

ORIGINAL RESEARCH



Large genomic introgression blocks of *Phaseolus parvifolius* Freytag bean into the common bean enhance the crossability between tepary and common beans

Santos Barrera¹ | Jorge C. Berny Mier y Teran² | Juan David Lobaton³ | Roosevelt Escobar⁴ | Paul Gepts² | Steve Beebe⁴ | Carlos A. Urrea¹

¹Department of Agronomy and Horticulture, University of Nebraska, Lincoln, Nebraska, USA

²Department of Plant Sciences, University of California, Davis, California, USA

³Department of Evolutionary Biology, National Australian University, Canberra, Australia

⁴International Center for Tropical Agriculture (CIAT), Palmira, Colombia

Correspondence

Santos Barrera Lemus, Department of Agronomy and Horticulture, University of Nebraska, Lincoln, NE, USA.
Email: sybarreral@huskers.unl.edu

Funding information

International Center for Tropical Agriculture (CIAT); Nebraska Dry Bean Commission

Abstract

The production of the common bean (*Phaseolus vulgaris* L.), one of the most important sources of protein and minerals and one of the most consumed grain legumes globally, is highly affected by heat and drought constraints. In contrast, the tepary bean (*Phaseolus acutifolius* A. Gray), a common bean-related species, is adapted to hot and dry climates. Hybridization to introduce complex traits from the tepary bean into the common bean has been challenging, as embryo rescue is required. In this study, we report three novel interspecific lines that were obtained by crossing lines from prior common bean × tepary bean hybridization with *Phaseolus parvifolius* Freytag in order to increase the male gametic diversity to facilitate interspecific crosses. These interspecific lines enhanced the crossability of the common bean and tepary bean species while avoiding the embryo rescue process. Crossing these three interspecific lines with tepary beans resulted in 12-fold more hybrid plants than crossing traditional common beans with tepary beans. Whole-genome sequencing analysis of these three interspecific lines shows large introgressions of genomic regions corresponding to *P. parvifolius* on chromosomes that presumably contribute to reproductive barriers between both species. The development of these lines opens up the possibility of increasing the introgression of desirable tepary bean traits into the common bean to address constraints driven by climate change.

KEYWORDS

embryo rescue, haplotype blocks, interspecific, introgressions, recombinant populations, whole-genome sequencing

1 | INTRODUCTION

The common bean (*Phaseolus vulgaris* L.) is one of the most highly consumed legumes globally, yet climate change threatens its ability to adapt to abiotic and biotic stressors (Beebe, 2012). In the Americas and Africa, the common bean is an essential source of protein,

carbohydrates, and minerals (Beebe, 2012; Broughton et al., 2003). The production of common beans relies mainly on small-farm agriculture, where farmers depend on crop production for their livelihood (Beebe, 2012; Broughton et al., 2003). However, due to abiotic and biotic constraints, common bean yield is relatively low in Latin America and Africa, where it seldom exceeds 900 kg ha⁻¹ (Beebe, 2012).

This is an open access article under the terms of the [Creative Commons Attribution](https://creativecommons.org/licenses/by/4.0/) License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2022 The Authors. *Plant Direct* published by American Society of Plant Biologists and the Society for Experimental Biology and John Wiley & Sons Ltd.

Furthermore, common bean production is negatively affected by heat and drought, which have increased due to climate change. These constraints further threaten common bean production in Latin America and Africa (Beebe et al., 2013; Ramirez-Cabral et al., 2016).

One potential solution to improve the common bean's ability to tolerate warmer and drier conditions is the introgression of tepary bean traits (*Phaseolus acutifolius* A. Gray: Dohle et al., 2019; Moghaddam et al., 2021). Of the five domesticated *Phaseolus* species, the tepary bean is the species with the strongest drought and heat tolerance (Dohle et al., 2019). It is a desert-adapted species, which is related to the common bean, and has a unique array of abiotic tolerance traits (Pratt & Nabhan, 1988). These traits include heat tolerance (Porch et al., 2021; Traub et al., 2018) and water use efficiency (Markhart, 1985; Polania et al., 2020). However, crossing common bean with tepary bean is challenging due to reproductive isolation barriers between the two species (Gaur et al., 2009; Waines et al., 1988). To obtain viable hybrid plants in common bean/tepar bean crosses requires numerous rounds of pollination, embryo rescue, tissue culture, and backcrosses to alternate parents (congruity backcrossing: Haghighi & Ascher, 1988; Mejía-Jiménez et al., 1994). Embryo rescue, an in vitro culture technique used to help developing plant embryos that will not survive in vivo, minimizes the hybridization to individual plants rather than large recombinant populations (Waines et al., 1988). In addition, alternating backcrosses limit the possibility of introducing significant variation from the tepary bean to the common bean (Mejía-Jiménez et al., 1994).

Although the interspecific hybridization of tepary and common beans has been conducted since 1956, few efforts have resulted in advantageous common bean/tepar bean lines. Most of the successful introgression studies have been focused on moving the resistance to common bacterial blight from the tepary bean to the common bean (McElroy, 1985; Singh et al., 2001; Thomas & Waines, 1984). Very few introgression studies have been focused on moving the drought tolerance from tepary beans to common beans (Souter et al., 2017), and no studies have reported a successful introduction of heat tolerance from tepary beans to common beans.

To address this gap, we crossed several tepary beans with elite common bean lines, but our initial effort was hindered by the high level of reproductive incompatibility between both species. However, one cross, which combines three species, *P. vulgaris*, *P. acutifolius*, and *P. parvifolius* resulted in three interspecific lines with enhanced crossability between tepary and common beans while avoiding embryo rescue; furthermore, they produced a large number of hybrids compared to earlier crosses.

In this research, we sought to increase the crossability of *vulgaris-acutifolius* crosses by broadening male gametic diversity. Earlier research by Wall and York (1960) and Pratt et al. (1985) had shown that gametic diversity obtained by using pollen of F1 or other hybrid plants instead of pure lines in *Phaseolus* and *Cucurbita* increased crossability in these two genera. To increase gametic diversity in *P. acutifolius*, we chose to hybridize it with *P. parvifolius*, a species closely related to the tepary bean. Indeed, its habitat overlaps partially with that of the tepary bean, and it is morphologically similar to

P. acutifolius var. *tenuifolius* (Freytag & Debouck, 2002). It belongs to the tepary bean's secondary gene pool (Dohle et al., 2019; Zink & Nagl, 1998). Thus, this species gives partially viable hybrids when it is crossed with *P. acutifolius* accessions (Freytag & Debouck, 2002). AFLPs and microsatellite molecular markers have shown a very close genomic relationship between *P. parvifolius* and *P. acutifolius* (Muñoz et al., 2006; Zink & Nagl, 1998).

In this study, we further confirm the close relationship between *P. acutifolius* and *P. parvifolius* using nuclear genome sequencing data. In addition, we describe the development of three interspecific lines resulting from a triple interspecific cross (*vulgaris-acutifolius-parvifolius* [VAP]) and the analysis of their whole-genome sequencing (WGS). The WGS results revealed large introgression blocks of *P. parvifolius* in the genomes of the three interspecific lines, which may explain the increased crossability between tepary and common beans, obviating the need for embryo rescue. Moreover, the results demonstrate why these three interspecific lines can be used potentially as bridge parents for introducing desirable, but genetically complex, traits such as heat and drought tolerance from tepary beans to common beans.

2 | MATERIALS AND METHODS

2.1 | Initial interspecific common-tepar bean hybridization

Initially, 11 Mesoamerican common bean lines were crossed with eight tepary bean genotypes and one *P. parvifolius* accession, which had been identified after a field screening under hot and humid conditions in northern Colombia (Supplementary Table S1). Crosses were carried out in the field at the International Center for Tropical Agriculture (CIAT) in Palmira, Colombia, in 2015, following the procedures described by Bliss (1980). Each parental combination included 50 manual pollinations. The common bean lines were used as female parents, whereas tepary beans and the *P. parvifolius* accession were used as male parents. Because early flower abscission and pod abortion occurred, embryos were rescued by removing immature pods from the plant before they abscised. Thirty-six immature pods from eight crosses were removed approximately 5–7 days after pollination and carried to the tissue culture laboratory at CIAT to conduct embryo rescue. All pods were approximately 3–5 cm long at the time of removal.

