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Chloroplast DNA as an evolutionary marker in the *Phaseolus vulgaris* complex

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Abstract We have analyzed the changes occurring in the chloroplast DNA (cpDNA) of taxa belonging to the Phaseolus vulgaris complex to help clarify relationships among species of this complex. Two restriction maps for 11 restriction enzymes comprising the whole chloroplast genome from a wild P. vulgaris and a wild P. coccineus accession were constructed. These maps allowed us to compare a total of 330 restriction sites between the two genomes in order to identify polymorphisms, assess the type of mutations detected, and identify regions of high variability. A region, located in the large single-copy region near the borders with the inverted repeats, accounted for a large portion of the variation. Most of the mutations detected were due to restriction sites gains or losses. Variable and conserved regions were then evaluated in 30 accessions belonging to taxa of the P. vulgaris complex. Phylogenetic analyses were made using parsimony methods. Conclusions obtained from such analyses were the following: (1) there was high cpDNA variability within P. coccineus but not in P. vulgaris. (2) P. coccineus subsp. glabellus showed a very distinct cpDNA type that strongly suggests that it actually belongs to a different but as yet undetermined section of the genus. Our cpDNA observations are supported by distinctive morphological traits and reproductive biology of this taxon. (3) In P. coccineus subsp. darwinianus (also classified as P. polyanthus), the cpDNA lineage was in disagreement with data obtained from nuclear markers and suggested a reticulated origin by hybridization between P. coccineus as the male parent and an ancestral P. polyanthus type, closely allied to P. vulgaris, as the seed parent. This initial cross was presumably followed by repeated backcrossing to P. coccineus. Our cpDNA studies illustrate the importance of molecular

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markers in elucidating phylogenetic relationships. They also indicate that accurate phylogenies will require analyses of both nuclear and cytoplasmic genomes.

Key words cpDNA · *Phaseolus vulgaris Phaseolus coccineus* · Phylogeny · Molecular evolution

Introduction

Phaseolus vulgaris (common bean), an annual, self-pollinating species, is economically the most important species of the genus. It is a major source of proteins for human consumption in many parts of the world (FAO 1980). The perennial, cross-pollinated *P. coccineus* (runner bean) is a source of variability for several agronomic traits for the improvement of the primary gene pool of P. vulgaris (Gepts 1981). In spite of marked differences in pollination systems and life cycles, both species are closely related as evidenced by data on genetic compatibility and reproductive biology (e.g., Mendel 1865; Hucl and Scoles 1985; Guo et al. 1991), morphology (Maréchal et al. 1978), seed protein immunology (Kloz and Klozová 1974), isozymes (Bassiri and Adams 1978), electrophoresis of seed proteins (Sullivan and Freytag 1986), and mitochondria DNA (mtDNA) polymorphism (Khairallah et al. 1991). Consequently, the two taxa have been grouped into the P. vulgaris complex.

P. coccineus and *P. vulgaris* are partially intercrossable, with strong reciprocal differences. When *P. coccineus* is used as the female parent, hybrid embryos do not develop beyond the late-cotyledonary stage. Viable plants are obtained only when *P. vulgaris* is used as the seed parent (Wall 1970; Shii et al. 1982; Hucl and Scoles 1985). Furthermore, a number of dwarf and other abnormal plants are observed in the progeny, and the viable hybrids have typically only 25% viable pollen (Gepts 1981; Shii et al. 1982).

Several additional taxa with a wide range of morphological and reproductive characteristics have been included in the *P. vulgaris* complex. They have been classified as

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 Table 1
 List of accessions

