Contrasting patterns of the 5S and 45S rDNA evolutions in the *Byblis liniflora* complex (Byblidaceae)

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Abstract To clarify the evolutionary dynamics of ribosomal RNA genes (rDNAs) in the *Byblis liniflora* complex (Byblidaceae), we investigated the 5S and 45S rDNA genes through (1) chromosomal physical mapping by fluorescence in situ hybridization (FISH) and (2) phylogenetic analyses using the nontranscribed spacer of 5S rDNA (5S-NTS) and the internal transcribed spacer of 45S rDNA (ITS). In addition, we performed phylogenetic analyses based on *rbcL* and *trnK* intron. The complex was divided into 2 clades: *B. aquatica*–*B. filifolia* and *B. guehoi*–*B. liniflora*–*B. rorida*. Although members of the complex had conservative symmetric karyotypes, they were clearly differentiated on chromosomal rDNA distribution patterns. The sequence data indicated that ITS was almost homogeneous in all taxa in which two or four 45S rDNA arrays were frequently found at distal regions of chromosomes in the somatic karyotype. ITS homogenization could have been prompted by relatively distal 45S rDNA positions. In contrast, 2–12 5S rDNA arrays were mapped onto proximal/interstitial regions of chromosomes, and some paralogous 5S-NTS were found in the genomes harboring 4 or more arrays. 5S-NTS sequence type-specific FISH analysis showed sequence heterogeneity within and between some 5S rDNA arrays. Interlocus homogenization may have been hampered by their proximal location on chromosomes. Chromosomal location may have affected the contrasting evolutionary dynamics of rDNAs in the *B. liniflora* complex.

Keywords *Byblis* · Chromosome · FISH · The 5S and 45S rDNAs · *rbcL* · *trnK* intron

Introduction The 5S and 45S ribosomal RNA genes (rDNAs) reside as independent tandem arrays in a plant genome. Their characteristic positions on chromosomes provide useful landmarks in chromosome study (Heslop-Harrison 2000). Physical mapping of the 5S and 45S rDNA provides universally applicable markers for chromosome and genome characterization in many plants. On the other hand, nucleotide sequences of the nontranscribed spacer of 5S rDNA (5S-NTS) and the internal transcribed spacer of 45S rDNA (ITS) are used globally as molecular markers for resolving species-level phylogenetic relationships (Persson 2000; Becerra 2003). Fluorescence in situ hybridization (FISH) detection and phylogenetic analysis allow us to learn the chromosomal-locus-level and nucleotide-sequence-level dynamics of these tandem arrays during diversification of the plant taxa. Chromosomal location of rDNA arrays has been suggested to affect the degree and speed of homogenization through concerted evolution...
(Wendel et al. 1995; Zhang and Sang 1999). Therefore, the combination of FISH and phylogenetic analyses may lead to a more sophisticated interpretation of rDNA evolution. However, chromosomal and molecular studies have often been carried out separately. This situation has masked evolutionary dynamics of rDNAs in plant taxa.

To trace the dynamics of 5S and 45S rDNA arrays in plant genomes, we investigated the chromosomal location and phylogenetic relationship of rDNAs in the *Byblis liniflora* complex. *Byblis* Salisb. (Byblidaceae) comprises seven species essentially native to Australia (Lowrie and Conran 2008). All members of the genus prey on small insects by aerial parts covered with secretory glands, which bear sticky droplets containing digestive enzymes (Bruce 1905; Hartmeyer and Hartmeyer 2005; Płachno et al. 2006). Their glistening appearance lends them the common name “the rainbow plants” (McPherson 2008). Currently recognized species fall into two complexes, the *B. gigantea* complex and the *B. liniflora* complex (Conran et al. 2002a). The former consists of temperate, perennial, shrubby plants distributed in South Western Australia, whereas the latter consists of tropical, annual, herbaceous plants mainly native to Northern Australia. Additionally, these complexes have been cytogenetically characterized by basic chromosome number: \(x = 9\) for the *B. gigantea* complex and \(x = 8\) for the *B. liniflora* complex (Conran et al. 2002a). The *B. liniflora* complex consists of three diploid species (*B. aquatica* Lowrie & Conran, *B. filifolia* Planch., and *B. rorida* Lowrie & Conran) and two tetraploid species (*B. liniflora* and *B. guehoi* Lowrie & Conran), whereas the *B. gigantea* complex comprises two diploids (*B. gigantea* Lindl. and *B. lamellata* Conran & Lowrie). *Byblis liniflora*, the representative member of the *B. liniflora* complex, is the most widespread species, being distributed up to New Guinea as well as to a wide area of northern Australia (Steenis 1971; Lowrie and Conran 1998). A seed fossil, which shows morphological resemblance to the *B. liniflora* complex, was discovered in clay lens from Eocene South America (Conran and Christophel 2004). Molecular phylogenetic studies have supported the placement of the family Byblidaceae in Lamiales (Albert et al. 1992; Müller et al. 2004). Despite the progress in phylogenetic investigations at higher taxonomic levels, the intrageneric relationships of *Byblis* have been less studied. The lack of a phylogenetic framework has made it difficult to understand the evolutionary history of this genus.

In this study, we investigated the 5S and 45S rDNA arrays in the *B. liniflora* complex through (1) chromosomal physical mapping by FISH and (2) phylogenetic analyses using 5S-NTS and ITS sequences. Furthermore, phylogenetic analysis using plastid sequences (*rbcL* and *tmK* intron) was used to gain further insight into the evolutionary history of the complex.