2.2 | Embryo rescue

One hundred fifty-two embryos were rescued using the method of Mejía-Jiménez et al. (1994), except that the gelling agent consisted of .2% Gelrite instead of .6% agarose. Briefly, the pods were sterilized for 1 min using 70% ethanol, followed by a 6-min treatment with 2.5% sodium hypochlorite. Next, the pods were washed three times with sterile deionized water and dissected to extract embryos from the seeds. The embryos were then transferred to a modified MS



medium (Murashige & Skoog, 1962) that was prepared as described by Mejía-Jiménez et al. (1994). Next, the embryos were moved first to a growth chamber ($25 \pm 1^\circ\text{C}$ and 75% relative humidity) for 2 weeks and then to a simulated photoperiod room with 12-h dark and 12-h light (1000 lux) until their primary leaves were fully expanded and their roots had initiated their development.

2.3 | Plant hardening and fertility recovering

Eighty-seven fully developed plants from the eight hybrid combinations were transplanted from in vitro tubes into pots with a soil-sand mixture in a 2:1 proportion by weight. The pots were placed in a growth chamber with a continuous nebulization water supply, where the maximum temperature did not exceed 20°C . One week later, the plants were moved to a greenhouse (26°C daytime and 19°C nighttime temperature). In the first round of crosses, only three hybrid plants from the cross between INB834 (female parent) and G40264 (*P. parvifolius*) (male parent) survived. These hybrids were self-sterile and were subsequently backcrossed with INB 834 or crossed with INB 841 (three-way cross) (Supplementary Figure S1). The two INB lines (INB834 and INB841) used in this study are interspecific common bean/teparty bean lines that resulted from fertile crosses between the interspecific lines INB605 and INB108, which have been reported as tolerant to drought stress (Beebe et al., 2012). INB605 and INB108 are interspecific lines that resulted from the congruity backcrosses and embryo rescue between *P. vulgaris* and *P. acutifolius* published by Mejía-Jiménez et al. (1994) and Muñoz et al. (2004). After the second round of crosses, early flower abscission was observed again, and only five pods could be recovered for embryo rescue. Twenty embryos were collected from these pods, but only three hybrid self-sterile plants were obtained. Since fertility was not fully recovered after the second round of crosses, a third round of alternate crosses with INB lines, used as male parents, was needed to fully recover the fertility (Supplementary Figure S1). After fertility was recovered, 132 seeds were advanced under greenhouse conditions for three generations of inbreeding (F_4) (Supplementary Figure S1).

2.4 | Identification of the VAP lines

One hundred thirty-two families from the F_4 generation were planted in 2017 in a net-covered screenhouse in Santander de Quilichao, Colombia (26°C daytime and 19°C nighttime temperature). To introduce a tepary bean background in these interspecific families, 16 of these families were crossed with four male tepary accessions (one cultivar: TARS-TEP22 [Porch et al., 2013]; three gene bank accessions: G40019, G40068, and G40119). Most of the pollinations either aborted or produced pods without seeds. However, in three families, the pollinations did not abort, and pods and seeds successfully matured on the plant. This result suggested that these families could be crossed directly with tepary beans, thereby avoiding the embryo rescue step. These families were named as experimental lines VAP1,

VAP2, and VAP3; VAP stands for *P. vulgaris*, *P. acutifolius*, and *P. parvifolius*.

To corroborate that these three VAP lines could be crossed with tepary beans without embryo rescue and used as bridge lines, we crossed them with 22 different wild and domesticated (landraces and improved lines) tepary beans (used as the males) (Supplementary Table S2). To compare the crossability of the VAP lines with other common bean lines, we crossed the same set of tepary beans with common beans (Supplementary Table S2). Approximately 8–12 pollinations were made per cross-combination. All pods from crosses with the VAP lines reached maturity. On the other hand, all pods from crosses with common bean genotypes exhibited early abscission and were, therefore, taken to the laboratory for embryo rescue. To confirm hybridization, hybrid plants from crosses with VAP lines as well as common beans were morphologically characterized using the traits listed in Supplementary Table S3.

2.5 | WGS study and reference genome mapping

To identify the genomic regions that enhance the crossability of the VAP lines with tepary beans, we conducted a WGS analysis of the VAP line parents (INB841, INB834, and G40264) and the three VAP lines (VAP1, VAP2, and VAP3). Due to high phenotypic variability, the WGS analysis was conducted in duplicate on the VAP lines. The WGS reads of G19833 (a bean line belonging to the Andean domesticated gene pool), G40001 (teparty bean), and RWR 719 (a Mesoamerican domesticated bean) were added as outgroup and introgression controls from the study published by Lobaton et al. (2018). The DNA extraction was performed following the methodology reported by Ariani et al. (2016). The WGS analysis was conducted at the genome center of the University of California, Davis, on an Illumina HiSeq 4000, with 100-bp single-end fragment length. FastQC (Andrews, 2010) was used to analyze the quality of the DNA sequence reads, and Trimmomatic (Bolger et al., 2014) was used to trim the adaptors and low-quality reads. The reads were mapped to the latest reference genomes *P. vulgaris* G19833 Andean (version 2.1, Schmutz et al., 2014; Phytozome Version 13: https://phytozome-next.jgi.doe.gov/info/Pvulgaris_v2_1) and *P. acutifolius* G40001 (Moghaddam et al., 2021; wild tepary: Phytozome Version 13: https://phytozome-next.jgi.doe.gov/info/PacutifoliusWLD_v2_0; domesticated tepary: Phytozome Version 13: https://phytozome-next.jgi.doe.gov/info/Pacutifolius_v1_0), using “Bowtie2” (Langmead & Salzberg, 2012).

2.6 | Interspecific introgression assignment

The identification of sequence variants was performed with the “Next Generation Sequencing Experience Platform” (NGSEP), using the “FindVariants” command with a Q40 quality score (Duitama et al., 2014). The resulting SNP matrix was filtered, leaving only sites where all the samples have SNPs data using the “VCFFilter” module

with the `-m` option. The genotype classification was calculated using the “DistanceMatrixCalculator” command, which uses identical-by-state (IBS) algorithms to address relatedness among individuals (Duitama et al., 2014; Tello et al., 2019). Genotypes were visualized with Flap-Jack (Milne et al., 2010). The interspecific introgression analysis was conducted with the “VCFIntrogression Analysis” module of NGSEP using 100 SNPs windows analysis, following the methodology described by Lobaton et al. (2018). Introgression was assigned if more than 80% of the SNPs were identical to one of the parental lines. In this analysis, we used G40264 as the *P. parvifolius* haplotype reference and G19833 as the *P. vulgaris* haplotype reference. The shared alleles were calculated based on the overall window similarities to G19833 and G40264 haplotypes. By doing this, we were able to determine the location of the interspecific introgressions and how these introgressions differed among the VAP lines and the INB lines.

3 | RESULTS

3.1 | Close phylogenetic relationship between *P. acutifolius* and *P. parvifolius* based on genome sequence data

The choice of the *P. parvifolius* accession (G40264) to increase gametic diversity was dictated by its phylogenetic closeness to *P. acutifolius* and its agronomic quality, namely, its tolerance to humid heat stress in a field trial on the Caribbean coast of Colombia (S. Barrera, unpubl. results). The percentage of the whole genome sequence reads of *P. parvifolius* mapped to the *P. acutifolius* genome reference was 97% (Table 1), whereas the percentage of whole genome sequence reads of *P. acutifolius* assigned to *P. parvifolius* was 98% (Table 2), confirming a close genomic relationship between these two species. Additionally, the genetic distance matrix tree (Figure 1)

shows that there was a closer genomic relationship between *P. acutifolius* (G40001) and *P. parvifolius* (G40264) than *P. acutifolius* and the interspecific lines (VAP and INB lines) and a common bean line (RWR719).

