The complete chloroplast genome of *Papaver setigerum* and comparative analyses in Papaveraceae

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**Abstract**

*Papaver setigerum* is an annual herb that is closely related to the opium poppy, *P. somniferum*. Genetic resources for *P. setigerum* are scarce. In the present study, we assembled the complete chloroplast (cp) genome of *P. setigerum* based on genome skimming data, and we conducted comparative cp genome analyses to study the evolutionary pattern in Papaveraceae. The cp genome of *P. setigerum* is 152,862 bp in length with a typical quadripartite structure. Comparative analyses revealed no gene rearrangement in the Papaveraceae family, although differences were evident in genome size, gene losses, as well as inverted repeats (IR) region expansion and contraction. The rps15 gene has been lost from the genomes of *Meconopsis racemosa*, *Coreanomecon hylomeconoides*, *P. orientale*, *P. somniferum*, and *P. setigerum*, and the ycf15 gene is found only in *C. hylomeconoides*. Moreover, 13 cpDNA markers, including psbA-trnH, rps16-trnQ, trnS-trnG, trnC-petN, trnE-trnT, trnL-trnF, trnF-ndhJ, petA-psbJ, ndhF-rps32, rpl32-trnL, ccsA-ndhD, ndhE-ndhG, and rps15-ycf1, were identified with relatively high levels of variation within *Papaver*, which will be useful for species identification in this genus. Among those markers, psbA-trnH is the best one to distinguish *P. somniferum* and *P. setigerum*.

**Keywords:** *Papaver setigerum*, chloroplast genomes, chloroplast hotspot, species identification.

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**Introduction**

*Papaver setigerum* DC., an annual herb of the poppy family (Kalis, 1979), occurs in the Mediterranean region, especially in southwestern Europe (Portugal, Spain, France, Italy, Greece) and North Africa (Pignatti, 1982). This plant is closely related to and sometimes treated as a variety or sub-species of opium poppy (*P. somniferum* L.) due to its similarity in flower-shape, color, fruit, and production of small amounts of morphine alkaloids (La Valva et al., 1985; Osalou et al., 2013). Of the 110 species of the genus *Papaver*, only *P. somniferum* and *P. setigerum* are controlled species in most countries (Choe et al., 2012). However, the cytological evidence shows that *P. somniferum* is diploid (2n = 22), while *P. setigerum* is tetraploid (2n = 44) (Fulton, 1944; Choe et al., 2012), indicating that *P. setigerum* is not likely the wild ancestral species of the cultivated *P. somniferum* (Farmilo et al., 1953). For *Papaver* species, inter-specific identification based only on morphological characteristics is difficult because of the similarities in appearance mentioned above (Osalou et al., 2013). Phytochemical methods (Zhang and Cheng, 2009; Osalou et al., 2013) and various molecular markers (Fan et al., 1987; Hosokawa et al., 2004; Choe et al., 2012; Zhang et al., 2015) have been used to identify *Papaver* species in previous studies. However, current studies involving *P. setigerum* have mostly focused on its chemical composition, largely ignoring its genetic background.

Chloroplasts (cp), the photosynthetic organelles of most green plants, are known to be derived from cyanobacteria through endosymbiosis and co-evolution (Dagan et al., 2012; Asaf et al., 2017). In most angiosperms, cp genomes have a typically circular and quadripartite structure. The genome size is usually from 115 to 165 kb in length, consisting of two regions of inverted repeats (IRs), separated by a large single-copy (LSC) region and a small single-copy (SSC) region (Wicke et al., 2011). Compared with nuclear and mitochondrial genomes, the cp genome is more conserved, not only in gene content and organization, but also in genome structure (Raubeson and Jansen, 2005). Due to its...
relatively conserved gene content and simple structure, small size, uniparental inheritance, and the fact that it is non-recombinant, the cp genome has been used as an ideal model for phylogenetic reconstruction (Liu et al., 2017), evolutionary and comparative genomic studies (Liu et al., 2018b), species identification (Thomson et al., 2010; Greiner et al., 2015) and markers development (Liu et al., 2018a). Currently, the rapid development and improvement of next-generation sequencing technology have made the assembly of the cp genome cheaper and more efficient compared with traditional sequencing (Alkan et al., 2011). In addition, the releases of many assembly processes or pipelines, such as SOAPdenovo2 (Luo et al., 2012), CLC Genomics Workbench (CLC Inc., Rarhus, Denmark) and GetOrganelle (Jin et al., 2018), have made cp genome reconstruction easier and much more effective.

