	Previously unpublished Corteva Agriscience sequence Bradi1g37410.1

2

1 Table 2. Transformation results with different developmentally regulated promoters driving *Cre* expression for auto-excision of
 2 morphogenic genes using construct design described in Figure 1. Data presents the T0 transformation frequency, qPCR detection of
 3 the number of excised events and the quality event frequency in two different inbreds, PH2RT and HC69.

4

Inbred	Promoter	Embryos transformed	T0 plants	T0 transformation frequency (% \pm SE)	Excised single copy, backbone-free events	Excision frequency (%)	Quality event (%)	Usable events (%)
PH2RT	<i>Ltp2</i>	229	75	32.8 (2.2) ^a	10	50	13.3	4.4
	<i>Ole</i>	228	59	27.2 (3.3) ^{ab}	8	40	13.6	3.5
	<i>Glb1</i>	280	38	13.6 (1.4) ^c	7	58.5	18.4	2.5
	<i>End2</i>	174	39	22.4 (2.6) ^b	3	100	7.7	1.7
	<i>Ubi</i>	440	40	9.1 (1.9) ^c	12	59.1	30.0	2.7
HC69	<i>Rab17</i>	121	35	28.9 (2.6) ^b	1	25	2.9	0.8
	<i>Ole</i>	151	49	37.1 (2.1) ^a	3	37.5	6.1	2.0
	<i>Glb1</i>	230	58	25.2 (1.8) ^b	5	38.5	8.6	2.2
	<i>End2</i>	178	48	27.0 (2.4) ^b	1	100	2.1	0.6
	<i>Ubi</i>	202	37	18.3 (1.2) ^c	3	13.6	8.1	1.5

5

6 Data from three independent transformers was used to determine T0 transformation frequency. The quality events (QE) were identified as single copy, backbone-
 7 free, and morphogenic gene-free (excised). The excision frequency was determined as the ratio of the number of excised single-copy events relative to the total
 8 single-copy events. The number QEs was divided by the total number of events recovered to calculate the QE frequency. The usable event (UE) frequency is a
 9 measure of the number of acceptable transgenic events per 100 embryos that was determined as the product of QE frequency and T0 transformation frequency.

10 Mean values followed by the same letter are not statistically different from each other at the significance level of 0.05.

1 Table 3. Transformation results from excision-activated marker gene selection using either the *Glb1_{pro}* or the *Ole_{pro}* driving *Cre*
 2 expression using construct design described in Figure 2. Data presents the T0 transformation frequency, qPCR detection of the number
 3 of excised events and the quality event frequency in maize inbred HC69.

Promoter	Embryos transformed	T0 plants	T0 transformation frequency (% ±SE)	Total single copy events	Excised single copy, backbone-free events	Excision frequency (%)	Quality event (%)	Usable events (%)
<i>Glb1</i>	126	57	44.7 (2.8) ^a	19	13	68.4	13.3	5.6
<i>Ole</i>	112	45	40.2 (1.9) ^a	15	8	53.3	8.8	3.6

4
 5 Data from two independent transformers was used to determine T0 transformation frequency. Quality events (QE) were identified as single copy, backbone-free,
 6 and morphogenic gene-free (excised). The excision frequency was determined as the ratio of the number of excised single-copy events relative to the total single-
 7 copy events. The number QEs was divided by the total number of events recovered to calculate the QE frequency. The usable event (UE) frequency is a
 8 measure of the number of acceptable transgenic events per 100 embryos that was determined as the product of QE frequency and transformation frequency.
 9 Mean values followed by the same letter are not statistically different from each other at the significance level of 0.05.

1 Table 4. Transformation results from excision-activated marker gene selection using *Glb_{pro}* driving *Cre* expression using construct
 2 design described in Figure 2. Data presents the T0 transformation frequency, qPCR detection of the number of excised events and the
 3 quality event frequency in three maize inbreds (HC69, PH85E, and PH84Z).

