Genetic diversity among Spanish pea (*Pisum sativum* L.) landraces, pea cultivars and the World *Pisum* sp. core collection assessed by retrotransposon-based insertion polymorphisms (RBIPs)

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Abstract

A total of 122 accessions of different wild and cultivated *Pisum* sp. were analysed using retrotransposon-based insertion polymorphisms (RBIP) markers. The *Pisum* materials included wild and cultivated (landraces and cultivars) materials from the World core collection of the John Innes Centre (JI) representing all generally recognized *Pisum* taxa, landraces materials from the Spanish core collection, and commercial pea cultivars largely sown in Spain. The overall polymorphism detected by RBIP marker was high and all accessions, except two pairs, could be distinguished by their marker pattern. Principal component and phylogenetic analyses clearly discriminated *P. fulvum* and *P. abyssinicum* samples from both each other and *P. sativum*, while *P. elatius* and *P. humile* samples were scattered among the other taxa clusters, supporting the existence of three well defined taxa in the genus *Pisum* (*P. abyssinicum*, *P. fulvum* and *P. sativum*). These results also suggest that the Spanish pea core collection of landraces maintains a relatively high variability which is only partially represented in cultivars generally sown in Spain. Thus, Spanish landraces are still a source of genetic variability for breeding new pea cultivars.

Additional key words: genetic resources, Pisum abyssinicum, Pisum fulvum, Pisum sativum, RBIP markers.

Resumen

Diversidad genética en variedades locales y cultivares españoles de guisante (*Pisum sativum* L.) y en la colección nuclear mundial de *Pisum* estimada mediante polimorfismo de inserción de retrotransposones (RBIP)

Se ha estudiado un total de 122 accesiones silvestres y cultivadas de *Pisum* sp. usando marcadores basados en polimorfismos de inserción de retrotransposones (RBIP). Las accesiones de *Pisum* incluyen materiales silvestres y cultivados (cultivares y variedades locales) de la colección nuclear mundial del John Innes Centre (JI) representando a todos los taxones generalmente reconocidos de *Pisum*, variedades locales de la colección nuclear española, y por último algunas variedades comerciales de guisante ampliamente cultivadas en España. Para el análisis genético se usaron 18 loci RBIP. El polimorfismo general detectado con los marcadores RBIP fue alto y todas las muestras, excepto dos pares, pudieron ser identificadas por un patrón particular de marcadores. Análisis de componentes principales y filogenéticos discriminaron claramente *P. fulvum* y *P. abyssinicum* entre ellas y de *P. sativum*, mientras que las muestras de *P. humile* y *P. elatius* se mezclaban con las de otros taxones en distintos grupos. Esto apoya la existencia de tres especies en el género *Pisum (P. abyssinicum, P. fulvum* y *P. sativum)*. Los resultados indican que la colección nuclear española de guisante mantiene una variabilidad relativamente elevada que está sólo parcialmente representada en los cultivares generalmente sembrados en España. Por tanto, las variedades locales españolas representan aún una fuente de variabilidad genética para la mejora de nuevos cultivares.

Palabras clave adicionales: marcadores RBIP, Pisum abyssinicum, Pisum fulvum, Pisum sativum, recursos genéticos.

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Abbreviations used: IRAP (inter-retrotransposon amplified polymorphism), ISSR (inter simple sequence repeats), JI (John Innes Centre), PCA (principal component analysis), RBIP (retrotransposon-based insertion polymorphisms), REMAP (retrotransposon-microsatellite amplified polymorphism), SSAP (sequence-specific amplification polymorphisms), ZP (ITACyL collection).

Introduction

Pea (Pisum sativum L.) is a major cool season legume crop for human consumption, as dry seeds or as vegetable, and for feeding livestock. Pea was also one of the first domesticated crops in the Old World and one of the first genetic research materials. The modern gene pool of cultivated Pisum is diverse, reflecting this early domestication and subsequent widespread cultivation. However, in spite of the extensive phenotypic and genetic variability, existing taxonomic classifications are confusing (Vershinin et al., 2003; Kosterin and Bogdanova, 2008). In addition to P. sativum, two other species are generally recognized within the genus Pisum, the wild Pisum fulvum Sibth. et Smith., which is almost reproductively isolated from P. sativum, and Pisum abyssinicum A. Br. represented by cultivated and some wild forms from South Arabia and Ethiopia. Other taxa once considered as species are in fact sensu lato representatives of P. sativum. Some of them are presently considered as subspecies, although the subspecies concept in the case of the pea remains quite vague, and according plastid, mitochondrial and nuclear markers all wild forms of P. sativum would better be considered within a fuzzy paraphyletic subspecies P. sativum ssp. elatius (Bieb.) Schmalh. sensu lato (Kosterin and Bogdanova, 2008). A similar phylogenetic organization of taxa was previously described by Maxted and Ambrose (2001) in which three species were recognized (P. abyssinicum, P. fulvum and P. sativum with two subspecies ssp. sativum and ssp elatius (Bieb.) Aschers. & Graebn., considering P. humile Boissier and Noe as a variety of *P. sativum* spp. *elatius*). According to plastid, mitochondrial and nuclear sequences (rbcL, coxI and SCA, respectively) Kosterin et al. (2010) pointed to the lineage B of P. sativum ssp. elatius as the origin of the cultivated P. sativum. A study of the genetic structure and evolutionary history of Pisum based on retrotransposon sequence-specific amplification polymorphisms (SSAP) revealed high polymorphism in all species, except P. abyssinicum. The results indicated a high contribution of recombination between multiple ancestral lineages compared to transposition within lineages, suggesting that the two independently domesticated pea species, P. abyssinicum and P. sativum, arose independently in contrasting ways via the common processes of hybridization, introgression, and selection (Vershinin et al., 2003).

