Developing a rapid and highly efficient cowpea regeneration, transformation and genome editing system using embryonic axis explants

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SUMMARY

Cowpea (Vigna unguiculata (L.) Walp.) is one of the most important legume crops planted worldwide, but despite decades of effort, cowpea transformation is still challenging due to inefficient Agrobacterium-mediated transfer DNA delivery, transgenic selection and in vitro shoot regeneration. Here, we report a highly efficient transformation system using embryonic axis explants isolated from imbibed mature seeds. We found that removal of the shoot apical meristem from the explants stimulated direct multiple shoot organogenesis from the cotyledonary node tissue. The application of a previously reported ternary transformation vector system provided efficient Agrobacterium-mediated gene delivery, while the utilization of spcN as selectable marker enabled more robust transgenic selection, plant recovery and transgenic plant generation without escapes and chimera formation. Transgenic cowpea plantlets developed exclusively from the cotyledonary nodes at frequencies of 4% to 37% across a wide range of cowpea genotypes. CRISPR/Cas-mediated gene editing was successfully demonstrated. The transformation principles established here could also be applied to other legumes to increase transformation efficiencies.

Keywords: cowpea transformation, embryonic axis, cotyledonary node (cot-node), shoot organogenesis, spectinomycin, Agrobacterium, CRISPR/Cas.

INTRODUCTION

Domesticated in Africa and widely cultivated in the tropical and subtropical zones of the world, cowpea (Vigna unguiculata (L.) Walp.), also known as black-eyed pea, is one of the most valuable grain legumes for high-quality dietary protein, carbohydrates, lipids, minerals and vitamins for people in developing countries of Africa and Asia (Abdu Sani et al., 2015; Phillips et al., 2003; Singh, 2014). It is estimated that over 200 million people consume cowpea daily in Africa (Phillips et al., 2003; Singh, 2014). Despite its high tolerance to heat, dry conditions and soil acidity, cowpea is highly susceptible to insect pests and pathogen infections, resulting in lower productivity (Abdu Sani et al., 2015; Boukar et al., 2016; Obembe, 2008; Singh, 2014; Solleti et al., 2008a). Due to limited genetic variability of cowpea and strong cross-incompatibility between wild Vigna species and cultivated cowpea, little progress has been made in genetic improvement through conventional breeding to achieve desirable agronomic traits (Abdu Sani et al., 2015; Fang et al., 2007; Gomathinayagam et al., 1998; Latunde-Dada, 1990; Wamalwa et al., 2016). Hence, plant biotechnology provides an alternative approach to overcome those constraints for improving the agronomic performance and developing improved cowpea cultivars with higher grain quality and yield (Carlos Popelka et al., 2004; Zaidi et al., 2005). The development of insect-resistant cowpea, unsuccessful through conventional breeding, was successfully achieved by introducing Bt genes through genetic transformation and is a good example of plant biotechnology application in an orphan crop (Bakshi et al., 2011; Bett
et al., 2017; Zaidi et al., 2005). Recently, significant progress has been made establishing genomic and gene expression data resources for two cowpea varieties, IT86D-1010 (Srippigs et al., 2018) and IT97K-499-35 (Lonardi et al., 2019; Munoz-Amatriain et al., 2017; Yao et al., 2016). However, the absence of an efficient genetic transformation and editing system (Popelka et al., 2006; Somers et al., 2003) has impeded the full utilization of these resources for cowpea functional genomic studies to elucidate the mechanisms of heat and drought stress tolerance and to improve the agronomic traits, such as insect and pathogen resistances and increased productivity.

Legumes, especially cowpea, are known to be recalcitrant for genetic manipulation (Manman et al., 2013; Popelka et al., 2006; Solleti et al., 2008b; Somers et al., 2003). This is mainly due to the inadequate Agrobacterium-mediated transfer DNA (T-DNA) delivery to the targeted tissue, the inefficient transgenic selection methods for viable transgenic plant recovery and the absence of an amenable in vitro shoot regeneration system. Although notable improvements have been made in recent years for Agrobacterium-mediated cowpea transformation using two types of explants, the cotyledonal node (cot-node) explants excised from germinated seedlings (Bakshi et al., 2011) and the cotyledon with attached embryonic axis (EA) from imbibed mature seeds (Bett et al., 2019), the published transformation frequencies (Bett et al., 2019) are still lower than 3.9% (Bett et al., 2019; Chaudhury et al., 2007; Manman et al., 2013; Mellor et al., 2012) and the tissue culture process generally requires excessive explant manipulation to remove the cotyledon, primary shoots and any regrown radicle and usually takes at least 5 to 8 months for generating fully developed plantlets after Agrobacterium infection (Chaudhury et al., 2007; Popelka et al., 2006). Besides different explant types, several selection systems have been reported for cowpea transformation (Manman et al., 2013), such as NPTII/kanamycin (Bett et al., 2019; Chaudhury et al., 2007), NPTII/G418 (Solleti et al., 2008b), PML/mannose (Bakshi et al., 2012), HPT/hygromycin (Kumar et al., 1996), BAR/glufosinate (Popelka et al., 2006) and ahasi/imazapyr (Citadin et al., 2013; Ivo et al., 2008). It was reported that incomplete selection and tissue necrosis were associated with those selection systems (Manman et al., 2013) and resulted in lower transgenic plant recovery (Bakshi et al., 2011; Chaudhury et al., 2007; Solleti et al., 2008b) and a higher percentage of chimera formation in cowpea (Das Bhowmik et al., 2019).

To increase the regeneration rate, enhance transgenic plant recovery and eliminate chimera formation under selection in this study, we first evaluated shoot regeneration using detached EA explants isolated from imbibed mature (dry) seeds and identified that only cot-node cells of the detached EA explants undergo the rapid cell division and dedifferentiation required to acquire organogenic competence for shoot regeneration. Based on this observation, we developed a rapid, robust and highly efficient Agrobacterium-mediated EA-based transformation using CTP-spcN as selectable marker and generated transgenic cowpea events without non-transgenic escapes and chimera formation. Finally, we applied this transformation technology to nine cowpea genotypes and achieved transformation frequencies in the range of 4% to 37% and demonstrated its potential to support efficient genome editing by creating inheritable knockouts in IT86D-1010 using CRISPR/Cas gene editing technology. The overall tissue culture process after Agrobacterium infection to generate fully developed plantlets was reduced to less than 3 months.