Inbred	Embryos transformed	T0 plants	T0 transformation (% ±SE)	Excised single copy, backbone-free events	Excision frequency (%)	Quality event (%)	Usable events (%)
HC69	393	196	49.9 (3.9) ^a	17	55.0	8.7	4.3
PH85E	363	47	12.9 (1.3) ^c	13	59.0	27.7	3.6
PH84Z	367	105	28.6 (2.5) ^b	7	61.0	6.7	1.9

4
 5 Data from two independent transformers was used to determine T0 transformation frequency. The quality events (QE) were identified as single copy, backbone-
 6 free, and morphogenic gene-free (excised). The excision frequency was determined as the ratio of the number of excised single-copy events relative to the total
 7 single-copy events. The number QEs was divided by the total number of events recovered to calculate the QE frequency. The usable event (UE) frequency is a
 8 measure of the number of acceptable transgenic events per 100 embryos that was determined as the product of QE frequency and transformation frequency.
 9 Mean values followed by the same letter are not statistically different from each other at the significance level of 0.05.

10

1 Table 5. Transformation results from screening of six different inducible promoters driving *Cre* expression for controlled gene
 2 excision. For this study, three different conditions were evaluated: two heat shock treatments (37°C for 1 day and 42°C, 2h/day for 3
 3 consecutive days) and no heat (control). Data presents the qPCR detection of the number of excised events and excision frequency
 4 across the different promoters, and a control construct without the *Cre* gene, in maize inbred HC69.

5

Promoter	Control				37°C, 1 day				42°C, 2h/day for 3 days			
	Embryos	T0	Excision		Embryos	T0	Excision		Embryos	T0	Excision	
		plants	QE	(%)		plants	QE	(%)		plants	QE	(%)
<i>Hsp17.7</i>	455	59	5	27.8	50	6	2	66.7	50	20	4	100
<i>Hsp26</i>	450	98	0	0.0	50	5	0	0	50	21	3	43
<i>Rab17</i>	455	127	1	3.4	50	10	0	0	50	18	0	0
<i>Rab21</i>	455	101	8	36.4	50	13	1	100	50	20	0	0
<i>Drp12</i>	450	79	2	11.1	50	16	0	0	50	22	2	66.7
<i>Drp1</i>	438	90	8	27.6	50	8	0	0	50	27	5	45.5
Control (no <i>Cre</i>)	450	182	0									

6

7 Data from two independent transformers was used to determine T0 transformation frequency. The quality events (QE) were identified as single copy, backbone-
 8 free, and morphogenic gene-free (excised). The excision frequency was determined as the ratio of the number of excised single-copy events relative to the total
 9 single-copy events.

1 Table 6. Transformation results optimizing the heat shock conditions for controlled gene excision using three inducible promoters
 2 driving *Cre* expression. The three different conditions evaluated were: no heat (control) and two heat shock treatments (42°C for 2h
 3 and 42°C, 2h/day for 3 consecutive days). The data presents the qPCR detection of the number of excised events and excision
 4 frequency across the different promoters in the study as compared to a control construct without the *Cre* gene in maize inbred HC69.

Promoter	Control					42°C, 2h					42°C, 2h/day for 3 days				
	Embryos	T0 Plants	QE	Excision frequency (%)	UE %	Embryos	T0 plants	QE	Excision frequency (%)	UE (%)	Embryos	T0 plants	QE	Excision frequency (%)	UE (%)
<i>Hsp17.7</i>	50	18	1	12.5	2	50	17	5	62.5	10	50	15	9	69.2	18.0
<i>Hsp26</i>	50	18	0	0	0	50	21	2	42.2	4.0	50	9	2	66.7	4.0
<i>Drp12</i>	50	11	1	25	2	50	9	1	50.0	2.0	50	14	1	20.0	2.0

5
 6 Data from two independent transformers was used to determine T0 transformation frequency. The quality events (QE) were identified as single copy, backbone-
 7 free, and morphogenic gene-free (excised). The excision frequency was determined as the ratio of the number of excised single-copy events relative to the total
 8 single-copy events. The number QEs was divided by the total number of events recovered to calculate the QE frequency. The usable event (UE) frequency is a
 9 measure of the number of acceptable transgenic events per 100 embryos that was determined as the product of QE frequency and transformation frequency.
 10 Mean values followed by the same letter are not statistically different from each other at the significance level of 0.05

1 Table 7. Optimizing heat shock conditions for controlled gene excision using heat shock promoters *Hsp17.7* and *Hsp26* driving *Cre*
 2 expression as shown in Figure 5. Four different conditions were evaluated side-by-side using split ears including no heat (control) and
 3 three heat shock treatments (42°C, 2h/d for 2d; 42°C/24h; and 45°C/2h). Transformation results and qPCR detection of the number of
 4 excised quality events, frequencies of excision and usable event are presented.