### Materials and methods

**Plant materials**

Six species were used in this study: one species from the *B. gigantea* complex (*B. gigantea*) and five species from all taxa in the *B. liniflora* complex (*B. aquatica*, *B. filifolia*, *B. guehoi*, *B. liniflora*, and *B. rorida*). Seeds and seedlings of *Byblis* species were purchased from commercial growers. Taxon names, cytotypes, voucher information, and GenBank accession numbers of DNA sequences presented in this study are given in Table 1. Voucher specimens were deposited in our laboratory of Tokai University. To test intraspecific variation, 3 to 5 accessions were analyzed for *B. aquatica*, *B. filifolia*, *B. liniflora* and *B. rorida*. *Byblis gigantea* and *B. guehoi* were sampled for single accession each. We sampled a total of 17 accessions from 6 species.

Seeds and shoot tips were surface sterilized with 1% benzalkonium chloride and with 1% sodium hypochlorite for 5 min each, repeatedly rinsed with sterilized water, and then sown on half-strength Murashige and Skoog medium (MS medium; Murashige and Skoog 1962) supplemented with 3.0% sucrose and 0.2% gellan gum. Established seedlings were maintained in vitro.

**DNA extraction, amplification, cloning, and sequencing**

Total genomic DNA was isolated from young leaves following modified cetyltrimethylammonium bromide method (Doyle and Doyle 1990). *rbcL* gene, *trnK* intron (including *matK* gene), ITS, and 5S-NTS (including 5S rDNA coding region) were amplified by polymerase chain reaction (PCR). The primers used are listed in Table 2. The cycle profile was an initial denaturation at 94°C (4 min); 35 cycles with 94°C (30 s), 57°C (30 s), and 72°C (60 s); and a final extension step at 72°C for 5 min. PCR products were column purified using QIAquick PCR Purification Kit (Qiagen, Tokyo, Japan) or gel purified using GeneClean III (BIO 101, Vista, CA, USA). The purified DNA was sequenced in both directions using ABI PRISM 310 DNA sequencer (Applied Biosystems, Foster City, CA, USA) with a BigDye Terminator Cycle Sequencing Kit (version 3.1, Applied Biosystems). When 3 or more arrays per somatic genomes were detected in rDNA-FISH analysis, the corresponding 5S-NTS and ITS were cloned into pGEM-T Easy vector (Promega Japan, Tokyo, Japan) and transformed into *Escherichia coli* DH5α competent cells. Recombinant clones were screened by colony direct PCR method, and were sequenced 3–10 clones per each sequence type using T7 (5’-TATACGACTCACCAGGG-3’) and SP6 (5’-TATTAGTGACACTAGAT-3’) primers. To exclude artifactual nucleotide variations generated in the PCR reaction, the polymorphism observed in
only 1 clone was removed from the analysis. The sequences reported in this paper were deposited in the GenBank database (Table 1).

DNA probes and labeling

To track the chromosomal locations of the 5S and 45S rDNA arrays, the entire region of the 5S rDNA unit and the 18S rDNA sequence were used as FISH probes, respectively. Primers used for PCR amplification are listed in Table 2. Sequence identities of the 5S and 18S rDNA fragments were confirmed by DNA sequencing (data not shown). To circumvent a contamination by dimeric or trimeric repeat sequences containing conservative 5S rDNA coding regions, 5S-NTS sequence type-specific probes were obtained by PCR amplification from clones. DNA fragments were biotin labeled through the random primed labeling technique (Feinberg and Vogelstein 1983) using Biotin-High Prime (Roche Applied Science, Indianapolis, IN, USA), or labeled in PCR with tetramethyl-rhodamine-5-dUTP (Roche Applied Science).

Chromosome preparation

Root tips that were 1–3 cm long were excised from in vitro plants and pretreated with 0.05% colchicine for 3–4 h at 18°C, fixed in 45% acetic acid, and then washed with distilled water. They were macerated in an enzymatic mixture containing 0.4% Cellulase Onozuka R-10 (Yakult Pharmaceutical Industry Co., Ltd., Tokyo, Japan) and 0.2% Pectolyase Y-23 (Kikkoman Co. Ltd., Tokyo, Japan) for 60 min at 37°C. After washing with distilled water, root tips were placed onto a glass slide and spread with ethanol–acetic acid (3:1). The slides were air dried for 40 h.

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**Table 1** Voucher accessions of *Byblis* species sampled in this study and GenBank accessions of *rbcL*, *trnK* intron, ITS and 5S-NTS sequences

| Species and cytotype | Voucher | GenBank accession |
|----------------------|---------|------------------|
|                      |         | *rbcL* | *trnK* intron | ITS | 5S-NTS |
| *Byblis aquatica*    | BaquKF01a, BaquKF02, BaquKF03 | AB546621b | AB546629b | GU810484c | AQU1: GU810492c |
| *B. filifolia*        | BfilKF03, BfilKF04a, BfilKF05, BfilKF01 | AB546622b | AB546630b | GU810485b | FIL-D1: GU810493c |
|                      |         |        |        |        | FIL-D2: GU810494c |
|                      |         |        |        |        | FIL-D3: GU810495c |
| *B. filifolia*        | BfilKF01a | AB546623b | AB546631b | GU810486c | FIL-T1: GU810496c |
|                      |         |        |        |        | FIL-T2: GU810497c |
|                      |         |        |        |        | FIL-T3: GU810498c |
| *B. guehoi*           | BgueKF01a | AB546624b | AB546632b | GU810488c | GUE1: GU810499c |
|                      |         |        |        |        | GUE2: GU810500c |
|                      |         |        |        |        | GUE3: GU810501c |
| *B. liniflora*        | BlinKF01, BlinKF02a, BlinKF03, BlinKF04 | AB546625b | AB546633b | GU810487b | LIN1: GU810502b |
|                      |         |        |        |        | LIN2: GU810503b |
|                      |         |        |        |        | LIN3: GU810504b |
| *B. rorida* cytotype A| BrorKF02a | AB546626b | AB546634b | GU810489b | ROR-A1: GU810505c |
| *B. rorida* cytotype B| BrorKF01, BrorKF04a | AB546627b | – | GU810490b | ROR-B1: GU810506c |
| *B. gigantea*         | BgigKF01a | AB546628b | AB546635b | GU810491c | GIG1: GU810507c |
| Antirrhinum majus      | –      | L11688e | AF051978e | – | – |
| Torenia baillonii     | –      | AB259805f | – | – | – |
| T. vagans             | –      | – | AF531812e |– | – |
| Buddleja asiatica     | –      | AJ001758h | – | – | – |
| B. alternifolia       | –      | – | AF531772e | – | – |
| Jasminum nudiflorum   | –      | NC_008407h | AF531779e | – | – |

