

The first complete chloroplast genome of the Genistoid legume *Lupinus luteus*: evidence for a novel major lineage-specific rearrangement and new insights regarding plastome evolution in the legume family

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• **Background and Aims** To date chloroplast genomes are available only for members of the non-protein amino acid-accumulating clade (NPAAA) Papilionoid lineages in the legume family (i.e. Millettoids, Robinoids and the 'inverted repeat-lacking clade', IRLC). It is thus very important to sequence plastomes from other lineages in order to better understand the unusual evolution observed in this model flowering plant family. To this end, the plastome of a lupine species, *Lupinus luteus*, was sequenced to represent the Genistoid lineage, a noteworthy but poorly studied legume group.

• **Methods** The plastome of *L. luteus* was reconstructed using Roche-454 and Illumina next-generation sequencing. Its structure, repetitive sequences, gene content and sequence divergence were compared with those of other Fabaceae plastomes. PCR screening and sequencing were performed in other allied legumes in order to determine the origin of a large inversion identified in *L. luteus*.

• **Key Results** The first sequenced Genistoid plastome (*L. luteus*: 155 894 bp) resulted in the discovery of a 36-kb inversion, embedded within the already known 50-kb inversion in the large single-copy (LSC) region of the Papilionoideae. This inversion occurs at the base or soon after the Genistoid emergence, and most probably resulted from a flip–flop recombination between identical 29-bp inverted repeats within two *trnS* genes. Comparative analyses of the chloroplast gene content of *L. luteus* vs. Fabaceae and extra-Fabales plastomes revealed the loss of the plastid *rpl22* gene, and its functional relocation to the nucleus was verified using lupine transcriptomic data. An investigation into the evolutionary rate of coding and non-coding sequences among legume plastomes resulted in the identification of remarkably variable regions.

• **Conclusions** This study resulted in the discovery of a novel, major 36-kb inversion, specific to the Genistoids. Chloroplast mutational hotspots were also identified, which contain novel and potentially informative regions for molecular evolutionary studies at various taxonomic levels in the legumes. Taken together, the results provide new insights into the evolutionary landscape of the legume plastome.

Key words: *Lupinus luteus*, European yellow lupine, legume, Genistoid clade, chloroplast genome evolution, structural plastid rearrangement, 36-kb inversion, inverted repeats, flip–flop recombination, lineage-specific marker, functional gene transfer, Papilionoideae, repeated plastid sequences, sequence divergence, plastome hotspots, Fabaceae phylogeny.

INTRODUCTION

Legumes (Fabaceae) are the third largest angiosperm family, with 727 genera and about 20 000 species (Lewis *et al.*, 2005). They are characterized by a wide biological and ecological diversity (Cronk *et al.*, 2006), and they are of great economic importance, particularly for human consumption or as animal forage. This family is composed of two main groups (Fig. 1): Caesalpinioideae *sensu lato* (*s.l.*; including Mimosoideae) and

Papilionoideae (Wojciechowski *et al.*, 2004; Cardoso *et al.*, 2012). The Papilionoideae is divided into six major clades: the Genistoids, Dalbergioids, Mirbelioids, Millettoids, Robinoids and the inverted repeat lacking clade (IRLC) (Cronk *et al.*, 2006). Within the Genistoids, *Lupinus* displays particular functional properties compared with other legumes, such as active nitrogen metabolism and the production of allelopathic substances of ecological and agronomical interest (Guillon and Champ, 2002; Magni *et al.*, 2004; Pilvi *et al.*, 2006). Additionally,

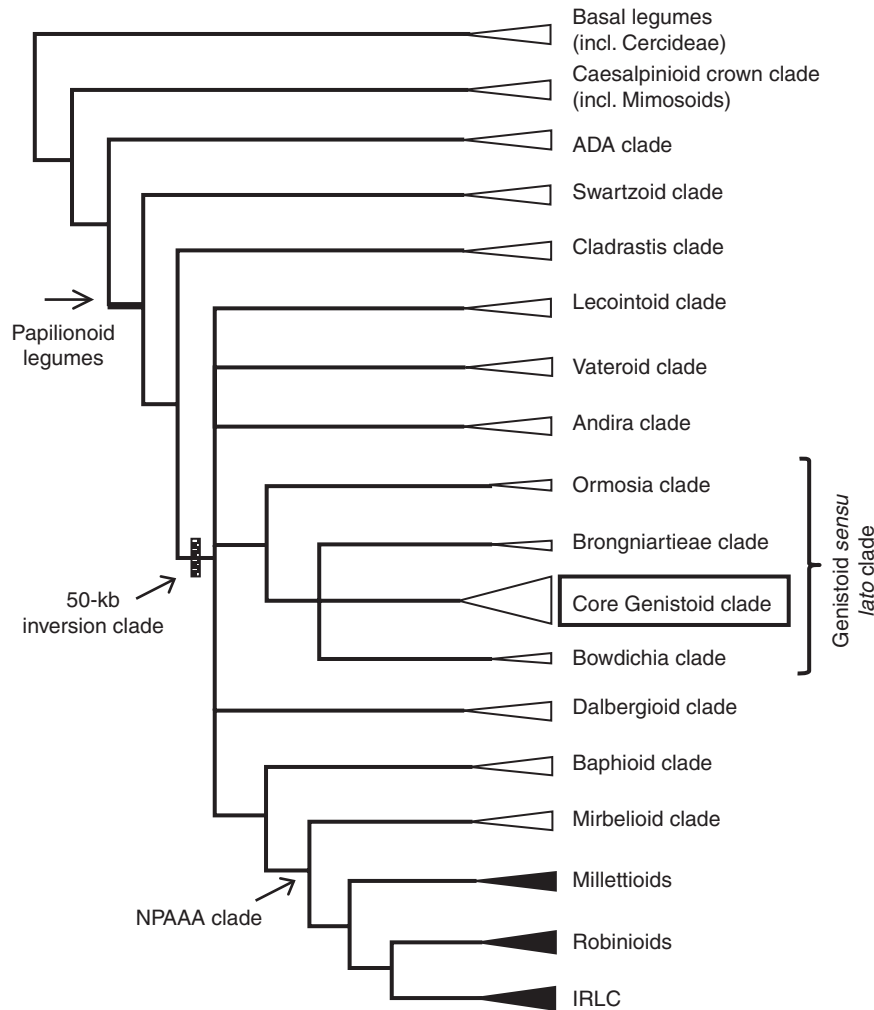


FIG. 1. Simplified phylogenetic tree representing the main clades circumscribed in the legume family [redrawn from Wojciechowski *et al.* (2004) and Cardoso *et al.* (2012)]. Filled triangles indicate lineages for which one or more whole chloroplast genome sequences are available, whereas open triangles indicate lineages for which no plastome sequence exists, including the core Genistoid clade (boxed) targeted in this study.

Lupinus is the only known legume that does not form mycorrhizal symbioses (Sprent, 2007). They also have considerable potential for phytoremediation due to their ability to metabolize nitrogen pollutants such as atrazine (Garcinuno *et al.*, 2003).

Plastome organization is highly conserved among most flowering plants (Jansen and Ruhlman, 2012), most having a quadripartite structure composed of two copies of an inverted repeat (IR) separated by large and small single-copy regions (LSC and SSC). However, a few angiosperm families, including the Fabaceae, present an unusual plastome structure and evolution. In this family, the loss of one IR in the Papilionoideae (Wojciechowski *et al.*, 2004), the presence of many repetitive sequences (Saski *et al.*, 2005; Magee *et al.*, 2010), the occurrence of relatively large inversions (Palmer and Herbon, 1988; Perry *et al.*, 2002; Magee *et al.*, 2010) and the presence of a localized hypermutable region (Magee *et al.*, 2010) have been detected. Aberrant DNA repair was inferred as a possible cause for these plastomic rearrangements and accelerated rates of nucleotide substitutions (Jansen *et al.*, 2007).

Although most photosynthetic angiosperm plastomes contain 79 protein-coding genes, various derived lineages exhibit slightly

fewer genes. Most of these rare chloroplastic gene losses occurred in species whose plastomes are highly rearranged relative to the ancestral angiosperm plastome (Jansen *et al.*, 2007). Since the emergence of the Fabaceae, there has been loss of five different chloroplastic genes: *accD*, *psaI*, *rpl23*, *rps16* and *ycf4* (Jansen *et al.*, 2007; Magee *et al.*, 2010). It is very likely that the genes lost from the plastome were previously functionally transferred to the nucleus or replaced by a nuclear gene of prokaryotic or eukaryotic origin. For example, the chloroplast *accD* gene was functionally transferred to the nucleus in *Trifolium* species (Magee *et al.*, 2010) and the plastidic *rps16* gene was functionally replaced by a nuclear-encoded *rps16* gene of mitochondrial origin in *Medicago truncatula* (Ueda *et al.*, 2008).