RESULTS AND DISCUSSION

De novo shoot organogenesis using EA as explants

A rapid, efficient and reproducible regeneration system is a prerequisite for establishment of an efficient cowpea genetic transformation system. Although several studies of in vitro regeneration of cowpea based on organogenesis have been reported (Aasim et al., 2010; Abdu Sani et al., 2015; Mamadou et al., 2008; Manman et al., 2013; Odutayo et al., 2005; Raveendrar et al., 2009; Sani et al., 2018; Tie et al., 2013; Yusuf et al., 2008), an efficient cowpea regeneration system that enables highly efficient transformation is still lacking (Manman et al., 2013). Soybean (Glycine max) transformation based on the pre-existing meristems of EA explants has been well established and provides a reliable and highly efficient means for introducing transgenes (Aragao et al., 2000; Liu et al., 2004; Turlapati et al., 2008). To test the regeneration efficiency of EA explants in cowpea, EA explants were isolated from imbibed mature seeds of cowpea variety IT86D-1010 by excising the cotyledons at the cot-node (Figure 1(a)). Those EA explants with the plumule excised (Figure 1(b)) were then cultured directly onto shoot induction medium (SIM) (Table S5) without selection in a vertical upright position with roots embedded in the media to induce shoot development. In most of the cases (>95%), a single primary shoot was developed per EA when the shoot apical meristem (SAM) of the EA explants was kept intact (non-decapitated EA explants) during regeneration (Figure 1(c)). However, multiple shoot development was observed occasionally for a small number of explants (<5%) (Figure 1(d)). Compared to the morphology of EA explants with single primary shoot development (Figure 1(c)), the EA explants with multiple shoots regenerated exclusively around cot-node (Figure 1(d)), were much shorter and lacked both epicotyls and SAMs. The lack of epicotyls and SAMs could be due to the accidental damage of those tissues during EA isolation and plumule excision. This finding indicated that the de novo organogenesis of shoots around the cot-nodes could be
inhibited by apical dominance. Indeed, removal of the SAM purposely by cutting through the middle of each epicotyl (decapitated EA explants without SAM and plumule) (Figure 1a,e)) induced 78% of explants to initiate multiple shoot regeneration in IT86D-1010 (Figure 1f,g) and Table S9). These results demonstrated that tissue culture and regeneration procedures can be applied to a wide collection of cowpea germplasm and extended to other legumes such as common bean.

Regeneration optimization under Agrobacterium-mediated transformation

Generally, the EA-based dicot transformation procedure consists of the following main steps: explant preparation, Agrobacterium infection, co-cultivation, shoot regeneration with selection and root induction (Figure S2 and the Experimental Procedures section). As described above, although EA explants per se have been described for soybean transformation, the cowpea EA-based regeneration system has a key difference. While the soybean EA transformation system relies on pre-existing apical meristematic tissue for regeneration and transformation (Aragão et al., 2000; Liu et al., 2004; Turlapati et al., 2008), the cowpea regeneration system shows that removal of the SAM stimulated multiple shoot organogenesis from the cot-node. This key difference raises the question of how well the decapitated EA explants will be able to survive and regenerate throughout the Agrobacterium-mediated transformation procedure.

To evaluate how the decapitation of EA explants affects the survival and regeneration capability, we conducted sonication, Agrobacterium infection and co-cultivation treatments (Figure S2 and the Experimental Procedures section) either with or without Agrobacterium, followed by regeneration on SIM. As shown in Figure 2a, the decapitated EA explants were extremely sensitive to the treatments and none of the EA explants survived on SIM without selection after mimicking all the treatment steps without Agrobacterium. On the contrary, all the non-decapitated EA explants survived and formed elongated epicotyls with a single primary shoot. The further decapitation of those primary shoots by cutting through the middle of
the elongated epicotyls after 4 days of regeneration stimulated multiple shoot organogenesis around the cot-nodes (Figure 2(b)). Collectively, those observations suggest that although the SAM negatively regulates multiple shoot organogenesis from cot-node tissues because of the apical dominance effect, the SAM is essential for EA explant survival through the transformation treatments before regeneration and the SAM should not be removed until the explants are fully recovered after 4 days of regeneration.

It has been reported that Agrobacterium-mediated infection leads to cell damage and tissue necrosis (Norkunas et al., 2018). To determine the survival rate of EA explants after infection and cocultivation with Agrobacterium, non-decapitated EA explants without plumules were subjected to the transformation procedure (Figure S2 and the Experimental Procedures section) using LBA4404 Th-y carrying the pPHP86170/pPHP71539 vector system as described below. As shown in Figure 2(c), about 70±10% EA explants (based on the average of three replicates and 75 EA explants) survived and formed elongated epicotyls after 4-day regeneration on SIM with selection (Table S5). Compared to the 100% survival rate without Agrobacterium inoculation (Figure 2(b)), the 30% mortality rate was most likely due to the sensitivity of EA explants to the Agrobacterium.

It is known that plumules interfere with shoot regeneration and need to be removed from soybean EA explants. The same is true for cowpea EA explants. No transgenic shoots can be regenerated without removing the plumule. However, removing plumules from isolated EAs, one by one very carefully without damaging the SAM, is the most time-consuming and labor-intensive step of this process. This is because EA isolation with an intact SAM is the critical step for maximizing explant survival during tissue culture inoculation as described above. To avoid SAM damage during EA explant preparation and for the purpose of developing a best practice, we simplified the EA isolation and subsequent transformation procedure as follows. Instead of removing the plumule at the beginning of EA isolation, as usually performed for soybean, both the SAM and the plumule are removed simultaneously on the fourth day on SIM by cutting through the middle of the epicotyl with a pair of surgical scissors (Figure S2 and the Experimental Procedures section). This improved procedure was implemented for all the subsequent transformation optimization experiments throughout this study.