In the present study, one P. setigerum individual was selected for genome skimming, and the complete chloroplast genome sequence was assembled and reported. We also compared the cp genomes among representatives of Papaveraceae and detected highly divergent regions of the cp genomes within the genus Papaver.

Material and Methods

Plant material, DNA extraction, and sequencing

We extracted whole-genomic DNA from silica-dried leaf tissue of one cultivated P. setigerum plant collected in Taizhou (Zhejiang, China), using modified CTAB reagent Plant DNAzol (Invitrogen, Shanghai, China) according to the manufacturer’s protocol. High quality DNA was sheared to yield fragments with length less than or equal to 800 bp. The quality of fragmentation was checked on an Agilent Bioanalyzer 2100 (Agilent Technologies). The 500 bp short-insert length paired-end library was prepared and sequenced by Beijing Genomics Institute (BGI, Wuhan, China). The library was run in one lane of an Illumina HiSeq X10 and obtained reads with length of 150 bp.

Chloroplast genome assembly and annotation

The raw reads were first screened for Phred score < 30 to remove low-quality sequences. In order to ensure the accuracy of cp genome assembly, we employed two different methods to assemble the cp genome. In the first method, all the remaining reads were assembled into contigs implemented in the CLC genome workbench (CLC Inc., Rarhus, Denmark). The parameters set in CLC were as follows: 200 bp for minimum contig length, 3 for deletion and insertion costs, bubble size of 98, 0.9 for length fraction and similarity fraction, and 2 for mismatch cost. Then, the principal contigs representing the cp genome were separated from the total contigs using a BLAST (NCBI BLAST V2.2.31) search, with the cp genome of P. somniferum set as the reference. The representative cp contigs were oriented and ordered on the basis of the reference cp genome, and the complete chloroplast genome of P. setigerum was reconstructed by connecting overlapping terminal sequences. In the second method, the cp genome of P. setigerum was de novo assembled using the GetOrganelle pipeline (Jin et al., 2018), with SPAdes 3.10.1 as assembler (Bankevich et al., 2012).

Geneious R11 (https://www.geneious.com) was used to annotate the cp genome of P. setigerum, and putative starts, stops, and intron positions were identified on the basis of comparisons with homologous genes of the P. somniferum cp genome. The tRNA genes were verified with tRNAscan-SE v1.21 (Schattner et al., 2005) with the default setting. We drew the circular chloroplast genome map of P. setigerum using the OrganellarGenomeDRAW program (OGDRAW, Lohse et al., 2013).

Comparative chloroplast genomic analyses

In order to study the sequence variation within Papaveraceae, we downloaded multiple publicly available cp genomes of the family from GenBank to compare the overall similarities, using Leontice incerta (Berberidaceae, MH940295) as the reference, according to the results of Kim and Kim (2016). The GenBank accession numbers for the Papaveraceae species are as follows (Table S1): P. orientale (NC_037832), P. rheas (NC_037831), P. somniferum (NC_029434), Meconopsis racemosa (NC_039625), Coreanomecon hyleomeconoides (NC_031446), and Macleaya microcarpa (NC_039623). The sequence identities of the seven Papaveraceae cp genomes were implemented in the mVISTA program with LAGAN mode (Frazer et al., 2004). The cp DNA rearrangement analyses of seven Papaveraceae cp genomes were based on Mauve Alignment (Darling et al., 2004).

Molecular markers development for Papaver

In order to screen variable characters within Papaver, multiple alignments of the four Papaver species cp genomes were carried out using MAFFT version 7.017 (Katoh and Standley, 2013). The nucleotide diversity (Pi) was determined by calculating the total number of mutations (Eta) and average number of nucleotide differences (K) using DnaSP v5.0 (Librado and Rozas, 2009).

