Genome Reports: Contracted Genes and Dwarfed Plastome in Mycoheterotrophic Sciaphila thaidanica (Triuridaceae, Pandanales)

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Abstract

With a reduced need for photosynthesis, the plastome of parasitic and mycoheterotrophic plants degrades. In the tiny, fully mycoheterotrophic plant Sciaphila thaidanica, we find one of the smallest plastomes yet encountered. Its size is just 12,780 bp and it contains only 20 potentially functional housekeeping genes. Thus S. thaidanica fits the proposed model of gene loss in achnorophyllous plants. The most astonishing feature of the plastome is its extremely compact nature, with more than half of the genes having overlapping reading frames. Additionally, intergenic sequences have been reduced to a bare minimum, and the retained genes have been reduced in length both compared with the orthologous genes in another mycoheterotrophic species of Sciaphila and in the autotrophic relative Carludovica.

Key words: gene loss, genome reduction, mycoheterotrophy, plastid genome evolution.

Introduction

Sciaphila Blume is a genus of fully mycoheterotrophic plants that parasitize mycorrhizal fungi. The need for carbon fixation has been eliminated and photosynthesis has been lost completely. The loss of photosynthesis is shared between fully mycoheterotrophic and holoparasitic plants, and a number of studies have investigated how plastid genomes (plastomes), housing a large number of genes related to photosynthesis, evolve when photosynthesis is no longer needed (reviewed in Krause 2011; Wicke et al. 2011; Graham et al. 2017). Not surprisingly, plastomes gradually degrade as genes are under relaxed or no selective pressure, have increased substitution rates, are pseudogenized and eventually lost. Simultaneously, the length of noncoding sequences is reduced and structural rearrangements are more frequent. Most changes lead to compaction of the plastome and potentially it may be lost completely, as have been suggested in Rafflesia R. Br. (Molina et al. 2014). As the plastome includes both photosynthesis genes and housekeeping genes, gene loss is not random, but follows a pattern, which seems to be shared among the different clades of achnorophyllous plants (Barrett and Davis 2012; Barrett et al. 2014; Wicke et al. 2016; Graham et al. 2017).

Sciaphila belongs in the monocot family Triuridaceae (Pandanales). With more than 50 species, the Triuridaceae constitute one of the largest clades of fully mycoheterotrophic plants, but only one complete plastome from a species of Sciaphila has been described previously (Lam et al. 2015). In order to further explore plastome evolution in response to loss of photosynthesis further, we here describe a complete plastome sequence from another species of Sciaphila, S. thaidanica K. Larsen (fig. 1), and compare it to the previously sequenced plastome of S. densiflora Schlr. We also compare both Sciaphila plastomes with the recently sequenced plastome of an autotrophic representative of the Pandanales, Carludovica palmata Ruiz. & Pav. (Cyclanthaceae) (Lam et al. 2015), and assess whether Sciaphila fits the pattern of plastome modification previously described for achnorophyllous plants (Barrett and Davis 2012; Barrett et al. 2014; Wicke et al. 2016; Graham et al. 2017).

Materials and Methods

DNA Extraction, Library Preparation, and Sequencing

DNA was extracted from a silica gel sample of Sciaphila thaidanica (Collected: Thailand, Phang Nga Province, Sra Nang...
Manora Forest Park, 17 Nov. 2014, specimen voucher Suddee et al. 4796; BKF) using the DNA Plant Minikit (Qiagen) according to the manufacturer’s instructions, with the addition of a Proteinase K treatment step, for 2 hours at 65°C, immediately after the bead-beating step (Qiashredder).