Agrobacterium-mediated gene delivery using a ternary vector system

A dramatic increase of T-DNA delivery efficiency was reported in cowpea by constitutive expression of additional vir genes in a resident pSB1 vector in Agrobacterium strain LBA4404 (Solleti et al., 2008b). Recently, we demonstrated that a newly designed ternary vector containing the T-DNA binary vector and the optimized pVir accessory (pPHP71539) plasmid with additional vir genes enhanced gene delivery and ultimately the transformation efficiency for both corn (Zea mays) and sorghum (Sorghum bicolor) (Anand et al., 2018; Che et al., 2018). This encouraged us to assess the gene delivery efficiency of the ternary vector for cowpea transformation.

To evaluate T-DNA delivery using the ternary vector system in cowpea, we transformed binary vector pPHP86170-10% EA explants (based on the average of three replicates and 75 EA explants) surviving EA explants with elongated epicotyls after 4 days of regeneration stimulated multiple shoot organogenesis around the cot-nodes (Figure 3(b)). This observation supported the hypothesis that only those cells within the cot-node tissue, especially around the cot-node tissue (Figure 3(b,c)), demonstrated highly efficient gene delivery in cowpea EA explants. Although gene delivery was efficient across the entire EA explant, only those fluorescent foci within the cot-node tissue showed subsequent development and substantially enhanced fluorescence intensity during regeneration (Figure 3(b,c)). This observation supported the hypothesis that only those cells within the cot-node region, but not any other tissues of the EA explant, actively undergo rapid cell division and dedifferentiation to acquire organogenic competence for shoot regeneration.

In vitro regeneration and transgenic selection of cowpea

Several selection systems have been reported for cowpea transformation with different explant types (Manman et al., 2013), such as NPTII/kanamycin (Bett et al., 2019;
Chaudhury et al., 2007), NPTII/G418 (Solleti et al., 2008b), PMI/mannose (Bakshi et al., 2012), HPT/hygromycin (Kumar et al., 1996), BAR/glufosinate (Popelka et al., 2006) and ahas/imazapyr (Citadin et al., 2013; Ivo et al., 2008). It was reported that incomplete selection and tissue necrosis were associated with those selection systems (Manman et al., 2013) and resulted in lower transgenic plant recovery (Bakshi et al., 2011; Chaudhury et al., 2007; Solleti et al., 2008b) and a higher percentage of chimeric tissue formation in cowpea (Das Bhowmik et al., 2019). To identify more efficient selection agents for the de novo organogenesis described above, we tested and compared the selection efficiency of CTP-NPTII/kanamycin, CTP-NPTII/G418 and CTP-spcN/SPEC (Anada et al., 2017) after Agrobacterium-mediated transformation using Agrobacterium strain LBA4404 Thy-. Ideally, the optimal concentration for effective selection is determined by testing different concentrations of selective reagents in SIM that completely inhibits the in vitro regeneration from the wild-type EA explants, but imposes minimal or no impact on transgenic shoot recovery and development. Since a CTP-spcN/SPEC selection system for EA-based soybean transformation has been well established and implemented for soybean transformation at a SPEC concentration of 25 mg L$^{-1}$ in Corteva, we decided to test three SPEC concentrations (15, 25 and 35 mg L$^{-1}$) for cowpea transformation and observed similar efficacy in terms of high transgenic shoot recovery with no transgenic escapes. Therefore, we chose a SPEC concentration of 25 mg L$^{-1}$ throughout the study. As described above, strong gene delivery was observed around the cot-node target tissue on the third day of regeneration as shown in Figure 3(b,c). Those fluorescent foci grew quickly, and single or multiple shoot buds emerged exclusively around the cot-nodes within 2 weeks following removal of the SAM on the fourth day of regeneration (Figure 3(d,e)). Transgenic shoots were fully developed from the buds within another 3 weeks in which all the shoots displayed strong fluorescence evenly across the entire regenerated shoot (Figure 3(f,g)) compared to the regenerated shoots from wild-type cowpea explants that showed no fluorescence at all (Figure 3(j,k)).

The elongated shoots were excised from the EA explants and transferred to root induction medium (RIM) (Table S6) for root development. Because of the stringent selection during shoot organogenesis, selection was not required for rooting. Approximately 95% (Table 1) of the elongated shoots fully rooted in the RIM within 2–3 weeks and all the regenerated shoots and roots displayed strong fluorescence (Figure 3(h,i)). The total time from inoculation of the EA explants to transplantation of a fully developed transgenic plantlet in the greenhouse took approximately 2–3 months. As shown in Table 1, the frequency of shoot formation was about 21% for IT86D-1010, of which about 23% of the events were single copy quality events (see the Experimental

Figure 3. Different stages of transgenic cowpea IT86D-1010 development using the CTP-spcN/SPEC selection system.
(a,b) T-DNA delivery, as determined by transient assay after 3 days on SIM.
(c) Enlarged view of the size, the intensity and the amount of fluorescent foci around the cot-node region. (d,e) Transgenic shoot budding after 2 weeks of regeneration on SIM with SPEC selection.
(f,g) Fully developed transgenic shoot after 5 weeks of regeneration with SPEC selection.
(h,i) Root development of regenerated transgenic shoot after 3 weeks of root induction on RIM.
(j,k) Autofluorescence evaluation of regenerated wild-type cowpea IT86D-1010. The arrows indicate the regenerated roots. (a,d,f,h,j) Bright field images. (b,c,e,g,i,k) Fluorescence images under RFP filter.

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Although the optimal concentration of G418 at 20 mg L⁻¹/C₀ was observed when a range of 17% to 33% (Table 3) was shown below in Table 3, a chimerism percentage in theogenic escapes nor chimera formation. Conversely, as more efficient and stringent transgenic selection and gen-

| # of EAs | # of transgenic shoots | # of chimeric shoots* | # of transgenic shoots with root developed | Rooting efficiency (%) | Transformation efficiency (%) | Quality events (%) |
|----------|------------------------|-----------------------|-------------------------------------------|------------------------|-------------------------------|-------------------|
| 30       | 7                      | 0                     | 6                                         | 86                     | 20                            | 23                |
| 30       | 7                      | 0                     | 7                                         | 100                    | 23                            |                   |
| 20       | 4                      | 0                     | 4                                         | 100                    | 20                            | 21±2**            |

*Chimeric shoots were identified by the presence of distinct sectors of cells exhibiting fluorescence, which were sharply demarcated from non-fluorescent tissue.