Promoter	Treatments	Embryos transformed	T0 plants	T0 transformation (% ±SE)	Quality events	Excision frequency (%)	Usable event (%)
<i>Hsp17.7</i>	none	102	56	54.9 (4.4) ^a	6	33.3	5.9
	42°C, 2h/d, 2d	102	39	38.2 (2.1) ^b	9	56.3	8.8
	42°C/24h	102	16	15.7 (1.8) ^c	6	75.0	5.9
	45°C/2h	102	50	49.0 (3.2) ^a	14	76.6	13.7
<i>Hsp26</i>	none	100	53	53.0 (4.0) ^a	1	5.6	1.0
	42°C, 2h/d, 2d	100	35	35.0 (1.2) ^b	12	66.7	12.0
	42°C/ 24h	100	41	41.0 (2.2) ^b	10	66.7	10.0
	45°C/2h	100	50	50.0 (2.3) ^a	13	61.9	13.0

5 Data from two independent transformers was used to determine T0 transformation frequency. The quality events (QE) were identified as single copy, backbone-
 6 free, and morphogenic gene-free (excised). The excision frequency was determined as the ratio of the number of excised single-copy events relative to the total
 7 single-copy events. The number QEs was divided by the total number of events recovered to calculate the QE frequency. The usable event (UE) frequency is a
 8 measure of the number of acceptable transgenic events per 100 embryos that was determined as the product of QE frequency and transformation frequency.
 9 Mean values followed by the same letter are not statistically different from each other at the significance level of 0.05.

10

1 Table 8. Transformation results and molecular event data using the *Hsp17.7* heat shock promoter for controlled excision of both
 2 morphogenic gene and marker gene in three maize inbreds (HC69, PH85E, and PH84Z). Two different SMGs were evaluated, *HRA*
 3 (resistance to the sulfonyleurea herbicide ethametsulfuron) and *NPTII* (resistance to antibiotic G418), using the same construct design
 4 with the same set of morphogenic genes as shown in Figure 5. Transformation results and qPCR detection of the number of excised
 5 quality events, frequencies of excision and usable event are presented.

Inbred	Selectable marker	Embryos transformed (number)	T0 plants (number)	T0 transformation (%)	Excised single copy, backbone-free events (number)	Excised single copy, backbone-free events (%)	Excision frequency (%)	Usable event (%)	Null (%)
HC69	<i>NPTII</i>	315	200	63.5	46	23.0	87.1	14.6	17.1
	<i>HRA</i>	407	281	69.0	45	16.0	82.3	11.1	37.3
PH85E	<i>NPTII</i>	219	64	29.2	23	35.9	96.7	10.5	15.3
	<i>HRA</i>	320	124	38.8	31	25.0	97.2	9.7	42.5
PH84Z	<i>NPTII</i>	356	145	40.7	19	13.1	50.4	5.3	14.2
	<i>HRA</i>	365	169	46.3	14	8.3	59.9	3.8	41.8

7 Data from two independent transformers was used to determine T0 transformation frequency. The quality events (QE) were identified as single copy, backbone-
 8 free, morphogenic and marker gene-free (excised). The number of QEs was divided by the total number of events analyzed to calculate the QE frequency. The
 9 excision frequency was determined as the ratio of the number of excised single-copy events relative to the total single-copy events. The usable event (UE)
 10 frequency is a measure of the number of acceptable transgenic events per 100 embryos that was determined as the product of QE frequency and transformation
 11 frequency.