 sampled for cpDNA restriction
 site variation

| Accession | Source ^a | Species | Subspecies | Status | Location |
|-------------|---------------------|-----------|-------------|------------|--------------------|
| BAT93 | 1 | vulgaris | | Cultivated | Breeding line |
| Jalo EEP558 | 1 | vulgaris | | Cultivated | Brazil |
| G12873 | 1 | vulgaris | | Wild | Mexico |
| PG11A | 1 | lunatus | | Cultivated | Colombia |
| ADS3 | 2 | coccineus | coccineus | Cultivated | Mexico (Puebla) |
| ADS4 | 2 | coccineus | coccineus | Cultivated | Mexico (Chiapas) |
| ADS5 | 2 | coccineus | coccineus | Cultivated | Mexico (Chiapas) |
| ADS11 | 2 | coccineus | formosus | Wild | Mexico (Oaxaca) |
| ADS14 | 2 | coccineus | glabellus | Wild | Mexico (Hidalgo) |
| Delgado286 | 3 | coccineus | coccineus | Wild | Mexico (Morelos) |
| Delgado550 | 2 | coccineus | coccineus | Cultivated | Mexico (Puebla) |
| Delgado553 | 2 | coccineus | darwinianus | Cultivated | Mexico (Puebla) |
| Delgado577 | 2 | coccineus | darwinianus | Cultivated | Mexico (Puebla) |
| Delgado578 | 2 | coccineus | glabellus | Wild | Mexico (Puebla) |
| Delgado584 | 2 | coccineus | darwinianus | Cultivated | Mexico (Puebla) |
| Delgado602 | 2 | coccineus | coccineus | Cultivated | Mexico (Puebla) |
| Delgado605 | 2 | coccineus | darwinianus | Cultivated | Mexico (Puebla) |
| Delgado608 | 2 | coccineus | glabellus | Wild | Mexico (Puebla) |
| DelgadoB | 2 | coccineus | coccineus | Wild | Mexico (Chihuahua) |
| DGD78/045 | 3 | coccineus | coccineus | Cultivated | Mexico |
| G35651 | 3 | coccineus | glabellus | Wild | Mexico |
| DGD2043 | 3 | coccineus | glabellus | Wild | Mexico |
| DGD2374 | 3 | coccineus | | Wild | Mexico (Puebla) |
| DGD2445 | 3 | coccineus | | Wild | Guatemala |
| DGD2462 | 3 | coccineus | | Wild | Guatemala |
| HSG22522 | 3 | coccineus | darwinianus | Cultivated | |
| DGD1661 | 3 | coccineus | darwinianus | Cultivated | Guatemala |
| DGD1684 | 3 | coccineus | darwinianus | Cultivated | Guatemala |
| OTCH1 | 3 | coccineus | coccineus | Cultivated | Colombia |
| DGD713 | 3 | coccineus | darwinianus | Cultivated | Peru (Cajamarca) |
| DGD1631 | 3 | coccineus | darwinianus | Wild | Guatemala |

^a Sources referred to are: 1, O. Tori, S. Singh, and M. Iwanaga, CIAT, Cali, Colombia; 2, A. Delgado Salinas, Universidad Nacional Autónoma de México; 3, D.G. Debouck, IBPGR, c/o CIAT, Cali, Colombia

separate species or subspecies of *P. coccineus* (Piper 1926; Maréchal et al. 1978; Delgado Salinas 1985). According to Delgado Salinas (1985), they include: (1) *P. coccineus* subsp. *coccineus* L., (2) *P. coccineus* subsp. *formosus* (Kunth) Maréchal, Mascherpa and Stainier (= *P. formosus* Kunth), (3) *P. coccineus* subsp. *glabellus* (Piper) A. Delgado Salinas (= *P. glabellus* Piper), (4) *P. coccineus* subsp. *griseus* (Piper) A. Delgado Salinas (= *P. griseus* Piper), and (5) *P. coccineus* subsp. *darwinianus* Hdez. X and Miranda C. (= *P. polyanthus* Greenman).

Chloroplast DNA (cpDNA) has been extensively used as a molecular marker in evolution analysis, and its advantages have been discussed elsewhere (e.g., Dowling et al. 1990; Crawford 1990; Clegg and Zurawski 1992). The disadvantages and risk of generalizations based on the use of cpDNA as the only marker to infer phylogenetic relationships in plant species have also been reviewed (Rieseberg and Brunsfeld 1992; Harris and Ingram 1991).

In this report we describe our analysis of changes occurring in the cpDNA among different taxa of the *P. vul*garis complex. The objectives of this analysis were: (1) to assess the mode of evolution of cpDNA based on the type and distribution of changes; (2) to use cpDNA as a molecular marker to clarify the systematics of the *P. vulgaris* complex; and (3) to evaluate the diversity of some of the taxa analyzed.