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a All accessions of the *B. liniflora* complex were analysed in FISH mapping of the 5S and 45S rDNAs whereas some accessions, followed by a, were used for DNA sequencing of *rbcL*, *trnK* intron, ITS and 5S-NTS

b The sequences were obtained by PCR-direct sequencing in this study

c The sequences were obtained by PCR following cloning in this study

de The sequences were retrieved from Olmstead et al. (1993) d, Young et al. (1999) e, Higashiyama et al. (2006) f, Müller et al. (2004) g, Oxelman et al. (1999) h, and Lee et al. (2007) i
Fluorescence in situ hybridization

Chromosome preparation was treated with 100 µg ml⁻¹ RNase A (NIPPON GENE Co., Ltd., Tokyo, Japan) for 120 min at 37°C in a humid chamber. After dehydration in a graded series of ethanol, a hybridization mixture containing 50% formamide, 10% dextran sulfate and DNA probes was dropped onto the slides. The preparation was sealed, denatured for 3 min at 78°C, and then incubated for 16 h at 37°C. Subsequently, the slides were rinsed in 2× SSC at 42°C for 10 min, 0.2× SSC at 42°C for 10 min, and 2× SSC/0.2% Tween-20 at room temperature for 10 min twice. The slides were blocked with 5% bovine serum albumin in 2× SSC/0.2% Tween-20 for 60 min at 37°C. Biotin-labelled probes were detected with streptavidin-Alexa Fluor 488 (Invitrogen, CA, USA) in 2× SSC for 2 h at 37°C in a humid chamber. The slides were washed in 2× SSC/0.2% Tween-20 for 10 min twice, and 2× SSC for 10 min twice at room temperature. The preparations were then mounted in Vectashield mounting medium containing 1.5 µl gm l⁻¹ 4',6-diamidino-2-phenylindole (DAPI) (Vector Laboratories, Inc., CA, USA). Chromosome images were taken by a digital camera (CoolSNAP: Roper Scientific, Inc., Chiba, Japan) on a microscope (Olympus BX51; Olympus, Tokyo, Japan).

Phylogenetic analyses

Species of *Antirrhinum*, *Buddleja*, *Torenia*, and *Jasminum* were selected as outgroup taxa of plastid gene analysis based on previous broadscale phylogenetic analysis in Lamiales (Müller et al. 2004); their GenBank accession numbers are listed in Table 1. *rbcL*, *trnK* intron, ITS, and 5S-NTS nucleotide sequences were aligned using the

### Table 2 Primers used for PCR amplification, DNA sequencing and FISH probe preparation

| Region     | Primer name | Primer sequence (5' to 3')                               | Design          |
|------------|-------------|----------------------------------------------------------|-----------------|
| *rbcL*     | aF<sup>a</sup> | ATGTCACCACAAACAGAGACTAAAGC | Hasebe et al. (1994) |
|            | aR<sup>b</sup> | CTTCTGCTCAAATAAGAATCTCCTCCCA | Hasebe et al. (1994) |
|            | cF<sup>b</sup> | TGAAAACGTGAATTCCCAACCGTTATGCG | Hasebe et al. (1994) |
|            | cR<sup>a</sup> | GCAGCAGCTGTGTCGGGCTCCA | Hasebe et al. (1994) |
|            | dF<sup>b</sup> | CTTTAGGCACTTTTTGGAAGGATCTGTT | This study |
|            | dR<sup>b</sup> | CCTCAAAGTTTACCTACGTTACCAGAG | This study |
| *trnK*      | MG1<sup>a</sup> | CTACTGCAAGAATGCGATAGAGAT | Liang and Hilu (1996) |
|            | MG15<sup>a</sup> | ATCTCGGTTGCTAATTCATAATG | Liang and Hilu (1996) |
|            | BtrnK-1<sup>b</sup> | ACAAGAAGTGGAGATATACCTAATTACGCAATTCG | This study |
|            | BtrnK-2<sup>b</sup> | GTTCTAGCACAAGAAGTCCGAGTGTTATATTGAT | This study |
|            | BtrnK-3<sup>b</sup> | CTGTCTGGGAGGTATTCTCTTATATTGAT | This study |
|            | BtrnK-4<sup>b</sup> | CTTTAGGCACTTTTTGGAAGGATCTGTT | This study |
| *ITS*       | ITS5<sup>a</sup> | GGAAGTAAAGTCTGTAACAGGAAGG | White et al. (1990) |
|            | ITS4<sup>a</sup> | TCCTCGCATTGATATGTCGATG | White et al. (1990) |
| 5S rDNA     | 5SF<sup>a,d</sup> | CGGTCATTAATGCTGGATAT | Hizume (1993) |
|            | 5SR<sup>a,d</sup> | CCATCGAAGACTCCGAGGTA | Hizume (1993) |
|            | FIL-T2-1<sup>d</sup> | AACCAGGCGCCATCATCATCATGATATGTCG | This study |
|            | FIL-T2-2<sup>d</sup> | GTGGACCGCCTGTCCTTACATGATATGTCG | This study |
|            | FIL-T3-1<sup>c,d</sup> | ATCCGTCGTCCGAGGTAATGTCG | This study |
|            | FIL-T3-2<sup>c,d</sup> | TACTGCGGGAAGGAGGATGTCG | This study |
|            | LIN2-1<sup>c,d</sup> | ACTAGTGGGATGTCGAGGGAAGG | This study |
|            | LIN2-2<sup>c,d</sup> | TCGGTTGCGCGGAGGAAATGTA | This study |
| 18S rDNA    | 18SF<sup>d</sup> | AACCTGTTGATCTCGAGGGAAGG | Sogin (1990) |
|            | 18SR<sup>d</sup> | TGATCCTTCTCGAGGTTACCAT | Sogin (1990) |