3.2 | Initial interspecific hybridization, embryo rescue, and plant hardening

To introduce genetic variation from the tepary bean to the common bean, we crossed 11 common bean elite experimental lines with eight *P. acutifolius* accessions and one *P. parvifolius* accession (Supplementary Table S1). Hybridization between *P. vulgaris*, *P. acutifolius*, and *P. parvifolius* was challenging as strong reproductive incompatibility was present. In eight out of 11 initial crosses, 93% of pollinated flowers abscised early or produced empty pods, and 136 embryos were rescued from 36 immature pods. From the 136 rescued embryos, only three plants from the cross with G40264 (*P. parvifolius*) survived the hardening step. These three plants represent a hybridization rate of .005 plants per pollination from the 550 pollinations that were made in the initial 11 crosses. The three plants were self-sterile, and it was necessary to make two consecutive crosses using INB lines as the male parent to fully recover fertility, which resulted in 132 inbred interspecific F_4 families (Supplementary Figure S1).

The initial challenges experienced while crossing tepary and common beans confirm the reproductive barriers observed previously between the two species (Andrade-Aguilar & Jackson, 1988; Mejía-Jiménez et al., 1994; Mok et al., 1978; Parker & Michaels, 1986; Smartt, 1970). Moreover, the observation that most pollinated flowers, from the cross between common bean and *P. parvifolius*, abscised early suggests a high level of reproductive incompatibility between these two species. Reproductive incompatibility between

TABLE 1 Alignment percentage of the whole genome sequence reads to the tepary bean reference genome (G40001) and Andean common bean reference genome (G19833)

Genotype	Genotype group	Total reads	Mapping percentage (%) to the common bean reference genome	Mapping percentage (%) to the tepary bean reference genome
VAP1_R1	Bridge line	41,280,074	96.6	88.8
VAP1_R2	Bridge line	48,043,157	97.2	89.0
VAP2_R1	Bridge line	44,831,917	96.1	89.8
VAP2_R2	Bridge line	34,774,461	96.7	89.7
VAP3_R1	Bridge line	40,358,828	96.7	88.5
VAP3_R2	Bridge line	34,909,203	96.0	89.2
INB841	Interspecific	36,755,518	97.1	88.9
INB834	Interspecific	41,323,904	97.1	88.4
RWR719	Common bean (Mesoamerican)	41,877,662	98.0	86.0
G19833	Common bean (Andean)	331,924,075	97.0	81.4
G40001	Tepary bean (<i>Phaseolus acutifolius</i>)	32,999,285	68.2	82.3
G40264	Tepary bean (<i>Phaseolus parvifolius</i>)	37,433,199	88.4	96.8



TABLE 2 Interspecific introgressions: Number of genomic regions and percentage assigned to either *Phaseolus parvifolius* or *Phaseolus vulgaris* whole genome sequence reads

Genotype	Genotype group	<i>P. parvifolius</i> Genome	<i>P. vulgaris</i> genome	<i>P. parvifolius</i> (%)	<i>P. vulgaris</i> (%)	Unmapped	Unmapped (%)	Hetero-zygous	Hetero-zygous (%)
G19833	<i>P. vulgaris</i> (Andean)	-	30,693	-	100	-	-	-	-
G40001	<i>Phaseolus acutifolius</i>	30,132	-	98.2	-	-	-	561	1.8
RWR719	<i>P. vulgaris</i> (Mesoamerican)	3	28,942	0.0	94.3	684	2.2	1064	3.5
VAP1_R1	Bridge line	1400	17,806	4.6	58.0	9318	30.4	2169	7.1
VAP1_R2	Bridge line	1658	18,605	5.4	60.6	8091	26.4	2339	7.6
VAP2_R1	Bridge line	1990	17,547	6.5	57.2	10,157	33.1	999	3.3
VAP2_R2	Bridge line	1948	11,704	6.3	38.1	16,291	53.1	750	2.4
VAP3_R1	Bridge line	1618	18,289	5.3	59.6	8381	27.3	2405	7.8
VAP3_R2	Bridge line	2264	12,432	7.4	40.5	12,948	42.2	3049	9.9
INB841	Interspecific	97	17,041	0.3	55.5	12,579	41.0	976	3.2
INB834	Interspecific	280	19,859	.9	64.7	9530	31.0	1024	3.3
G40264	<i>P. parvifolius</i>	30,693	-	100.0	-	-	-	-	0.0

common bean and *P. parvifolius* had previously been overcome by embryo rescue and congruity backcrosses (Singh et al., 1998). However, in a study reported by Rao et al. (2013), these interspecific lines obtained with congruity backcrosses did have a comparative advantage in their drought tolerance response because of the dilution of the *P. parvifolius* contribution caused by the congruity backcrossing and the use of the drought-susceptible common bean parent, ICA Pijao.

3.3 | Identification of the VAP lines

We observed early flower abscission and empty pods when we crossed 16 interspecific F₄ families with four male tepary beans. However, in three families (which we named VAP1, VAP2, and VAP3), three pollinated flowers per family did not abscise early, and the pods reached maturity in the plant, leading to 12 mature seeds in total. The hybrid seeds were viable when planted and produced hybrid self-sterile plants, which confirmed the hybridization. This phenomenon was unusual because we had not obtained mature seeds in any of the previous direct crosses with tepary beans. Only five studies have reported the obtention of hybrid mature F₁ seeds between common and tepary beans (Ferwerda et al., 2003; Haghghi et al., 1984; Park et al., 1986; Smartt, 1970; Thomas et al., 1983). The ability to produce hybrid seeds from crosses between tepary beans and lines with a high level of common bean ancestry represents a vital breakthrough to overcoming reproductive barriers between the two species.

To confirm that VAP lines allowed fertile crossing between common and tepary beans without the need for embryo rescue, we created two sets of crosses under greenhouse conditions. One set of crosses used VAP lines (as female parents) and tepary beans (as male parents), and the other set used common beans (as female parents) and tepary beans (as male parents). Unsurprisingly, common beans that were fertilized with pollen from tepary beans showed premature shedding of flowers, and pods did not develop seeds (Figure 2a). From 278 pollinations between common and tepary beans, no pods reached maturity, and 40 flowers produced immature pods. Moreover, only 20 hybrid plants were obtained with embryo rescue (Figure 3). Conversely, when we crossed the VAP lines with tepary beans, a substantial hybridization success was observed. From 198 pollinations, 139 pods reached maturity in the plant, and 243 hybrid seeds were harvested from these pods (Figure 2b, Figure 3). From the hybrid seeds obtained, 39% corresponded to VAP1, 35% to VAP2, and 26% to VAP3. Overall, a substantial difference in the hybridization rate was observed for crosses between common bean and VAP lines with tepary beans. Specifically, common beans had a hybridization rate of .071 plants per pollination, whereas the hybridization rate for VAP lines was 1.23 plants per pollination.

All the interspecific hybrids obtained from crosses with VAP lines and common bean lines were self-sterile and showed intermediate common-tepary bean morphological characteristics (Supplementary Table S3). The fertility was fully recovered when the hybrids were crossed once with pollen of common bean lines

Tree scale: 0.1

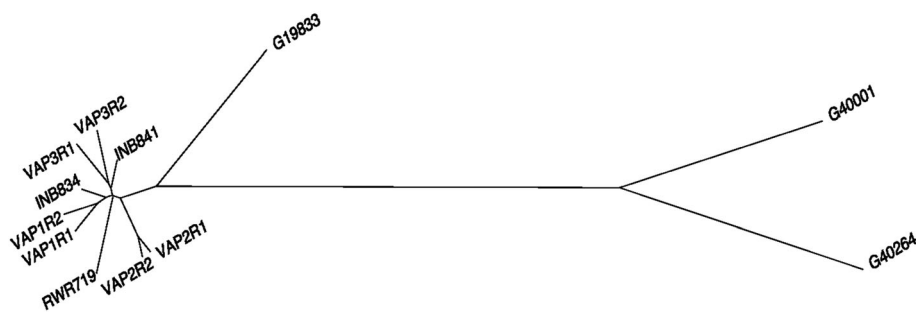


FIGURE 1 Relatedness tree based on the distance matrix between the interspecific lines (VAP and INB lines), the tepary bean genotype (G40001), the common bean genotype (RWR719), and the *Phaseolus parvifolius* genotype (G40264). The distance matrix was calculated, and the plot was created in the software TASSEL v.5.0 (Bradbury et al., 2007).