The extracted DNA was sheared using a Bioruptor (Diagenode). A 100-µl dilution at a concentration of 10 ng/µl DNA was sheared using the following conditions: 5 cycles of 15 seconds ON, 90 seconds OFF. The sheared DNA was then run on a 2% agarose gel with a 100-bp DNA ladder (Thermo Scientific), to check the size variation of the DNA fragments, which were in the range of 200–600 bp. To prepare the fragments for inserting the adapter sequences and unique barcodes, the NEBNext DNA Sample Prep Master Mix Set 2 (New England Biolabs, Ipswich, Massachusetts) using blunt-end adapters specified by Meyer and Kircher (2010) was used. In order to determine the number of cycles needed for library construction, qPCRs were performed on a 1:40 dilution of the DNA sample, using the SYBR Green Master Mix 1 (Agilent Technologies) on an Mx3500p qPCR machine (Agilent Technologies). Libraries were amplified 13 cycles in a total volume of 100 µl, containing 5 U AmpliTaq Gold polymerase (Applied Biosystems, Foster City, CA), 1× AmpliTaq Gold buffer, 2.5 mM MgCl₂, 0.4 mg/ml bovine serum albumin (BSA), 0.2 mM of each dNTP, 0.2 µM IS4 forward primer, 0.2 µM indexed reverse primer, and 20 µl DNA library template. Following amplification, libraries were purified using a QIAquick PCR Purification kit (Qiagen, Hilden, Germany), according to manufacturer’s instructions. DNA was eluted in 32 µl EB buffer and the column was incubated for 10 minutes at 37°C prior to centrifugation. The libraries were first quantified on a Qubit 2.0 (Life Technologies, Carlsbad, CA) using dsDNA high sensitivity assay, and then run on a TapeStation 2200 using the high sensitivity tapes (Agilent, Santa Clara, CA) to determine the average insert size and molarity of each library. About 100-bp paired-end sequencing was performed using an Illumina Hiseq 2500 at the National High-Throughput DNA Sequencing Centre.

Genome Assembly, Annotation, and Analysis

Raw reads were trimmed for quality, adapters, and unidentified nucleotides (Ns) using Adapter Removal (Lindgreen 2012). To roughly assess the gene content, reads were initially mapped to the complete plastome sequences of Sciaphila densiflora (NC027659) and Carludovica palmata (NC026786) using the Map to Reference option of Geneious version 11.0.2 (Biomatters Ltd) under default settings. As sequence reads only mapped to few genes, suggesting the plastome to be very small, we subsequently followed a procedure based on repeated reference mapping and de novo assembly for assembling a complete plastome: Consensus sequences calculated from the reads mapped to reference plastome genes were extracted and extended using several rounds of Map to Reference with custom settings using a higher sensitivity (maximum mismatches per read 5%; allow only 5% gaps per read) and up to 10 iterations where reads are automatically mapped to the consensus of the previous iteration. After each round of reference mapping, assemblies were inspected for errors and potential places where reads could split in two directions. Cases of the latter were not observed. Accepted new consensus sequences were used for de novo assembly using default options of Geneious version 11.0.2 (Biomatters Ltd). This procedure was continued until all contigs were assembled into a circular structure. The plastome genes were manually annotated in Geneious version 11.0.2 (Biomatters Ltd) following comparison to reference organellar genomes from Pandanales. Open reading frames of more than half the length of orthologous genes from autotrophic relatives were accepted as potentially functional genes.

Plastome sequences may be transferred to the mitochondrial genome, and this may confound plastome assembly. Thus, in order to roughly assess genomic location of sequences, we mapped the reads from Sciaphila thaidanica to five mitochondrial protein coding genes from Carludovica palmata available in GenBank (AF197734, AF197707, DQ406948, DQ508954, GU3511605) to determine mitogenome coverage.

Sequence similarity of the coding sequences retained in Sciaphila and those in Carludovica palmata were calculated from pairwise alignments using MUSCLE version 3.8.425 (Edgar 2004) as implemented in Geneious version 11.0.2 (Biomatters). For protein coding genes, we applied the translation alignment option.

Results and Discussion

From approximately 42 million paired-end reads, the plastome of Sciaphila thaidanica was assembled into a circular structure of 12,780 bp (fig. 2; GenBank accession number MG757197) with an average coverage of ~452×. Coverage is rather even across the plastome with differences being related to GC content. Coverage of the mitochondrial loci is ca. 60×, thus it is not likely that potentially transferred plastome sequences have been confounding for assembly of the plastome.

Like the previously sequenced plastome of S. densiflora (Lam et al. 2015), the S. thaidanica plastome lacks one of the inverted repeat regions. Although higher coverage of two regions of the S. densiflora plastome suggested possible duplications, this is not the case for S. thaidanica. The plastome is largely colinear with the Carludovica palmata plastome except for one inversion spanning a region from rps12 exon1 to rps2. Thus, colinearity is better conserved in S. thaidanica than in S. densiflora, which differed from Carludovica by two inversions (Lam et al. 2015). The inversion in S. thaidanica is not similar to any of those in S. densiflora indicating that they represent three separate structural rearrangement events. Although plastome structure and gene order is usually

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highly conserved among angiosperms, achlorophyllous plants are particularly prone to structural changes (Wicke et al. 2016).