Materials and methods

DNA extraction

Plants from the accessions listed in Table 1 were grown in a greenhouse. Total DNA was isolated from young leaves according to the method described by Gepts et al. (1992). Harvested leaves (2–5 g per plant) were ground in liquid nitrogen and placed in 10 ml extraction buffer (50 mM TRIS, 700 mM NaCl, 1 mM Na₂EDTA, 1% SDS, 1% β -mercaptoethanol, 50 mg/ml polyvinyl polypyrrolidone, pH 9.0). The suspension was then incubated at 60 °C for 1 h, followed by two chloroform:isoamyl alcohol (24:1) extractions. DNA was precipitated with isopropanol, washed with 70% ethanol and resuspended in TE (10 mM TRIS, 1 mM Na₂EDTA). DNA concentration was estimated using a dedicated fluorometer TK-100 (Hoefer, San Francisco, Calif.).

DNA samples from some accessions presented digestibility problems and were further purified by ultracentrifugation in a cesium chloride-ethidium bromide gradient basically according to Sambrook et al. (1989). Samples were set in a 1.55 g/ml cesium chloride solution containing 250 g/ml ethidium bromide. After ultracentrifugation at 400,000 g for 12 h, the DNA was removed and ethidium bromide was extracted with TE-saturated butanol. Samples were dialyzed against TE (1000:1) for 10 h and then precipitated with ethanol. Purified DNA was resuspended in TE at a final concentration of 300 µg/ml.

Restriction mapping

The wild accession 'G12873' belonging to the Middle American gene pool (Gepts 1990) was used for the *P. vulgaris* map. The cpDNA map

for *P. coccineus* subsp. *coccineus* was constructed using a wild accession collected in Chihuahua, Mexico ('DelgadoB'). Additionally, partial maps were made from a wild *P. coccineus* subsp. *glabellus* accession ('DGD2043') and from a cultivated Andean *P. lunatus* accession ('PG11A').

In each case, 3 µg of total DNA was digested separately with the following 6-base restriction endonucleases: SalI, PstI, PvuII, SmaI, KpnI, XhoI, SstI, HindIII, BamHI, DraI, and EcoRI (Bethesda Research Laboratories; New England Biolabs; Boehringer Mannheim). The reaction conditions were those specified by the supplier. In addition, 50 ng/µl RNase was added to the reaction mixture. Samples were incubated for 4–5 h using an eightfold enzyme excess to assure total digestion. In addition to the 11 single digestions, 44 double digestions were made. When optimal conditions were different for each of the 2 enzymes in a double digestion, the digestion with lower salt concentration was accomplished first, followed by addition of the second enzyme and KCl or NaCl to the recommended concentration.

For mapping, restriction fragments were separated by electrophoresis in a 20 to 25 cm-long, 0.5%-0.6% horizontal agarose gels in $1 \times TAE$ (40 mM TRIS, 20 mM glacial acetic acid, 10 mM Na₂-ED-TA), running at 1 V/cm, as described by Sambrook et al. (1989). Lambda phage DNA cut with *Hind*III was used as the size standard. For comparative analysis among accessions, gels and DNA transfer were made according to the same protocols. However, gels for comparative analysis were only 10 cm long and consisted of 0.8% agarose in $1 \times TAE$.

Hybridization was carried out initially according to the procedure described in Gepts et al. (1992). The procedure was subsequently modified as follows: DNA fragments from the agarose gels were denatured and transferred by capillarity onto nylon (Zetabind) membranes according to Sambrook et al. (1989) and then crosslinked with 150 mJ UV light using a Bio-Rad UV linker. Nylon membranes were prehybridized at 42 °C in 5 × SSPE, 1% SDS, 50% formamide for 30 minutes. The probe (10^5 cpm/cm²) was then added to the same solution and incubated at 42 °C for another 20 h. The membrane was washed briefly in 2 × SSC, 0.1% SDS at 60 °C. An X-ray film was subsequently exposed to the membrane for from 3–4 h to 1 day. This method gave excellent results and was inexpensive and faster than the previous method. Membranes were reused approximately 15 times. Prior to reuse, the probe was removed by washing in 0.4 *N* NaOH at 42 °C for 30 min, followed by a 30-min wash in 0.5 *M* TRIS, 0.1×SSC, 0.1% SDS at 42 °C.