FIGURE 2 Interspecific hybridization between common bean and tepary bean. (a) Flowers absconded at a very early stage and pods without seeds, from crosses between tepary beans and common beans. (b) Mature pods and seeds from crosses between the VAP lines and tepary beans

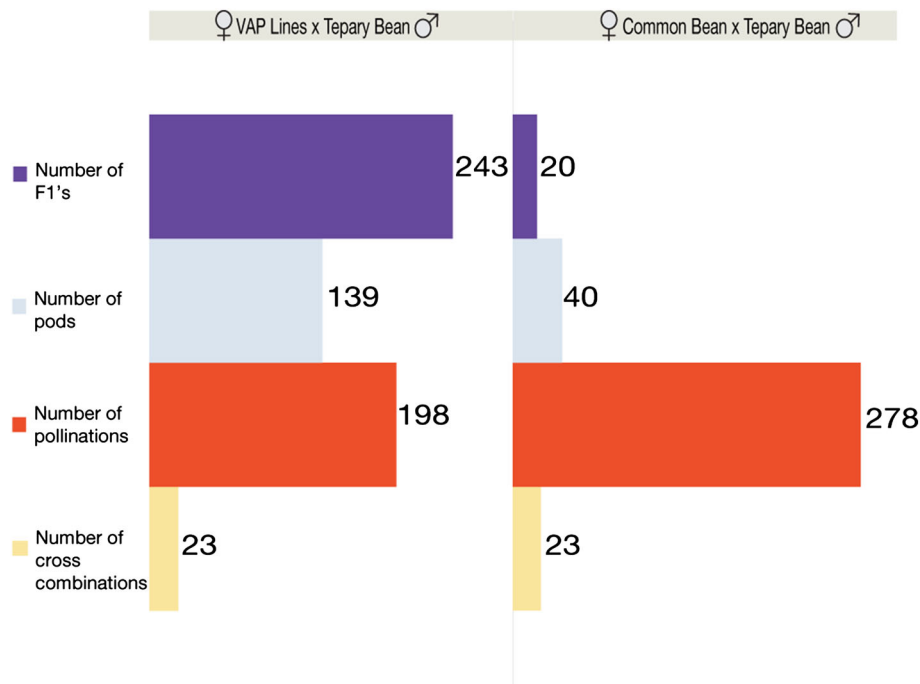
(Supplementary Figure S2). Although embryo rescue was not required for the hybridization between the VAP lines and tepary beans, self-sterility in the hybrids indicates that there is still a limitation in using these lines, and there is a need for an additional cross with a fertile common bean parent to recover full fertility.

3.4 | WGS and sequence read mapping

To elucidate the genomic constitution of the VAP lines and identify genomic regions that could be responsible for enhancing VAP lines'

ability to cross with tepary beans, we conducted WGS. We aligned the whole genome sequences of VAP lines with the tepary bean (*P. acutifolius* = G40001) and common bean (*P. vulgaris* = G19833) reference genomes and determined differences in the mapping percentage. The VAP lines, interspecific parents, and common bean controls generally had a higher alignment rate with the common bean reference (97%) than the tepary bean reference (88%) (Table 1). This result suggests that, overall, the VAP lines and interspecific parents are more related to the common bean than the tepary bean. Given that the VAP lines were backcrossed to INB lines that generally have over 90% common bean background (Lobaton et al., 2018), we

FIGURE 3 Comparison of the ability to cross tepary beans with traditional common bean lines and the interspecific VAP lines. Left) 22 crosses of VAP lines (used as female parents) with tepary beans (used as male parents). Right) 23 crosses of traditional common beans (female parents) with tepary beans (male parents)



expected a higher percentage alignment between VAP lines and the common bean reference compared to the tepary bean reference.

The *P. parvifolius* parental accession (G40264) had an alignment rate of 88% with the common bean reference genome and 97% with the tepary reference genome (Table 1). Due to the close phylogenetic relationship between G40264 and the tepary bean reference genome (G40001) (Freytag & Debouck, 2002), a higher percentage alignment between G40264 with tepary bean compared to common bean was also expected. The VAP lines had, on average, a 3% higher overall alignment with the tepary bean reference (G40001) than the common bean control (RWR719), and an 8% higher overall alignment than the Andean common bean reference (G19833) (Table 1).

3.5 | Interspecific introgressions mapping

To identify the genomic introgressions that could be critical in enhancing the crossability between the VAP lines and tepary beans, we mapped the genomic regions that corresponded to *P. parvifolius* and *P. vulgaris*. We found that the VAP lines had more genomic regions corresponding to the *P. parvifolius* genome than the INB parental lines and the common bean controls. While less than 1% of the *P. parvifolius* genomic regions were found in the INB lines, 4.6%–7.4% of these genomic regions were found in the VAP lines (Table 2), which indicates that the VAP lines have more interspecific introgressions in their genomes than the INB lines. Also, the VAP lines had on average 930 more heterozygous genomic regions than the INB lines and the common bean controls, indicating that those introgressions need to be fixed or recombined (Table 2). We also found that *P. parvifolius* and *P. acutifolius* share 98% of the 30,693 mapped

genomic regions (Table 2), confirming the close genetic relationship between both species (Freytag & Debouck, 2002).

The introgression analysis also revealed large *P. parvifolius* haplotype blocks in the VAP lines that are not present in the INB lines and the common bean controls. Specifically, VAP1 has large heterozygous introgression blocks spanning over 28 Mb, and small homozygous haplotype blocks spanning over 1.2 Mb, both on chromosome Pv03. It also has smaller heterozygous as well as homozygous introgression blocks on chromosomes Pv04 and Pv08 (Figure 4). VAP2 has a large homozygous *P. parvifolius* haplotype block of more than 39 Mb on chromosome Pv01 and a small (35 K) homozygous introgression block on chromosome Pv08 (Figure 4). It also has heterozygous introgression blocks on chromosomes Pv04, Pv08, and Pv011 (Figure 4).

VAP3 is the bridge line that has the most heterozygous introgression blocks of the three VAP lines (Figure 5). Interestingly, nearly the entire chromosome Pv09 (~36 Mb) is composed of heterozygous SNPs from *P. parvifolius* and *P. vulgaris* (Figures 4 and 5). VAP3 also has a large heterozygous block (54 Mb) in replication 2 (R2) on chromosome Pv04, which is not present in replication 1 (R1) (Figure 3), indicating that this line is not entirely fixed. Interestingly, the introgressions we found on chromosome Pv08 were also found by Lobaton et al. (2018) in the VAX and INB lines, indicating that these tepary introgressions are conserved across multiple interspecific lines. However, unlike the VAP lines, the INB lines do not have more interspecific tepary haplotype blocks on the other chromosomes. Also, as expected, the unintrogresed control common bean line RWR719 did not show any introgressions (Figure 4). Still, three genomic regions were assigned to the *P. parvifolius* group (Table 2), which indicates that small genomic regions assigned to *P. parvifolius* cannot necessarily be considered interspecific introgressions but reflect a common ancestry in the genus *Phaseolus*.