12

13

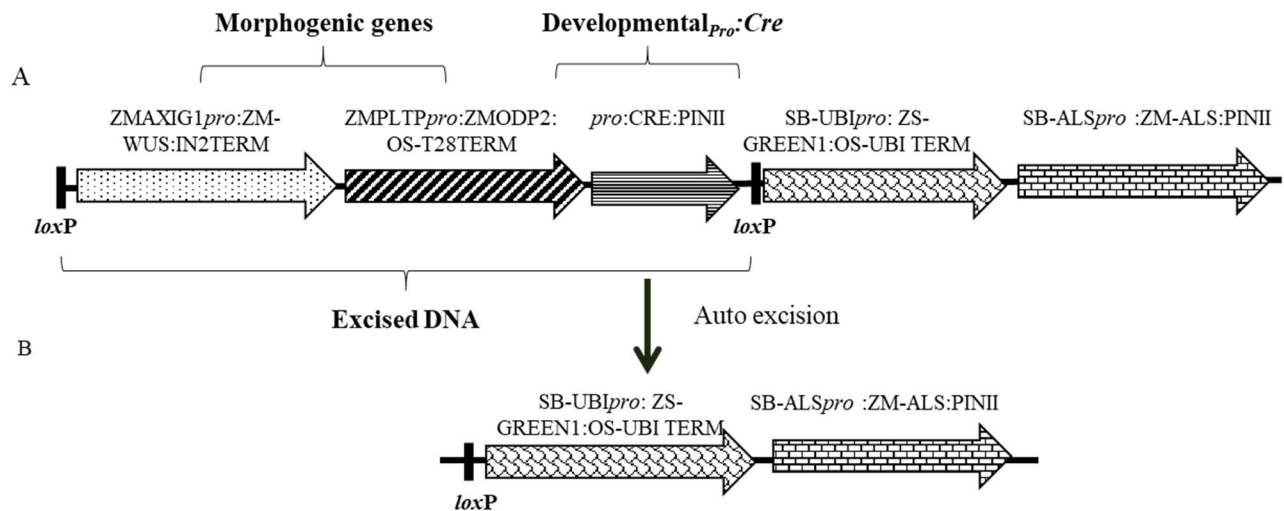
- 1 Table 9. Observed and expected number of homozygous, hemizygous and null plants for T-DNA integration copy number in in T1
- 2 generation of 13 SC excised quality events across two maize inbreds (PH84Z and HC69).

Inbred	Event ID	Total Plants	Homozygous	Hemizygous	Null	Chi-square	P-value*
PH84Z	ZMYF66.001.83A	23	7	11	5	0.39	0.82
	ZMCJK9.001.74A	31	10	13	8	0.76	0.68
	ZMCJK9.001.13A	30	8	18	4	2.03	0.36
	ZMCJK9.001.96A	32	6	17	9	0.69	0.71
	ZMCJK9.001.34A	30	10	12	8	1.5	0.47
	ZMCJK9.001.77A	24	5	10	9	2	0.36
	ZMCJK9.001.3A	27	4	17	6	2.07	0.35
HC69	ZMNW4W.001.72A	23	11	7	5	6.65	0.03
	ZMNW4W.001.30A	31	11	13	7	1.83	0.39
	ZMNW32.001.49A	32	4	17	11	3.19	0.2
	ZMNW32.001.58A	31	8	14	9	0.35	0.84
	ZMNW32.001.43A	32	9	10	13	5.5	0.06
	ZMNW32.001.65A	32	9	14	9	0.5	0.78

- 3
- 4 * No statistically significant deviations identified from expected 1:2:1 (homozygous:hemizygous:null) segregation at 5% level

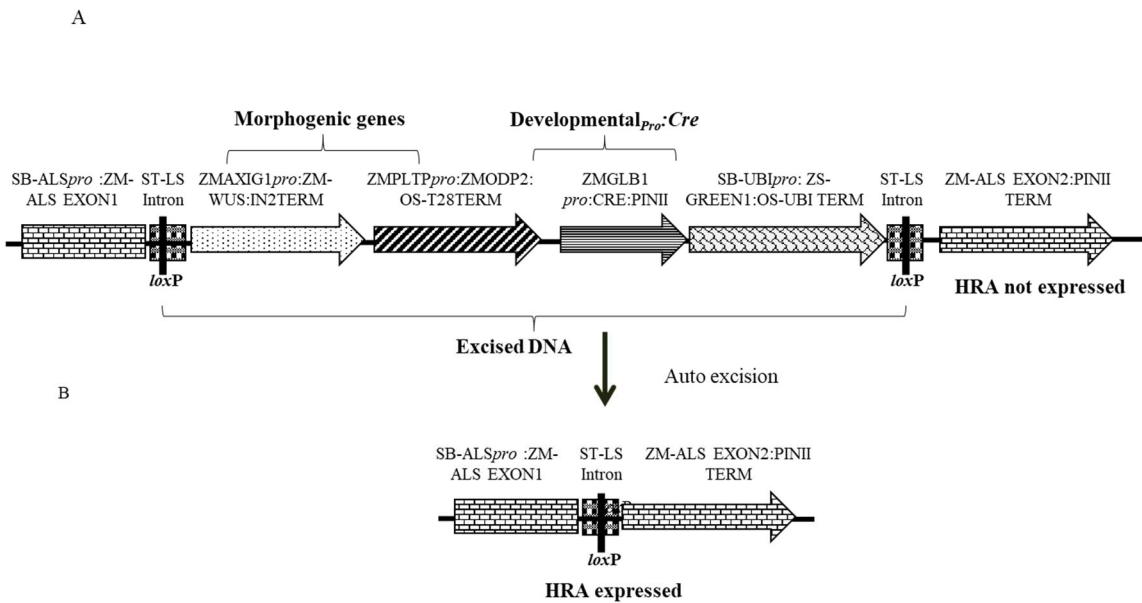
1 **Figure 1.** Schematic representation of an auto-excision construct design used for testing different
2 developmentally regulated or stress-inducible promoters to achieve excision of morphogenic
3 genes. A) The excision construct with different promoter combinations driving *Cre* expression
4 (represented by *pro:CRE*) and the DNA fragment to be excised flanked by two directly oriented
5 *loxP* recombination sites. B) The excised product following auto-excision. Refer to Table S-1 for
6 description of construct components used in T-DNA construction.