DNA probes and labeling

Plasmids containing *Vigna radiata* cpDNA fragments (kindly provided by Dr. J.D. Palmer, University of Indiana at Bloomington; Palmer and Thompson 1981) were amplified in competent cells of *Escherichia coli* strain DH5 α prepared according to the CaCl₂ method and transformed with the plasmids containing the fragments. Plasmids were isolated by the boiling miniscreen procedure (Sambrook et al. 1989) and then labeled with [³²P]-dCTP by nick translation (Rigby et al. 1977) or by random priming (Feinberg and Vogelstein 1983).

Phylogenetic analysis

Phylogenetic analysis was performed with the program PAUP (Phylogenetic Analysis Using Parsimony) Version 3.0 provided by D.L. Swofford (Illinois National History Survey, Champaign, Ill.). Mapped restriction sites were analyzed in a cladistic fashion using Wagner (Farris 1970), Dollo (LeQuesne 1974; Farris 1977), and weighted parsimony (Albert et al. 1992). For the weighted method we used a restriction site gain:loss ratio equal to 1:2.3, which is considered to be optimal for cpDNA restriction site analysis (Albert et al. 1992).

Initially, *P. lunatus* was used as the outgroup in order to polarize the mutations because it is phylogenetically distant from the *P. vul*-

garis complex within the genus (Delgado Salinas et al. 1993; Mok et al. 1978). However, as explained later, *P. lunatus* turned to be an inadequate outgroup for our analyses and was subsequently included as an additional accession in the unrooted trees.

We estimated phylogenies using the whole set of accessions or representative accessions from each taxa. For the whole set of accessions (30 samples and 48 synapomorphies), we used heuristic searches using Tree Bisection-Reconnection (TBR) and ACCTRAN optimization in PAUP 3.0. For smaller phylogenies (e.g., 6 taxa and 46 synapomorphies), we were able to use the Branch-and-Bound search (for weighted parsimony) and the exhaustive search modes (for Dollo and Wagner parsimonies) and ACCTRAN optimization. For bootstrapping, we made 100 repetitions in the Phylip 3.01 programs Boot and Dolboot (Felsenstein 1988).

For the estimation of phylogeny under weighted parsimony, we used the Stepmatrix option of PAUP 3.0, with heuristic search using Tree Bisection-Reconnection (TBR) and ACCTRAN optimization. Random data acquisition sequence was used with heuristic search and Tree Bisection-Reconnection (TBR) swapping to find the most parsimonious trees. A 10:23 step matrix was employed as suggested by Albert et al. (1992).

Results

Development of a *Phaseolus vulgaris* cpDNA restriction map

Single and double digestions of total genomic DNA from the wild *P. vulgaris* accession 'G12873' were performed using 11 endonucleases. cpDNA was visualized by sequential Southern hybridization of total DNA to radioactively labeled heterologous cpDNA probes from mung bean (*V. radiata*), a species from a related genus (Palmer and Thompson 1981). Mubumbila et al. (1983) and Palmer et al. (1983) previously constructed a *P. vulgaris* cpDNA map for 3 enzymes (*PstI, XhoI* and *SalI*) and also demonstrated the colinearity and high similarity between this genome and that of *V. radiata* cpDNA.

A total of 11 simple and 44 double digestions were carried out. The mung bean cpDNA probes comprised 96% of the complete cpDNA genome. Three hundred and thirty 6-bp restriction sites were assigned to a restriction map. This corresponds to 1.3% of the total cpDNA sequence (on average, 1 site every 460 bp). The restriction enzymes *Sal*I, *PstI*, *SmaI*, *KpnI*, *XhoI*, *PvuII*, and *SstI* cut *P. vulgaris* cpDNA at a low frequency (5–20 sites), whereas the enzymes *DraI*, *HindIII*, *Bam*HI, and *Eco*RI cut it at high frequencies (40–100 sites/genome) (Fig. 1).

The complete *P. vulgaris* cpDNA map is presented in Fig. 1. The map presents only one of two possible isomeric conformations present in similar proportions in *P. vulgaris* as demonstrated by Palmer (1983). The map is complete for 9 enzymes. For 2 high-frequency cutting enzymes (*Eco*RI and *Dra*I), there were some ambiguous regions that have been omitted in the map. The ambiguity is due to the presence of many small fragments and the absence of overlapping fragments in small non-probed regions.