<sup>a</sup> Primers used for PCR amplification prior to DNA sequencing  
<sup>b</sup> Internal primers used to fill gaps between sequences which were too short to overlap  
<sup>c</sup> Primers used to obtain cryptic 5S-NTS paralogues which were not represented in 5S-NTS clones come from PCR amplification using 5SF and 5SR primer pair  
<sup>d</sup> Primers used for FISH probe preparation
Clustal X program (Thompson et al. 1994) implemented in MEGA4 (Tamura et al. 2007) and then refined by eye. Gaps and ambiguous sites were eliminated from datasets. Following phylogenetic analyses were separately performed for rbcL, trnK intron, ITS and 5S-NTS matrices. In addition, to enhance the reliability of plastid tree topology, the conjugate matrix of rbcL and trnK intron was analyzed. The congruence between rbcL and trnK matrices was confirmed using the incongruence length difference test ($P = 0.22$, Farris et al. 1994).

To independently test the reliability of the tree topology, phylogenetic analyses were performed under maximum parsimony (MP) analysis, maximum likelihood (ML) analysis, and Bayesian inference (BI). The MP tree was constructed using the close-neighbor-interchange search implemented in MEGA4, in which the search level was 3 and the initial trees were obtained with the random addition of sequences (10 replicates). The bootstrap percentages (BP; Felsenstein 1985) for the MP trees were calculated for 1,000 replicates. For ML and BI searches, a best-fit model of nucleotide substitution was determined based on the evaluation of each dataset under AIC and hLRT criteria using MrModeltest (Nylander 2004). For rbcL, trnK intron, rbcL + trnK intron, ITS, and 5S-NTS datasets, HKY85 + I+G, GTR+I+G, GTR+I+G, GTR+I+G, and K80+I+G models were optimized, respectively. ML trees were obtained using PhyML 3.0 (Guindon and Gascuel 2003). Topology search was carried out under the “Best of NNIs and SPRs” option with 10 random additions. Support for nodes was estimated by bootstrapping with 1,000 replications. BI was carried out by Metropolis-coupled Markov-chain Monte Carlo (MC$^3$) analysis using MrBayes 3.1.2 (Ronquist and Huelsenbeck 2003). Topology search was performed by running four simultaneous chains initiated from random trees for 1,000,000 generations (heating parameter = 0.2), in which trees were sampled every 100 generation. At the end of the run, convergence was evaluated by visual inspection of a graph of likelihood as a function of generation. Because a conservative burn-in period was identified for 250,000 generations, the first 2,500 trees were discarded. A majority-rule consensus tree with posterior probabilities (PP) calculated for each clade was yielded from the 7,500 post-burn-in trees.

Results

Chromosome number

Table 3 summarizes the chromosomal characteristics of the B. liniflora complex at mitotic metaphase. Intraspecific variations of chromosome number were observed among accessions of B. filifolia: 4 accessions had a chromosome number of 2n = 16 (diploid cytotype), whereas 1 accession exhibited 2n = 32 (tetraploid cytotype). The other chromosome numbers are 2n = 16 for B. aquatica and B. rorida, and 2n = 32 for B. liniflora and B. guehoi.

Chromosomal distribution of 5S and 45S rDNAs

Figure 1 shows the results of double-target FISH using the 5S and 45S rDNA probes to mitotic metaphase of the B. liniflora complex. All species were distinguishable in terms of FISH patterns of rDNAs. The number and distribution of 5S and 45S rDNAs are summarized in Table 3. Two signals of the 45S rDNA probe were observed in B. filifolia diploid cytotype, B. liniflora, and B. rorida, whereas four arrays were detected in B. aquatica, B. filifolia tetraploid cytotype, and B. guehoi. Two distal 45S rDNA signals of B. aquatica were strong as seen in those of other species, but the signals on the other two interstitial arrays were very weak. The 45S rDNA signals were located at secondary constrictions, except for 2 minor signals in B. aquatica. The 5S rDNA arrays were detected in 2 chromosomes of B. aquatica, 4 chromosomes of B. filifolia diploid cytotype, 8 chromosomes of B. filifolia tetraploid cytotype, 10 chromosomes of B. liniflora, and 12 chromosomes of B. guehoi. In B. rorida, 1 accession had 2 arrays whereas the other 2 accessions harbored 3 arrays. Hereafter, the former was designated as cytotype A, and the latter as cytotype B. In terms of proximo-distal (centromeric–telomeric) distribution of rDNA arrays, 5S and 45S rDNA arrays tended to occupy the proximal and distal positions, respectively.