During the last decade, knowledge of the organization and evolution of legume plastomes has rapidly expanded with the development of next-generation sequencing (NGS) technologies. Ten legume plastomes have now been sequenced: *Cicer arietinum* (Jansen *et al.*, 2008), *Glycine max* (Saski *et al.*, 2005), *Lathyrus sativus* (Magee *et al.*, 2010), *Lotus japonicus* (Kato *et al.*, 2000), *M. truncatula* (NC_003119), *Millettia pinnata* (Kazakoff *et al.*, 2012), *Phaseolus vulgaris* (Guo *et al.*, 2007), *Pisum sativum*

(Magee *et al.*, 2010), *Trifolium subterraneum* (Cai *et al.*, 2008) and *Vigna radiata* (Tangphatsornruang *et al.*, 2010). The sequencing of these plastomes confirmed previous observations of major rearrangements in this family, including a 50-kb inversion present in most papilionoids (Palmer and Thompson 1982; Lavin *et al.*, 1990; Doyle *et al.*, 1996; Wojciechowski *et al.*, 2004; Jansen *et al.*, 2008) and the loss of one copy of the IR region in one of the papilionoid clade, called the IRLC (Palmer and Thompson, 1982; Lavin *et al.*, 1990; Wojciechowski *et al.*, 2004; Jansen *et al.*, 2008). However, all the Papilionoideae plastomes sequenced to date belong to three clades (Millettoids, Robinoids and IRLC) within the non-protein amino acid-accumulating clade (NPAAA clade; according to Cardoso *et al.*, 2012). Thus it is essential to investigate representatives from other Papilionoid lineages to better understand plastome evolution within the Papilionoideae, and more broadly within legumes. In this context, the genus *Lupinus* is a good candidate to represent the core Genistoids (Wojciechowski *et al.*, 2004; Cronk *et al.*, 2006; Cardoso *et al.*, 2012) that is one of the poorly studied legume lineages. Although considerable strides have been made in elucidating the evolutionary history of the Fabaceae using plastid DNA sequence-based phylogenies (Wojciechowski *et al.*, 2004; Cardoso *et al.*, 2012), there is still a great need to elucidate more accurately relationships at other taxonomic levels among and within lineages of the 50-kb-inversion Papilionoid clade, including within the Genistoids and in the genus *Lupinus* (Ainouche and Bayer, 1999; Ainouche *et al.*, 2004; Hughes and Eastwood, 2006; Drummond, 2008; Mahé *et al.*, 2011a, b). Therefore, the lupine plastome sequence not only provides the raw material to extend understanding of legume genome organization and evolution, but also provides an important source of phylogenetically informative plastid molecular markers, which have the advantage of being uniparentally (maternally) inherited and generally non-recombinant (Jansen *et al.*, 2007; Moore *et al.*, 2007, 2010).

Here we report the complete sequence of the chloroplast genome of *Lupinus luteus*, the first sequenced in the core Genistoids. After reconstruction and annotation, this genome has been compared with other Fabaceae and extra-Fabales plastomes, allowing the identification of a noteworthy 36-kb inversion. A PCR and sequencing survey of this inversion across various legume representatives provided evidence that this inversion represents a novel genomic rearrangement, characterizing the core Genistoids. The gene content within the *L. luteus* plastome has also been compared with that of other Fabaceae and closely related species in order to identify chloroplast genes lost from the *L. luteus* plastome. We verified that the chloroplast genes missing in the *Lupinus* plastome were functionally transferred to the nucleus. Finally, we evaluated the sequence divergence between the lupine and other Fabaceae plastomes at different levels (exon, intron and intergenic) in order to better understand the unusual plastome evolution and to suggest potentially useful plastid regions for molecular phylogenetic analyses in Fabaceae.

MATERIALS AND METHODS

DNA extraction, high-throughput sequencing and isolation of chloroplast sequences

Genomic DNA was extracted from fresh leaves of an individual sample (Lab. collection ref. number: M6=EGSM6Llu2) from a

natural population of *Lupinus luteus* collected at Bou Tlelis, Oran in Algeria, North Africa. DNA extraction was performed using a NucleoSpin® Plant II kit (Macherey Nagel) following the manufacturer's instructions. The genomic DNA was subjected to two high-throughput methods of sequencing: one run using pyrosequencing with the GS-FLX (454 Life Science, Roche) platform (OSUR/biogenouest; Université de Rennes-1) that generated 799 732 reads of approx. 400 bases, and one flow cell lane performed with an Illumina HiSeq 2000 platform (BGI, Hong Kong) that yielded 11.46 million 2×100 -base paired-end reads from a library of approx. 500 base DNA fragments. Reads corresponding to plastome sequences were extracted from the Roche-454 data set using a blast similarity search (e-value 10^{-6} , 90 % identity) against the fully sequenced plastomes of *G. max* (NC_007942), *M. truncatula* (NC_003119), *L. japonicus* (NC_002694), *C. arietinum* (NC_011163), *P. sativum* (NC_0147057), *T. subterraneum* (NC_011828), *L. sativus* (NC_014063), *M. pinnata* (NC_016708), *V. radiata* (NC_013843), *P. vulgaris* (NC_009259), *Populus trichocarpa* (NC_009143) and *Arabidopsis thaliana* (NC_000932). A total of 21 018 reads corresponding to plastid sequences were obtained from the 454 sequencing and 509 962 paired-end reads from Illumina.

Plastome assembly and annotation

De novo assembly was performed from filtered Roche-454 reads using Newbler (v. 2.5.3, 454 Life Science). A total of 45 contigs ranging from 450 to 25 000 bases were obtained and organized using the *G. max* plastome as a reference. Illumina paired-end reads having at least one mate mapping with Bowtie (Langmead *et al.*, 2009) on the 45 contigs were extracted from the Illumina data set. The draft plastome sequence as well as the junctions between contigs were verified and corrected with the 509 962 paired-end Illumina reads extracted using Mira v. 3.4.0 (Chevreux *et al.*, 1999) and Bowtie (Langmead *et al.*, 2009). The 454 and Illumina data sets allowed a $73 \times$ (s.d. 53) and $884 \times$ coverage (s.d. 466) of the newly reconstructed *L. luteus* plastome, respectively.

Plastome annotation was conducted in four steps. (1) Identification of protein-coding sequences by aligning (blastp, e-value threshold 10^{-5}) *G. max* protein-coding sequences obtained from the ChloroplastDB (Cui *et al.*, 2006) against chloroplastic open reading frames (ORFs) extracted from the *Lupinus* plastome sequence using the perl script *get_orf.pl* designed by Paul Stothard (University of Alberta). (2) Identification of rRNA and tRNA sequences by direct alignment of *G. max* tRNAs and rRNAs against the *Lupinus* plastome sequence. (3) Verification of the identification of all plastomic genes using DOGMA (Wyman *et al.*, 2004). (4) Verification of the annotation by performing manual alignment using BioLign and multiple contig editor (v. 4.0.6.2). A graphical representation of the chloroplast genome was performed using the CIRCOS software (Krzywinski *et al.*, 2009).

To determine the presence of codon bias, the number of codons ending with A–T or C–G was tallied and a χ^2 test was performed for each amino acid. These tests were subjected to a Bonferroni correction for multiple testing performed with the R software package (<http://www.r-project.org>).

Identification of repeat elements

The number and location of repeated elements (tandem, palindrome, dispersed direct and dispersed inverted repeats) in the *L. luteus* plastome were determined using REPuter (Kurtz et al., 2001). We used the same parameters as previously used for other Fabaceae species (Saski et al., 2005; Cai et al., 2008; Tangphatsornruang et al., 2010). More precisely, we searched for repeated elements with a minimum size of 30 bp and a Hamming distance of 3 (sequence identity of ≥ 90 %). One copy of the IR was removed before performing the analysis.

Identification and origin of the 36-kb inversion by PCR screening and sequencing

In order to identify the putative presence of large structural variation (>1 kb) within the *L. luteus* plastome, breaks of synteny were searched between plastomes of *L. luteus*, other legumes and two outgroup taxa (*Cucumis sativus* from the Cucurbitales and *Prunus persica* from the Rosales) by performing dot plots using the Gepard software (Krumstiek et al., 2007).