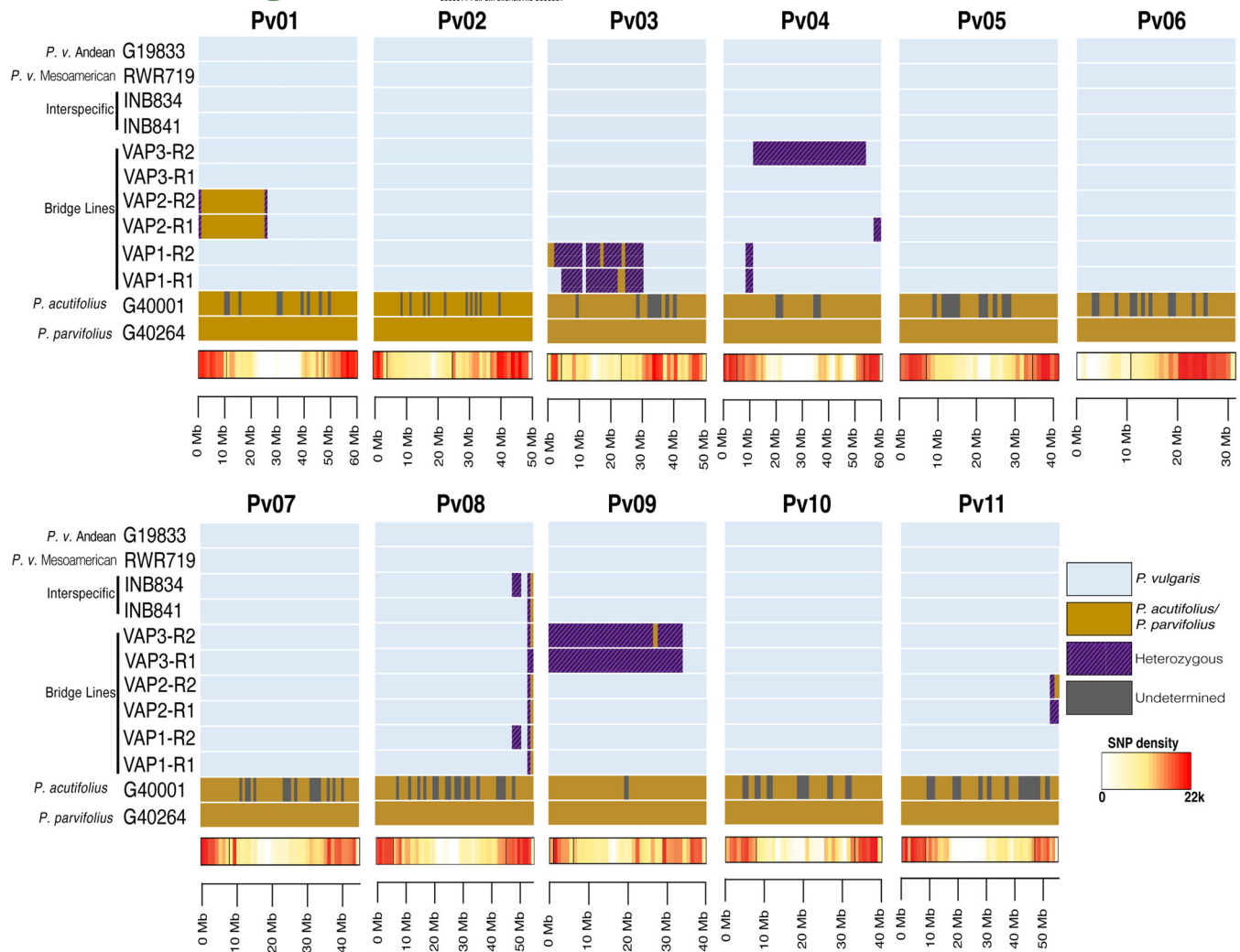


FIGURE 4 Mapping of the interspecific introgressions in the VAP lines, INB lines, two common bean genotypes (Andean and Mesoamerican), and two tepary bean genotypes in the 11 chromosomes. Light blue indicates the common bean background. Brown indicates the tepary bean background. Purple indicates heterozygous SNPs. The heat map shows the SNP density. G40001, which is the tepary bean reference genome is presented in this figure only to compare *Phaseolus parvifolius* and *Phaseolus acutifolius* genomes.

4 | DISCUSSION

In this study, we have confirmed that crossing the traditional common bean (i.e., without tepary introgressions) with tepary bean is challenging. Although hybridization is possible with embryo rescue, it limits efficient interspecific introgression. Similar challenges have been reported in previous common and tepary bean hybridization studies (Andrade-Aguilar & Jackson, 1988; Mok et al., 1978; Mejía-Jiménez et al., 1994; Parker & Michaels, 1986; Pratt et al., 1985; Rabakoarihanta et al., 1979; Souter et al., 2017; Thomas et al., 1983). Although different hybridization strategies such as alternating female genotypes (Parker & Michaels, 1986; Rabakoarihanta et al., 1979) or congruent backcrossing have been attempted (Anderson et al., 1996; Haghghi & Ascher, 1988), only a few hybrid plants are usually obtained. Here, we demonstrate that it is possible to improve the crossability of common bean and tepary bean by using three novel interspecific genotypes (VAP lines). These interspecific genotypes

allowed us to obtain a large number of interspecific common–teparty bean hybrids without embryo rescue (Figure 3). Strikingly, we were able to achieve up to 12-fold more hybrids when we crossed these VAP lines with tepary beans than when we crossed traditional common beans with tepary beans and used embryo rescue (Figure 3). We needed 21% fewer pollinations when we crossed VAP lines with the tepary beans than when we crossed common beans with tepary beans (Figure 3).

Five other studies have also reported successful hybridization between common bean and tepary bean without embryo rescue but did not report finding a large number of hybrid seeds (Ferwerda et al., 2003; Haghghi et al., 1984; Park et al., 1986; Smartt, 1970; Thomas et al., 1983). Producing large recombinant populations is fundamental to introducing genetically complex traits, such as heat and drought tolerance, from the tepary to the common bean. Additionally, this is the first study that has achieved interspecific crosses using a large number of different forms of tepary beans, such as wild

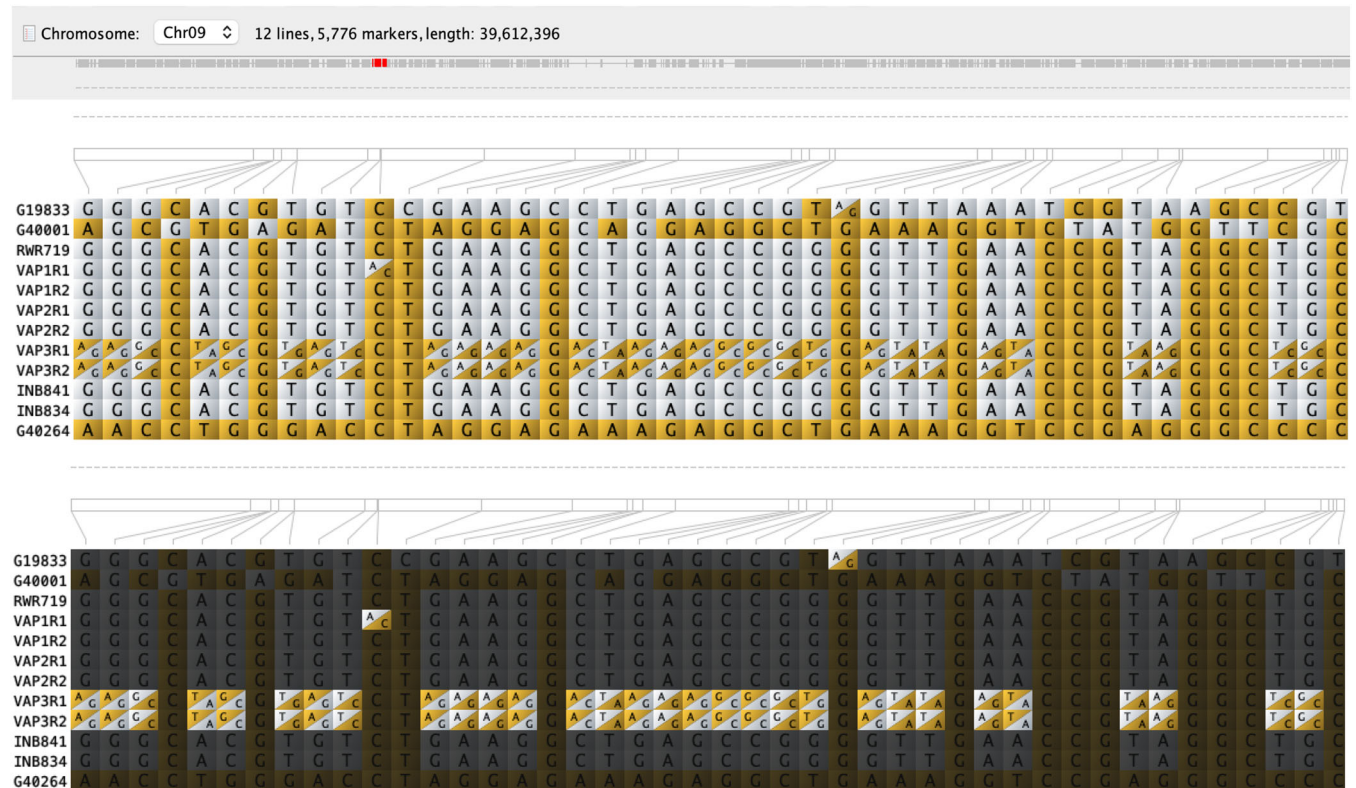


FIGURE 5 Flapjack visualization of introgressions from *Phaseolus parvifolius* (G40264) in the VAP lines in chromosome Pv09. Highlighted heterozygous introgression regions in the line VAP3 in replication 01 (R1) and replication 02 (R2). Heterozygous regions are defined as regions that share nucleotides from both reference haplotypes (G19833 and G40264). G40001, which is the tepary bean reference genome is presented in this figure only to compare *P. parvifolius* and *Phaseolus acutifolius* genomes.

accessions, landraces, and improved varieties. Our success in increasing crossability may also be due to an increase in gametic diversity, as *P. parvifolius*, a close relative of *P. acutifolius* (Freytag & Debouck, 2002; Figure 1), was included in the crossing scheme. Gametic diversity has been credited with (or has been suggested to achieve) an increase in interspecific crossability in *Phaseolus* and *Cucurbita* (Gepts, 1981; Paris, 2016; Pratt et al., 1985; Wall & York, 1960), rye-wheat (Meister & Tjumjakoff, 1928), and eggplant (Schaff et al., 1982).