**Data are presented as the average rooting and transformation efficiencies ± SD of three biological replicates.

Procedures section for the definition and determination of ‘quality events’.

Chimeric tissue formation (a single plant tissue containing a mixture of transformed and non-transformed sections) during tissue culture transformation is a prevalent issue in legumes, including cowpea (Das Bhowmik et al., 2019). Chimeric plants affect the segregation of the transgene to the subsequent generation and reduce the efficiency of recovering stable transgenic lines (Das Bhowmik et al., 2019). Reporter genes, such as uidA (encoding GUS) (Das Bhowmik et al., 2019), GFP (Dutt et al., 2007) and Ds-RED (Xu et al., 2020), are often used to determine the uniformity of gene expression on the regenerated shoots for chimera formation. In this study, the formation of chimerism was evaluated for all the transgenic events generated using binary vector pPHP86170 (Figure S1(a)) containing the proDMMV:TagRFP as the visual marker and pPHP92782 (Figure S1(c)) containing the proGM-EF1A2:Ds-RED as the visual marker. In addition to the 18 transgenic shoots reported in Table 1, an additional 234 regenerated shoots generated using pPHP86170 (Figure S1(a)) and 59 regenerated shoots generated using pPHP92782 (Figure S1(c)) were screened and showed no signs of chimerism. Therefore, the utilization of the CTP:spcN/SPC selection system provided more efficient and stringent transgenic selection and generated transgenic cowpea events with neither non-transformed escapes nor chimera formation. Conversely, as shown below in Table 3, a chimerism percentage in the range of 17% to 33% (Table 3) was observed when a hypervirulent Agrobacterium strain, AGL1 (Lazo et al., 1991), and an alternate SPEC selection system, CTP:aadA SPEC (Martinell et al., 2017), were utilized.

In contrast to SPEC as the selection reagent, cowpea IT86D-1010 EA explants showed a high degree of resistance to kanamycin and no selection pressure could be built up by culturing wild-type EA explants on SIM containing kanamycin at concentrations as high as 600 mg L⁻¹. Although the optimal concentration of G418 at 20 mg L⁻¹ for selection of transformed shoots was established by culturing non-transformed EA explants on SIM containing different concentrations of G418 (10-40 mg L⁻¹), the G418 selection was not as stringent as for SPEC. As indicated by the uneven and partial fluorescence of the regenerated shoot in Figure S5, all five events generated from 110 EA explants were chimeric events using the binary vector pPHP94518 (Figure S1(b)) containing proGM-EF1A2:Ds-RED as a visual marker and proGM-UBQ:CTP-NPTII as the selectable marker. Because of the low transformation efficiency and high chimera formation rate under G418 selection, we stopped conducting further optimization and concluded that CTP-NPTII/kanamycin and CTP-NPTII/G418 were not efficient selection systems for cowpea EA-based transformation.

Transgene inheritance in the progeny

To evaluate the inheritance of T-DNA integration events, we selected nine independent single copy quality T0 transgenic events (lines) in the IT86D-1010 background transformed with construct pPHP92782 (Figure S1(c)) containing proGM-EF1A2:Ds-RED as a visual marker gene and proGM-UBQ:CTP-spCN as the selectable marker gene. Those T0 plants with chimera formation based on Ds-RED expression were self-pollinated in the greenhouse and the resultant T1 seeds displayed various levels of red color (Figure S6), because the overall intensity of the seed color is largely represented by the color of cotyledons passing through the semi-translucent seed coat, as described by Nishizawa et al. in soybean carrying the Ds-RED transgene (Nishizawa et al., 2006). Therefore, the variation of seed color intensity in T1 cowpea seeds could represent the level of transgene expression in the cotyledon and indicate transmission and segregation of the transgene in the progeny. To further determine the segregation ratio, T1 seeds from each of the nine independent events were randomly chosen regardless of seed color and advanced to the T1 generation. The zygosity (homozygous, hemizygous and null) of segregated T1 plants was characterized by determining the copy number of the integrated T-DNA based on the assays described in the Experimental Procedures section. As shown in Table 2, the segregation pattern of eight out of nine transgenic events showed a typical 1:2:1 Mendelian ratio in the T1 generation based on chi-square test
analysis, except event 125739949, which had a $P$-value of 0.04, which was barely lower than the threshold at 0.05. Nonetheless, 28 homozygous plants were still identified from 98 segregated T1 plants in this event. These results demonstrate that all the transgenic events analyzed possessed stably integrated T-DNA without obvious chimera formation and the T-DNA was successfully passed on to the cot-node region and, in most of the cases, no more than two transgenic shoots per explants were developed (Figure S7). These results demonstrate that the transformation protocol developed for IT86D-1010 described herein is transferable to other genotypes even with an alternative Agrobacterium-mediated transformation system. However, a high percentage of chimerism (in the range of 17% to 33%) (Table 3) was observed using this quick transformability assay, most likely attributable to the hypervirulent AGL1 Agrobacterium strain and/or the CTP-aadA/SPEC selection system.

In general, a better shoot organogenesis response tends to produce a higher transgenic shoot regeneration frequency, but this is not always the case. As shown in Tables 3 and S9, although all five genotypes, IT86D-1010, PI 527675, PI 580227, PI 582835 and TVu 79, showed very good and comparable shoot organogenesis rates (in the range of 78% to 81%), only IT86D-1010, PI 527675 and PI 582835 demonstrated significant high transformability efficiencies (19±7.5%, 31% and 37%, respectively), in contrast to PI 580227 and TVu 79 (5.6% and 4.5%, respectively). In contrast, all three genotypes, TVu 3562, TVu 9693 and PI 583259, had relatively low shoot organogenesis rates (68%, 61% and 56%, respectively) (Table S9), but their transformability efficiency was relatively high (18%, 22% and 26%, respectively) (Table 3). Therefore, the transformability of those germplasm lines is determined not only by the shoot organogenesis capability, but also by the combination of the susceptibility to Agrobacterium-mediated T-DNA delivery and sensitivity to Agrobacterium infection and sonication-related damage.