Although the plastome of *S. densiflora* is drastically reduced compared with *Carludovica palmata* (Lam et al. 2015), reduction has proceeded even further in *S. thaidanica* (table 1). The size has been reduced dramatically and the gene content is also substantially lower. All the 20 genes included in the *S. thaidanica* plastome are present in *S. densiflora*, but in addition, the latter includes putatively functional copies of *clpP*, *matK*, *rpl20*, *rpl36* and four more tRNA genes (*trnC*, *trnI*, *trnQ*, *trnW*) (Lam et al. 2015). The genes retained in *S. thaidanica* (*accD*, *rps12*, *rrn*, *trn*), which are all housekeeping genes, fit a model of gene loss from achlorophyllous plants perfectly (Barrett and Davis 2012; Barrett et al. 2014; Wicke et al. 2016), and the presence of *trnE* as one of only two tRNA genes supports the importance of this gene (Barbrook et al. 2006). The only nonribosomal protein coding gene, *accD*, is retained as in most other mycoheterotrophs yet sequenced (Lam et al. 2016; Graham et al. 2017). Only in some mycoheterotrophic Ericaceae has this gene possibly been functionally lost (Braukmann et al. 2017).

Two of the ribosomal protein genes (*rpl2*, *rps12*) retained in the *S. thaidanica* plastome contain a group IIA intron supposed to be spliced by the *matK* gene product (Zoschke et al. 2010). However, this gene has been lost from the plastome.

Loss of *matK* coupled with retention of *rpl2* and *rps12* can be observed in a remarkable high number of parasitic and mycoheterotrophic plants suggesting that intron splicing may occur by alternative means (Graham et al. 2017). Alternatively, *matK* could be functionally transferred to the nuclear genome.

Reduction of the plastomes in *Sciaphila* is associated with compaction of the remaining genes. In *Carludovica palmata* genes occupy 70.2% of the plastome, in *Sciaphila densiflora*, the gene proportion is increased to 78.0% and in *S. thaidanica* to 91.9% leaving just 1.030 bp of intergenic sequence (table 1). This compaction of the *S. thaidanica* plastome has led to an unusual overlap of reading frames. Eleven genes have short overlapping reading frames in the 3′- or 5′-end, and among them an array of five genes (*rpl14-rpl16-rps3-rps19-rpl2*) has no intergenic sequence at all (fig. 2). In addition, two ribosomal RNA genes (*rrn23*, *rrn4.5*) are just touching. Although reduction of the length of noncoding sequence, both through intron loss and deletions in intergenic regions, is well-known for plastomes from parasitic plants (Funk et al. 2007; McNeal et al. 2007; Petersen et al. 2015; Naumann et al. 2016), none of the previously described plastomes have attained a stage of gene compaction as observed here. To further understand how compaction has occurred a denser taxon sampling is crucial. Provided that the overlapping genes are functional (see below), they may be
cotranscribed, but this possibility will need to be tested in live plant material.

The genes retained in the Sciaphila plastomes are also shorter in length; both with regard to coding and noncoding sequence. Intron coverage of the complete genomes drops from 12.7\% in Carludovica palmata to 7.0\% in Sciaphila thaidanica (table 1), but these differences are partly caused by loss of intron containing genes, thus a more precise picture is provided by the few remaining intron-containing genes in Sciaphila (table 2). Compared with Carludovica palmata, intron length of the three relevant genes (*rpl2, rpl16, rps12*) is reduced to 75.3\% in *S. densiflora* and 41.6\% in *S. thaidanica*. In the latter, the *rpl16* intron is lost completely. The length of the coding sequence of the genes present is reduced, too, although less drastically. The difference between *Carludovica palmata* and *S. densiflora* is hardly significant, but in *S. thaidanica*, the length of coding sequence has been reduced to 88.5\% of the former, affecting most of the retained genes (table 2). Although intron loss is not uncommon, consistent reduction of coding sequence appears unusual. In *Thismia*

### Table 1
Comparison of Complete Plastomes of *Carludovica* and *Sciaphila*

|                      | *Carludovica palmata* | *Sciaphila densiflora* | *Sciaphila thaidanica* |
|----------------------|-----------------------|------------------------|------------------------|
| Complete plastome (bp) | 158,545               | 21,485                 | 12,780                 |
| Coding (% of total)   | 91,183 (57.5)         | 14,769 (68.7)          | 10,856 (84.9)          |
| Introns (% of total)  | 20,174 (12.7)         | 1,998 (9.3)            | 894 (7.0)              |
| Genes (% of total)    | 111,357 (70.2)        | 16,767 (78.0)          | 11,750 (91.9)          |
| Intergenic (% of total) | 47,189 (29.8)   | 4,718 (22.0)           | 1,030 (8.1)            |