Retrotransposons are ubiquitous in plant genomes and they vary in copy number and chromosomal location within and between species, and play significant roles in genome evolution (Flavell *et al.*, 1992; Baucom *et al.*, 2009; Hawkins *et al.*, 2009). Particular retrotransposon families can vary greatly in abundance and chromosomal location even between closely related species (Pearce *et al.*, 1996; Kubis *et al.*, 1998; Hill *et al.*, 2005), or contribute to novel satellite repeats (Macas *et al.*, 2009). Because of this ubiquity and diversity, retrotransposons based markers are powerful tools for the assessment of genetic diversity, and have shown their usefulness as genetic markers and in biodiversity and phylogenetic analyses (Ellis *et al.*, 1998; Flavell *et al.*, 1998; Vershinin *et al.*, 2003; Jing *et al.*, 2007; Martín-Sanz *et al.*, 2017; Agarwal *et al.*, 2008; Tam *et al.*, 2009; Jing *et al.*, 2010).

Several types of genetic markers have been derived from retrotransposons, including retrotransposonbased insertion polymorphisms (RBIP), inter-retrotransposon amplified polymorphism (IRAP), retrotransposon-microsatellite amplified polymorphism (REMAP), and SSAP (Waugh et al., 1997; Syed and Flavell, 2006); and they have been used in genetic analyses such as gene mapping (Ellis et al., 1998), genotyping, and pea cultivar fingerprinting (Smýkal, 2006). RBIP markers detect presence or absence of individual retrotransposon insertions in the genome; the method requires flanking sequence information for primer design and yields co-dominant markers, where the different allelic states at a locus can be revealed (Flavell et al., 1998). RBIP markers have been recently used for both broad diversity analysis and variety discrimination in pea (Smýkal et al., 2008b; Jing et al., 2010) and have proved to be the most robust and easy to score retrotransposon-based marker method in comparison to IRAP and other marker systems (Smýkal et al., 2008a).

The estimation of the genetic variability within pea collections and the relationships between accessions using molecular markers has been carried out in numerous works. Recently, evaluations of genetic diversity among European pea materials using isozyme, protein and PCR markers and among Spanish materials using inter simple sequence repeats (ISSR) markers have been published (Baranger *et al.*, 2004; Lázaro and Aguinagalde, 2006). Zong *et al.* (2009) analyzed two very wide collections of Chinese and world accessions (this later collection included some wild materials), respectively, using microsatellite markers. They found that the genetic diversity of *P. sativum* within China appears to be quite different to that detected in the global gene pool, these genetically distinct gene pools within domestic field pea has significant implications in broadening the available variability for further genetic improvement.

Here we described the use of RBIP markers in the analysis of genetic variability and relationships of the Spanish *P. sativum* landrace core collection and pea cultivars currently sown in Spain in relation to a representative set of the *Pisum* World core collection provide by the John Innes Centre, which includes wild and cultivated accessions. The results obtained will contribute to gain a better knowledge of *Pisum* resources genetic variability and be useful for future breeding purposes to improve pea cultivars.

Material and methods

The Pisum accessions used are summarized in Table 1. They included wild and cultivated materials, some of them are included in the World core collection of the John Innes Centre (JI), another set of materials represent the Spanish core collection of landraces (Caminero et al., 2001; Ramos, 2003) and traditional varieties (ZP) conserved in the Spanish gene bank, and finally some commercial cultivars largely cultivated in Spain, currently or in the past. Spanish landraces were collected by the Plant Genetic Resource Center (INIA, Spain) from 1971 to 2000 in pea-growing areas with different agroclimatic conditions. These materials have been traditionally cultivated by local farmers in conventional and organic farms. The JI accessions included in this study coincide with accessions included in previous works (Jing et al., 2005, 2007); this JI collection represents the diversity of the genus Pisum (Jing et al., 2005). All materials will be referred as accessions in this paper. A total of 122 accessions were analyzed. A summary on the species and the type of material (landrace, cultivar) is shown in Table 1, additional information on their collection numbers, names, origin, etc., is compiled in Appendix 1. Irrespective of the mainly accepted taxonomic classification of *Pisum* species and subspecies (Maxted and Ambrose, 2001), the binomial nomenclature will be used in this work for simplicity.

Genomic DNA was isolated from young leaf tissue using the Quiagen (Valencia, CA) Dneasy 96 plant kit method according manufacturer recommendations. A single individual per accession was analyzed. The RBIP technique was performed as described by Flavell *et al.* (1998), using as primers the flanking sequences of 25 insertions of the retrotransposon *PDR1* defined by Jing *et al.* (2005). A total of 25 RBIP markers were assayed, but monomorphic and those amplifying multiple bands were not considered. The primer sequences and size of the 18 RBIP loci scored is summarized in Appendix 2. Each RBIP was considered as a locus, defining one or more alleles depending of the amplicon fragment size, or the absence of any PCR product corresponding to primer site mutation (Jing *et al.*, 2005) (Fig. 1). Expec-



Figure 1. Agarose gel (1.5%) showing three different alleles of the RBIP locus 281x1. Numbers indicate the reference size markers in base pairs.

Species	World collection	Spanish landrace collection	Cultivars	Others	Total
P. sativum					
ssp. sativum	22	43	27	1	93
ssp. elatius	12				12
ssp. humile	2				2
P. fulvum	10				10
P. abyssinicum	1			4	5
Total	47	43	27	5	122

 Table 1. Pisum materials^a used in the retrotransposon-based insertion polymorphisms (RBIP) analysis

^a Figures indicate the number of accessions analyzed.

Collection	Average	Standard deviation	Maximum value	Minimum value
			(locus)	
Whole collection ^a	0.451	0.167	0.640 (<i>45-x31</i>)	0.079 (2385-x64)
P. fulvum	0.144	0.220	0.580(95-x2)	0.0 (12 loci)
P. elatius	0.367	0.224	0.653 (399-80-46)	0.0 (3 loci)
<i>P. sativum</i> Wc ^b	0.370	0.202	0.628 (64-x45)	0.0 (3 loci)
<i>P. sativum</i> Sc ^c	0.373	0.208	0.623(281-x1)	0.0 (2 loci)
<i>P. sativum</i> cc ^d	0.222	0.220	0.590 (45-x29)	0.0 (6 loci)

Table 2. Average of expected heterozygosity (H_e) over loci

^a Including *P. abyssinicum* and *P. elatius*. These two species were not included in the subset analyses due to the low number of accessions. ^b Wc: World collection. ^c Sc: Spanish landrace collection. ^d cc: commercial cultivars.

ted heterozygosity was estimated as $H_e = 1 - \Sigma p_i^2$, were p_i are the allelic frequencies. Shared allele distance (Chakraborty and Jin, 1993) was calculated from the proportion of shared alleles P_{AS} as $D_{AS} = 1 - P_{AS}$. Nei distance was also calculated as $1 - I_S$. Both distances were calculated by the program MSAT2 (http://hpgl. stanford.edu/projects/microsat/). Bootstrap was carried out, and Neighbour-Joining and UPGMA methods were used for generating trees. Principal component analysis was performed using «Proportion-of-shared-alleles distance» genetic distance.