An average size of *P. vulgaris* cpDNA of 152.4 kb was calculated using the 9 restriction endonucleases with the lowest cutting frequency because of the possibility of large amounts of undetected small fragments present in the high-



Fig. 1 High-resolution restriction map of *P. vulgaris* cpDNA. The *top row* shows the *Vigna radiata* cpDNA probes used in the Southern hybridizations. The *second row* is a composite restriction map showing all restriction sites detected with the 11 enzymes. The *following rows* are restriction maps for individual enzymes. The two *heavy horizontal bars* are the inverted repeats. The three *boxes* marked by *stippled lines* represent regions with a high frequency of polymorphism. *Arrows* indicate restriction site differences with the *P. coccineus* subsp. *coccineus* map. The *blank region* in the *Eco*RI map could not be mapped because of the high frequency of restriction sites.

frequency restriction digestions with *Dra*I and *Eco*RI. Minor discrepancies among size calculations for each enzyme were probably due to small fragments (0.3-0.05 kb) not being detected under our conditions. Estimates for some of the fragments sizes differed slightly (0.5-3%) from those previously published (Mubumbila et al. 1983). This is probably due to differences in the procedures followed in both works. We visualized restriction fragments after Southern hybridization of total DNA fragments separated in ethidium bromide-agarose gels instead of using purified cpDNA. Agarose gel overloading and the presence of ethidium bromide in the gel may retard DNA fragments in gel.

The small single copy region was approximately 19 kb long. Each repeat had a size of at least 25 kb, whereas the large single-copy region was approximately 83 kb long. These values are consistent with those obtained for *P. vulgaris* and other angiosperms with cpDNA containing inverted repeats (reviewed in Crouse et al. 1986; Palmer and Stein 1986; Palmer 1987). Comparison among cpDNA restriction maps

A second cpDNA map was constructed for wild accession 'DelgadoB' of P. coccineus subsp. coccineus using the same restriction endonucleases as above. Similar results were obtained in P. coccineus subsp. coccineus and P. vulgaris with respect to the frequency of restriction sites recognized by the various restriction enzymes, but a total of 16 differences were found between their respective genomes, which corresponds to 5% of the sites analyzed. Figure 1 shows the differences found between P. coccineus subsp. coccineus and P. vulgaris cpDNA. All of the detected differences except one are probably simple restriction site gains or losses. The exception is most likely due to a small insertion-deletion mutation in region p11.1. No major rearrangements (large insertions, deletions, inversions, or translocations) were detected. However, additional insertions and inversions of up to 300 bp may have been overlooked by our methods in some cases.

All mutations except 3 were located in the large singlecopy region. The distribution of mutations in this region was uneven: 1 single 10-kb region in s13.3 contained 6 site differences, that is, 40% of the total observed. The presence of monomorphic sites between polymorphic sites suggests that the former sites are not the product of a single event (e.g., insertion, deletion or inversion), but have been produced independently. An uneven distribution of mutations has been found in other species (Close et al. 1989). In our subsequent analyses, all mutations, except those of *Bam*HI p16.2 and *Bam*HI p17.2, were considered to be independent. The *Bam*HI p16.2 and *Bam*HI p17.2 mutations are located in the inverted repeat, one in each, and may have resulted from sequence homogenization, as has been inferred in other plant species (Palmer 1985).

Partial maps based on 4 enzymes (*DraI*, *EcoRI*, *Bam*HI and *Hin*dIII) were also generated for *P. lunatus* (PG11A) and *P. coccineus* subsp. *glabellus* ('DGD2043'). Such maps were required in order to assign the position of sites in regions where inference from the other maps was not possible because of the presence of multiple changes.

Phylogenetic analysis

We screened several regions of cpDNA in 30 accessions of the *P. vulgaris* complex. The samples included wild and cultivated accessions of (1) *P. coccineus* subsp. coccineus, (2) *P. coccineus* subsp. darwinianus (= *P. polyanthus*), (3) *P. coccineus* subsp. formosus, (4) *P. coccineus* subsp. glabellus, (5) the Middle American and Andean *P. vulgaris* gene pools, and (6) *P. lunatus*. The latter was included originally as the outgroup to polarize mutations within the *P. vulgaris* complex.