To determine the origin of the large inversion observed in *L. luteus*, its presence/absence was surveyed by PCR in *Lupinus* and in representatives of various genera more or less closely related to the lupines in the core Genistoids: *Argyrolobium uniflorum*, *Chamaecytisus mollis*, *Crotalaria saharae*, *Echinopartum boissieri*, *Genista florida*, *Genista tricuspidata*, *Laburnum anagyroides*, *Lupinus microcarpus*, *Retama sphaerocarpa*, *Sophora japonica*, *Thermopsis rhombifolia* and *Ulex minor*. Outgroup taxa were also screened for the presence/absence of this inversion, for instance: *Cercis siliquastrum* that is basal in the legume family; *Acacia dealbata* that belongs to the Mimosoids; and *Cladrastis lutea*, a Papilionoid that is sister to the 50-kb-inversion clade. A PCR strategy using primer pairs diagnostic for the presence or absence of the inversion was conducted. The primer pairs were designed in either conserved *ycf3* and *psbI*, or *rps4* and *ycf3* protein-coding sequences, which are flanking the inversion endpoints, to allow the assessment of the presence or absence of the inversion.

Each PCR amplification was performed in a total volume of 50 μ L containing 10 μ L of $5 \times$ Go taq green buffer (Promega), 5 μ L of 2 mM deoxyribonucleotide mix, 4 μ L of each primer (5 mM), 0.2 μ L of Go Taq polymerase (5 U μ L⁻¹) and 20 ng of template DNA. Cycling conditions were 94 °C for 2 min, followed by 35 cycles of 94 °C for 45 s, 55 °C for 30 s and 72 °C for 90 s, and a final extension of 72 °C for 7 min. The primer pairs used to detect the absence or presence of the 36-kb inversion were: *rps4-bef-F* (5'-CAATCAAATAATAGATAGTAAATGGGTTG-3') and *ycf3-bef-R* (5'-GGAATTATTCGTAATAATATATTGGCTAC-3'); and *ycf3-inv-F* (5'-CGTAATAAGATATTGGCTAC-3') and *psbI-int-R* (5'-CTCTTTTCATCTTCGGATTC-3'). The PCR products were then purified using the NucleoSpin Gel and PCR Clean-up purification kit (Macherey-Nagel) and sequenced directly in both directions (Macrogen Europe, Amsterdam).

Evolution of the gene content in the Fabaceae plastome and identification of genes functionally transferred to the nucleus in *Lupinus*

In order to determine whether *L. luteus* has recently lost chloroplastic genes, its plastome was compared with those of ten other

legume species (*M. pinnata*, *V. radiata*, *G. max*, *P. vulgaris*, *T. subterraneum*, *M. truncatula*, *L. japonicas*, *C. arietinum*, *P. sativum* and *L. sativus*) and two outgroup species available in GenBank. During Fabaceae evolution, five chloroplastic genes (*accD*, *psaI*, *rpl22*, *rpl23* and *rps16*) have been lost from the plastome of various lineages, of which three (*accD*, *rpl22* and *rps16*) were shown to have been independently functionally relocated to the nucleus or replaced by a nuclear gene in different Fabaceae (Gantt et al., 1991; Millen et al., 2001; Ueda et al., 2008; Magee et al., 2010). We searched for putative functional transfer to the nucleus (functional relocation or intermediate stage) of these five plastid genes within transcriptomic data available from our laboratory for *Lupinus mariae josephi* (unpubl. data). The identification of these putative functional transfers was performed by blasting (e-value threshold: 10^{-10}) the following sequences against the transcripts of *L. mariae josephi*: the *Trifolium repens* nuclear *accD* (Magee et al., 2010) and the *L. luteus* plastidic *accD* genes; the *P. sativum* nuclear *rpl22* sequence (Gantt et al., 1991); the plastidic *psaI*, *rpl23* and *ycf4* genes from various Fabaceae (*L. luteus*, *L. japonicus* and *P. vulgaris*); and the *M. truncatula* nuclear-encoded *rps16* genes of mitochondrial origin (Ueda et al., 2008). The presence of a transit peptide-encoding sequence within the identified chloroplastic genes functionally replaced in the nucleus was then predicted using BaCelLo (Pierleoni et al., 2006), Predotar (Small et al., 2004) and TargetP (Emanuelsson et al., 2000) software programs. To confirm that the nuclear *rpl22* gene identified in *L. mariae josephi* results from an early functional transfer to the nucleus in the common ancestor of all flowering plants, as demonstrated with *P. sativum* by Gantt et al. (1991), we aligned these sequences (*Lupinus* and *Pisum*) with the *rpl22* amino acid sequences from eubacteria, algae, bryophytes and land plants using the Geneious package (<http://www.geneious.com/>). After excluding the extreme 5' and 3' ends of the sequences, a data matrix of 98 amino acids was subjected to phylogenetic analyses using PHYML (Guindon and Gascuel, 2003) and Neighbor-Joining (Saitou and Nei, 1987). The tree was rooted using the eubacteria *Mycoplasma*. Bootstrap values were performed with 1000 replicates (Felsenstein, 1985).

Evaluating sequence divergence between the complete lupine plastid genome and those from other legumes and Fabids

Sequence divergence between *L. luteus* and ten other Fabaceae plastomes was evaluated independently for each homologous regions aligned with MUSCLE (Edgar, 2004). Pairwise distances were calculated with the *ape* R-cran Package (Paradis et al., 2011, available at <http://cran.r-project.org/web/packages/ape/ape.pdf>) using the Kimura-2-parameters (K2p) evolution model (Kimura, 1980). The mean sequence divergence rate of the different genetic categories [i.e. intergenic spacers (IGSs), introns, rRNA and tRNA, and exons] was compared using Mann-Whitney test with Bonferroni correction. Additionally, sequence divergence of coding-protein genes (exons) was estimated using the synonymous (Ks) and non-synonymous (Ka) nucleotide substitution rates with the yn00 method (Yang and Nielsen, 2000) from the PAML package (Yang, 2007). Finally, fast-evolving sequences were identified. Only the protein-coding, intronic or intergenic regions presenting a higher evolutionary rate than the regions most commonly used for

evolutionary studies in Fabaceae (*rbcL* and *matK* genes, the 5'*trnK* and *trnL* introns, and the *trnK_trnF*, *trnL_trnT* and *trnS_trnG* IGSs) and a minimum size of about 300 bp were retained.

A list of all the software programs used in this study, their purpose and availability can be found in the Supplementary Data Table S1.

RESULTS

Organization, gene content and characteristics of the *L. luteus* plastome

The *Lupinus luteus* plastome (deposited in GenBank: KC695666) has a length of 151 894 bp, with a quadripartite structure composed of two IRs (25 860 bp) separated by an SSC (17 847 bp) and an LSC (82 327 bp) region (Fig. 2). It contains 111 different genes, including 77 protein-coding genes, 30 tRNA genes and four rRNA genes (Table 1). Protein-coding genes, tRNA and rRNA represent, respectively, 51.6, 1.8 and 6.0% of the plastome. Non-coding DNA, including IGSs and introns, represents 40.6% of the genome. The overall GC content of the *L. luteus* plastome is 36.6%. It is higher in tRNA and rRNA (53.3 and 55.3%, respectively), slightly higher in protein-coding genes (37.3%), similar in introns (36.3%) and lower in IGSs (30.3%).

The *L. luteus* plastome contains 18 different intron-containing genes (of which six are tRNA), as in most Fabaceae species. All intronic genes contain one intron, apart from two genes (*clpP* and *ycf3*) that contain two introns. Within the IR, four rRNA, seven tRNA and five protein-coding genes are repeated. Only the 5' end of the *ycf1* gene (519 bp) is present in the IR, and the gene *rps12* is trans-spliced, with the 5' exon in the LSC and the remaining two exons in the IR.

TABLE 1. *Lupinus luteus* plastome characteristics

Plastome characteristics	
Size (bp)	151 894
LSC size in bp (%)	82 327 (54.2)
SSC size in bp (%)	17 847 (11.7)
IR length in bp (%)	25 860 (34.1)
Size in bp (%) of coding regions	90 217 (59.4)
Size in bp (%) of protein-coding regions	78 363 (51.6)
Size in bp (%) of introns	19 136 (12.6)
Size in bp (%) of rRNA	9056 (6)
Size in bp (%) of tRNA	2798 (1.8)
Size in bp (%) of IGSs	42 541 (28)
No. of different genes	111
No. of different protein-coding genes	77
No. of different tRNA genes	30
No. of different rRNA genes	4
No. of different genes duplicated by IR	17
No. of different genes with introns	18
Overall % GC content*	36.6
% GC content in protein-coding regions*	37.3
% GC content in introns*	36.3
% GC content in IGSs*	30.3
% GC content in rRNA*	55.3
% GC content in tRNA*	53.3

*The sequence of the two inverted repeats were taken into account for this analysis.

Thirty different tRNA are present in the *L. luteus* plastome. They correspond to 28 different codons, at least one for each amino acid. Seven of the 28 different anticodon tRNAs encoded in the *Lupinus* plastome correspond to the most common codon (where synonymous codons exist). The codon usage is biased towards a high representation of A and T at the third position (Supplementary Data Table S2).