Results

A total of 122 accessions were analyzed using 18 RBIP loci. A total of 56 alleles were observed in the 18 loci (2 to 4 alleles per locus). As expected in highly self-pollinated materials the homozygosity was predominant, only 13 cases of heterozygosity were detected in the 2,196 accession × loci combinations, approximately a 0.6% of observed heterozygosity which agree with the predominantly self-pollination mating system of *Pisum*. Twelve accessions and seven loci showed at least a case of heterozygosity. The RBIP alleles used indicated a relatively high level of polymorphism. The expected heterozygosity (H_e) for the whole set of accessions and loci was 0.658, the average heterozygosity among loci was 0.451 ranging from 0.079 (locus 2385-x64) and 0.640 (locus 45-x31). For those set of accessions represented by at least 10 accessions the heterozygosity values are indicated in Table 2. The levels of polymorphism maintained within the pea World collection, the Spanish landrace collection and *P. elatius* were similar (H_e approximately 0.370) while the set of cultivars showed a significantly (p < 0.05) lower average value (0.222).

This high polymorphism of RBIP markers allowed identifying all accessions by a RBIP pattern combination, except P. fulvum accessions JI-2519 from JI-2544 and cultivars Lucy from Messire (both French commercial cultivars), respectively. Thus RBIPs are suitable markers for the identification of pea materials. Some species-specific alleles due to retrotransposon insertion were observed in P. abyssinicum (loci 2385-x64 and 95-x19) and P. sativum (loci 45-x31, Birte-x34, and 281-x1). Since the probability of detecting rare alleles increases as sample size increases, it is possible that the species-specific RBIP alleles observed in the most represented sample of P. sativum (93 accessions) represent rare Pisum alleles, but in a sample of only five P. abyssinicum accessions those alleles most be «true» species-specific alleles, frequent in a species but rare or absent in related species.

The average distances within and between taxa are shown in Table 3. On the basis of RBIP polymorphisms

Table 3. Average distances^a within and between *Pisum* species (standard deviation)

	P. fulvum	P. abyssinicum	P. humile	P. elatius	P. sativum
P. fulvum	0.172 (0.083)				
P. abyssinicum	0.570 (0.096)	0.307 (0.161)			
P. humile	0.460 (0.052)	0.561 (0.076)	0.426 (0.091)		
P. elatius	0.556 (0.073)	0.562 (0.090)	0.444 (0.133)	0.389 (0.124)	
P. sativum	0.640 (0.094)	0.647 (0.098)	0.498 (0.120)	0.485 (0.139)	0.388 (0.130)

^a Proportion-of-shared-alleles distance.

and H_e , *P. fulvum* seemed to be the less diverse taxon, while *P. elatius* and *P. sativum* showed similar level of internal diversity, and *P. abyssinicum* was lower than these two taxa. The number of accession of *P. humile* was too low in our study to draw valid conclusions. On the other hand, *P. fulvum* and *P. abyssinicum* showed the greatest and similar average distances to *P. sativum*.

Clustering methods showed low bootstrap confidence values for tree nodes. Although these values were too low to be significant, the two distances used (proportion-of-shared-alleles and Nei) and two clustering methods (Neighbour-Joining and UPGMA) generated very similar or identical topologies (data not shown), which conferred robustness to the results. The shown data were obtained with the proportion-of-shared-alleles distance and Neighbour-Joining method. The unrooted tree (Fig. 2) showed that *P. fulvum* formed a well differentiated cluster (1 in Fig. 2); *Pisum abyssinicum* formed a second cluster with some *P. elatius* accessions (2). The remaining *P. elatius* and *P. fulvum* accessions were mainly grouped with pea landraces. All the pea cultivars were grouped in a big cluster (3) with some pea landraces. Pea cultivars included in this group belong to two sets of accessions, to materials sown in Spain and to cultivars included in the JI collection (JI-321 is cultivar Alaska from Canada, JI-399 is Cenia-The Netherlands, JI435 is Wisconsin Perfection-USA, JI-516 is Maro-UK, and JI-113 is an unnamed cultivar from Russia). The remaining pea landraces were scattered in several small clusters irrespective their origin, Spanish landraces or landraces from JI collection.

Principal component analysis (PCA) also clearly discriminated between *P. fulvum* and *P. abyssinicum* and these two taxa from the remaining *Pisum* accessions (Fig. 3). Each of the three first components explained percentages of the total variance higher than 10%



Figure 2. Neighbour joining unrooted tree for *Pisum* accessions deduced from 18 RBIP retrotransposon markers. Names indicate the accession register as follow: Pa, *P. abyssinicum;* Pe, *P. elatius;* Pf, *P. fulvum,* Ph, *P. humile* from the John Innes World Core Collection (JI), all other accessions are registered as *P. sativum.* Numbers indicate the register number at the JI and the ITACyL (ZP) collections, respectively. Accessions indicated with names are cultivars from the ZP collection. 1, 2 and 3 indicate the clusters in which *P. fulvum, P. abyssinicum* and the pea cultivars are grouped, respectively. Branch length units are shown at the bottom.



Figure 3. Plot of the principal component analysis. Abbreviations as in Figure 2. Ps, *Pisum sativum* landraces and traditional materials; PsC, *pea cultivars. P. fulvum, P. abyssinicum* and pea cultivars are highlighted within circles.

(17.4, 13.7 and 10.7, respectively; 41.7% accumulated). The first component differentiated *fulvum-abyssinicum* from the remaining *Pisum* taxa, while the second discriminated between *P. fulvum* and *P. abyssinicum*. The other three *Pisum* taxa were not clearly discriminated among them. The Neighbour-Joining clustering method at the species level is shown in Fig. 4. Bootstrap values support the conclusion that *P. sativum* and *P. elatius* are sister taxa. The rest of relationships were not clearly supported by bootstrap values.