Table 3 Karyotype characteristics in the B. liniflora complex

| Species and cytotype       | Chromosome number | rDNA number and distribution$^a$ |
|----------------------------|-------------------|----------------------------------|
|                            |                   | 5S | 45S                          |
| B. aquatica                | 2n = 16           | 2P | 2D + 2I                      |
| B. filifolia diploid cytotype | 2n = 16           | 4P | 2D                           |
| B. filifolia tetraploid cytotype | 2n = 32           | 4P + 4I | 4P                        |
| B. guehoi                  | 2n = 32           | 12P | 4D                        |
| B. liniflora               | 2n = 32           | 10P | 2D                        |
| B. rorida cytotype A       | 2n = 16           | 2P | 2D                        |
| B. rorida cytotype B       | 2n = 16           | 3P | 2D                        |

$^a$ rDNA position on a chromosome is classified as follow: P, proximal location; I, interstitial location; D, distal location
Because an odd number of 5S rDNA arrays was observed in *B. rorida* cytotype B, assigning pairs of homologous chromosomes became difficult. Hence, the number of rDNA arrays per somatic chromosome complement but not at the haploid level is mentioned in the following sections.
Plastid dataset and tree topology

Cytogenetic screening revealed interspecies and intraspecies karyotype variations that guided the successive phylogenetic analyses. A representative 1 accession for each species and cytotype was analyzed in phylogenetic analyses. Unfortunately, we failed to obtain the PCR product of *trnK* intron from *B. rorida* cytotype B. Thus, this cytotype was excluded from the analyses of *trnK* intron and conjugated matrices. Figure 2 shows plastid gene phylogenies of *Byblis* and related genera. Although separate analyses of *rbcL* and *trnK* intron datasets did not show any topological incongruence in ingroup taxa, these datasets gave rise to less-resolved topology (Fig. 2a, b). The conjugated matrix of *rbcL* and *trnK* intron sequences resulted to a single most parsimonious tree with improved resolution (Fig. 2c). ML and BI trees resulting from the conjugated matrix exhibited essentially the same topology in comparison with the MP tree (data not shown).

Plastid trees supported monophyly of not only *Byblis* but also the *B. liniflora* complex, with high confidence. In ingroup taxa, *B. gigantea* was the sister to five species of the *B. liniflora* complex. In the conjugated tree, the *B. liniflora* complex was divided into two clades, *B. aquatica*–*B. filifolia* and *B. guehoi*–*B. liniflora*–*B. rorida*, although the latter clade received low supports (BP in MP/BI in ML/PP in BI = 62/-/50). *Byblis filifolia* diploid and tetraploid cytotypes formed a well-supported clade (100/100/100), and this pair was sister to *B. aquatica* (100/100/100).

Two tetraploid species, *B. guehoi* and *B. liniflora*, formed a clade with high support values (100/100/100). *Byblis rorida* cytotype A and B were indicated as sister lineage with a weak support in ML analysis of the *rbcL* dataset (BP = 51, data not shown), but MP and BI analyses did not resolve this clade.

Characterization of 5S rDNA sequences

By PCR amplification with universal primers (5SF and 5SR) following cloning, 1 sequence type of 5S-NTS was detected in *B. aquatica* (AQU1), *B. gigantea* (GIG1), *B. rorida* cytotype A (ROR-A1), and *B. rorida* cytotype B (ROR-B1); 2 sequence types in *B. filifolia* diploid cytotype (FIL-D1 and FIL-D2) and *B. guehoi* (GUE1 and GUE2); and 3 sequence types in *B. filifolia* tetraploid cytotype (FIL-T1, FIL-T2, and FIL-T3) and *B. liniflora* (LIN1, LIN2, LIN3) (Fig. 3b). Since considerable polymorphisms within and among species were observed, we carried out a detailed characterization of 5S-NTS. We designed sequence type-specific primer pairs, FIL-T2-1 and FIL-T2-2 for FIL-T2, FIL-T3-1 and FIL-T3-2 for FIL-T3, and LIN2-1 and LIN2-2 for LIN2 (Fig. 3a; Table 2). The presence of FIL-T2-like and FIL-T3-like sequences in

Fig. 2 Plastid gene phylogenies of *Byblis* and its related taxa. a Strict consensus tree of 30 most parsimonious trees generated by analysis of the *rbcL* data matrix composed of 1,209 nucleotide positions (45 sites are parsimony informative). The consistency index (CI) is 0.750 and the retention index (RI) is 0.843. b Strict consensus tree of 2 most parsimonious trees yielded by the *trnK* intron dataset consisted of 2,266 positions (263 sites are parsimony informative). CI and RI are 0.797 and 0.863, respectively. c Single most parsimonious tree resulting from the *rbcL* + *trnK* intron conjugate dataset composed of 3,475 positions (308 sites are parsimony informative). CI and RI are 0.782 and 0.853, respectively. Numbers above branches are bootstrap percentages in MP/ML analyses. Bayesian posterior probabilities are indicated below the branches. A minus sign (−) denotes a clade not supported in the respective analysis.
cytotype as FIL-D3. In addition, the LIN2-like sequence in *B. guehoi* was distinguishable from GUE1/GUE2 and therefore referred to as GUE3.

To localize specific 5S-NTS sequence types on chromosomes, we then carried out FISH analysis using FIL-T2-specific probe (FIL-T2sp), FIL-T3-specific probe (FIL-T3sp), and LIN2-specific probe (LIN2sp) (Fig. 3c–i). Based on sequence similarity, FIL-T2sp is predicted to recognize FIL-D1 and FIL-D2 in addition to FIL-T2 (Fig. 3a). Similarly, FIL-T3sp presumptively detects FIL-D3 in addition to FIL-T3, and LIN2sp seems to recognize GUE3, LIN2, LIN3, ROR-A1, and ROR-B1 (Fig. 3a).
These sequence type-specific probes were hybridized to chromosome spreads of *B. aquatica* as a negative control, showing no apparent signal (Fig. 3c). FIL-T2sp hybridized to all arrays of *B. filifolia* diploid and tetraploid cytotypes (Fig. 3d), while FIL-T3sp specifically recognized 2 out of 4 and 4 out of 8 arrays, respectively (Fig. 3e). FIL-T3sp signals in the diploid cytotype were detected on 2 chromosomes carrying secondary constriction (the 45S rDNA sites) (Fig. 3d, arrows and arrowheads). Contrarily, in the tetraploid cytotype, the signals were observed on 4 chromosomes without secondary constriction (Fig. 3e, arrows and arrowheads). These data suggested interlocus and intralocus heterogeneity of the 5S rDNA arrays in *B. filifolia* diploid and tetraploid cytotypes. LIN2sp was detected as two major sites along with some faint signals in *B. guelhoi* and *B. liniflora* (Fig. 3f, g), suggesting interlocus heterogeneity. LIN2sp also recognized the 5S rDNA arrays of *B. rorida* cytotypes A and B (Fig. 3h, i).