Further genomic analysis of the VAP lines, conducted to identify genomic regions that are key in enhancing their crossability, revealed large introgression blocks from *P. parvifolius* to the VAP lines. The inheritance of these large introgression blocks seen in the VAP lines suggests a lack of recombination events in those genomic regions. Some of the introgression blocks found here have been reported in genomic regions where considerable rearrangements between common and tepary beans have occurred (Moghaddam et al., 2021). For instance, a major QTL has been reported, which contributes to the reproductive incompatibility between the two species on chromosome Pv09 (Soltani et al., 2020), which is one of the chromosomes with large common-tepary bean rearrangements (Gujaria-Verma et al., 2016; Moghaddam et al., 2021). In this study, we saw that chromosome Pv09 was one of the chromosomes with the largest *P. parvifolius* introgression events, suggesting that cross-compatibility

between tepary beans and the VAP3 lines may have been enhanced by this interspecific introgression. In contrast, different introgressions present on chromosomes Pv01, Pv03, Pv04, Pv08, and Pv11, found in the VAP1 and VAP2 lines (Figure 3), could have enhanced their cross-compatibility with tepary beans since they lacked the Pv09 introgression event.

VAP 3 had a reduced ability to cross with tepary beans (only 26% of the crosses were obtained with VAP3), which could be explained by the high level of heterozygous introgressions of this line. A high number of heterozygous introgression regions, such as the introgressions on chromosome Pv09 (Figures 4 and 5), suggest that it might be challenging to maintain the ability to cross the VAP lines with tepary beans since the lines could segregate in favor of common bean alleles. Therefore, there is a need to continue breeding the VAP lines and fix the heterozygous introgressions.

The only shared introgressed segment between the two INB and three VAP lines, but absent in the two common bean lines, was the short terminal segment observed on chromosome Pv08. This introgression had already been observed by Lobaton et al. (2018) in three VAX lines (VAX 3, 4, and 6), resulting from interspecific hybridization with tepary beans and pyramiding multiple sources of resistance to common bacterial blight, including tepary beans (Singh et al., 2001; Singh & Muñoz, 1999). Given the frequent position of disease resistance genes or clusters in distal chromosome positions (e.g., Kelly

et al., 2003; Schmutz et al., 2014), this segment could be a carrier of part of the bacterial resistance exhibited by the VAX lines mentioned. Although the two INB lines included in this study (INB834 and INB841) carried the terminal Pv08 introgression, this is not always the case as Lobaton et al. (2018) showed that INB827 did not carry this introgression. Thus, any putative crossability-enhancing property attributable to the terminal Pv08 introgression cannot be assigned with certainty to all INB lines. Nevertheless, it is possible that this segment could carry both common bacterial blight resistance and enhanced crossability, either by pleiotropy or tight linkage.

Heterozygosity is still high in the VAP lines, based on the sequence data obtained in this study. Although the pursuit of fixation is highly desirable for the maintenance of the VAP lines, it remains to be seen whether the increased crossability with tepary beans can be maintained in a homozygous state or if the heterozygous state has to be maintained, perhaps by a marker-assisted selection of heterozygous introgressed chromosome segments.

The development of the novel VAP lines, which allows hybridizing common beans with tepary beans and obtaining large common bean/tepar bean populations, may open up the possibility of efficiently exploiting the tertiary gene pool to move heat and drought tolerance from tepary beans to common beans without the need for embryo rescue and the associated cost and infrastructure.

ACKNOWLEDGMENTS

This study was financially supported by the International Center for Tropical Agriculture (CIAT) and the Nebraska Dry Bean Commission. The funders did not influence the result reported in this study. The authors acknowledge colleagues from CIAT for their support in the data collection in the field, greenhouses, and laboratory. A special acknowledgment is given to Gersain Galarza from CIAT for his support in the interspecific crosses.

CONFLICTS OF INTEREST

The authors declare that they have no conflict of interest related to this manuscript.

AUTHOR CONTRIBUTIONS

Santos Barrera Lemus developed the VAP lines, aligned the reads to the reference genomes, and wrote the manuscript draft. Jorge C. Bery Mier y Teran conducted the DNA extraction, coordinated the WGS analysis, and edited the manuscript. Juan David Lobaton performed de bioinformatic analysis of the WGS data and introgression analysis. Roosevelt Escobar performed the embryo rescue. Steve Beebe, Paul Gepts, and Carlos A. Urrea supervised the study and edited the final version of the manuscript.

DATA AVAILABILITY STATEMENT

The Whole Genome Sequence data for this study are available online at the National Center for Biotechnology Information (NCBI) repository <https://submit.ncbi.nlm.nih.gov/subs/sra/SUB10611384/>. (BioProject ID:PRJNA778000)

ORCID

Santos Barrera  <https://orcid.org/0000-0001-6452-4979>

Roosevelt Escobar  <https://orcid.org/0000-0002-4661-6199>

Paul Gepts  <https://orcid.org/0000-0002-1056-4665>

REFERENCES

- Anderson, N. O., Ascher, P. D., & Haghghi, K. (1996). Congruity backcrossing as a means of creating genetic variability in self pollinated crops: Seed morphology of *Phaseolus vulgaris* L. and *P. acutifolius* A. Gray hybrids. *Euphytica*, 87(3), 211–224. <https://doi.org/10.1007/BF00023748>
- Andrade-Aguilar, J. A., & Jackson, M. T. (1988). Attempts at interspecific hybridization between *Phaseolus vulgaris* L. and *P. acutifolius* A. Gray using embryo rescue. *Plant Breeding*, 101(3), 173–180. <https://doi.org/10.1111/j.1439-0523.1988.tb00285.x>
- Andrews, S. (2010). A quality control tool for high throughput sequence data. <https://www.bioinformatics.babraham.ac.uk/projects/fastqc/>
- Ariani, A., Bery Mier y Teran, J. C., & Gepts, P. (2016). Genome-wide identification of SNPs and copy number variation in common bean (*Phaseolus vulgaris* L.) using genotyping-by-sequencing (GBS). *Molecular Breeding*, 36(7), 1, 87–11. <https://doi.org/10.1007/s11032-016-0512-9>
- Beebe, S. E. (2012). Common bean breeding in the tropics. *Plant Breeding Reviews*, 36, 357–426. <https://doi.org/10.1002/9781118358566.ch5>
- Beebe, S. E., Rao, I., Cajiao, C. H., & Grajales, M. (2012). Mining the *Phaseolus* genus: How many species can you get into a cross? *Annual Report Bean Improvement Cooperative*, 55, 61–62.
- Beebe, S. E., Rao, I., Mukankusi, C., & Buruchara, R. (2013). Improving resource use efficiency and reducing risk of common bean production in Africa, Latin America, and the Caribbean. *Eco-Efficiency: From Vision to Reality*, January 2015, 117–134. http://ciat.cgiar.org/wp-content/uploads/2013/04/eco-efficiency_book.pdf
- Bliss, F. A. (1980). Common Bean. In W. R. Fehr & H. H. Hadley (Eds.), *Hybridization of crop plants* (pp. 273–284). American Society of Agronomy. <https://doi.org/10.2135/1980.hybridizationofcrops>
- Bolger, A. M., Lohse, M., & Usadel, B. (2014). Trimmomatic: A flexible trimmer for Illumina sequence data. *Bioinformatics*, 30(15), 2114–2120. <https://doi.org/10.1093/bioinformatics/btu170>
- Bradbury, P. J., Zhang, Z., Kroon, D. E., Casstevens, T. M., Ramdoss, Y., & Buckler, E. S. (2007). TASSEL: Software for association mapping of complex traits in diverse samples. *Bioinformatics*, 23, 2633–2635. <https://doi.org/10.1093/bioinformatics/btm308>
- Broughton, W. J., Hernández, G., Blair, M., Beebe, S. E., Gepts, P., & Vanderleyden, J. (2003). Beans (*Phaseolus* spp.)—Model food legumes. *Plant and Soil*, 252(1), 55–128. <https://doi.org/10.1023/A:1024146710611>
- Dohle, S., Bery Mier y Teran, J. C., Ashley, E., Kisha, T., & Khoury, C. K. (2019). Wild Beans (*Phaseolus* L.) of North America. In S. L. Greene, K. A. Williams, C. K. Khoury, L. F. Marek, & M. B. Kantar (Eds.), *North American crop wild relatives: Important species* (Vol. 2). Springer. <https://doi.org/10.1007/978-3-319-97121-6>
- Duitama, J., Quintero, J. C., Cruz, D. F., Quintero, C., Hubmann, G., Foulquié-Moreno, M. R., Verstrepen, K. J., Thevelein, J. M., & Tohme, J. (2014). An integrated framework for discovery and genotyping of genomic variants from high-throughput sequencing experiments. *Nucleic Acids Research*, 42(6), e44. <https://doi.org/10.1093/nar/gkt1381>
- Ferwerda, F. H. M., Bassett, J., & Beaver, J. (2003). Viability of seed of reciprocal interspecific crosses between *Phaseolus vulgaris* L. and *Phaseolus acutifolius* A. Gray. *Bean Improvement Cooperative Annual Report*, 29–30. <http://handle.nal.usda.gov/10113/IND43757103>