Discussion

RBIPs have proved to be suitable markers for genetic diversity evaluation, evolutionary analysis and variety discrimination in *Pisum* (Smýkal *et al.*, 2008a,b;



Figure 4. Neighbour joining unrooted tree at species-subspecies level. Abbreviations as in previous figures. Only bootstrap support values over 50% are indicated.

Jing *et al.*, 2010). We have evaluated here the RBIP variability in a Spanish pea collection in relation to an accession set of the World *Pisum* core collection.

In relation to diversity at species level, the results on *P. fulvum* and *P. abyssinicum* contrast with the results described by Vershinin *et al.* (2003) using SSAP markers in which *P. fulvum*, *P. elatius* and *P. sativum* shared a similar high level of polymorphism, while *P. abyssinicum* showed a significant lower level of polymorphism. We suggest that these contrasting results are mainly due to the different accession sets of *fulvum* and *abyssinicum* used in both studies, and less to the markers since both RBIP and SSAP are based on retrotransposon insertion polymorphisms.

The fact that pea cultivars are grouped within a single cluster (Fig. 2, cluster 3) which represents a subset within the cultivated pea accessions set, and that their polymorphism level, estimated as He, is lower than pea landraces points to that only a part of the genetic variability available in traditional crop materials have been used to bred modern pea cultivars. Zong et al. (2009), in a comparative analysis of Chinese and global wide pea collections, stressed the importance of genetically distinct pea gene pools for future breeding programs. But even less «exotic» pea landrace collections can have additional genetic variability to be exploited in cultivar improvement. The advantage of local materials is that they are probably better adapted to the local environmental conditions than «exotic» materials. For instance, resistance to the race 6 of Pseudomonas syringae pv. pisi has been found in Spanish pea landraces (Elvira-Recuenco and Taylor, 2001; Martín-Sanz, 2008), while resistance to this race was previously only described in P. abyssinicum (Schmit et al., 1993).

Ellis *et al.* (1998) have also described that pea cultivars, and some «landraces», formed a separate cluster from other pea materials and wild species using an almost completely different set of JI accessions from the one included here. The cluster (Ellis *et al.*, 1998) included two classes of cultivars which were bred separately as the crop has different requirements: some were bred to be harvested as immature seeds for vegetable use, and other to be harvested as dry seeds. Smýkal *et al.* (2008b) used RBIP and SSR markers to analyze a collection of 164 Czech and Slovak pea accessions, and the cluster analysis of the molecular data not fully separated fodder pea types from other pea types, and they suggested that no global genomic differences exist between the two pea types. There was no apparent relationship among clusters and geographical origin of cultivated pea accessions, either between those from the Spanish collection or from the World collection. Previous data on a Spanish collection of 120 pea landraces indicated that the groups formed on the basis of ISSR markers were not related to agro-climatic regions within Spain (Lázaro and Aguinagalde, 2006).

Previous PCA analysis based on SSAP transposon polymorphisms (Vershinin *et al.*, 2003) have pointed out similar close relationships among *P. elatius*, *P. humile* and *P. sativum* to the observed in our work (Fig. 3). Likewise, clustering methods (Fig. 4) agreed with the generally accepted hypothesis that *P. fulvum* is the most distant species from cultivated *Pisum sativum* (Maxted and Ambrose, 2001; Vershinin *et al.*, 2003; Kosterin and Bogdanova, 2008). The representation of *P. humile* in our accession collection was too low to obtain valid conclusions.

Studies based in the analysis of different set of markers (retrotransposon based markers; AFLPs, mitochondrial-chloroplast-nuclear markers, etc.) agree that three main taxonomic groups of Pisum can be distinguished; these are P. abyssinicum, P. fulvum and P. sativum, other taxa once considered as species are presently considered as subspecies of *P. sativum*, although the subspecies concept in pea remains quite vague (Ellis et al., 1998; Maxted and Ambrose, 2001; Kosterin and Bogdanova, 2008). Our result with RBIP markers agree with this species differentiation. Both principal component analysis and clustering method distinguished two groups including each one the P. abyssinicum and P. fulvum accessions, and clearly segregated from the remaining Pisum accessions. On the basis of RBIP markers, the elatius and humile accessions were scattered among other Pisum cluster, some near P. abyssinicum and most interposed with pea landraces. The same set of P. fulvum accessions also formed a differentiated cluster apart from other Pisum materials in previous works based on the use of 39 gene segments or 54 PDR1 SSAP retrotransposon markers (Jing et al., 2005; 2007). Some P. elatius accessions formed clusters in the phylogenetic trees described in these two previous works, while others were clustered with P. sativum accessions as observed in our results with RBIP markers, but in our case all P. elatius accessions (almost an identical set to the used by Jing et al., 2007) were grouped within cluster with other Pisum taxa. Likewise the two P. humile accessions were clustered with other Pisum materials. In these previous works

by Jing *et al.* (2005, 2007), *P. abyssinicum* was represented only by accession JI 2385, also included in this work.

Like SSAP markers (Vershinin et al., 2003), all the RBIP markers analyzed here were polymorphic, with unique and species-specific markers making up only a small proportion of them. This situation is consistent with the possibility that introgression, segregation, and small rearrangements, rather than transposition itself, are the dominant modes of diversity generation in Pisum (Vershinin et al., 2003). Likewise, according to Vershinin et al. (2003) the absence of common markers shared exclusively by P. abyssinicum and P. sativum strongly supports the idea that both species were brought into cultivation independently and bayesian structure analysis of RBIP data provide a plausible model for how this occurred (Jing et al., 2010). Our result would support this hypothesis since, in spite that additional markers and accessions are included in the comparison, only a single common allele was shared exclusively by these two species.

The analysis of shared alleles also supported the close relationships between the *sativum* (92 accessions), *elatius* (12 accessions), and *humile* (two accessions) taxa. The numbers of alleles detected in these taxa were 50, 40 and 26, respectively. The shared alleles between *sativum* and *elatius* were 37 (over a possible maximum of 40) and between *sativum* and *humile* 23 (over a possible maximum of 26). Among these shared pairs seven were shared exclusively by *sativum* and *elatius* and one by *sativum* and *humile*, in spite of the small number of *humile* accessions considered.