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### Table 2
Comparison of Plastome Genes Shared by *Sciaphila* and *Carludovica*

| Locus | *Carludovica* | *Sciaphila densiflora* | *Sciaphila thaidanica* |
|-------|---------------|------------------------|------------------------|
| accD  | 1,476         | 1,401                  | 1,179                  |
| rpl2  | 1,479         | 1,381                  | 1,173                  |
| rpl2 CDS | 825          | 810                    | 765                    |
| rpl14 | 369           | 369                    | 345                    |
| rpl16 | 1,358         | 976                    | 375                    |
| rpl16 CDS | 411          | 411                    | 375                    |
| rps2  | 714           | 714                    | 555                    |
| rps3  | 657           | 657                    | 612                    |
| rps4  | 606           | 594                    | 507                    |
| rps7  | 468           | 450                    | 321                    |
| rps8  | 399           | 402                    | 405                    |
| rps11 | 417           | 381                    | 357                    |
| rps12 | 915           | 850                    | 873                    |
| rps12 CDS | 369          | 369                    | 387                    |
| rps14 | 303           | 306                    | 316                    |
| rps18 | 306           | 252                    | 240                    |
| rps19 | 279           | 279                    | 291                    |
| rnr4.5| 103           | 108                    | 102                    |
| rnr5  | 121           | 121                    | 116                    |
| rnr16 | 1,491         | 1,486                  | 1,440                  |
| rnr23 | 2,811         | 2,816                  | 2,656                  |
| trnE  | 73            | 78                     | 75                     |
| trnFM | 74            | 74                     | 73                     |

*For Sciaphila, the percentage relative to Carludovica is given in parentheses.*

*The sum is 10,957, but due to overlap of several genes 101 bp are shared between two genes.*
tentaculata K. Larsen & Aver. (Thismiaceae), having retained only 12 plastome genes, a 5% overall reduction in coding length can be observed, but the reduction applies to only five genes while another five have actually increased in length (Lim et al. 2016).

The reduction of coding sequence in the retained genes, coupled with extensive sequence divergence (see Table 2), may indicate that some genes are no longer functional. However, if they are pseudogenized the reading frames would be expected to be even shorter. Most dubious in terms of functionality is rps14, which has been reduced to almost half the length compared with Carludovica palmata, and has a very low sequence similarity. With the exception of rps18, the genes of Sciaphila thaidanica are consistently more divergent than those of S. densiflora when compared with Carludovica palmata (Table 2) indicating a further progressed state of degradation.

It is, however, remarkable that the pairwise similarity between genes from the two species of Sciaphila is just as low as between genes from S. thaidanica and Carludovica palmata (Table 2) suggesting a very high degree of randomness in the evolution of the retained genes. Sequence data from additional species of Sciaphila are strongly needed for understanding this extraordinary divergence in an evolutionary perspective.

In S. densiflora Lam et al. (2015) calculated that most genes were still under strong purifying selection. Only three genes were under significantly relaxed selection (rps7, rpl14, clpP), but of these only clpP has been deleted in S. thaidanica and none of the other genes missing from the S. thaidanica plastome (matK, rpl20, rpl36) showed any sign of relaxation in S. densiflora. This may indicate that gene loss follows different routes in different evolutionary lineages of Sciaphila. The pronounced sequence divergence of S. thaidanica is unfortunately a hindrance to meaningful analysis of selection pressure on individual genes, as such analyses are very sensitive to correct alignment. Neither is it possible to test transcription as we do not have access to live plants or suitably preserved tissue from Sciaphila.

With a maximum of 20 functional genes, the plastome of S. thaidanica is in the final stage of degradation. Only in the mycoheterotroph Thismia tentaculata and the holoparasitic genus Pilostyles Gull. (Apodanthaceae) have plastomes with even fewer genes been described (Bellot and Renner 2015; Lim et al. 2016). The plastome of Sciaphila thaidanica is also the second smallest yet described; only the 11,348-kb plastome of Pilostyles aethiopica Welw. is smaller. With more than 50 fully mycoheterotrophic species of Triuridaceae and a crown age of at least 50 Ma (Mennes et al. 2013), this clade represents one of the oldest examples of photosynthesis loss. Hence, the Triuridaceae offers an outstanding framework for evolutionary studies of the final stages of plastome degradation.

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