- Freytag, G. F., & Debouck, D. G. (2002). Taxonomy, distribution, and ecology of the genus *Phaseolus* (Leguminosae-Papilionoideae) in North America, Mexico, and Central America. In Research Institute of Texas (BRIT); Forth Worth, TX; USA. 298 p. (SIDA; *Botanical Miscellany* no. 23).
- Gaur, P. M., Mallikarjuna, N., Knights, T., Beebe, S. E., Debouck, D., Mejía, A., Malhotra, R. S., Imtiaz, M., Sarker, A., Tripathi, S., & Gowda, C. L. L. (2009). Gene introgression in grain legumes. In International Conference on Grain Legumes: Quality Improvement, *Value Addition and Trade*. Indian Society of Pulses Research and Development.
- Gepts, P. (1981). Hibridaciones interespecíficas para el mejoramiento de *Phaseolus vulgaris* L. (Seminarios internos. Serie SE-10-81). In: SE-10-81 SiS (ed). Centro Internacional de Agricultura Tropical (CIAT), Cali, Colombia.
- Gujaria-Verma, N., Ramsay, L., Sharpe, A. G., Sanderson, L. A., Debouck, D. G., Tar'an, B., & Bett, K. E. (2016). Gene-based SNP discovery in tepary bean (*Phaseolus acutifolius*) and common bean (*P. vulgaris*) for diversity analysis and comparative mapping. *BMC Genomics*, 17(1), 1–17. <https://doi.org/10.1186/s12864-016-2499-3>
- Haghighi, K., Zuluaga, S., & Ascher, P. D. (1984). Hybridization and rapid asexual propagation of *Phaseolus vulgaris*-*P. acutifolius*. *Bean Improvement Cooperative Annual Report*, 27, 42–43.
- Haghighi, K. R., & Ascher, P. D. (1988). Fertile, intermediate hybrids between *Phaseolus vulgaris* and *P. acutifolius* from congruity backcrossing. *Sexual Plant Reproduction*, 1(1), 51–58. <https://doi.org/10.1007/BF00227023>
- Kelly, J. D., Gepts, P., Miklas, P. N., & Coyne, D. P. (2003). Tagging and mapping of genes and QTL and molecular marker-assisted selection for traits of economic importance in bean and cowpea. *Field Crops Research*, 82, 135–154. [https://doi.org/10.1016/S0378-4290\(03\)00034-0](https://doi.org/10.1016/S0378-4290(03)00034-0)
- Langmead, B., & Salzberg, S. L. (2012). Fast gapped-read alignment with bowtie 2. *Nature Methods*, 9(4), 357–359. <https://doi.org/10.1038/nmeth.1923>
- Lobaton, J. D., Miller, T., Gil, J., Ariza, D., Hoz, J. F., Soler, A., Beebe, S., Duitama, J., Gepts, P., & Raatz, B. (2018). Resequencing of common bean identifies regions of inter-gene pool introgression and provides comprehensive resources for molecular breeding. *The Plant Genome*, 11(2), 170068. <https://doi.org/10.3835/plantgenome2017.08.0068>
- Markhart, A. H. (1985). Comparative water relations of *Phaseolus vulgaris* L. and *Phaseolus acutifolius* Gray. *Plant Physiology*, 77(1), 113–117. <https://doi.org/10.1104/pp.77.1.113>
- McElroy, J. B. (1985). *Breeding dry beans, Phaseolus vulgaris* L., for common bacterial blight resistance derived from *Phaseolus acutifolius* A. Gray. Cornell University.
- Meister, N., & Tjumjakoff, N. A. (1928). Rye-wheat hybrids from reciprocal crosses. *Journal of Genetics*, 20, 233–245. <https://doi.org/10.1007/BF02983142>
- Mejía-Jiménez, A., Muñoz, C., Jacobsen, H. J., Roca, W. M., & Singh, S. P. (1994). Interspecific hybridization between common and tepary beans: Increased hybrid embryo growth, fertility, and efficiency of hybridization through recurrent and congruity backcrossing. *Theoretical and Applied Genetics*, 88(3–4), 324–331. <https://doi.org/10.1007/BF00223640>
- Milne, I., Shaw, P., Stephen, G., Bayer, M., Cardle, L., Thomas, W. T. B., Flavell, A. J., & Marshall, D. (2010). Flapjack-graphical genotype visualization. *Bioinformatics*, 26(24), 3133–3134. <https://doi.org/10.1093/bioinformatics/btq580>
- Moghaddam, S. M., Oladzad, A., Koh, C., Ramsay, L., Hart, J. P., Mamidi, S., Hoopes, G., Sreedasyam, A., Wiersma, A., Zhao, D., Grimwood, J., Hamilton, J. P., Jenkins, J., Vaillancourt, B., Wood, J. C., Schmutz, J., Kagale, S., Porch, T., Bett, K. E., ... McClean, P. E. (2021). The tepary bean genome provides insight into evolution and domestication under heat stress. *Nature Communications*, 12(1), 1–14. <https://doi.org/10.1038/s41467-021-22858-x>
- Mok, D. W., Mok, M. C., & Rabakoarihanta, A. (1978). Interspecific hybridization of *phaseolus vulgaris* with *P. lunatus* and *P. acutifolius*. *Theoretical and Applied Genetics*, 52(5), 209–215. <https://doi.org/10.1007/BF00273891>
- Muñoz, C. G., Blair, M. W., Duque, M. C., Tohme, J., & Roca, W. (2004). Introgression in common bean × tepary bean interspecific. *Crop Science*, 44(2), 637–645. <https://doi.org/10.2135/cropsci2004.6370>
- Muñoz, L. C., Duque, M. C., Debouck, D. G., & Blair, M. W. (2006). Taxonomy of tepary bean and wild relatives as determined by amplified fragment length polymorphism (AFLP) markers. *Crop Science*, 46(4), 1744–1754. <https://doi.org/10.2135/cropsci2005-12-0475>
- Murashige, T., & Skoog, F. (1962). A revised medium for rapid growth and bio assays with tobacco tissue cultures. *Physiologia Plantarum*, 15(3), 473–497. <https://doi.org/10.1111/j.1399-3054.1962.tb08052>
- Paris, H. S. (2016). Genetic resources of pumpkins and squash, *Cucurbita* spp. In R. Grumet, N. Katzir, & J. Garcia-Mas (Eds.), *Genetics and genomics of Cucurbitaceae* (pp. 111–154). Springer International Publishing. https://doi.org/10.1007/7397_2016_3
- Park, S. J., Chesney, M., & Dhanvantari, B. N. (1986). Reaction to common bacterial blight in selfed progeny from *Phaseolus vulgaris* × *P. acutifolius*. *Bean Improvement Cooperative Annual Report*, 107–108.
- Parker, J. P., & Michaels, T. E. (1986). Genetic control of abnormal seedling development in *Phaseolus vulgaris* L. and *P. vulgaris* × *P. acutifolius* A. Gray hybrids. *Plant Breeding*, 104(3), 212–217. <https://doi.org/10.1111/j.1439-0523.1990.tb00425.x>
- Polania, J. A., Chater, C. C. C., Covarrubias, A. A., & Rao, I. M. (2020). *Phaseolus* species responses and tolerance to drought. In M. Hasanuzzaman, S. Araujo, & S. S. Gill (Eds.), *The plant family Fabaceae: Biology and physiological responses to environmental stresses* (pp. 319–336). Springer Singapore. https://doi.org/10.1007/978-981-15-4752-2_12
- Porch, T., Barrera, S., Berny, J., Díaz-Ramírez, J., Pastor-Corrales, M., Gepts, P., Urrea, C., & Rosas, J. C. (2021). Release of tepary bean TARS-Tep 23 germplasm with broad abiotic stress tolerance and rust and common bacterial blight resistance. *Journal of Plant Registrations*, 16, 109–119. <https://doi.org/10.1002/plr2.20180>
- Porch, T. G., Beaver, J. S., & Brick, M. A. (2013). Registration of tepary germplasm with multiple-stress tolerance, TARS-Tep 22 and TARS-Tep 32. *Journal of Plant Registrations*, 7(3), 358–364. <https://doi.org/10.3198/jpr2012.10.0047crg>
- Pratt, R. C., Bressan, R. A., & Hasegawa, P. M. (1985). Genotypic diversity enhances recovery of hybrids and fertile backcrosses of *Phaseolus vulgaris* L. × *P. acutifolius* A. Gray. *Euphytica*, 34, 329–344. <https://doi.org/10.1007/BF00022926>
- Pratt, R. C., & Nabhan, G. P. (1988). Evolution and diversity of *Phaseolus acutifolius* genetic resources. In P. Gepts (Ed.), *Genetic resources of Phaseolus beans* (pp. 409–440). Kluwer. https://doi.org/10.1007/978-94-009-2786-5_17
- Rabakoarihanta, A., Mok, D. W. S., & Mok, M. C. (1979). Fertilization and early embryo development in reciprocal interspecific crosses of *Phaseolus*. *Theoretical and Applied Genetics*, 54(2), 55–59. <https://doi.org/10.1007/BF00265469>
- Ramirez-Cabral, N. Y. Z., Kumar, L., & Taylor, S. (2016). Crop niche modeling projects major shifts in common bean growing areas. *Agricultural and Forest Meteorology*, 218–219, 102–113. <https://doi.org/10.1016/j.agrformet.2015.12.002>
- Rao, I., Beebe, S., Polania, J., Ricaurte, J., Cajiao, C., Garcia, R., & Rivera, M. (2013). Can tepary bean be a model for improvement of drought resistance in common bean. *African Crop Science Journal*, 21(4), 265–281. <https://doi.org/10.4314/acsj.v21i4>
- Schaff, D. A., Jelenkovic, G., Boyer, C. D., & Pollack, B. L. (1982). Hybridization and fertility of hybrid derivatives of *Solanum melongena* L. and