Jing et al. (2007) pointed out that recombination has been very effective in shuffling genetic diversity between major Pisum lineages, and data based on transposable elements support the extensive introgression and intermixing among lineages. Even the homogeneous P. abyssinicum appears to have a hybrid origin (Vershinin et al., 2003). And probably this is also true in relation to modern cultivars originated from the recombination and introgression of genetically distant gene pools. As consequence the use of single genetic distance to represent the genetic structure and diversity in Pisum is inadequate (Jing et al., 2007), and this fact has implications for the management of plant genetic resources and the selection of germplasm for plant breeding. Sample pairs that are very closely related in single distance analysis may nevertheless carry distantly related gene alleles and vice versa (Jing et al., 2007). This is the situation also observed in our results. For instance, a relatively infrequent allele in locus 2055-x29 is shared by a Spanish cultivar (Ucero) and a Spanish landrace (JI1831), which differ in the alleles present in other seven loci but, on other hand, the allele is present in several cultivars or landraces from distant countries such as Canada or India. All these accessions sharing this infrequent 2055-x29 allele are scattered in cluster 3 and in other clusters of Fig. 2. A similar situation can be observed in relation to geographic information, which is also important in the design of germplasm collections. Cultivars Blizzar, Cea, Esla, Fortune, Raffale, Victor, and Iceberg and landrace ZP806 were close related by genetic distance (Fig. 2) in spite that they were bred in different countries. Thus, Jing et al. (2007) suggest that multilocus haplotype analysis of germplasm collections will be required to provide the solution to these problems. The importance of considering multilocus analysis in plant genetics and breeding was stressed by Allard (1999) and its importance in relation to germplasm collections was indicated (Pérez de la Vega et al., 1994).

This work has provided additional information on pea germplasm collections and on the use of RBIP markers in the evaluation of genetic diversity in these collections. Data are complementary to previous data on the same or similar materials and, in general, agree and support previous conclusions on *Pisum* taxa relationships and genetic variability distribution. Results also point to that the Spanish pea core collection of landraces maintains a relatively high variability which is only partially represented in modern bred pea cultivars adapted to Spanish conditions.

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References

- AGARWAL M., SHRIVASTAVA N., PADH H., 2008. Advances in molecular marker techniques and their applications in plant sciences. Plant Cell Rep 27, 617-631.
- ALLARD R.W., 1999. Principles of plant breeding, 2nd ed. John Wiley & Sons, New York.

- BARANGER A., AUBERT G., ARNAU G., LAIN A.L., DENIOT G., POTIER J., WEINACHTER C., LEJEUNE-HÉNAUT I., LALLEMAND J., BURSTIN J., 2004. Genetic diversity within *Pisum sativum* using protein and PCR-based markers. Theor Appl Genet 108, 1309-1321.
- BAUCOM R.S., ESTILL J.C., CHAPARRO C., UPSHAW N., JOGI A., DERAGON J.M., WESTERMAN R.P., SAN-MIGUEL P.J., BENNETZEN J.L., 2009. Exceptional diversity, non-random distribution, and rapid evolution of retroelements in the B73 maize genome. PLoS Genet 5, 732 (e1000732).
- CAMINERO C., CAMPO L., GONZÁLEZ R., RODRÍGUEZ M., GARCÍA A., RIBAS M.J., LAGUNA R., RAMOS A., 2001. Advances in the formation of the Spanish pea (*Pisum sativum* L.) core collection. Proc 4th European Conference on Grain Legumes, Cracow. pp. 10-11.
- CHAKRABORTY R., JIN L., 1993. Determination of relatedness between individuals using DNA fingerprinting. Hum Biol 65, 875-895.
- ELLIS T.H.N., POYSER S.J., KNOX M.R., VERSHININ A.V., AMBROSE M.J., 1998. Polymorphism of insertion sites of *Ty1*-copia class retrotransposons and its use for linkage and diversity analysis in pea. Mol Gen Genet 260, 9-19.
- ELVIRA-RECUENCO M., TAYLOR J.D., 2001. Resistance to bacterial blight (*Pseudomonas syringae* pv. *pisi*) in Spanish pea (*Pisum sativum*) landraces. Euphytica 118, 305-311.
- FLAVELL A.J., DUNBAR E., ANDERSON R., PEARCE S.R., HARTLEY R., KUMAR A., 1992. *Ty1*-copia group retrotransposons are ubiquitous and heterogeneous in higher plants. Nucleic Acids Res 20, 3639-3644.
- FLAVELL A.J., KNOX M.R., PEARCE S.R., ELLIS T.H.N., 1998. Retrotransposon-based insertion polymorphisms (RBIP) for high throughput marker analysis. Plant J 16, 643-650.
- HAWKINS J.S., PROULX S.R., RAPP R.A., WENDEL J.F., 2009. Rapid DNA loss as a counterbalance to genome expansion through retrotransposon proliferation in plants. Proc Natl Acad Sci USA 106, 17811-17816.
- HILL P., BURFORD D., MARTIN D.M.A., FLAVELL A.J., 2005. Retrotransposon populations of *Vicia* species with varying genome size. Mol Genet Genomics 273, 371-381.
- JING R., KNOX M.R., LEE J.M., VERSHININ A.V., AMBROSE M., ELLIS T.H.N., FLAVELL A.J., 2005. Insertional polymorphism and antiquity of PDR1 retrotransposon insertions in *Pisum* species. Genetics 171, 741-752.
- JING R., JOHNSON R., SERES A., KISS G., AMBROSE M.J., KNOX M.R., ELLIS T.H.N., FLAVELL A.J., 2007. Gene-based sequence diversity analysis of field pea (*Pisum*). Genetics 177, 2263-2275.
- JING R., VERSHININ A., GRZEBYTA J., SHAW P., SMYKAL P., MARSHALL D., AMBROSE M.J., ELLIS T.H.N., FLAVELL A.J., 2010. The genetic diversity and evolution of field pea (*Pisum*) studied by high throughput retrotransposon based insertion polymorphism (RBIP) marker analysis. BMC Evol Biol 10, 44.