Phylogenetic inferences

The phylogenetic relationships of Papaveraceae were inferred using the whole chloroplast genome sequences of seven species; two species from Ranunculaceae (Ranunculus macranthus) and Berberidaceae (Leontice incerta) were chosen as the outgroups, according to the results of Kim and Kim (2016). The phylogeny inferences were conducted using Bayesian inference (BI) and maximum likelihood (ML) methods. ML analysis was performed with RAxML-HPC v8.1.11 on the CIPRES cluster (Miller et al., 2010) with GTR + I + G set as the best-fit nucleotide substitution model. BI analysis was implemented in MrBayes v3.2.3 using the same substitution model mentioned above (Ronquist and Huelsenbeck, 2003).
Results

P. setigerum cp genome assembly, organization and gene content

The complete cp genomes of P. setigerum assembled from two different assembly strategies were identical. However, using GetOrganelle to assemble the cp genome of P. setigerum was much faster and more effective than using CLC genome workbench (< 1h vs. > 6h). The cp genome size was 152,862 bp, and had a typical quadripartite structure that was similar to the majority of land plant cp genomes, consisting of an 83,022 bp large single copy region (LSC), a 17,944 bp small single copy region (SSC) and two 25,948 bp inverted repeats. The P. setigerum cp genome contains 113 unique genes, including 79 protein-coding genes, 30 tRNA genes and four ribosomal RNA genes (Figure 1 and Table 1). Eight protein-coding, seven tRNA, and four rRNA genes are duplicated and located in the IR regions. Among the 113 genes, nine protein-coding genes and six tRNA genes contain one intron; three protein-coding genes (clpP, ycf3 and rps12) contain two introns. We submitted the cp genome of

P. setigerum to GenBank with the accession number MK820043.

Genome comparison of Papaveraceae

The chloroplast genomes of the seven Papaveraceae species were relatively conservative, and the IR region is more conserved compared to the LSC and SSC regions (Figure 2). No rearrangements, such as translocations or inversions, occurred in gene organization after verification in this family (Figure 3). However, differences existed in genome size, gene losses, and IR expansion and contraction.

In terms of the cp genome size observed among the representative Papaveraceae species, the four Papaver species were the smallest and had similar genome sizes ranging from 152,799 bp to 152,931 bp (Figure 4). Of the other species, Macleaya microcarpa (161,124 bp) exhibited the largest cp genome, while Meconopsis racemosa (153,763 bp) had the smallest one.

The rps15 gene has been lost from the genomes of M. racemosa, C. hylomeconoides, P. orientale, P. somniferum, and P. setigerum, although it is present in P. rhoeas and the reference genome. In addition, the ycf15 gene occurred only

Figure 1 - Chloroplast genome map of Papaver setigerum. Genes inside the circle are transcribed clockwise, genes outside are transcribed counterclockwise. The light gray inner circle corresponds to the AT content, the dark gray to the GC content. Genes belonging to different functional groups are shown in different colors.
in *C. hylomeconoides* compared to the other analyzed cp genomes.

In addition, we compared the exact IR border positions and their adjacent genes between the seven Papaveraceae cp genomes and the reference genome (Figure 4). The results showed that the *ycf1* gene spanned the SSC/IRA region and the pseudogene fragment of *ycf1* varied from 912 bp to 1379 bp. The *ndhF* gene shares some nucleotides (25 bp) with the *ycf1* pseudogene in *Meconopsis racemosa* but is separated from *ycf1* by spacers in the other analyzed species. The *trnH-GUG* gene was located in the LSC region of all genomes, but varied from 5 bp to 117 bp apart from the IR/IRA junctions. In addition, the *rps19* pseudogene appeared in all the representative Papaveraceae species due to the *rps19* gene extending to the IR region.