Repeat elements in the *L. luteus* plastome

All repeat sequences that present a minimum size of 30 bp and with a sequence identity $\geq 90\%$ were identified in the *L. luteus* plastome using REPuter (Kurtz et al., 2001). A total of 31 repeats were found (Supplementary Data Table S3), including 13 dispersed short inverted repeats (sIRs), nine palindromes, six dispersed direct repeats and three tandem repeats.

All the palindromic repeats observed in the *L. luteus* plastome are localized in IGSs (except one in the *ycf1* coding sequence) and tandem repeats are mainly found in coding sequences (*ycf2*).

Most repeats (93.55%) are 30–50 bp long. The largest repeat in the plastome is a 288 bp dispersed direct repeat corresponding to a fragment of *ycf2* duplicated in each IR, between *rpl23* and *trnI_CAU*. This repeated element is absent from extra-Fabales plastomes (*C. sativus* and *P. persica*) or from IRLC plastomes in the Papilionoideae, but is present in other non-IRLC Papilionoideae plastomes (*G. max*, *L. japonicus*, *P. vulgaris* and *V. radiata*), as previously observed by Guo et al. (2007).

Origin of a 36-kb inversion detected in *L. luteus*

Global alignment and comparison of gene order between the plastomes of *L. luteus* and other legumes, as well as with outgroups, revealed an inversion of about 36 kb between the *trnS-GCU* and *trnS-GGA* genes in *L. luteus* (Fig. 3). This inversion is highlighted by dot plot analyses that compared the plastome of *L. luteus* with that of either *G. max* or *C. sativus* (Supplementary Data Fig. S1). This unique 36-kb inversion is embedded within the 50-kb inversion that occurred earlier in the Papilionoideae after the divergence between the *Cladrastis* clade and the rest of the more derived papilionoid legumes (Doyle et al., 1996).

To verify the existence of this inversion in *Lupinus* and to screen other Genistoids and legume species for the presence or absence of this 36-kb inversion, two diagnostic primer pairs were designed. The localization of these primers is indicated in Fig. 3. PCR amplification was expected from the primers located in the *rps4* and *ycf3* protein-coding sequences only for the species without the inversion, whereas PCR amplification using primers within the *ycf3* and *psbI* genes was only expected in species with the inversion. Since the 36-kb inversion identified in *L. luteus* is not present in the plastomes available for extra-Fabaceae taxa or in the derived Fabaceae (representative of the Millettoid, Robinoid and IRLC clades), it most probably occurred after the emergence of the Genistoids. For the 11 core Genistoid species screened here (including representatives of the Sophoreae, Thermopsidae and Genisteae tribes), amplifications were only successful when using the diagnostic primers pair for the presence of the inversion. In contrast, all non-core Genistoids tested gave amplification only when using the diagnostic primers pair for the absence of the 36-kb inversion (Fig. 4). In both cases, the results were confirmed by sequencing of the PCR products (deposited in GenBank:

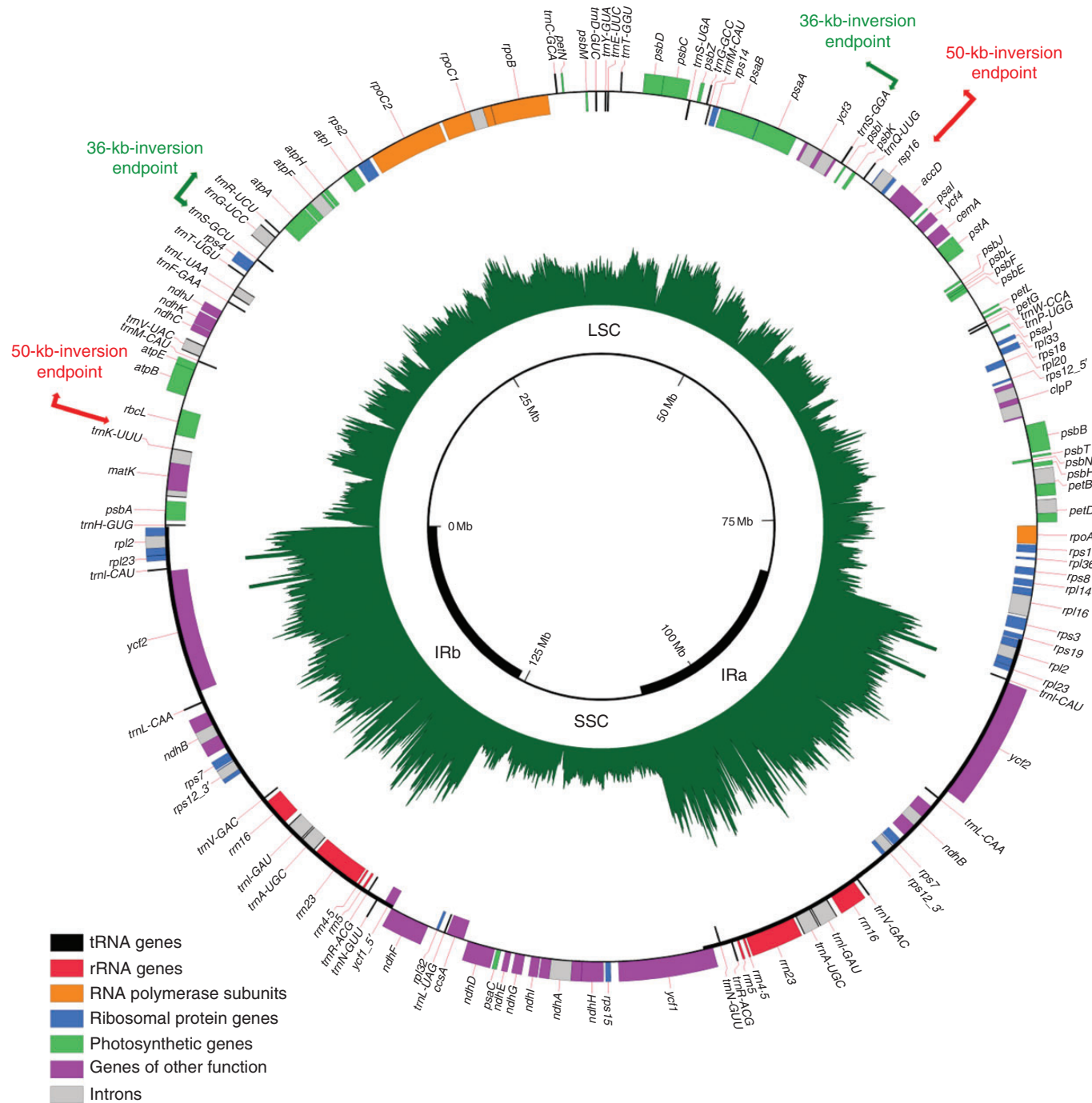


FIG. 2. Circular gene map of the *Lupinus luteus* (Genistoid; Fabaceae) plastid genome. Genes are represented with boxes inside and outside the first circle to indicate a clockwise or counterclockwise transcription direction, respectively. Genes belonging to different functional groups are colour coded. Read depth coverage of the plastome is represented by proportional radial lines in the inner green circle. The locations of the different main plastomic regions (inverted repeats, large single copy and small single copy) are indicated in the inner circle. The endpoints of the 50-kb and 36-kb inversions are represented by red and green arrows, respectively.

KC695667–KC695681) and alignment of the sequences with the homologous plastidic region from other Fabaceae plastomes, including *L. luteus*.

Several multiple alignments of the sequences surrounding the endpoints of the 36-kb inversion using Papilionoideae species with or without the 36-kb inversion allowed determination of

the exact location of the inversion (Supplementary Data Fig. S2). It occurred between the 3' end of the *trnS-GGA* and the *trnS-GCU* genes that are identical for the last 29 bp and are in inverse orientation (Fig. 5; Supplementary Data Table S3). A similar sIR also exists between *trnS-GGA* and *trnS-UGA* that are 9 kb distant.

Gene content and gene transfers to the nucleus in *Lupinus* compared with other Fabaceae

The protein-coding gene content of the *L. luteus* plastome was compared with those of ten other Fabaceae and two outgroup taxa. The aim of this comparison was to examine whether the lupines (representing the core Genistoids) have lost or retained the chloroplast genes known to have lost their functionality in the plastome of various lineages during legume evolution (reviewed in Magee et al., 2010) such as: *accD*, *psaI*, *rpl22*, *rpl23*, *rps16* and *ycf4* (as indicated in Fig. 6). Out of these six plastidic genes lost from legume lineages, only *rpl22* is missing in the plastome of *L. luteus*. The functional transfer of this gene to the nucleus, already demonstrated in *P. sativum* (Gantt et al., 1991), was verified in a lupine species (*L. mariae josephi*) by the identification of a nuclear *rpl22* transcript that is similar to the nuclear *rpl22* transcript found in *P. sativum*. The presence of a chloroplast target peptide was predicted in the *L. mariae josephi* nuclear *rpl22* transcript using a variety of software (data not shown). The alignment and phylogenetic analysis of nuclear and chloroplastic *rpl22* sequences (Supplementary Data Fig. S3) showed that the nuclear *rpl22* gene observed in *Lupinus* and *Pisum* derives from the same transfer event, which occurred in the common ancestor of all flowering plants (Gantt et al., 1991). Concerning the other chloroplast genes lost during Fabaceae evolution, investigations were performed to determine whether they could be at an intermediate stage of functional transfer to the nucleus. We identified nuclear *rps16* transcripts in *L. mariae josephi* that were similar to the *M. truncatula* nuclear *rps16* genes (Ueda et al., 2008), but no nuclear *accD*, *psaI*, *rpl23* or *ycf4* transcripts could be detected.