- KOSTERIN O.E., BOGDANOVA V.S., 2008. Relationship of wild and cultivated forms of *Pisum* L. as inferred from an analysis of three markers, of the plastid, mitochondrial and nuclear genomes. Genet Resour Crop Evol 55, 735-755.
- KOSTERIN O.E., ZAYTSEVA O.O., BOGDANOVA V.S., AMBROSE M.J., 2010. New data on three molecular markers from different cellular genomes in Mediterraneam accessions reveal new insights into phylogeography of *Pisum sativum* L. subsp. *elatius* (Bieb.) Schmalh. Genet Resour Crop Evol 57, 733-739.
- KUBIS S.E., HESLOP-HARRISON J.S., DESEL C., SCHMIDT T., 1998. The genomic organization of non-LTR retrotransposons (LINEs) from three *Beta* species and five other angiosperms. Plant Mol Biol 36, 821-831.
- LÁZARO A., AGUINAGALDE I., 2006. Genetic variation among Spanish pea landraces revealed by Inter Simple Sequence Repeat (ISSR) markers: its application to establish a core collection. J Agric Sci 144, 53-61.
- MACAS J., KOBLIZKOVA A., NAVRATILOVA A., NEUMANN P., 2009. Hypervariable 3' UTR region of plant LTR-retrotransposons as a source of novel satellite repeats. Gene 448, 198-206.
- MARTÍN-SANZ A., 2008. Bacteriosis en guisante (*Pisum sativum* L.): situación en Castilla y León, caracterización de los patógenos implicados y búsqueda de fuentes de resistencia. Ph D dissertation. Universidad de León, Spain. [In Spanish].
- MARTÍN-SANZ A., GILSANZ-GONZÁLEZ S., SYED N.H., SUSO M.J., CAMINERO C., FLAVELL A.J., 2007. Genetic diversity analysis in *Vicia* species using retrotransposon-based SSAP markers. Mol Genet Genomics 278, 433-441
- MAXTED N., AMBROSE M., 2001. Peas (*Pisum* L.). In: Plant genetic resources of legumes in the Mediterranean (Maxted N., Bennett S.J., eds). Kluwer Academic Publishers, Dordrech. pp. 181-190.
- PEARCE S.R., HARRISON G., LI D., HESLOP-HARRISON J.S., KUMAR A., FLAVELL A.J., 1996. The *Ty1*-copia group retrotransposons in *Vicia* species: copy number, sequence heterogeneity and chromosomal localisation. Mol Gen Genet 250, 305-315.
- PÉREZ DE LA VEGA M., GARCÍA P., SÁENZ DE MIERA L.E., VENCES F.J., 1994. Genetic diversity in inbreeding species. Proc Eucarpia Genetic Resource Section Meeting (Balfourier F., Perretant M.R., eds). Clermont-Ferrand. pp 83-90.

- RAMOS A., 2003. Estudio de la variabilidad en la colección de variedades locales españolas de guisante (*Pisum sativum* L.). Ph D dissertation. Universidad Politécnica de Madrid. [In Spanish].
- SCHMIT J., TAYLOR J.D., ROBERTS S.J., 1993. Sources of resistance to pea bacterial bligth (*Pseudomonas syringae* pv. *pisi*) in pea germplasm. Proc 6th Internacional Congress of Plant Pathology, Montreal. p. 180.
- SMÝKAL P., 2006. Development of an efficient retrotransposon-based fingerprinting method for rapid pea variety identification. J Appl Genet 47, 221-230.
- SMÝKAL P., HORÁÈEK J., DOSTÁLOVÁ R., HÝBL M., 2008a. Variety discrimination in pea (*Pisum sativum* L.) by molecular, biochemical and morphological markers. J Appl Genet 49, 155-166.
- SMÝKAL P., HÝBL M., CORANDER J., JARKOVSKÝ J., FLAVELL A.J., GRIGA M., 2008b. Genetic diversity and population structure of pea (*Pisum sativum* L.) varieties derived from combined retrotransposon, microsatellite and morphological marker analysis. Theor Appl Genet 117, 413-424.
- SYED N.H., FLAVELL A.J., 2006. Sequence-specific amplification polymorphisms (SSAPs): a multi-locus approach for analyzing transposon insertions. Nat Protoc 1, 2746-2752.
- TAM S.M., LEFEBVRE V., PALLOIX A., SAGE-PALLOIX A.M., MHIRI C., GRANDBASTIEN M.A., 2009. LTRretrotransposons Tnt1 and T135 markers reveal genetic diversity and evolutionary relationships of domesticated peppers. Theor Appl Genet 119, 973.989.
- VERSHININ A.V., ALLNUTT T.R., KNOX M.R., AMBROSE M.J., ELLIS T.H.N., 2003. Transposable elements reveal the impact of introgression, rather than transposition, in *Pisum* diversity, evolution, and domestication. Mol Biol Evol 20, 2067-2075.
- WAUGH R., MCLEAN K., FLAVELL A.J., PEARCE S.R., KUMAR A., THOMAS B.T., POWELL W., 1997. Genetic distribution of BARE-1 retrotransposable elements in the barley genome revealed by sequence-specific amplification polymorphisms (S-SAP). Mol Gen Genet 253, 687-694.
- ZONG X., REDDEN R.J., LIU Q., WANG S., GUAN J., LIU J., XU Y., LIU X., GU·J., YAN L., ADES P., FORD R., 2009. Analysis of a diverse global *Pisum* sp. collection and comparison to a Chinese local *P. sativum* collection with microsatellite markers. Theor Appl Genet 118, 193-204.