### 5S-NTS dataset and tree topology

In this study, a total of 83 5S-NTS clones were sequenced and condensed into the 16 sequence types from 8 taxa. 5S-NTS length was quite variable from 294 bp in AQU1 to 793 bp in GIG1, owing to the presence of large indels that markedly reduced the amount of sequence available for comparison (Fig. 3a). A large fraction of NTS was inevitably discarded in phylogenetic analyses. The 5S-NTS dataset used for phylogenetic analyses contained partial 5S rRNA coding region (65 positions) and 5S-NTS (77 positions). Because of the difficulty in sequence alignment, outgroup taxa in plastid dataset could not be included to the 5S-NTS and ITS datasets. Alternatively, the trees were rooted with *B. gigantea* based on of plastid tree topology.

Maximum parsimony analysis generated a single most parsimonious tree (Fig. 4). MP, ML, and BI analyses yielded roughly congruent trees (data not shown). 5S-NTS sequences were aligned, condensed into the 16 sequence types from 8 taxa. 5S-NTS length was quite variable from 294 bp in AQU1 to 793 bp in GIG1, owing to the presence of large indels that markedly reduced the amount of sequence available for comparison (Fig. 3a). A large fraction of NTS was inevitably discarded in phylogenetic analyses. The 5S-NTS dataset used for phylogenetic analyses contained partial 5S rRNA coding region (65 positions) and 5S-NTS (77 positions). Because of the difficulty in sequence alignment, outgroup taxa in plastid dataset could not be included to the 5S-NTS and ITS datasets. Alternatively, the trees were rooted with *B. gigantea* based on of plastid tree topology.

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sequence types in the *B. liniflora* complex were divided into two major clades as seen in plastid phylogenies: *B. aquatica*– *B. filifolia* (73/84/92) and *B. guehoi*– *B. liniflora*– *B. rorida* (97/51/99). In the MP tree, AQU1 was nested among 5S-NTS sequence types from *B. filifolia* with a low support (54/-/-); however, this clade did not appear in the ML and BI trees. Alternatively, in ML and BI analyses, 5S-NTS sequence types from *B. filifolia* diploid and tetraploid cytotypes formed a monophyletic clade (BP in ML/PP = 52/67; not shown). FIL-D3, which was isolated as a FIL-T3-like sequence, was predictably sister to FIL-T3 (91/87/93). On the other hand, ROR-A and ROR-B were nested in 5S-NTS sequence types from *B. guehoi* and *B. liniflora* with low confidences (61/40/52). Despite its original isolation as a LIN2-like sequence, GUE3 was sister to LIN3 (77/85/99).

### ITS dataset and tree topology

The length of ITS sequences in datasets ranged from 602 to 654 bp. *Byblis aquatica*, *B. filifolia* tetraploid cytotype, and *B. guehoi*, which harbored four 45S rDNA arrays, did not show any polymorphic sequence in an examination of clones (approximately 20 clones each were examined in colony PCR, and 5, 8, and 12 clones were sequenced, respectively). These data indicate an almost completion of interlocus homogenization of the 45S rDNA arrays in these taxa.

A single most parsimonious tree was obtained from this set of sequences (Fig. 4). MP, ML, and BI tree topologies were in full accordance with each other except for a collapsed clade of *B. guehoi* and *B. liniflora* in Bayesian analysis. Relationships in the *B. liniflora* complex indicated by the ITS tree were essentially identical to those of plastid and 5S-NTS phylogenies (Figs. 2, 4).

### Discussion

#### Phylogenetic framework and morphological traits in the *B. liniflora* complex

In this study, we used maternally inherited plastid sequences (*rbcL* and *trnK* intron; Fig. 2) and biparentally inherited nuclear rDNA sequences (ITS and 5S-NTS; Fig. 4) for phylogenetic analyses. Comparing plastid and rDNA phylogenies, we found the topologies to be largely congruent. *Byblis filifolia* diploid and tetraploid cytotypes were suggested to be sister lineages. Morphological variations in *B. filifolia* are known to exist, for instance, flower size and petal color (Lowrie and Conran 1998; McPherson 2008). Among samples we analyzed, plants of the diploid cytotypes had navy-colored stigma and mauve-striped yellowish abaxial petal. In contrast, the tetraploid plants had white-colored stigma and whitish to yellowish abaxial petal. Extended sampling is needed to clarify the relationships between these morphological and cytogenetic characteristics in the *B. filifolia* lineage.

An affinity of *B. filifolia* and *B. aquatica* was suggested by both plastid and rDNA phylogenies. Their reproductive isolation was previously verified by a crossing experiment (Meyers-Rice 1993). Previous reports have suggested that seed morphology is a useful characteristic to define the different taxa in *Byblis* (Lowrie and Conran 1998, 2008; Conran et al. 2002b). These two species share relatively smooth anticlinal sculpturing of seeds presumably as a synapomorphy.

On the other hand, an association of *B. guehoi* and *B. liniflora* was observed. These two species share a putative synapomorphic structure—pedicels that are longer than leaves (Lowrie and Conran 2008). *Byblis rorida* was sister to the *B. liniflora*–*B. guehoi* clade in plastid and rDNA trees. The three species share a morphological trait—denticulate sculpturing of seed coat.