Sequence divergence between the plastome of *L. luteus* and other Fabaceae

A comparison of pairwise distances (K2p) calculated for non-coding regions between *L. luteus* and other legumes (Supplementary Data Table S4) revealed that, as expected, IGSs evolve significantly more rapidly than introns. The slowest evolving regions are tRNAs and rRNAs (Supplementary Data Table S4). For introns (Fig. 7A), the mean sequence divergence ranged from 0.028

(for the *rps12* intron) to 0.270 (for *clpP* intron1). The two main introns previously used for phylogenetic inference in legumes showed relatively low rates of variation: 0.100 for the *trnL* intron (501 bp length) and 0.148 for the *trnK* 5' intron (318 bp length). Among introns, seven exhibited a higher level of divergence (Fig. 7A): the *trnG-UCC* intron (K2p = 0.183; 698 bp), *rpoC1* intron (0.164; 766 bp), *clpP* intron 2 (0.230; 739 bp), *clpP* intron 1 (0.270; 655 bp), *petD* intron (0.224; 743 bp), *rpl16* intron (0.195; 1155 bp) and *ndhA* intron (0.215; 1171 bp). The highest mean sequence divergence of IGS regions corresponds to the *accD-psaI* region (0.473). Among IGSs and in comparison with the IGS regions most used for legume phylogeny (*trnL-trnF*, mean K2p = 0.255; *trnL-trnT*, 0.352; *trnS-trnG*, 0.316), five IGSs >300 bp showed divergence rates slightly or significantly higher than *trnL-trnT*, i.e. *ycf4_cemA* (0.357; 317 bp), *rpl36-rps8* (0.357; 453), *psbZ-trnG-GCC* (0.357; 345 bp), *trnV-UAC-ndhC* (0.355; 497 bp) and *accD-psaI* (0.473; 293 bp (Fig. 7A). For protein-coding regions (Fig. 7B; Supplementary Data Table S5), the evolutionary rates have been evaluated by comparison of their synonymous (Ks) nucleotide substitution rates (Fig. 7B). The mean divergence rate between *L. luteus* and the other legume genes ranged from 0.072 (for *rpl23*; 282 bp) to 0.667 (for *rps16*; 47 bp), with most loci presenting mean Ks values lower than those of the two protein-coding genes used for phylogenetic inference in legumes, *matK* (0.235; 1521 bp) and *rbcL* (0.367; 1428 bp). Fourteen genes displayed Ks values higher than *rbcL*, of which nine are >300 bp: *rpoC2* (0.419; 4149 bp), *rps16* (0.667; 407 bp), *accD* (0.538; 1497 bp), *ycf4* (0.659; 555 bp), *rps8* (0.388; 405 bp), *rpl14* (0.441; 369 bp), *ycf1* (0.518; 5296 bp), *ndhH* (0.394; 1182) and *ndhF* (0.454; 2241 bp). For most loci (65/77), the non-synonymous nucleotide substitution (Ka) values calculated between *Lupinus* and the other legumes were <0.1 (Supplementary Data Table S6). Among the 12 remaining loci, only five displayed higher values than the reference *matK* gene (mean Ka = 0.132): *rpl32* (0.147), *rps16* (0.169), *accD* (0.184) and particularly *ycf1* (0.306) and *ycf4* (0.398). Regarding these low values of Ka, the Ka/Ks ratio calculated for each protein-coding gene (Supplementary Data Table S7) was <1 and even <0.5 for almost all loci, indicating that plastidic genes are under a high negative (i.e. purifying) selective constraint (Kimura, 1977; Messier and Stewart, 1997). Nevertheless, it can be noted that

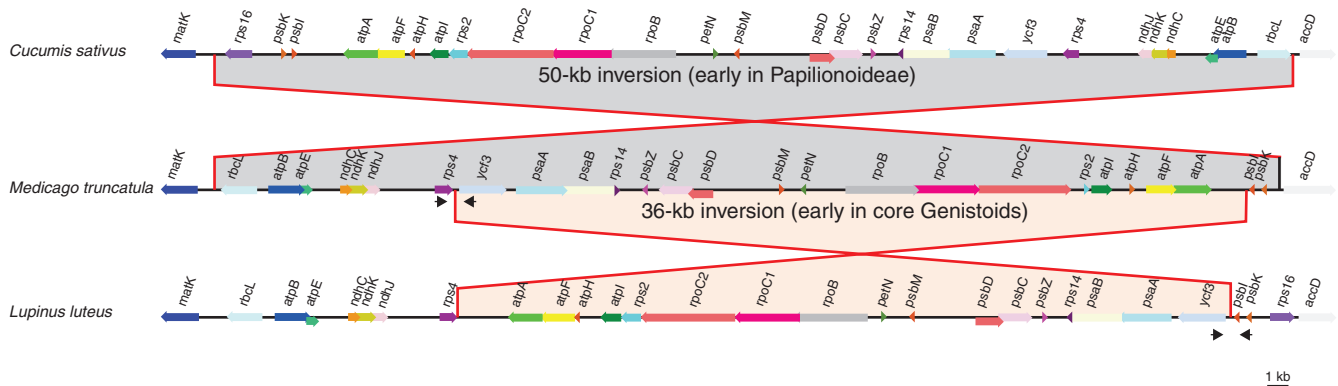


FIG. 3. Comparative plastomic maps showing the endpoints of the large 50-kb inversion present in most Papilionoideae (Fabaceae) and of a new 36-kb inversion detected in most core Genistoids surveyed in this study. The plastomes of *Cucumis sativus*, *Medicago truncatula* and *Lupinus luteus* were used to represent the structural patterns observed in most flowering plants, in most Papilionoid legumes and in the core Genistoids, respectively. The partial plastomic maps are drawn to scale, and only protein-coding genes are mapped. Approximate positions of diagnostic primer pairs used to detect the presence or absence of the 36-kb inversion are designated by black arrows (not to scale).

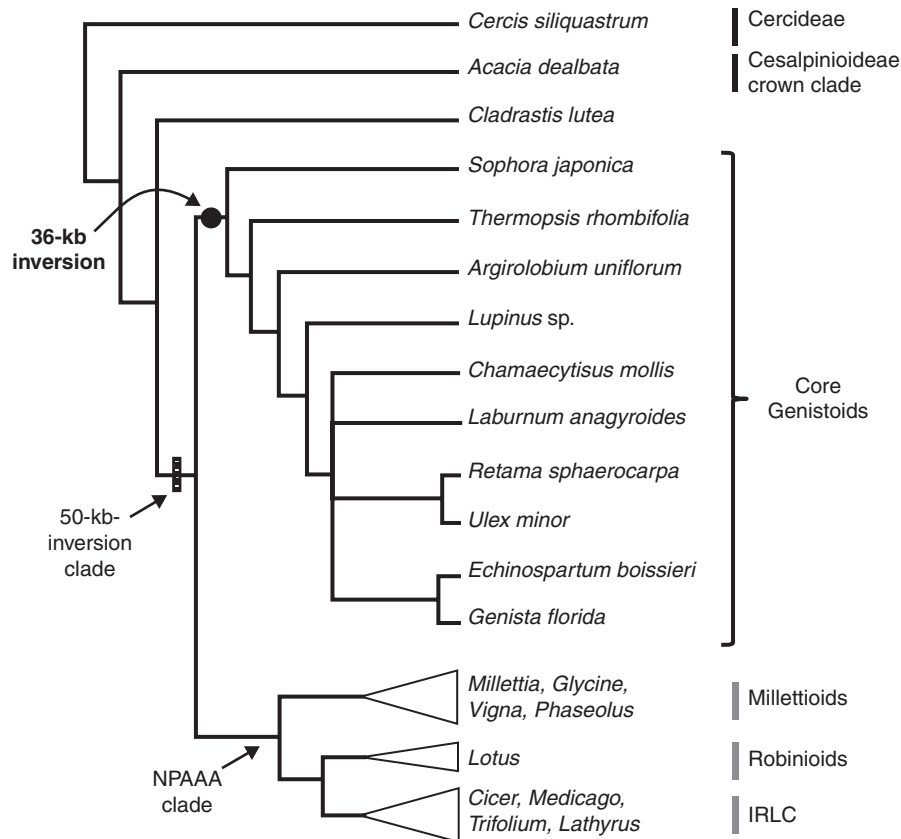


FIG. 4. Phylogenetic position of the 36-kb inversion rearrangement (solid black circle) detected in the plastomes of *Lupinus* and representatives of the core Genistoid clade (Papilionoideae; Fabaceae). All taxa screened by PCR and sequencing for the presence or absence of this inversion have their names labelled in bold. The taxa for which the plastome sequence is publicly available and for which the 36-kb inversion is absent belong to the Millettoids, Robinioids and IRLC (labelled in grey). The phylogenetic tree is redrawn from Cardoso *et al.* (2012).

the three *ycf* genes (*ycf1*, *ycf2* and *ycf4*) exhibit remarkably higher Ka/Ks values (0.601, 0.682 and 0.649, respectively) than all the other genes, indicating an increase in their sequence evolutionary rate.