Accesion number/ Cultivar name	Species	Status	Country	Place	County	Complementary information ^a
JI0130	P. abvssinicum	Landrace	Palestine			ZP1237
JI1640	P. abvssinicum	Landrace	Ethiopia			ZP1246
JI2202	P. abvssinicum	Landrace	Ethiopia			ZP1254
JI2202/1	P. abyssinicum	Landrace	Ethiopia			Derived from JI2202
JI2385	P. abyssinicum	Landrace	Yemen			ZP1525
JI0261	P. elatius	Wild	Turkey			
JI1075	P elatius	Wild	Turkey			
JI1094	P elatius	Wild	Greece			
JI1703	P elatius	Wild	Unknown			
JI2078	P elatius	Wild	Unknown			
JI2105	P elatius	Landrace	Iran			
JI3147	P elatius	Wild	Turkey			
II3149	P elatius	Wild	Turkey			
II3151	P elatius	Wild	Turkey			
113155	P elatius	Wild	Turkey			
II3156	P elatius	Wild	Turkey			
JI0224	P fulvum	Wild	Israel			
JI1006	P fulvum	Wild	Israel			
II1000	P fulvum	Wild	Iran			
II1796	P fulvum	wild	Israel			
112473	P fulvum	Wild	Israel			
II2517	P fulvum	Wild	Svria			
II2519	P fulvum	Wild	Svria			
112523	P fulvum	Wild	Svria			
JI2520	P fulvum	Wild	Svria			
JI2530	P fulvum	Wild	Svria			
JI0241	P humile	Wild	Israel			
II1704	P humila	Wild	Israel			
II0113	P satiyum	Cultivar	Russia			Spontaneus mutant
JI0115	P satiyum	Cultivar	Sweden			Spontaneus mutant
JI0267	P sativum	Landrace	Greece			
JI0281	P sativum	Landrace	Ethionia			
JI0201	P sativum	Cultivar	Canada			Alaska
110300	P sativum	Cultivar	Netherlands			Cennia
JI0435	P sativum	Cultivar				Wisconsin Perfection
JI0435	P sativum	Cultivar	UK			Maro
11086	P sativum	Landrace	A fahanistan			maro
110975	P sativum	Landrace	Costa Rica			
II1267	P sativum	Wild	India			
J11207 J11308	P satiyum	Wild	China			
II1544	P satiyum	Landrace	China			
II1775	P satiyum	Landrace	Chile			
J11775 II1831	P satiyum	Landrace	Spain			
J110J1 J11844	P satiyum	Landrace	Mexico			
112200	D satiyum	Landrace	Pussio			
J12200 J12262	P. satiyum	Wild	Tunicio			
J12203 J12376	1. suuvum P satiyum	Landrago	Tullisla			
112383	1. suuvum P satiyum	Landrace	Zambia			
J12303 II2421	1. Sullvum P sativum	Landrace	Latvia			
J12721 112546	1. sullvuill P sativum	Wild	Georgia			P transcaucasiour
J12340 112605	r. suuvum D. satiyere	Wild	Libyo			r. transcaucastcum
7D0064	r. suuvum P satiyum	Wild Landrage	Spain		Dalanaia	r. speciosum BGE022221 CC
ZP0004 ZP0074	P. sativum P. sativum	Landrace	Spain	Santibáñez de Vidriales	Zamora	BGE052251-CC BGE004041-CC

Appendix 1. *Pisum* materials used in the RBIP analysis

Accesion number/ Cultivar name	Species Status		Country Place		County	Complementary information ^a
ZP0076	P. sativum	Landrace	Spain	Paradela del Río	León	BGE004043-CC
ZP0104	P. sativum	Landrace	Spain	Vall d'Alba	Castellón	BGE001034
ZP0109	P. sativum	Landrace	Spain	Maguilla	Badajoz	BGE001100
ZP0115	P. sativum	Landrace	Spain	M. de la Salud	Baleares	BGE001414-CC
ZP0126	P. sativum	Landrace	Spain	Jerez de los Caballeros	Badajoz	BGE001646-CC
ZP0138	P. sativum	Landrace	Spain	Boracan de San Cristóbal	Oviedo	BGE002088-CC
ZP0150	P. sativum	Landrace	Spain	Garellas	Pontevedra	BGE002165-CC
ZP0152	P. sativum	Landrace	Spain	Santibáñez de Vidriales	Zamora	BGE002167-CC
ZP0156	P. sativum	Landrace	Spain	Castiñeiro	La Coruña	BGE003046-CC
ZP0168	P. sativum	Landrace	Spain	Pola de Somiedo	Asturias	BGE003303-CC
ZP0171	P. sativum	Landrace	Spain	La Riera	Asturias	BGE003306-CC
ZP0177	P. sativum	Landrace	Spain	Hospital	Asturias	BGE003312-CC
ZP0180	P. sativum	Landrace	Spain	Llanos de Somerón	Asturias	BGE003315-CC
ZP0181	P. sativum	Landrace	Spain	Jomezana Baja	Asturias	BGE003316-CC
ZP0202	P. sativum	Landrace	Spain	Corvelle	Lugo	BGE003436-CC
ZP0206	P. sativum	Landrace	Spain	Corvelle	Lugo	BGE003440-CC
ZP0213	P. sativum	Landrace	Spain	Tagarabuena	Zamora	BGE003690-CC
ZP0344	P. sativum	Landrace	Spain	Cervera de Pisuerga	Palencia	BGE032241-CC
ZP0516	P. sativum	Landrace	Spain	Fontecha de la Peña	Palencia	BGE005515-CC
ZP0535	P. sativum	Landrace	Spain	Cumbres San Bartolomé	Huelva	BGE001662-CC
ZP0593	P. sativum	Landrace	Spain	Béjar	Salamanca	BGE001519-CC
ZP0798	P. sativum	Landrace	Spain	Albunan	Granada	BGE032226-CC
ZP0799	P. sativum	Landrace	Spain	Colomera	Granada	BGE030152-CC
ZP0806	P. sativum	Landrace	Spain	Alhama de Granada	Granada	BGE030158-CC
ZP1261	P. sativum	Landrace	Spain	Puntallana	Tenerife	BGE019598-CC
ZP1262	P. sativum	Landrace	Spain	Lena	Asturias	BGE019600-CC
ZP1263	P. sativum	Landrace	Spain	Gijón	Asturias	BGE019778-CC
ZP1264	P. sativum	Landrace	Spain	Pina de Ebro	Zaragoza	BGE020326-CC
ZP1278	P. sativum	Landrace	Spain	Zas	La Coruña	BGE023269-CC
ZP1282	P. sativum	Landrace	Spain	Luanco	Asturias	BGE023273-CC
ZP1290	P. sativum	Landrace	Spain	Zas	La Coruña	BGE023282-CC
ZP1294	P. sativum	Landrace	Spain	Vélez Rubio	Almeria	BGE023644-CC
ZP1300	P. sativum	Landrace	Spain	Peñarrubia	Cantabria	BGE024375-CC
ZP1301	P. sativum	Landrace	Spain	La Fuente	Cantabria	BGE024376-CC
ZP1311	P. sativum	Landrace	Spain	Mirandilla	Badajoz	BGE025269-CC
ZP1315	P. sativum	Landrace	Spain	Villamanín	León	BGE025273-CC
ZP1318	P. sativum	Landrace	Spain	Santiesteban del Puerto	Jaen	BGE025728-CC
ZP1338	P. sativum	Landrace	Spain	Torrepacheco	Murcia	BGE027119-CC
ZP1347	P. sativum	Landrace	Spain	Ordes	La Coruña	BGE028986-CC
ZP1355	P. sativum	Landrace	Spain	Xinzo de Limia	Orense	BGE028998-CC
ZP1358	P. sativum	Landrace	Spain	Lousame	La Coruña	BGE029002-CC
ZP1366	P. sativum	Landrace	Turkey			Vavilov Institute 2274
Aravalle	P. sativum	Breeding line	Spain			ZP1460
Baccara	P. sativum	Cultivar	France			ZP1457
Badmington	P. sativum	Cultivar	France			ZP1454
Blizzard	P. sativum	Cultivar	France			ZP1672
Burbia	P. sativum	Breeding line	Spain			ZP1664
Cea	P. sativum	Cultivar	Spain			ZP0866
Chevenne	P. sativum	Cultivar	France			ZP1456
Chicarrón	P. sativum	Cultivar	Spain			ZP1666
Coomonte	P. sativum	Cultivar	Spain			ZP1406
Dove	P. sativum	Cultivar	France			ZP1667
Esla	P. sativum	Cultivar	Spain			ZP0864