An analysis of 190 restriction sites revealed 60 mutations. All of these mutations except 9 were shared by at least 2 or more taxa, and consequently they were phylogenetically informative. Most of the mutations (56) could be explained by restriction site gain/loss events, whereas 4 were probably occasioned by small (ca. 0.1 kb) insertions or deletions. Length mutations were not included in our

Fig. 2 Cladogram showing relationships from the restriction site data among accessions of the *P. vulgaris* complex. This is one of ten most parsimonious cladograms for the 30 separated accessions, with a length of 63 steps and a consistency index of 780 (Homoplasy index = 220). The *first three letters* of the label for each accession represent the taxon to which it belongs: (*coc P. coccineus* subsp. *coccineus*, *dar P. coccineus* subsp. *darwinianus*, *for P. coccineus* subsp. *formosus*, *gla P. glabellus*, *lun P. lunatus*, *vul P. vulgaris*). The *last letter* indicates whether the accession is wild-growing (*w*) or cultivated (*c*)



phylogenetic analysis. As observed in Fig. 2, cpDNA data analysis showed *P. lunatus* to be more closely related to *P. coccineus* subsp. *coccineus* and *P. coccineus* subsp. *darwinianus* than to *P. coccineus* subsp. *glabellus*. Surprisingly, the latter taxon was distantly related to all of the other taxa analyzed in this study.

Discussion

Levels of polymorphism

We detected significant levels of polymorphism among species and within species in the P. vulgaris complex. Recent work indicates that moderate variation at the interpopulational (and even intrapopulational) level is common in at least several other species (see Soltis et al. 1992 and Crawford 1990 for a review). Our results show taxon-dependent differences in polymorphism. P. coccineus subsp. coccineus showed the highest number of polymorphic sites. On the other hand, Phaseolus coccineus subsp. darwinianus did not show polymorphism in any of the wild and cultivated accessions analyzed. Only 1 polymorphism was observed among the 3 accessions of P. vulgaris in spite of the marked divergence of nuclear DNA sequences between Middle American (e.g., 'BAT93') and Andean (e.g., 'Jalo') genotypes (Nodari et al. 1992). This polymorphism differentiates 'BAT93' and other genotypes from race Mesoamerica (Singh et al. 1991) from other Middle American genotypes and Andean genotypes (V. Llaca, V. Becerra, and P. Gepts unpublished results).

Phylogenetic implications for *P. coccineus* subsp. *darwinianus*

cpDNA from all of the accessions belonging to *P. coccineus* subsp. *darwinianus* was more similar to that of *P. vulgaris* than to that of *P. coccineus* subsp. *coccineus*. According to our most parsimonious trees, *P. vulgaris* is separated from *P. coccineus* subsp. *darwinianus* by only 3 changes, whereas at least 14 changes are required to explain the differences between the closest *P. coccineus* subsp. *coccineus* subsp. *darwinianus*. Similar results were obtained by Schmit (1992).

These results suggest a discrepancy between the phylogenetic relationships suggested by cpDNA and nuclear DNA of *P. coccineus* subsp. *darwinianus*. Data from previous studies using nuclear markers such as enzymes and antibodies against seed proteins (Piñeiro and Eguiarte 1989; Kloz and Klozova 1974) and our own observations using restriction fragment length polymorphism (RFLP) analysis of random single-copy and highly repetitive nuclear clones as well as the polymerase chain reaction (PCR) amplification of phaseolin genes (V. Llaca and P. Gepts unpublished results) indicate a close relationship between *P*. coccineus subsp. darwinianus and P. coccineus subsp. coccineus, whereas both are readily discernible from P. vulgaris. The similarity in the nuclear genomes of P. coccineus subsp. darwinianus and P. coccineus subsp. coccineus is consistent with the reduced reproductive isolation between these two subspecies (Gepts 1981).