7



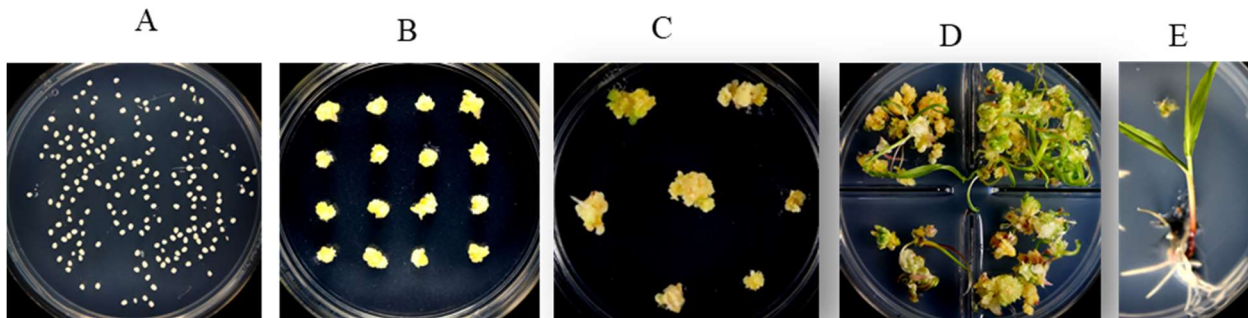
8

- 1 **Figure 2.** Schematic representation of an auto-excision construct design used for testing
- 2 developmentally regulated promoters driving *Cre* expression (represented by *pro:CRE*) for
- 3 excision-activated SMG expression. A) An excision-activated selectable marker construct design
- 4 with the DNA fragment to be excised flanked by two directly oriented *loxP* recombination sites.
- 5 B) Following excision, the *HRA* gene is activated and events are selected on a media
- 6 supplemented with 0.1 mg/L imazapyr. Refer to Table S-1 for description of construct
- 7 components used in T-DNA construction.



8

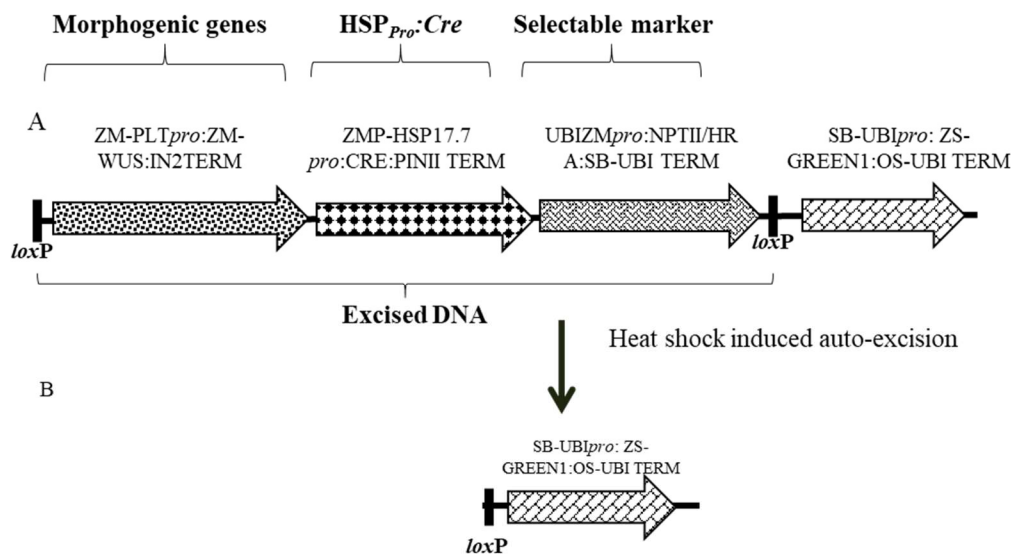
- 1 **Figure 3.** The different stages in rapid maize transformation and heat shock treatment. A)
- 2 immature zygotic embryos are isolated and infected with *Agrobacterium tumefaciens*, (B)
- 3 transgenic somatic embryos are placed for 3 weeks on selection media based on selectable
- 4 marker used (*HRA* or *NPTII*), (C) somatic embryos are heat shock treated and transferred to
- 5 maturation media, (D) transgenic plants are regenerated without selection pressure for 2 weeks
- 6 and, (E) regenerated plants are placed on a rooting media for 2-3 weeks.