Appendix 1 (cont.). Pisum materials used in the RBIP analysis

Accesion number/ Cultivar name	Species	Status	Country	Place	County	Complementary information ^a
Fortune	P. sativum	Cultivar	U.K.			ZP1524
Gracia	P. sativum	Cultivar	Spain			ZP0028
Hardy	P. sativum	Cultivar	France			ZP1669
Iceberg	P. sativum	Cultivar	Denmark			ZP1458
Ideal	P. sativum	Cultivar	France			ZP1463
Lobos	P. sativum	Breeding line	Spain			ZP1665
Lucy	P. sativum	Cultivar	France			ZP1670
Luna	P. sativum	Cultivar	Spain			ZP1231
Messire	P. sativum	Cultivar	France			ZP1468
Rafalle	P. sativum	Cultivar	France			ZP1013
Sidney	P. sativum	Cultivar	France			ZP1671
Speleo	P. sativum	Cultivar	France			ZP1673
Talanda	P. sativum	Breeding line	Spain			ZP1461
Ucero	P. sativum	Cultivar	Spain			ZP1414
Víctor	P. sativum	Cultivar	U.S.A.			ZP1465
Volcano	P. sativum	Cultivar	France			ZP1668

Appendix 1 (cont.). *Pisum* materials used in the RBIP analysis

^a Reference number in other germplasm bank collections. BGE: Spanish germplasm bank, these accessions are included in the Spanish pea core collection. ZP: ITACyL collection. Alaska, Cennia, Wisconsin Perfection, and Maro are cultivar names. CC: included in the Spanish core collection.

Appendix 2. RBIP markers used, primer sequences and linkage group in which they have been located

Marker name and specific primers	Annealing temperature	No occupied size (bp)	Occupied size (bp)	Linkage group
Birte-B1 5' CCCATTGATTCTCGTCTCAAGAC 5' TCACGAGGGTGTGATAGTAACTCA	55	244	281	
Birte-x16 5' CTTACCACCAAGCGCGCGAC 5' AGGCTTCTGATCCAACCAG	55	134	226	II
Birte-x34 5' GTTACTGCGGACGGTGGTC 5' GGCTGAAATCTCACTTTTGC	55	591	183	
281x1 5' TAATTATTATGGTATTCTGTG 5' CATATATTCACCCAAATCTTAAAG	55	267	240	IV
281x44 5' GATCAGAGAATCATGTCCAG 5' TCGAGGTGTGACAAAGTGC	55	339	237ª	II
281x5 5' GTAAATATGGACGTAAGATATC 5' CGATACCCTATTCCCAAAAG	55	361	215	
1794-2 5' GGGCCATGTACGACACATTC 5' GAGGAAATAAGAATGGTAGAGCATC	55	247	181	
399-80-46 5' GTTCTACTTCCTCTGAGTCA 5' CGATACGAAGGAGGAGTTAG	55	89	167	V

Marker name and specific primers	Annealing temperature	No occupied size (bp)	Occupied size (bp)	Linkage group
2055x29 5' CGATCATGATAAATATATTTAAT 5' CGAAGCATTAATGTATTAGAAC	55	311	183	
2055nr23 5' ATATGTGATTACGACAATAGG 5' CGACAGTGTAAATCTTTTTACA	60	256	181	
2055nr53 5' TGGATAGGGTATTGGAGTTC 5' ATAGCAGTAATTATGAACATG	55	379	434	
2385x64 5' GAAACATGATAGTAAGTTGCTC 5' CTTCCCTAAGCATTTTAATTGATC	55	556	234	
95x19 5' GGCGAGTATGTGCGCATG 5' CGACACCAGTCCCGTATTC	55	599	241	
95x2 5' CTGCAAAGGGTGCATATAG 5' GTTTTACAGGTGAAGAATCGTG	55	390	213	VI
64x45 5' CAAAGTATAACGTGTATCAAG 5' GTCATCGTCCAAACACACTC	55	641	601ª	
45x15 5' CGAAGTCAATATAATTGGCG 5' TCAACATGTCACTCCCATTAC	55	320	285	III
45x29 5' TGATGAACAGCATCCTGG 5' CGAAACTGGCTAGTTGCAAG	55	412	197	VII
45x31 5' GACTACTAGTTGGAACTCTTG 5' CATCTGGTTAGACAAGAAGAG	60	289	340ª	II

Appendix 2 (cont.). RBIP markers used, primer sequences and linkage group in which they have been located

^a These bands when the retrotransposon is inserted were obtained with the general primer 5' GGGCTTTGACTAATGGACCTC, the rest were obtained with the general primer 5' TAAGGTCCATTAGTCAAAGCCC.