The lack of coherence between nuclear and chloroplast lineages may be accounted for by a reticulated origin of P. coccineus subsp. darwinianus. The intermediate morphology of P. coccineus subsp. darwinianus compared with that of P. vulgaris and P. coccineus subsp. coccineus was first noted by Hernandez-Xolocotzin et al. (1959), who proposed a hybrid origin for this taxon, with *P. coccineus* subsp. *coccineus* as the seed parent and *P. vulgaris* as the pollen parent. Alternate views considered a gradual differentiation of P. vulgaris from P. coccineus subsp. darwinianus more likely (Smartt 1973, Schmit and Debouck 1991). Our work supports the first point of view but suggests that P. vulgaris or a taxon closely related to it would be the seed parent due to maternal inheritance of cytoplasmic DNA in Phaseolus (H. Bannerot personal communication). Recurrent unidirectional backcrosses with P. coccineus plants as male parents could explain the similarity of the nuclear genomes. Hybridization would have occurred before the split of the two major gene pools (Middle American and Andean) in P. vulgaris (Gepts 1990; Gepts and Debouck 1991) because all P. coccineus subsp. darwinianus accessions share the same apomorphies with the P. vulgaris accessions representing the two gene pools. According to the distribution of both the cultivated and few wild P. coccineus subsp. darwinianus accessions collected (Schmit and Debouck 1991; R. Ramirez personal communication), this taxon originated in Middle America (Mexico and Central America) and has only recently been introduced into South America.

Phaseolus glabellus cpDNA

The results obtained in this work strongly suggest that the chloroplast of *P. coccineus* subsp. *glabellus* is very distant from that of the other taxa analyzed. *P. coccineus* subsp. *glabellus* had been included in the *P. vulgaris* complex because of the presence of scarlet flowers, cross-pollination, and a perennial life cycle, which are all also observed in *P. coccineus* subsp. *coccineus* (Piper 1926; Maréchal 1978; Delgado-Salinas 1985). In this study, *P. coccineus* subsp. *glabellus* cpDNA restriction sites showed a considerable divergence from those of any of the other taxa. Furthermore *P. lunatus* cpDNA was intermediate between that of *P. coccineus* subsp. *glabellus* and that of the other taxa surveyed. These results confirm those obtained by Schmit (1992).

Several morphological traits, the presence of manifest reproductive barriers, and the great divergence at the molecular level suggest the reclassification of *P. glabellus* as a different species outside the *P. vulgaris* complex. Inflorescence number and structure, leaf structure, and seed morphology (Maréchal et al. 1978, Delgado-Salinas 1985) as well as the absence of natural or artificial hybrids (Delgado Salinas 1988; Sousa-Peña et al. unpubl. results) also argue against a close relationship with *P. coccineus*. Variability of cultivated P. coccineus subs. coccineus

P. coccineus subsp. *coccineus* showed a high level of intraspecific polymorphism. All of the cultivated accessions were grouped in the same cluster, together with a wild accession from the Mexican state of Puebla, which suggests a single center of domestication. The reduction in diversity observed between wild and cultivated *P. coccineus* subsp. *coccineus* confirms observations made previously for cpDNA in other wild ancestor-cultivated descendant combinations (Doebley 1992). It also confirms observations made in other species of the *Phaseolus-Vigna* group with other molecular markers (*P. vulgaris*: Gepts et al. 1986; *P. acutifolius*: Schinkel and Gepts 1988, 1989; *Vigna unguiculata*: Panella and Gepts 1992).

We were unable to distinguish between the cpDNA of *P. coccineus* subsp. *formosus* and *P. coccineus* subsp. *coc*cineus. Furthermore, some wild P. coccineus subsp. coccineus accessions were closer to P. coccineus subsp. formosus than to other P. coccineus subsp. coccineus accessions. Therefore, a very close relationship among accessions of both subspecies is inferred. Nevertheless, P. coccineus subsp. formosus seems to include a heterogenous group of populations. For example, individuals with high reciprocal compatibility to P. vulgaris and lilac flowers have been described within P. coccineus subsp. formosus, whereas common wild, red-flowered wild plants are also included in the same group (Delgado Salinas 1988). The P. coccineus subsp. formosus accession included in this work corresponds to the second type, that similar in morphology to wild P. coccineus subsp. coccineus. The analysis of cpDNA in lilac-flowered, *P. vulgaris*-compatible populations could show if *P. coccineus* subsp. *formosus* is in fact a monophyletic group or whether it consists of a heterogeneous group of populations.

Overall, our results set the stage for further analysis of phylogenetic relationships in the genus *Phaseolus* and related genera. They underscore the need to analyze cytoplasmic and nuclear genomes to establish more reliables phylogenies.

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