**Targeted cowpea genome editing with CRISPR/Cas9**

The lack of mutation resources and efficient means for gene inactivation has dramatically hampered the genetic improvement of cowpea for breeding. Targeted genome editing using the CRISPR/Cas system has proven to be a

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### Table 2 Segregation analysis of self-fertilized IT86D-1010 transgenic cowpea plants in the T1 generation

| Event ID   | Total # of plants analyzed | # of two copy plants | # of single copy plants | # of null plants | Expected segregation ratio | Chi-square test | P-value $^2$ | H$_0$ hypothesis $^3$ |
|------------|----------------------------|----------------------|-------------------------|-----------------|-----------------------------|-----------------|--------------|---------------------|
| 125739938  | 100                        | 17                   | 54                      | 29              | 1:2:1                       | 3.52            | 0.17         | Accepted            |
| 125739949  | 98                         | 28                   | 37                      | 33              | 1:2:1                       | 6.39            | 0.04         | Rejected            |
| 125739950  | 190                        | 41                   | 106                     | 43              | 1:2:1                       | 2.59            | 0.27         | Accepted            |
| 125739952  | 84                         | 19                   | 49                      | 26              | 1:2:1                       | 1.21            | 0.55         | Accepted            |
| 125739954  | 92                         | 25                   | 44                      | 23              | 1:2:1                       | 0.88            | 0.88         | Accepted            |
| 135577762  | 84                         | 19                   | 46                      | 19              | 1:2:1                       | 0.76            | 0.68         | Accepted            |
| 135577763  | 82                         | 21                   | 41                      | 20              | 1:2:1                       | 0.024           | 0.99         | Accepted            |
| 135577764  | 88                         | 28                   | 39                      | 21              | 1:2:1                       | 2.25            | 0.32         | Accepted            |
| 135577766  | 73                         | 15                   | 45                      | 13              | 1:2:1                       | 4.06            | 0.13         | Accepted            |

$^1$Copy numbers were determined by PSB1, PSA2, Ds-RED and spcN-SO qPCR assays (Experimental Procedures).

$^2$The observation ratio is considered to fit the expected segregation ratio of 1:2:1 if $P > 0.05$.

$^3$H$_0$ hypothesis: The segregated T1 population fits the 1:2:1 genotype ratio.

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powerful tool for crop engineering and has been successfully applied to maize, sorghum, soybean, rice \((\text{Oryza sativa})\) and numerous other plant species to generate stable genome-edited lines with targeted modifications (e.g., insertions, deletions and replacements). Recently, CRISPR/Cas-mediated genome editing in cowpea was demonstrated in non-inheritable mutated hairy roots \(\text{(Ji et al., 2019)}\) and stable, heritably edited, mutated plants \(\text{(Juranic et al., 2020)}\). To establish the CRISPR/Cas-mediated genome editing system in cowpea using the transformation protocol described herein and to test the feasibility for highly efficient editing, we designed a new cowpea gene editing construct, \(\text{pPHP96249}\), carrying \(\text{spCas9}\) driven by a soybean elongation factor \(\text{EF1A2}\) promoter \(\text{(Li et al., 2015)}\) and the \(\text{Vu-SPO11-CR2}\) single guide RNA \(\text{(sgRNA)}\) driven by the cowpea \(\text{U6}\) promoter \(\text{(Vu-U6.1)}\) \(\text{(Juranic et al., 2020)}\) to knock out the cowpea meiosis gene \(\text{Vu-SPO11-1}\) \(\text{(Juranic et al., 2020)}\) \(\text{Figure 4(b–e)}\). A total of 35 independent T0 transgenic plants were obtained from 250 EAs in the \(\text{IT86D-1010}\) background. All the T0 plants were analyzed by deep sequencing to identify mutations at the target site \(\text{Table S12 and the Experimental Procedures section}}\). As shown in Table 4 and Table S12, a genome editing efficiency of 68.6% was observed, in which 25.7% of the edited plants showed an edited read percentage higher than 30% of at least one allele. A higher allele read percentage generally indicates higher allele recovery potential in the next generation. This large population of T0 plants with high editing efficiency provided us with the opportunity to investigate the types of edit mutations in cowpea for the first time. As shown in Table 4 and Table S12, 70.8% of edited plants carried a 1-bp G deletion and 33.3% of the edited plants carried a multiple-bp deletion. The only type of insertion detected by deep sequencing was a 1-bp G insertion, at a frequency of 37.5%. Thus, deletions are the predominant type of edited mutation, especially the single base pair G deletion (70.8%), which results in a frameshift or knockout of the gene. Among the edited plants, 41.6% carried more than one edited allele forming chimeric mutations (Table 4). Remarkably, two biallelic knockouts (plants #1 and 9) were identified from 35 T0 transgenic plants by deep sequencing (Table 4 and Table S12). Plant #1 contained a 1-bp G deletion on both alleles (94% of allele reads) and plant #9 had two types of mutated alleles, a 1-bp G insertion (42% of allele reads) and a 22-bp deletion (52% of allele reads), all leading to a null mutation of \(\text{Vu-SPO11}\) \(\text{(Table S12)}\). Those two biallelic knockouts allowed us to characterize mutant phenotypes in the T0 generation. As shown in Figure 4(b–e), both T0 biallelic mutants showed complete infertility with no seed pods developed on the plants. On the contrary, all the other T0 heterogeneous mutations were totally fertile and produced pods and seeds. To demonstrate that the edited allele can be stably transmitted and to further confirm the fertility phenotype of \(\text{Vu-SPO11}\) mutation in the

| Germplasm | # of EAs | # of EAs with fluorescent shoots at 2 weeks | Transformability\(^1\) (%) | # of EAs with fluorescent shoots at 4 weeks | # of chimeric shoots\(^2\) | Chimerism rate\(^3\) (%) |
|-----------|----------|---------------------------------------------|--------------------------|---------------------------------------------|--------------------------|--------------------------|
| Experiment 1 | | | | | |
| \(\text{IT86D-1010}\) | 123 | 14 | 11 | 24 | 7 | 29 |
| \(\text{PI 527675}\) | 166 | 52 | 31 | 34 | 9 | 26 |
| \(\text{PI 580227}\) | 178 | 10 | 6 | 12 | 4 | 33 |
| \(\text{TVu 9693}\) | 160 | 35 | 22 | 43 | 9 | 21 |
| PI 80670 | | | | | |
| PI 582359 | 117 | 22 | 19 | 43 | 11 | 26 |
| TVu 79 | 157 | 7 | 4 | 22 | 6 | 27 |
| TVu 8670 | 125 | 11 | 9 | 13 | 3 | 23 |
| Experiment 3 | | | | | |
| \(\text{IT86D-1010}\) | 102 | 27 | 26 | NA | NA | NA |
| \(\text{PI 58235}\) | 90 | 33 | 37 | NA | NA | NA |
| \(\text{TVu 3562}\) | 177 | 32 | 18 | NA | NA | NA |