#### Tetraploidizations in the *B. liniflora* complex

The chromosome counts in this study were essentially consistent with previous works (Peng and Kenton 1982; Conran et al. 2002a; Hoshi et al. 2007; Lowrie and Conran 2008; Fukushima et al. 2008). However, *B. filifolia*, which is thus far recognized as a diploid species, showed a tetraploid chromosome number along with the diploid cytotype in the accessions we examined (Table 1). FISH analysis indicated the autoploidal karyotype for the tetraploid cytotype (Fig. 1), providing a further support for polyploidization within the lineage of *B. filifolia*. Inverted positional relationship between the 5S and 45S rDNA arrays on a chromosome arm of *B. filifolia* diploid and tetraploid cytotypes suggested the occurrence of a paracentric inversion at this chromosome arm (Fig. 1d, f). Although compositional discordance of 5S-NTS between the loci of diploid and tetraploid cytotypes observed in FISH with FIL-T3sp indicated unexplained complexity of the karyotype evolution in the taxa (Fig. 3d and e), the chromosome rearrangement would occur before tetraploidization because all four chromosomes carrying both 5S and 45S rDNA arrays in the tetraploid cytotype exhibited identical FISH pattern (Fig. 1f).

Peng and Kenton (1982) found the lack of multivalent formation during meiosis in *B. liniflora*, suggesting a functionally diploidal nature of this tetraploid species. Ribosomal DNA distribution in *B. liniflora* did not represent the exact doubling of any diploid species (Fig. 1) and might help hamper multivalent formation. In phylogenetic analyses, *B. guehoi* and *B. liniflora* formed a clade, and the
pair was sister to *B. rorida* (Figs. 2, 4). A candidate for at least 1 of the parental genome donor of these tetraploids is *B. rorida*-like species or its ancestor.

### 5S and 45S rDNA variabilities at the chromosomal distribution level

The copy number and chromosomal location of rDNA could be rapidly changed in plant genomes (Schubert and Wobus 1985; Raina and Mukai 1999). Schubert and Wobus (1985) postulated that the mobility of rDNAs in *Allium* is rendered by an adjacent transposable element or by recombination hot spots in the terminal heterochromatin blocks. In some members of Poaceae, *Enhancer/Supressor-mutator*-like (*En/Spm*-like) transposons and Ty3-gypsy retrotransposons localize inside rDNA cluster (Raskina et al. 2004; Belyayev et al. 2005; Altinkut et al. 2006). Transposase activity of *En/Spm*-like transposons increases during meiosis and seems to contribute to rDNA migration in *Aegilops speltoides* (Raskina et al. 2004; Altinkut et al. 2006). Similarly, in *Nemesia* and *Oryza*, the key mechanism facilitating diversification of rDNA distribution patterns is considered as transpositions rather than chromosome rearrangements (Datson and Murray 2006; Chung et al. 2008). FISH mapping revealed interspecies- and intraspecies-specific rDNA distribution in the *B. liniflora* complex (Fig. 1) without deviation of chromosome morphology (chromosome length and arm ratio; data not shown). Taking conservative karyotypes into account, a large contribution of transpositions to diversification of rDNA distribution patterns in the complex is conceivable. The copy number of migrated rDNA repeats would be amplified by unequal crossing over to the extent that these new sites can be detected by FISH. The array number of 5S rDNA (2–12 arrays) was more flexible compared to that of 45S rDNA (2–4 arrays) (Fig. 1 and Table 3). In the *B. liniflora* complex, generation of a novel locus via transposition may be more frequent in 5S rDNA than in 45S rDNA. Alternatively, amplification or maintenance of copy number within a novel array is more effective in the 5S rDNA.

Chromosomal rDNA distribution often varies among closely related taxa, even within species (Garrido et al. 1994). *Byblis rorida* exhibited intraspecies polymorphisms of the 5S rDNA distribution: cytotype A with 2 arrays and cytotype B with 3 arrays (Fig. 11, n). We could not find any 5S-NTS polymorphism in *B. rorida* cytotype B (12 clones were examined in colony PCR and 8 clones were sequenced) (Fig. 4). Thus, most of the units in a surplus 5S rDNA array in cytotype B seem to be homogenized with the units of the other 2 arrays. Vaio et al. (2005) observed chromosomal polymorphisms in a population of *Paspalum quadrifarium*, and attributed the heterozygous karyotype with the odd number of 45S rDNA arrays to self-incompatibility in this species. The odd number of rDNA arrays was also found in *Brachyscome lineariloba* complex cytode E, which transmits specific chromosomes to progeny through microspores but not through megaspores (quasi-diploid; Carter et al. 1974; Adachi et al. 1997), similar to the permanent odd polyploid *Rosa canina* (Lim et al. 2005; Kovarik et al. 2008). Karyotype screening among populations and an investigation on reproductive strategy of *B. rorida* can provide an insight into the odd number of the 5S rDNA arrays.

In polyploid species, parental rDNA loci follow different evolutionary paths. The sum of the parental 45S rDNA arrays has been observed in various allopolyploid species, such as *Oryza punctata* (Chung et al. 2008). On the other hand, the decrease in the expected number of arrays has been observed in *O. minuta* and *O. latrifolia* (Chung et al. 2008). Likewise, a number of the 5S rDNA loci were conservative in some taxa (e.g., *Sanguisorba*, Mishima et al. 2002) but variable in others (e.g., *Iris versicolor*, Lim et al. 2007). At a glance, rDNA arrays seem to behave arbitrarily during the evolutionary history of the polyploid species. Differing from the estimation based on its tetraploid chromosome number, two 45S rDNA arrays were detected in *B. liniflora* (Fig. 1j), suggesting complete or almost complete deletion of at least 2 arrays (Fukushima et al. 2008). On the other hand, *B. filifolia* tetraploid cytotype and *B. guehoi* retained 4 arrays (Fig. 1f, h). Although an apparent decrease of the 5S rDNA array number was not observed in 3 polyploids (Fig. 1f, h, j), the lack of information on parental genome obscured the evolutionary dynamics of rDNA site number in polyploid species.