- Solanum macrocarpon* L. *Theoretical and Applied Genetics*, 62, 149–153. <https://doi.org/10.1007/bf00293348>
- Schmutz, J., McClean, P. E., Mamidi, S., Wu, G. A., Cannon, S. B., Grimwood, J., Jenkins, J., Shu, S., Song, Q., Chavarro, C., Torres-Torres, M., Geffroy, V., Moghaddam, S. M., Gao, D., Abernathy, B., Barry, K., Blair, M., Brick, M. A., Chovatia, M., ... Jackson, S. A. (2014). A reference genome for common bean and genome-wide analysis of dual domestications. *Nature Genetics*, 46, 707–713. <https://doi.org/10.1038/ng.3008>
- Singh, S. P., Debouck, D. G., & Roca, W. M. (1998). Interspecific hybridization between *Phaseolus vulgaris* L. and *P. parvifolius* Freytag. *Bean Improvement Cooperative Annual Report*, 4, 7–8. <https://doi.org/10.2307/2408534>
- Singh, S. P., & Muñoz, C. G. (1999). Resistance to common bacterial blight among *Phaseolus* species and common bean improvement. *Crop Science*, 39, 80–89. <https://doi.org/10.2135/cropsci1999.0011183X003900010013x>
- Singh, S. P., Muñoz, C. G., & Terán, H. (2001). Registration of common bacterial blight resistant dry bean germplasm VAX 1, VAX 3, and VAX 4. *Crop Science*, 41, 275–276. <https://doi.org/10.2135/cropsci2001.411275x>
- Smartt, J. (1970). Interspecific hybridization between cultivated American species of the genus *Phaseolus*. *Euphytica*, 19(4), 480–489. <https://doi.org/10.1007/BF01902923>
- Soltani, A., Walter, K., & Lowry, D. B. (2020). A major reproductive isolation qtl is associated with F1 sterility in common bean × tepary bean hybrids. *Bean Improvement Cooperative Annual Report*, 63, 153–154.
- Souter, J. R., Gurusamy, V., Porch, T. G., & Bett, K. E. (2017). Successful introgression of abiotic stress tolerance from wild tepary bean to common bean. *Crop Science*, 57(3), 1160–1171. <https://doi.org/10.2135/cropsci2016.10.0851>
- Tello, D., Gil, J., Loaiza, C. D., Riascos, J. J., Cardozo, N., & Duitama, J. (2019). NGSEP3: Accurate variant calling across species and sequencing protocols. *Bioinformatics*, 35(22), 4716–4723. <https://doi.org/10.1093/bioinformatics/btz275>
- Thomas, C. V., Manshardt, R. M., & Waines, J. G. (1983). Teparies as a source of useful traits for improving common beans. *Desert Plants*, 5, 43–48.
- Thomas, C. V., & Waines, J. G. (1984). Fertile backcross and allotetraploid plants from crosses between tepary beans and common beans. *Journal of Heredity*, 75, 93–98. <https://doi.org/10.1093/oxfordjournals.jhered.a109901>
- Traub, J., Porch, T., Naeem, M., Urrea, C. A., Austic, G., Kelly, J. D., & Loescher, W. (2018). Screening for heat tolerance in *Phaseolus* spp. using multiple methods. *Crop Science*, 58(6), 2459–2469. <https://doi.org/10.2135/cropsci2018.04.0275>
- Waines, J. G., Manshardt, R. M., & Wells, W. C. (1988). Interspecific hybridization between *Phaseolus vulgaris* and *P. acutifolius*. In P. Gepts (Ed.), *Genetic resources of Phaseolus beans. Current plant science and biotechnology in agriculture* (Vol. 6). Springer. https://doi.org/10.1007/978-94-009-2786-5_20
- Wall, J. R., & York, T. L. (1960). Gametic diversity as an aid to interspecific hybridization in *Phaseolus* and in *Cucurbita*. *Proceedings American Society for Horticultural Science*, 75, 419–428.
- Zink, D., & Nagl, W. (1998). A taxon identified by microsatellite-primed PCR and southern hybridization in the secondary gene pool of the tepary bean. *Bean Improvement Cooperative Annual Report*, 41, 107–108.

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

How to cite this article: Barrera, S., Berny Mier y Teran, J. C., Lobaton, J. D., Escobar, R., Gepts, P., Beebe, S., & Urrea, C. A. (2022). Large genomic introgression blocks of *Phaseolus parvifolius* Freytag bean into the common bean enhance the crossability between tepary and common beans. *Plant Direct*, 6(12), e470. <https://doi.org/10.1002/pld3.470>