Altogether, these analyses allow circumscription of fast-evolving regions in the legume plastomes, as inferred from pairwise comparison of *Lupinus* with the other available legume plastomes (highlighted in Supplementary Data Fig. S4). Among these regions, three are remarkable: one in the SSC, between *ycf1* and the *ndhA* intron (9043 bp); two in the LSC, around the *rpl36-rpl16* genes (3178 bp), and the *accD-ycf4_cemA* region (2968 bp) that exhibits the highest rates of sequence divergence for genes (*rps16*, *accD* and *ycf4*) and IGSs (*accD_psaI* and *ycf4_cemA*). The latter region, which includes the *ycf4* gene, was shown to have a dramatic increase in its evolutionary rate in the NPAAA clade (including Millettoids, Robinioids, and IRLC) and most particularly in *Lathyrus* (Magee *et al.*, 2010). To investigate whether such acceleration also occurred in the lupine lineage, maximum likelihood phylogenetic analyses using legume *ycf4* gene sequences (including *L. luteus ycf4*) and based on Ks and Ka substitution rates were performed. Our results (Supplementary Data Fig. S5) do not provide evidence of such acceleration in the *ycf4* gene in *Lupinus*, in accordance with the previous results obtained by Magee *et al.* (2010) using a few Genistoid members (*Crotalaria*, *Goodia* and *Laburnum*).

The other regions showing peaks of divergence when comparing *Lupinus* with other legume plastomes include some isolated genes

(*rpoC2* and *ndhF*), introns (*trnG-UCC*, *rpoC1* and *petD*) and IGSs (*trnV_ndhC* and *psbZ_trnG-GCC*) that were not previously detected as fast-evolving regions in NPAAA clade members.

DISCUSSION

In this work the plastome of *Lupinus luteus* has been sequenced using NGS technologies. Its size and gene content are within the range found in plastomes containing two IRs (Raubeson and Jansen, 2005). It is AT rich (with the exception of rRNA and tRNA genes) and the codon usage is biased toward a high representation of A and T at the third position as previously observed by Clegg *et al.* (1994). This sequence, which represents the first plastome sequenced in the core Genistoids, is of major interest because all legume plastomes sequenced so far belong to only three Papilionoid clades, the Millettoids, the Robinioids and the IRLC, which derive from within the NPAAA clade (Cardoso *et al.*, 2012). Thus it was essential to sequence plastomes from representatives of other Papilionoid lineages in order to have a better understanding of the unusual plastome evolution observed in legumes (Jansen and Ruhlman, 2012). Most photosynthetic angiosperms have a highly conserved plastome organization, except the Campanulaceae, Fabaceae and Geraniaceae families that exhibit remarkable and extensive rearrangements (Jansen and Ruhlman, 2012). Within the Fabaceae, one of the most remarkable inversions that occurred after the emergence of the

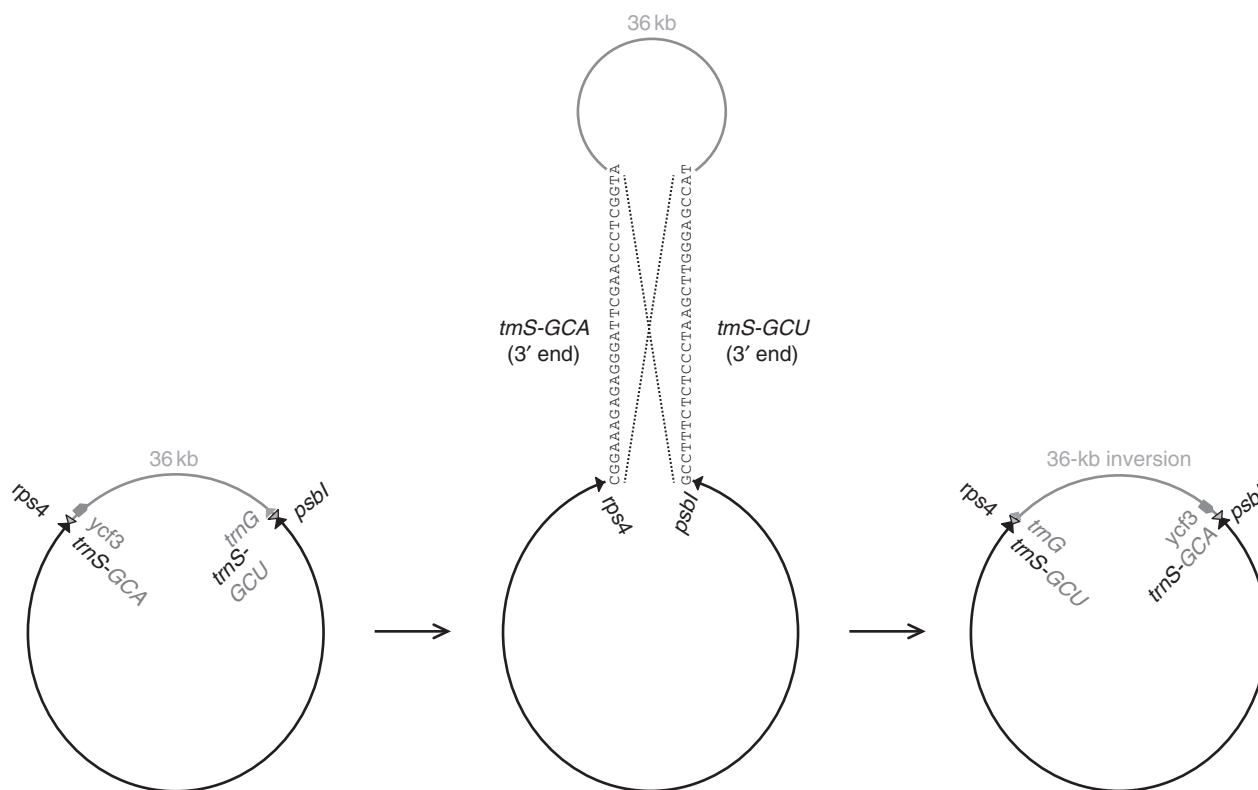


FIG. 5. Illustration of the suggested flip–flop recombination event that occurred after the divergence of the Genistoids from the other Fabaceae clades, resulting in a 36-kb inversion (grey region). This recombination event was most probably caused by the presence of inverted repeat sequences (29 bp shown) at the 3' end of *trnS-GGA* and *trnS-GCU* that are 36 kb apart.

family is the 50-kb inversion, which is shared by most papilionoid taxa (Doyle *et al.*, 1996). The plastome sequences of three IRLC species, *L. sativus* (Magee *et al.*, 2010), *P. sativum* (Palmer and Herbon, 1988) and *T. subterraneum* (Cai *et al.*, 2008), revealed that, relative to the ancestral angiosperm plastome organization, and after the 50-kb inversion event, six, eight and fifteen inversions occurred, respectively (Magee *et al.*, 2010). Within the 50-kb clade, our study reveals that the *L. luteus* plastome experienced an additional 36-kb inversion internal to the 50-kb inversion, which most probably occurred at the origin of the core Genistoids. Previous molecular characterization of large plastomic inversion endpoints in a few plant families or genera, including the 50-kb inversion present in most Papilionoideae (Doyle *et al.*, 1996), the 22-kb inversion in Asteraceae (Kim *et al.*, 2005), the 42-kb inversion in *Abies* (Tsumura *et al.*, 2000) or the 21-kb inversion in *Jasminae* (Lee *et al.*, 2007), showed that these large plastomic inversions were often associated with sIRs present within, or adjacent to, a tRNA. The detailed survey of the regions surrounding the 36-kb inversion endpoints in core Genistoids allowed us to determine that this inversion is most likely to be due to the presence of sIR motifs (29 identical nucleotides) at the 3' end of *trnS-GGA* and *trnS-GCU*. The role of repeated elements present in inverse orientation in promoting flip–flop recombination resulting in inversions has been previously demonstrated using tobacco transplastomic lines (Rogalski *et al.*, 2006). Such repeated elements can promote plastid DNA inversions which may vary in size from a few base pairs to several kilobases (reviewed in Downie and Palmer, 1992).