\(^1\)Transformability is defined as the number of EAs with fluorescent shoots divided by the total number of EA explants.

\(^2\)Chimeric shoots were identified by the presence of distinct sectors of cells exhibiting fluorescence, which were sharply demarcated from non-fluorescent tissue.

\(^3\)Chimerism rate is defined as the number of chimeric shoots divided by the total number of regenerated shoots with fluorescence at 4 weeks after transformation.

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Edits are defined as mutagenesis at target sites. In the subsequent generation, we characterized the target gene mutation in the progenies from three selected heterogeneous T0 events (#26, #28 and #29) (Table S12 and Table 5). As shown in Table 5, T1 heterozygous mutations and null mutations were recovered in the progenies of all three selected T0 events, and T1 homozygous mutations were identified in the progenies of both event #26 and event #28. These results demonstrate that the edited alleles are heritable and can be segregated to form heterozygous, homozygous and null genotypes in the next generation. Consistent with the observation of the two T0 biallelic knockouts (Figure 4(b–e)), all T1 homozygous identified in Table 5 were completely infertile with aborted seed pod development through plant maturity (Figure 4(g)). Contrarily, all the segregated null mutants and heterozygous plants were fertile with normal seed pod development and not distinguishable from the wild-type (Figure 4(a,f,h,i)). These observations further supported the functionality of Vu-SPO11 in meiosis, as characterized by Juranic et al. (Juranic et al., 2020). Overall, these results demonstrated that we have established a highly efficient CRISPR/Cas9-mediated editing system for selectively altering genome DNA sequences in cowpea.

In summary, we have developed a rapid, robust, flexible and highly efficient cowpea transformation and CRISPR/Cas9-mediated genome editing system using EA as explants. The principles established in this study have the potential to increase the transformation efficiencies of other legume species and, potentially, other dicot crops. With recent progress establishing cowpea genetic and genomic resources (Lonardi et al., 2019; Munoz-Amatriain et al., 2017; Spriggs et al., 2018; Yao et al., 2016), we believe that the broad application of this cowpea transformation and editing system will have an immediate and far-reaching impact on cowpea research that will improve cowpea productivity.

**EXPERIMENTAL PROCEDURES**

**Agrobacterium strains and vectors**

Two *Agrobacterium* strains, the auxotrophic strain LBA4404 Thy- and AGL1, were used in this study. *Agrobacterium* auxotrophic strain LBA4404 Thy- was used with the ternary vector transformation system for cowpea IT86D-1010 transformation. The ternary vector system contains the T-DNA binary vector and the optimized pVIR accessory (pPH71539) plasmid as previously described by Che et al. (Che et al., 2018) and Anand et al. (Anand et al., 2018). The T-DNA binary plasmid pPHP86170 (Figure S1(a)) contains the PUC ORI, the NPTII bacterial selectable marker, the TagRFP reporter gene and spcN (Anada et al., 2017) as the plant selectable marker gene. The binary plasmid pPHP94518 (Figure S1(b)) contains the PVS1 ORI, the SPEC bacterial selectable marker, the De-RED reporter gene and NPTII as the plant selectable marker gene. The

### Table 4 CRISPR/Cas9-mediated gene editing efficiency and types for Vu-SPO11 in IT86D-1010

| Total # of T0 edited plants | # of T0 edited plants with allele read percentage > 30% | # of T0 edited plants with biallelic knockouts | # of T0 edited plants with chimeric edits (≥2 edited alleles) | # of T0 edited plants with multiple-bp deletion | # of edited T0 plants with 1-bp G deletion | # of edited T0 plants with 1-bp G insertion |
|-----------------------------|---------------------------------------------|-----------------------------------------------|-------------------------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| 35                          | 24 (68.6%)^4                               | 9 (25.7%)^4                                   | 2 (5.7%)^4                                       | 10 (41.6%)^5                                 | 8 (33.3%)^5                                 | 17 (70.8%)^5                                 | 9 (37.5%)^5                                 |

^1Edits are defined as mutagenesis at target site.
^2See Table S12 for the allele read percentage in each plant.
^3Biallelic knockout is defined as targeted mutagenesis that results in a frameshift mutation for both alleles.
^4The number in parentheses represents the frequency of the mutation type in the total T0 plants.
^5The number in parentheses represents the frequency of the mutation type in the total edited plants detected.

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(a) The fertile phenotype of wild-type cowpea IT86D-1010 with mature pod development. (b–e) The infertile phenotype of T0 biallelic Vu-SPO11 mutants, plant #1 (b) and plant #9 (d). (c,e) Enlarged view of a portion of biallelic Vu-SPO11 mutants, plant #1 (b) and plant #9 (c).

(g–i) Fertility phenotypes of the T1 segregants, homozygous (g), heterozygous (h) and null mutant (i), of edited T0 plant #26. Seed pods are indicated by green arrows. Floral sectors with pod abortion are indicated by yellow arrows.
production binary plasmid pHPR2782 (Figure S1(c)) contains the PVS1 ORI, the NPTII bacterial selectable marker, the Ds-RED reporter gene and spcN (Anada et al., 2017) as the plant selectable marker gene. The ternary design was assembled by first mobilizing the accessory plasmid pHPR71539 in the LBA4404 Thy- containing the accessory plasmid pPHP71539 (Figure S1(e)). The spCas9 is driven by a soybean elongation factor EF1A2 promoter (Li et al., 2015) and the sgRNA cassette fused with CRISPR RNAs (crRNA) and trans-activating crRNA is driven by a cowpea U6 promoter (Vu-U6). (Juranic et al., 2020). The N20 region of the crRNA NA hybridization region represents the RNA sequence used to target the genomic sequence upstream of the DNA triplet ‘NGG’ in the VU-SPO11 gene (Figure S8) that is recognized by the spCas9 enzyme. The guide RNA Vu-SPO11-CR2 (Figure S8) is the same as SPO11-sg3 as described by Juranic et al. (2020).