### Table 1 - Genes contained in *P. setigerum* chloroplast genome (113 genes in total).

| Category                         | Group of gene                  | Name of gene       |
|----------------------------------|--------------------------------|--------------------|
| Self-replication                 | Ribosomal RNA genes            | *rrn*4.5*<sup>a</sup> | *rrn*5<sup>a</sup> | *rrn*16<sup>a</sup> | *rrn*23<sup>a</sup> |
|                                  | Transfer RNA gene               | *trn*-UGC<sup>a</sup> | *trn*-C-GCA | *trn*-D-GUC | *trn*-E-UUC |
|                                  | *trn*-F-GAA               | *trn*-G-M-CAU | *trn*-G-GCC<sup>a</sup> | *trn*-G-UCC |
|                                  | *trn*-H-GUG               | *trn*-I-C-AU<sup>a</sup> | *trn*-I-GAU<sup>a</sup> | *trn*-K-UUU<sup>a</sup> |
|                                  | *trn*-L-CAA<sup>a</sup> | *trn*-L-UAA<sup>a</sup> | *trn*-L-UAG | *trn*-M-CAU |
|                                  | *trn*-N-GU<sup>a</sup> | *trn*-P-UUG | *trn*-Q-UUG | *trn*-R-ACG<sup>i</sup> |
|                                  | *trn*-R-UCU               | *trn*-S-GCU | *trn*-S-GGA | *trn*-S-UCA |
|                                  | *trn*-T-GGU               | *trn*-T-UUG | *trn*-V-GAC<sup>a</sup> | *trn*-V-UAC<sup>a</sup> |
|                                  | *trn*-W-CCA               | *trn*-Y-GUA |
|                                  | Small subunit of ribosome   | *rps*2 | *rps*3 | *rps*4 | *rps*7<sup>a</sup> |
|                                  |                             | *rps*8 | *rps*11 | *rps*12<sup>44</sup>| *rps*14 |
|                                  |                             | *rps*15 | *rps*16<sup>a</sup> | *rps*18 | *rps*19<sup>a</sup> |
|                                  | Large subunit of ribosome  | *rpl*2<sup>**</sup> | *rpl*14 | *rpl*16<sup>a</sup> | *rpl*20 |
|                                  |                             | *rpl*22 | *rpl*23<sup>a</sup> | *rpl*32 | *rpl*33 |
|                                  |                             | *rpl*36 |            |            |           |
| Photosynthesis                    | RNA polymerase subunits     | *rpo*A | *rpo*B | *rpo*C | *rpo*C<sup>1</sup> | *rpo*C2 |
|                                  | Subunits of photosystem I   | *psa*A | *psa*B | *psa*C | *psa*I |
|                                  |                             | *psa*I | *ycb*2<sup>**</sup> |            |           |
|                                  | Subunits of photosystem II  | *psb*A | *psb*B | *psb*C | *psb*D |
|                                  |                             | *psb*E | *psb*F | *psb*H | *psb*I |
|                                  |                             | *psb*J | *psb*K | *psb*L | *psb*M |
|                                  | Subunits of cytochrome     | *pet*A | *pet*B<sup>a</sup> | *pet*D<sup>a</sup> | *pet*G |
|                                  |                             | *pet*L | *pet*N |            |           |
|                                  | Subunits of ATP synthase    | *atp*A | *atp*B | *atp*E | *atp*7<sup>a</sup> |
|                                  |                             | *atp*H | *atp*I |            |           |
|                                  | Large subunit of Rubisco    | *rbc*L |            |            |           |
|                                  | Subunits of NADH           | *ndh*A<sup>a</sup> | *ndh*B<sup>a</sup> | *ndh*C | *ndh*D |
|                                  | Dehydrogenase              | *ndh*E | *ndh*F | *ndh*G | *ndh*H |
|                                  |                             | *ndh*I | *ndh*J | *ndh*K |           |
| Other genes                      | Translational initiation factor | *inf*A | |            |           |
|                                  | Maturase                    | *mat*K |            |            |           |
|                                  | Envelope membrane protein   | *cem*A |            |            |           |
|                                  | Subunit of acetyl-CoA       | *acc*D |            |            |           |
|                                  | C-type cytochrome synthesis gene | *ccr*A | |            |           |
|                                  | Protease                    | *clp*P<sup>**</sup> |            |            |           |
| Unknown function                 | Conserved open reading frames | *ycf1*(part) | *ycf2<sup>a</sup> | *ycf4* |

<sup>a</sup> Two gene copies in IRs; <sup>b</sup> gene divided into two independent transcription units; one and two asterisks indicate one- and two-intron containing genes, respectively.
The cp genome of *Papaver setigerum*

Figure 2 - Visualization of alignment of the seven Papaveraceae chloroplast genome sequences, with *Leontice incerta* as the reference. The horizontal axis indicates the coordinates within the chloroplast genome. The vertical scale indicates the percentage of identity, ranging from 50 to 100%. Genome regions are color coded as protein coding, intron, mRNA, and conserved non-coding sequences (CNS).