Minor inversions are more common than major ones and mainly occur in non-coding regions, IGSs and introns (Palmer, 1985). Interestingly the sIR motif in *trnS-GGA* and *trnS-GCU*, which caused the 36-kb inversion in the core Genistoids, is present in almost all Rosids (Supplementary Data Table S8) and is separated by at least 30 kb. Thus, even though this 36-kb inversion was only observed in the core Genistoids, it could have occurred in any other rosoid species.

The *L. luteus* plastome contains fewer repeats (31) than other Fabaceae species, such as *V. radiata*, *L. japonicus*, *G. max* or *M. truncatula* that have 50, 67, 104 and 191 repeats, respectively (Saski *et al.*, 2005; Tangphatsornruang *et al.*, 2010). Most *L. luteus* repeats are relatively small in size (90% are 30–50 bp in size). The longest repeat observed in *L. luteus* is a 288 bp direct repeat (within the IR) that is also present in the non-IRLC Papilionidae (*G. max*, *L. japonicus*, *P. vulgaris* and *V. radiata*) but not in the IRLC or outgroup taxa (*C. sativus* and *P. persica*). The low number of repeats observed in *Lupinus* is in stark contrast to the *T. subterraneum* plastome that contains a high number of large repeats and shows a high rate of rearrangement: 14 inversions occurred since its divergence with other IRLC species (Cai *et al.*, 2008; Magee *et al.*, 2010). The number of large repeats was demonstrated to be positively correlated to the degree of plastome rearrangements in plants (Maul *et al.*, 2002; Pombert *et al.*, 2005; Guisinger *et al.*, 2011). Within the repeats observed in *Lupinus*, 42% (13/31) are dispersed sIRs (6–66 kb distant) that could promote inversions. However, apart from the dispersed sIR at the origin of the 36-kb inversion in the core Genistoid and

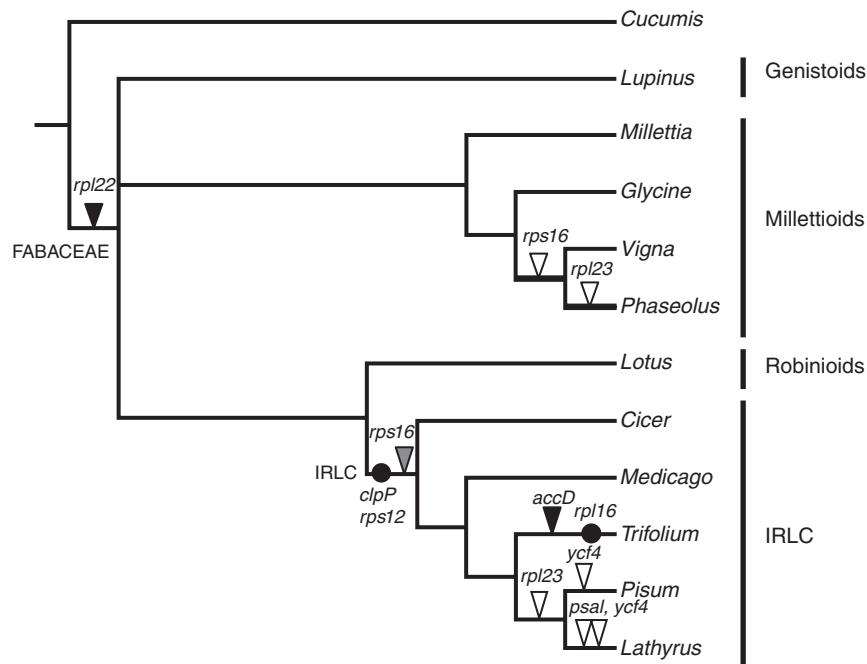


FIG. 6. Chloroplast genes and introns lost during Fabaceae evolution. All Fabaceae species whose plastome has been fully sequenced are presented in the phylogenetic tree (redrawn from Cardoso *et al.*, 2012). Black circles on branches indicate intron losses, whereas triangles show the genes recently lost from the plastome in various Fabaceae species. Black triangles indicate genes that were shown to be functionally transferred to the nucleus (Gantt *et al.*, 1991; Magee *et al.*, 2010), including the transfer of *rpl22* in *Lupinus luteus* detected in this study. The grey triangle indicates the functional replacement of the plastomic *rps16* gene by a mitochondrial gene functionally transferred to the nucleus (Ueda *et al.*, 2008). White triangles show the plastid genes lost during Fabaceae evolution and for which no functional replacement in the nucleus has been observed. The independent losses of *ycf4* in *Lathyrus sativus* and *Pisum sativum* were inferred from the results of Magee *et al.* (2010) who showed the presence of intact *ycf4* genes in some *Lathyrus* species.

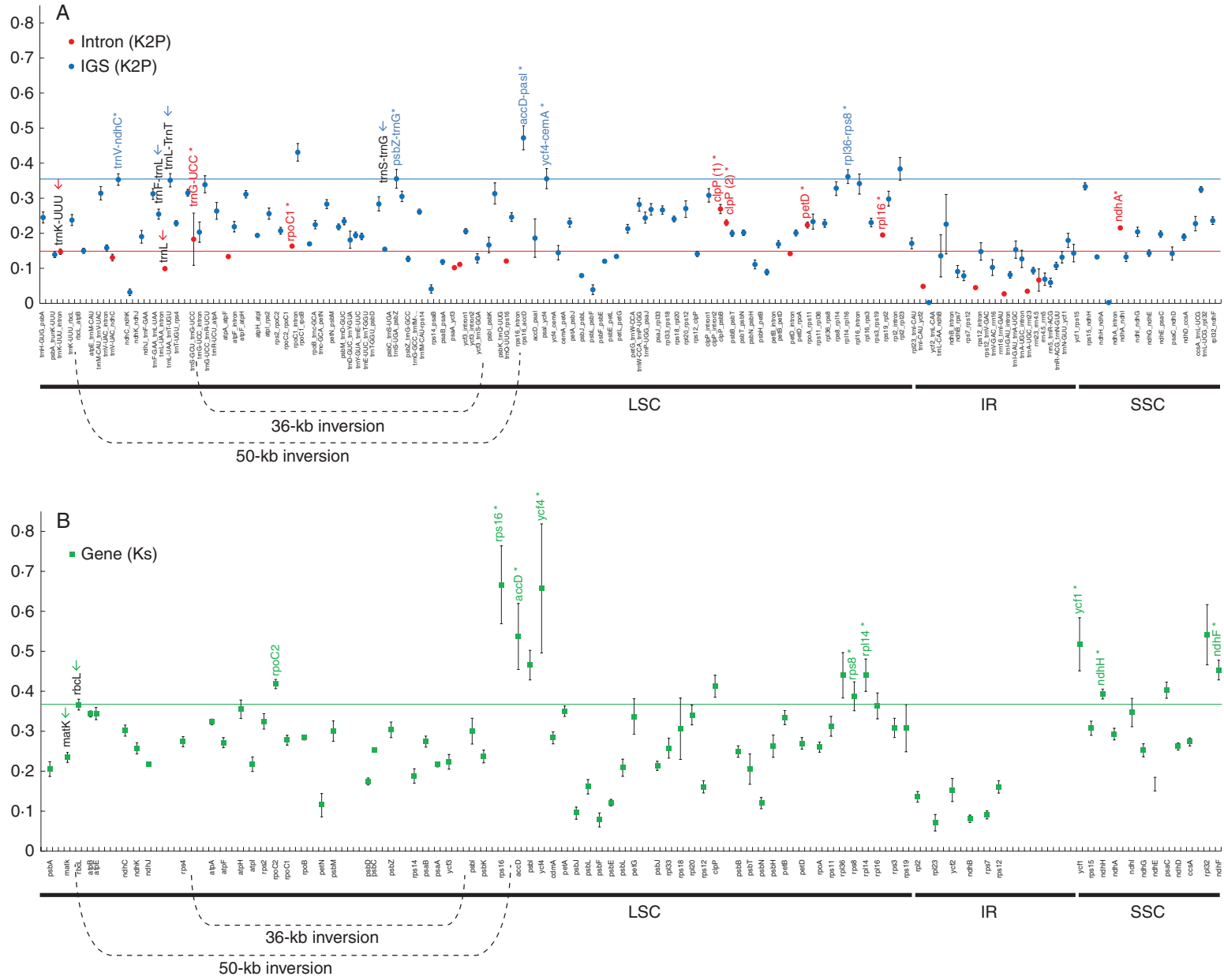
another repeat between *trnS-UGA* and the *trnS-GGA* (9 kb distant), all the others would presumably lead to the loss of functionality of one or several genes in the case of an inversion event and thus may be deleterious (Ruf *et al.*, 1997; Drescher *et al.*, 2000). Whilst the above 9-kb region is potentially prone to inversion, to date no evidence of such an event has been observed from this or previous studies in other Fabaceae species.