7

1 **Figure 4.** Schematic representation of an auto-excision construct design used for testing
2 elimination of a morphogenic gene and a marker gene using heat shock promoter driving *Cre*
3 expression for controlled gene excision. A) Construct design depicting the order of cassettes
4 including morphogenic genes, *Hsp17.7_{pro}* driving *Cre* expression, and the selectable marker
5 (*HRA* or *NPTII*) flanked by directly oriented *loxP* sites (a) which will be excised upon *Cre*
6 expression. B) Following excision, the DNA piece containing the ZS-GREEN expression
7 cassette is left in the T0 event for visual confirmation of excision. Refer to Table S-1 for
8 description of construct components used in T-DNA construction.

9



10

1 Table S-1. Construct components used in T-DNA construction.

2

Component type	Label	Description	References
Promoters	<i>Sb-Als_{pro}</i>	The sorghum ALS promoter	SB-ALS promoter and 5'UTR, DOE-JGI Sbi v3.1, SBChr04, bases 49239164-49240031. DOE-JGI Sbi v3.1 corresponds to Sorghum bicolor BTx623 assembly v3.0.1 and gene annotation v3.1 available from phytozome (http://phytozome.jgi.doe.gov/). Chromosome 4 of Sbi v3.1 is registered as NCBI accessions NC_012873.2 and CM000763.3
	<i>Pltp_{pro}</i>	Maize phospholipid transferase promoter	See GenBank sequence (MN380778)
	<i>Axig1_{pro}</i>	The maize Axig1 promoter	(Garnaat et al., 2002)
	<i>Sb-Ubipro</i>	The sorghum Ubiquitin promoter	(Shane, 2007)
3' Sequences	<i>In2-2</i>	The maize IN2-2 terminator	(Hershey and Stoner, 1991)
	<i>PINII</i>	The potato proteinase inhibitor II (pinII) 3' sequence	(An et al., 1989)
	<i>Os-Ubi 3'</i>	The rice Ubiquitin terminator	Terminator region of the rice Ubiquitin (Os06g46770.1), unpublished
	<i>Sb-Ubi 3'</i>	The sorghum Ubiquitin terminator	(Shane, 2007)
	<i>Os-T28 3'</i>	The T28 3' regulatory sequence from <i>Oryza sativa</i>	(Bhyri et al., 2014)
Marker genes	<i>NPTII</i>	Maize codon-optimized Neomycin Phosphotransferase II	Previously unpublished Corteva Agriscience sequence

	<i>HRA</i>	The maize ALS double mutant gene conferring herbicide resistance	(Green et al., 2009)
	<i>Zs-YELLOW</i>	The Zs-Yellow1 N1 gene encoding a yellow fluorescent protein from <i>Zoanthus sp</i>	(Matz et al., 1999)
Maize morphogenic genes	<i>Zm-Wus2</i>	The maize <i>Wuschel2</i> (<i>Wus2</i>) gene	(Lowe et al., 2007)
	<i>Zm-Bbm</i>	The maize <i>Baby boom</i> gene (<i>Bbm</i>)	(Gordon-Kamm et al., 2005)
Recombinase Expression Cassettes	<i>Cre</i>	A maize-optimized <i>Cre</i> recombinase gene (originally from the P1 bacteriophage), with an inserted potato LS1 intron	(Odell et al., 1990)
Recombinase Target Sites	<i>loxP</i>	The recombinase target site for the Cre recombinase from <i>E. coli</i>	(Odell et al., 1990)

1