### 5S and 45S rDNA variabilities at the nucleotide sequence level

A growing body of evidence indicates that there are three evolutionary patterns of different rDNA sequences separated by chromosomal location: (1) be retained and evolve independently without interaction, (2) be lost either through loss of entire array or through interlocus homogenization, and (3) be replaced by another rDNA type (Bao et al. 2010). We found one ITS sequence in *Byblis* species with two or four 45S rDNA arrays (Fig. 4). In contrast, multiple 5S-NTS sequence types were detected in species harboring 4 or more arrays (Fig. 4). Although orthology-paralogy relationships of 5S-NTS were largely obscured by extensive sequence modification in the *B. liniflora* complex (Fig. 3a), phylogenetic relationships suggested by the 5S-NTS tree were largely congruent with plastid and ITS phylogenies (Fig. 4). Based on our phylogenetic analysis, 5S-NTS polymorphisms within a genome would occur after the separation of *B. aquatica*–*B. filifolia* and
B. guehoi–B. liniflora–B. rorida clades. Although network or phylogenetic analysis of entire sequences is not applicable due to a large amount of missing positions (Fig. 3a), 5S-NTS variants within a genome seem to be generated via recombination events between different sequence types. For example, the 5' side of LIN2 is similar to that of LIN1, whereas the 3' side of LIN2 resembles LIN3 (Fig. 3a). Likewise, the 5' and 3' sides of GUE2 had similarity with GUE3 and GUE1, respectively (Fig. 3a). These data indicated 5S-NTS sequence-type diversification in a genome via recombination events. FISH analysis suggested that sequence types recognized by FIL-T2sp and FIL-T3sp coexisted within an array (Fig. 3d, e). Colocalization of sequence types recognized by FIL-T2sp and FIL-T3sp occurred due to a large amount of missing positions (Fig. 3a), whereas the 3S-NTS variants within a genome seem to be generated via recombination between different sequence types. FISH analysis suggested that sequence types recognized by FIL-T2sp and FIL-T3sp coexisted within an array (Fig. 3d, e). Colocalization of polymorphic sequences may facilitate the generation of novel sequence types in the B. liniflora complex.

PCR selection and variation in copy number among polymorphic sequences present in a genome lead to a biased representation of different paralogues (Wagner et al. 1994; Alvarez and Wendel 2003). Initially, 2 sequence types of 5S-NTS were found in B. filifolia diploid cytotype (15 clones were examined in colony PCR and 6 clones were sequenced), whereas 3 types were observed in the tetraploid cytotype (15 clones were examined in colony PCR and 13 clones were sequenced) (Fig. 3b). Using sequence type-specific primers, we found additional sequence type, FIL-D3, in the diploid cytotype (Fig. 3b). Sequence type-specific FISH analysis indicated an abundance of FIL-D3 in the 5S rDNA arrays of the diploid cytotype (Fig. 3d). GUE3, which was not represented in 5S-NTS pool of B. guehoi amplified with universal primers (36 clones were examined in colony PCR and 12 clones were sequenced) (Fig. 3b), was also detected by FISH using LIN2sp (Fig. 3f). Our results underline the risk in sequence analysis exclusively depending on universal primers.

Relationships between chromosomal distribution and sequence variation of 5S and 45S rDNAs

In cytogenetically well-characterized Gossypium and Nicotiana, 45S rDNA but not 5S rDNA is homogenized between loci in polyploid species (Wendel et al. 1995; Cronn et al. 1996; Fulneek et al. 2002; Kovarik et al. 2004;). The 5S rDNA loci are located at the proximal region, whereas the 45S rDNA loci occupy the terminal locations in these two genera (Hanson et al. 1996; Fulneek et al. 2002; Kovarik et al. 2004). The mechanism facilitating interlocus homogenization would be crossing over between rDNA repeats located on different chromosomes (Wendel et al. 1995). When crossing over occurs between proximally located arrays, inviable gametes could be generated more frequently. In contrast, telomeric or subtelo-meric locations may allow interlocus crossing over without deleterious effect. After interlocus crossing over is established, intralocus homogenization would be completed via successive intralocus unequal crossing over. Under these assumptions, the chromosomal location has a substantial impact on the tempo of interlocus concerted evolution of tandemly repeated DNA elements (Zhang and Sang 1999). Our results in the B. liniflora complex appear to meet the above hypothesis. In all taxa we examined, other than B. filifolia tetraploid cytotype, the 45S rDNA arrays were more distal relative to the 5S rDNA arrays (Fig. 1; Table 3). ITS sequences seem to be well homogenized in the genome harboring four 45S rDNA arrays (Fig. 4). In B. guehoi and B. aquatica, ITS homogenization could have been prompted by relatively distal 45S rDNA positions. We believed that ITS uniformity in B. filifolia tetraploid cytotype may stem from a presumptive autotetraploid origin. In B. liniflora, ITS can be homogenized through the entire locus loss of the 45S rDNA array. In contrast, the genomes harboring 4 or more 5S rDNA arrays surely had polymorphic 5S-NTS sequence types (Fig. 4). FISH with sequence type-specific probes indicated interlocus and intralocus heterogeneity in some 5S rDNA tandem repeats (Fig. 3d–g). Together, these data indicate that, in the B. liniflora complex, the homogenization of the 5S rDNA could not be completely accomplished within the genomes with multiple 5S rDNA arrays. Interlocus homogenization may have been hampered by their proximal location on chromosomes. Chromosomal position may have affected the contrasting evolutionary dynamics of rDNAs in the B. liniflora complex.

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