The rarity of plastomic rearrangements in flowering plants makes these characters powerful phylogenetic markers (Kim *et al.*, 2005) since they present an extremely low level of homoplasy (Cosner *et al.*, 2004). The 36-kb inversion identified in this study is present in all core Genistoid species surveyed (12) and therefore provides a robust additional synapomorphy supporting monophyly of the core Genistoids (Crisp *et al.*, 2000). Further screening of representatives from Brongniartieae and Bowdichia clades, shown to be closely related to the core Genistoids (Cardoso *et al.*, 2012), will determine whether this 36-kb inversion is strictly specific to the core Genistoids or whether it occurred earlier or at the base of the large Genistoid *s.l.* assemblage (includes Brongniartieae and Bowdichia clades). Thus, after the 50-kb inversion that is shared by a majority of Papilionideae (Doyle *et al.*, 1996), and the 78-kb inversion supporting the Papilionoid subtribe Phaseolinae (Bruneau *et al.*, 1990) in legumes, this 36-kb inversion represents an additional example highlighting the phylogenetic usefulness of plastidic inversions. Such clade-demarkating inversions were also detected in other Angiosperm families. Within the Asteraceae, a 22-kb inversion allowed identification of the subtribe Barnadesiinae as the most primitive lineage in the family (Jansen and Palmer, 1987). In the Campanulaceae, which also have highly rearranged plastomes, reliable phylogenetic relationships could be reconstructed within the

family only based on the use of the numerous rearrangements (including inversions) as characters (Cosner *et al.*, 2004). Interestingly, there is also evidence of specific mutational and restructuring events that affected the nuclear genome of *Lupinus* (Mahé *et al.*, 2011a), which suggests that the Genistoids experienced noteworthy genomic changes, in both the plastid and the nuclear genomes, after their divergence from the NPAAA papilionoid lineages (approx. 50–56 million years ago).

Gene content is highly conserved among photosynthetic angiosperm plastomes (Timmis *et al.*, 2004). However, within the Fabaceae, several chloroplastic genes (*accD*, *psal*, *rpl22*, *rpl23*, *rps16* and *ycf4*) have been lost recently and independently in various lineages (Magee *et al.*, 2010). However, within the *L. luteus* plastome, only the *rpl22* gene is missing, which is in accordance with the previous finding of Gantt *et al.* (1991) who demonstrated that the functional transfer of this gene from the chloroplast to the nucleus occurred in a common ancestor of all flowering plants, and thus preceded its loss from the chloroplast genome by about 100 million years (Supplementary Data Fig. S3). Among the chloroplast genes that have been lost in legume lineages following their divergence from the common ancestor with lupines, we found that the *rps16* gene is at an intermediate stage of functional replacement in the *Lupinus* nuclear genome, as it is still represented by a functional copy in the chloroplast genome while another is in the nucleus. This nuclear-encoded *rps16* gene targeted to the plastid is of mitochondrial origin and was transferred prior to the monocot–dicot divergence (Ueda *et al.*, 2008).

The evaluation of sequence divergence between *Lupinus* and the other sequenced legumes allowed identification of fast-evolving



sequences (Fig. 7; Supplementary Data Fig. S4). This information is essential for a better understanding of the dynamic nature of plastome evolution in legumes and for improving legume phylogeny, especially within the Genistoids and the genus *Lupinus* (Eastwood *et al.*, 2008; Mahé *et al.*, 2011a; Cardoso *et al.*, 2012). As expected, most coding regions are well conserved, particularly in the IR region, and in most cases IGSs are evolving faster than introns, in accordance with previous observations (Clegg and Zurawski, 1991; Raubeson *et al.*, 2007). Compared with the plastid sequences used in legume evolutionary studies, we have detected several sequences (Fig. 7) that exhibit higher rates of divergence: (1) seven introns (*trnG-UCC*, *rpoC1*, *rpl16*, *ndhA*, *petD* and *clpP* introns 1 and 2); (2) five IGSs (*ycf4_cemA*, *rpl36_rps8*, *psbZ_trnG-GCC*, *trnV-UAC_ndhC* and *accD_psaI*); and (3) eight protein-coding genes (*rpoC2*, *accD*, *ycf4*, *rps8*, *rpl14*, *ycf1*, *ndhH* and *ndhF*). Interestingly, most of these variable regions have not been or have rarely been employed in legume phylogeny [e.g. the *trnS-trnG* region in *Lupinus* by Drummond (2008); *ycf1* in *Astragalus* by Bartha *et al.* (2013)], and thus represent a new set of markers to explore evolutionary relationships within legumes. Each of these sequences needs to be tested in order to evaluate at which taxonomic level and in which lineages they could be more informative and useful. As an example, the remarkable increase in the evolutionary rate observed in the *ycf4* gene is specific to the IRLC, Robinoid and Millettoid lineages (NPAAA clade), and occurred after the divergence of the latter from the other legumes (Magee *et al.*, 2010). Thus, this region is most probably a good candidate for the NPAAA clade but seems less interesting for phylogenetic inference within the Genistoids and earlier legume lineages. In contrast, we have detected several loci exhibiting an increase in their evolutionary rate that is specific to the lupine/genistoid lineage. These loci include the *rpoC2* and *ndhF* genes, the *trnG-UCC*, *rpoC1* and *petD* introns, and the *trnV_ndhC* and *psbZ_trnG-GCC* IGSs. Regardless of the specificity and degree of utility of each locus, altogether these variable sequences constitute an important source of novel characters for single- or multigene-based reconstruction of evolutionary patterns in legumes at various taxonomic levels.

These variable sequences (mentioned above) are distributed in well-circumscribed fast-evolving regions that shape the legume plastome landscape (Fig. 7; Supplementary Data Fig. S4). Interestingly, three of these variable regions are located at boundaries of the 50-kb inversion (*rps16_ycf4* region), the 36-kb inversion (*trnS-GCU_trnG-UCC* region) and the IR region (*ycf1* region) (Fig. 7). As previously pointed out by Magee *et al.* (2010), these fast-evolving regions include gene and intron losses, such as genes lost from the *rps16_ycf4* region (*rps16*, *accD*, *psaI*, *rpl23* and *ycf4*) and introns lost from the *clpP_rps12* and the *rpl16* regions (Fig. 7). This suggests that these regions are most probably involved in structural rearrangements and thus represent unstable regions or hotspots that contribute significantly to the evolutionary dynamics of legume plastomes. Future research on the efficiency of the four classes of nuclear-encoded genes that are involved in chloroplast DNA repair and the maintenance of plastome stability (Maréchal and Brisson, 2010; Guisinger *et al.*, 2011) may reveal whether one or several of these four genes are implicated in legume plastome evolution. Additionally, this study demonstrates that it is essential to sequence plastomes from other papilionoid and earlier legume lineages that remain unexplored to date in order to

have a better understanding of the atypical plastome evolution observed in this family.

SUPPLEMENTARY DATA

Supplementary data are available online at www.aob.oxfordjournals.org and consist of the following. Table S1: list of software used in this paper. Table S2: codon usage bias. Table S3: repeated elements in the *Lupinus luteus* chloroplast genome. Table S4: sequence divergence (K2p) between *L. luteus* and ten other Fabaceae plastomes. Table S5: synonymous mutation rate between *L. luteus* and ten other Fabaceae plastome protein-coding sequences. Table S6: non-synonymous mutation rate between *L. luteus* and ten other Fabaceae plastomes. Table S7: Ka/Ks ratio between *L. luteus* and ten other Fabaceae plastomes. Table S8: identification of the presence of inverted repeated elements in *trnS-GGA* and *trnS-GCU* genes within rosid plastomes. Fig. S1: dot matrix plots showing the presence of a 36-kb inversion in the *Lupinus luteus* plastome. Fig. S2: comparative plastomic maps showing the presence of a 36-kb inversion in *Lupinus luteus* in comparison with other Papilionoideae. Fig. S3: phylogenetic analysis of plastidic and nuclear *rpl22* protein sequences. Fig. S4: pairwise distance between *Lupinus luteus* and other Fabaceae orthologous plastomic regions. Fig. S5: synonymous and non-synonymous divergence in legume chloroplast *ycf4* gene.

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