Plant materials and growth conditions

Cowpea varieties IT86D-1010, PI 527675, PI 580227, PI 582835, PI633259, TVu 8670, TVu 3562, TVu 9693 and TVu 79, originally obtained from the U.S. NPGS (https://www.ars-grin.gov/npgs/), were used for this study. Mexican cowpea accessions TPC1-001 and MRS-001 were collected from local farming communities in the Mexican states Tabasco and Morelos, respectively; the common bean varieties black bean (CBB-001) and pinto bean (CBP-001) were obtained from local producers in the Mexican state Guanajuato. Those varieties were maintained in the greenhouse to collect mature seeds for EA explant isolation.

Cowpea transformation procedure

The main steps of cowpea transformation mediated by Agrobacterium are illustrated in Figure S2. The detailed protocol is described below. Unless otherwise specified, all the chemicals used for medium preparation were obtained from Sigma-Aldrich.

Agrobacterium preparation.

1 Master plate preparation: Streak Agrobacterium from glycerol stock on the master plate medium (Table S1) containing different antibiotics based on the Agrobacterium strains and the constructs that bacterium carries to make master plates. Incubate the master plates at 28°C for 3-4 days. The master plates can be kept in the refrigerator to make working plates and last for a month.

2 Working plate preparation: Streak a working plate on the working plate medium (Table S2) using a loop of bacteria from the master plate prepared above and incubate the working plate at 28°C overnight or for 20 h for LBA4404 Thy- and AGL1, respectively.

3 Incubation cycle: Collect five to seven full loops of bacteria from the working plate using a sterile loop, suspend bacteria in 30 ml infection medium (IM) (Table S3) with acetosyringone (1 m stock in DMSO protected from light, final concentration, 200 µM) and dithiothreitol (DTT, 1 m stock, final concentration, 1 mM) freshly added in a sterile 50-ml centrifuge tube. Adjust OD to 0.5.
Cowpea EA explant preparation.
1 Seed sterilization: Cowpea seeds are surface-sterilized using chlorine gas made by mixing in 3.5 ml of 12 M HCl and 100 ml bleach (5.25% sodium hypochlorite) for 16 h.
2 Seed pre-treatment: Soak sterilized cowpea I790D-1010 seeds in bean germination medium (BGM) (Table S4) with approximately 45 ml of water for approximately 16 h. For other cowpea varieties transformed with AGL1, 30 ml 0MS medium (Table S8) is used to replace BGM.
3 EA explant isolation: Isolate embryonic axis (EA) explants with plumules by removing the seed coats and cotyledons and placing them into sterile water in a Petri dish until infection.

Agrobacterium inoculation.
1 Infection: Remove water from the Petri dish (as much as possible), and add 15 ml inoculum and 50 µl sterile Poloxamer 188 10% solution. Wrap the plate with parafilm and sonicate (VWR Motel 50T or FS30H, Fisher Scientific, 120 V, 1 A) for 3 sec. After sonication, add an additional 10 ml of inoculum (total 25 ml in Petri dish) to the mix and gently shake on a shaker at approximately 60 rpm for 1.5 h at room temperature.
2 Co-cultivation: Remove bacteria and transfer EAs to filter paper (VWR Cat No. 28320-020) blotted with 700 µl sterile Poloxamer 188 10%. Wrap the plate with parafilm and sonicate (VWR Motel 50T or FS30H, Fisher Scientific, 120 V, 1 A) for 3 sec. After sonication, add an additional 10 ml of inoculum (total 25 ml in Petri dish) to the mix and gently shake on a shaker at approximately 60 rpm for 1.5 h at room temperature.

Cowpea regeneration.
1 Regeneration with selection: Insert the roots of EAs vertically into SIM (Table S5) with cot-node and SAM above the medium. Incubate the EAs on SIM at 26°C under 24 h light conditions. Remove the SAM and plumules together by cutting through the middle of the epicotyl using a pair of surgical scissors after 4–5 days of culturing on SIM to promote axillary shoot formation at the cot-node region.
2 Rooting: After 3–5 weeks of regeneration, harvest shoots larger than 3 cm by cutting at the base of the shoot, and place into RIM (Table S6).
3 Shoot elongation (optional): If shoots do not reach 3 cm after 3–5 weeks of regeneration, they should be transferred to shoot elongation medium (SEM) (Table S7) for 2–4 weeks before transfer to RIM.

Microscopy and imaging
Images were taken using a dissecting Leica M165 FC stereo-epifluorescence microscope, with RFP and Ds-RED filters for the detection of fluorescence, using the PLANAPo 1.0 × objective, 0.63 × zoom and Leica Application Suite V4.7 acquisition software. The autofluorescence of the wild-type regenerated cowpea was evaluated using the same system. For testing transformation on the eight additional cowpea germplasm lines from NPGS, transgenic shoots expressing TdTomato were monitored with a Stemiview 77 dissemination system equipped with an HBO illuminator (Carl Zeiss, Thornwood, NY) and a Ds-RED filter (excitation: 545/25 nm, emission: 605/70 nm, Chroma Technology, Bellows Falls, VT). Images were taken with an AxioCam camera (Carl Zeiss, Oberkochen, Germany) and AxioVision LE4 software and composed using Photoshop CC (Adobe, San Jose, CA).