Figure 3 - MAUVE alignment of seven Papaveraceae chloroplast genomes. *Leontice incerta* is shown at the top as the reference. Within each of the alignments, local collinear blocks are represented by blocks of the same color connected by lines.
In order to explore the divergence hotspot regions in *Papaver*, we divided the genome alignment of the four *Papaver* species into non-coding regions, coding genes, and intron regions. We eventually identified 125 loci (53 coding genes, 55 inter-genic spacers, and 17 intron regions) within *Papaver* having more than 200 bp in length (Figure 5). Of these 125 regions, nucleotide variability ($\pi$) values ranged from 0.0003 (*rrn16*) to 0.0474 (*psbA-trnH*). Thirteen of these variable loci ($\pi > 0.02$), including *psbA-trnH*, *rps16-trnQ*, *trnS-trnG*, *trnC-petN*, *trnE-trnT*, *trnL-trnF*, *trnF-ndhJ*, *petA-psbH*, *ndhF-rpl32*, *rpl32-trnL*, *ccsA-ndhD*, *ndhE-ndhG*, and *rps15-ycf1*, showed high levels of intrageneric variation. Within the 13 regions, seven varied between *P. setigerum* and *P. somniferum* (Table S2), and these can be candidate markers to identify these two species. Among those markers, *psbA-trnH* is the best one to distinguish *P. somniferum* and *P. setigerum*, which are different by seven site mutations.

**Phylogenetic inferences**

The tree topologies from both ML and Bayesian analyses were consistent with each other (Figure 6). All but one node within Papaveraceae have full support (maximum likelihood bootstrap, MLBS = 100%, Bayesian inference posterior probabilities, BIPP = 1). The four *Papaver* species formed one clade with full support and is sister to *Meconopsis racemosa*. The remaining two species, *Macleaya microcarpa* and *Coreanomecon hylomeconoides*, formed another clade.

**Discussion**

In the last decades, the rapid development of high throughput sequencing technologies have greatly reduced sequencing cost. Considering the large number of copies of
the plastid genome in a single cell, it is easy to get enough reads to reconstruct a complete cp genome from low-coverage, whole-genome sequencing data (Twyford and Ness, 2017), viz. genome skimming data (Straub et al., 2012). Since the first complete nucleotide sequence of the cp ge-

With the publication of many cp genome assembly pipelines (Luo et al., 2012; Jin et al., 2018), cp genome reconstruction by these protocols is more effective than the Sanger method. Since the first complete nucleotide sequence of the cp ge-

**Figure 5** - Comparative analysis of the nucleotide variability (Pi) values among the four *Papaver* species.

**Figure 6** - Phylogenetic tree reconstruction of Papaveraceae using maximum likelihood (ML) based on whole chloroplast genome sequences. Numbers above the branches represent bootstrap values from maximum likelihood analyses and posterior probabilities from Bayesian inference, respectively.
nome was generated (Nicotiana tabacum; Shinozaki et al., 1986), more than 3000 cp genomes have been submitted to GenBank (Jin et al., 2018). In this study, we tried to assemble the cp genome sequence of Papaver setigerum using two different pipelines, the CLC Genomics Workbench (CLC Inc., Rarhus, Denmark) and GetOrganelle (Jin et al., 2018). Cp genome sequences produced by the two pipelines were completely identical in terms of both genome size and base information. However, the GetOrganelle pipeline is faster and more effective in assembling a circular cp genomes of P. setigerum or other species (we are preparing to publish a comparative study separately), especially for the low coverage data of the whole genome.