Evaluation of transgenic plants
The integrated copy number of the T-DNA of the binary vector in the transgenic plants was determined by a series of quantitative PCR (qPCR) analyses based on the method previously described by Wu et al. (2014). In this study, an endogenous control qPCR assay (LBS) (Table S10) was developed using the house-keeping gene 3-isopropylmalate dehydrogenase (Vignaung5g298700), which is involved in the leucine biosynthetic pathway in cowpea (Misra et al., 2017). Five qPCR assays (PINII_TERM, PSJ, spcN_SO, CTP and UBQ14_TERM) for pHPP86170 (Figure S1(a) and Table S10) and five qPCR assays (Ds-RED, spcN_SO, PSJ, PSA2 and PSB1) for pHPP92782 (Figure S1(c) and Table S10) were developed to determine the T-DNA copy number by normalizing with the endogenous control LBS assay. Two qPCR assays, PSA2 and PSB1, were specially designed for pHPP92782 just within the right and left border regions to determine not only the copy number, but also the integrity of the integrated T-DNA (Figure S1(c)).

Outside the border integration sites, PCR backbone-specific assays were developed to check for any border read-through (Wu et al., 2014). The presence or absence of Agrobacterium vector backbone integration of the binary vector was detected based on screening for sequences from three regions outside of the T-DNA integration sites for each vector, such as SPC, LEFTBORDER and NPTIII for plasmid pHPP92782 and HYG, VIRG and HYGROMYCIN for plasmid pHPP86170 (Figure S1(a) and Table S10).

Stable T-DNA integration was confirmed by copy number determination using genomic DNA extracted from the putative T0 transgenic events. The T0 transgenic plants carrying a single copy of the intact T-DNA integrations without vector backbone for all assays described were defined as quality events (Che et al., 2018; Zhi et al., 2015). The percentage of quality events was divided by the total number of events analyzed to calculate the quality event frequency. Only quality events were advanced to the greenhouse for the next generation. The zygosity of the T1 plants was established by determining the copy number of the T-DNA for all the event quality assays (Figure S1 and Table S10). Chi-square analysis was performed to determine whether the difference between the observed and the expected ratio was statistically significant.

Amplicon deep sequencing
CRISPR/Cas edits were characterized by amplicon sequencing from DNA extracted from a single fresh leaf punch from each plant as per the manufacturer recommendations via the seabdex™ tissue extraction system (LGC Limited, UK). DNA concentration was adjusted to 10 ng ml⁻¹ and 20-cycle target region PCR was performed on 50 ng of genomic DNA with Phusion™ Flash 2× master mix (Thermo Scientific, Waltham, MA) as per the manufacturer’s recommendations. Primary PCR product (5 µl) was used for 20-cycle secondary amplification containing primers to attach individual sample indices and sequencing components, again with Phusion™ Flash 2× master mix. Paired-end sequencing was performed on an Illumina MiSeq®, with 150 cycles per read following the Illumina standard operating procedure. Sequence reads were aligned to the wild-type reference sequence via Bowtie2. The allele return threshold was set to 5%. The primers used to amplify VU-SPO11 genomic loci are listed in Table S11. Edited sequences are reported in Table S12.

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**AUTHOR CONTRIBUTIONS**

PC, SC, MS, MA and TJ conceptualized the methods. PC, SC, MS, ZZ, JV, PO, MA and TJ designed research, analyzed the data and wrote the paper. AS conducted cowpea transformation and collected the data for IT86D-1010. ZZ, YG and KM conducted transformation and genotyping for other cowpea genotypes collected from NPGS. JO and DO developed event quality assays. MT conducted regeneration experiments for Mexican cowpeas and beans.

**CONFLICT OF INTEREST**

The authors have no conflict of interest to declare.

**DATA AVAILABILITY STATEMENT**

All relevant data supporting the findings of this work are available within the manuscript and the supporting materials. Novel biological materials described in this publication may be available to the academic community and other not-for-profit institutions solely for non-commercial research purposes upon acceptance and signing of a material transfer agreement between the author’s institution and the requester. In some cases, such materials may originally contain genetic elements described in the manuscript that were obtained from a third party (e.g., TagRFP, Ds-RED and Cas9), and the authors may not be able to provide materials including third-party genetic elements to the requester because of certain third-party contractual restrictions placed on the author’s institution. In such cases, the requester will be required to obtain such materials directly from the third party. The authors and the authors’ institution do not grant any express or implied permission(s) to the requester to make, use, sell, offer for sale or import third-party proprietary materials. Obtaining any such permission(s) will be the sole responsibility of the requester. In order to protect Corteva Agriscience™ proprietary germplasm, such germplasm will not be made available except at the discretion of Corteva Agriscience™ and then only in accordance with all applicable governmental regulations.

**SUPPORTING INFORMATION**

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

**Figure S1.** Schematic representation of the constructs used in this study.

**Figure S2.** Diagram of the cowpea EA-based Agrobacterium-mediated transformation process. The detailed transformation procedure is described in the Experimental Procedures section.

**Figure S3.** Dry mature seeds of selected accessions of cowpea and common bean.

**Figure S4.** Shoot organogenesis of selected accessions of cowpea and common bean.

**Figure S5.** Development of chimeric events using the CTP-NPTII/G418 selection system. (a) Bright field image. (b) Fluorescence image under RFP filter.

**Figure S6.** Transgene segregation in the progeny. (a) Mature wild-type cowpea IT86D-1010 seeds. (b) Segregated T1 seeds in the IT86D-1010 background harvested from T0 plants containing proGM:EF1A2:Ds-RED as visual marker.

**Figure S7.** Formation of transgenic shoots expressing TdTomato on the EA explants of nine cowpea germplasm lines after 14-day culture on SIM. Bar = 1 mm.

**Figure S8.** Diagram of the DNA sequence of the VU-SP011 target site in exon 3.

**Table S1.** Master plate medium.

**Table S2.** Working plate medium

**Table S3.** Infection medium (IM).

**Table S4.** Bean germination medium (BGM).

**Table S5.** Shoot induction medium (SIM).

**Table S6.** Root induction medium (RIM).

**Table S7.** Shoot elongation medium (SEM).

**Table S8.** OMS.

**Table S9.** Shoot organogenesis of selected accessions of cowpea and common bean.

**Table S10.** Primers used for event quality assays.

**Table S11.** Primers for the CRISPR/Cas target site.

**Table S12.** Sequence changes in Cas9-edited plants.

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