In recent years, comparative studies of cp genomes have been applied to a number of focal species (Young et al., 2011), genera (Greiner et al., 2008; Liu et al., 2018b), or plant families (Daniell et al., 2006; Liu et al., 2017). Comparative analyses of cp genomes are useful for phylogenetic inference at higher taxonomic levels (Moore et al., 2010; Li et al., 2017), as well as for understanding the evolution of genome size variations, gene and intron losses, and nucleotide substitutions. In the present study, multiple complete cp genomes of representative Papaveraceae species provide an opportunity to compare the sequence variation within the family. No rearrangement, such as translocations and inversions, occurred in gene organization in this family. However, we identified differences in genome size, gene losses, and IR expansion and contraction. The rps15 gene is found in most cp genomes in land plants (Tsujii et al., 2007). However, comparative analysis revealed that this gene was found in P. rhoes and the reference genome Leonice incerta, but was not present in other Papaveraceae species. Previous studies have confirmed that the rps15 loss has also appeared in other families (Tsujii et al., 2007; Krause, 2012). Similarly, the function of the ycf15 gene has attracted the attention of previous workers (Raubeson et al., 2007; Shi et al., 2013), and it has apparently been lost in other taxa (Liu et al., 2017; Liu et al., 2018). The ycf15 gene, which displays a small open reading frame (ORF), is located immediately downstream of the ycf2 gene (Dong et al., 2013). In our study, the ycf15 gene occurred only in Coreanomecon hylomeconoides, located immediately downstream of the ycf2 gene but absent in other analyzed cp genomes. These findings suggest that parallel losses of particular genes have occurred during the evolution of land plant cp genomes.

In the genus Papaver, almost all of the species are similar in their flower-shapes (two sepalsthat fall off as the bud opens and four to six petals), colors, and fruits, complicating species identification based on morphological characteristics alone (Osalou et al., 2013; Zhou et al., 2018). Previous studies have identified Papaver species using physicochemical methods, including discrete stationary wavelet transform (Zhang et al., 2009), amplified fragment length polymorphism (Lu et al., 2008), as well as phytochemical methods (Osalou et al., 2013). Hosokawa et al. (2004) identified Papaver species using the plastid gene rpl16 and rpl16-rpl14 spacer sequences. Zhang et al. (2015) had verified that trnL-trnF can be considered a novel DNA barcode to identify the Papaver genus, and ITS, matK, psbA-trnH, and rbcL can be used as combined barcodes for identification. Zhou et al. (2018) screened five hypervariable regions, including rpoB-trnC, trnD-trnT, petA-psbJ, psbE-psrL, and ccsA-ndhD, as specific DNA barcodes. In this study, except for the regions mentioned above, we developed nine additional regions (ps16-trnQ, trnS-trnG, trnC-petN, trnE-trnT, trnF-ndhJ, ndhF-rpl32, rpl32-trnl, ndhE-ndhG and rps15-ycf1) with relatively high levels of intragenic variation, which can be used for identifying Papaver species in the future. Moreover, P. setigerum was formerly treated as a variety or subspecies of P. somniferum due to the similar morphological appearance and chemical signature (La Valva et al., 1985; Osalou et al., 2013). However, the cytological evidence rejects this perspective (Fulton, 1944). Besides, there are seven cp regions varied between P. setigerum and P. somniferum (Table S2). In the phylogenetic tree of the present study, P. setigerum is sister to P. somniferum with full support within the Papaver clade, which cannot be applied for determining the phylogenetic relationship of these two species due to lack of population sampling. Therefore, more samples for each species in subsequent studies will help us to resolve the genetic relationship between P. setigerum and P. somniferum.

Conclusion

In the present study, we assembled the complete chloroplast genome sequence of Papaver setigerum based on genome skimming data. The chloroplast genome of P. setigerum had a typical quadripartite structure with similar size and organization to other sequenced angiosperms. The evolutionary pattern of cp genomes in Papaveraceae was also detected utilizing seven representative species. Moreover, we screened additional cp hotspots regions for the genus Papaver, which will contribute to identification of species in this genus. The inter-genic region psbA-trnH is the best marker to distinguish P. somniferum and P. setigerum.

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Conflict of Interest

The authors declare that they have no potential conflict of interest.
The cp genome of *Papaver setigerum*  

Author Contributions  
PL and JL designed the study. PL conducted the sampling. LXL, YXD and CS produced and analyzed the data. LXL, YXD and RL wrote the manuscript. PL and JL revised the manuscript. All authors approved the final manuscript.

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Supplementary Material
The following online material is available for this article:
Table S1 - The GenBank accession numbers for other seven species used in comparative chloroplast genome analyses.
Table S2 - The variable sites information of seven chloroplast (cp) regions between P. somniferum and P. setigerum.

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