Research Article

Conflicting phylogenetic signals in genomic data of the coffee family (Rubiaceae)

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Received 30 August 2019; Accepted 2 January 2020; Article first published online 3 January 2020

Abstract Reconstructions of phylogenetic relationships in the flowering plant family Rubiaceae have up until now relied heavily on single- or multi-gene data, primarily from the plastid compartment. With the availability of cost- and time-efficient techniques for generating complete genome sequences, the opportunity arises to resolve some of the relationships that, up until now, have proven problematic. Here, we contribute new data from complete 58 plastid genome sequences, representing 55 of the currently 65 recognized tribes of the Rubiaceae. Also contributed are new data from the nuclear rDNA cistrons for corresponding taxa. Phylogenetic analyses are conducted on two plastid data sets, one including data from the protein coding genes only, and a second where protein coding data are combined with non-coding regions, and on a nuclear rDNA data set. Our results clearly show that simply adopting a “more characters” approach does not resolve the relationships in the Rubiaceae. More importantly, we identify conflicting phylogenetic signals in the data. Analyses of the same plastid data, treated as nucleotides or as codon-degenerated data, resolve and support conflicting topologies in the subfamily Cinchonoideae. As these analyses use the same data, we interpret the conflict to result from erroneous assumptions in the models used to reconstruct our phylogenies. Conflicting signals are also identified in the analyses of the plastid versus the nuclear rDNA data sets. These analyses use data from different genomic compartments, with different inheritance patterns, and we interpret the conflicts as representing “real” conflicts, reflecting biological processes of the past.

Key words: coffee family, conflicting signals, phylogenomics, plastid DNA, rDNA, Rubiaceae.

1 Introduction

The flowering plant family Rubiaceae, or the coffee family, comprises close to 14 000 species, and recent classifications recognize somewhere around 600 genera, 65 tribes, and three different subfamilies (Bremer et al., 1999; Bremer & Eriksson, 2009; Davis et al., 2009; Rydin et al., 2009a; Kainulainen et al., 2013; Mouly et al., 2014; Govaerts et al., 2018). Representatives of this species-rich family can be found in all regions of the world, even on the subantarctic islands, but most occur in tropical and subtropical regions across the globe (Govaerts et al., 2018). Also from ecological, morphological, and functional perspectives, the diversity of the family is considerable: life forms span from small annual and perennial herbs to large tropical trees; flowers are adapted to a broad range of different pollinators, including bats (Sazima et al., 1999); fruit types include animal dispersed fleshy fruits to dry fruits such as nuts and capsules, the latter often with small and wind dispersed seeds (Robbrecht, 1988; Eriksson & Bremer, 1991); epiphytes, lianas, succulents, rheophytes, and aquatic life forms can be found; and members of the family inhabit a wide range of different habitats, from dry dessert like conditions to wet tropical rain forests (Robbrecht, 1988).

Over the last 20–30 years, a large number of papers have been published with the primary focus of resolving phylogenetic relationships in Rubiaceae at different taxonomic levels (see Robbrecht & Manen, 2006; Bremer, 2009 for reviews of work up until 2009). This work has resulted in a reasonably well-resolved and coherent picture of the phylogenetic relationships in the family (Fig. 1). Two early diverging groups are consistently recognized and supported (e.g., Bremer et al., 1995; Andersson & Rova, 1999; Robbrecht & Manen, 2006; Bremer & Eriksson, 2009; Rydin et al., 2009a; Wikström et al., 2015). The first corresponds to the subfamily
Fig. 1. Continued
Rubioideae, the largest of three subfamilies, with almost 8000 species and 27 different tribes. The second comprises the other two subfamilies: Cinchonoideae with about 1700 species and 9 (or 10) recognized tribes; and Ixoroideae with about 4000 species and 26 recognized tribes. The latter two subfamilies are sometimes united as one (Cinchonoideae s. lat. Robbrecht & Manen, 2006). More detailed analyses of the Rubioideae resolve the subfamily in two informal alliances: the Spermacoceae alliance (with 10 tribes recognized by Rydin et al. (2009b) and two additional tribes described more recently by Wen & Wang (2012) and Ginter et al. (2015)); and the Psychotrieae alliance (with 9 tribes recognized by Razafimandimbison et al., 2014, 2017), each alliance including more than 3000 species. Sister to the two alliances are Coussareaeae (414 species) and a grade of tribes including Lasiantheae (291 species), Colletocemateae (3 species), Urophyllaeae (251 species), and Ophiorrhizeae with around 420 species (Bremer & Eriksson, 2009; Rydin et al., 2009a, 2009b; Wikström et al., 2015). Also, a fifth tribe, Perameaeae (14 species), a neotropical tribe including the single genus Perama, should probably be included here and has been resolved together with Lasiantheae in several analyses (Andersson & Rova, 1999; Robbrecht & Manen, 2006; Smedmark et al., 2014). In the subfamily Cinchonoideae, four groups have emerged as well supported (Antonelli et al., 2009; Manns & Bremer, 2010; Manns et al., 2012; Wikström et al., 2015): the first includes the two predominantly paleotropical tribes, Hymenodictyeae (25 species) and Naucleeae (196 species), the second includes the “quinine-tribe” Cinchoneae (125 species) and Isertieae (17 species), the third includes Guettardeae (176 species) and Rondeletiaceae (192 species), and the last group includes the tribes Hamelieae (181 species), Hillieae (30 species), and Chicocceae (218 species). A fourth tribe Strumpfieae should probably be recognized in this last group (Paudyal et al., 2014). Two informal alliances are commonly recognized also in the subfamily Ixoroideae: the Coffeeeae alliance with more than 2000 species and 10 tribes (as recognized by Kainulainen et al. (2013) and Moly et al. (2014)); and the Vangerieae alliance with about 1200 species and eight different tribes (Kainulainen et al., 2013), and a grade of Retiniphyllaeae (21 species), Steenisieae (5 species), Mussaendeae (216 species) together with Sipaneeae (161 species), Condamineae (310 species), and a clade comprising Henriquezeea (21 species), Posoquerieae (23 species), and Sipaneeae with about 43 species (Kainulainen et al., 2010, 2013; Razafimandimbison et al., 2011).

There are of course uncertainties with respect to detailed relationships in many of the recognized tribes, but with respect to the overall relationships among the recognized tribes, a few problems stand out as having been particularly difficult to resolve (Fig. 1). The relationships of the two small tribes Luculieae, with a single genus and a handful of species, and Coptosapelteae, with two genera and 50–60 species, for example, are not well understood. They show inconsistent and often unsupported relationships in most analyses (Robbrecht & Manen, 2006; Bremer & Eriksson, 2009; Rydin et al., 2009a; Manns et al., 2012; Wikström et al., 2015), and they remain unclassified with respect to their subfamily placement. In the subfamily Rubioideae, relationships among early diverging lineages are problematic. Usually, the tribe Lasiantheae is grouped sister to a group comprising Coussareaeae and the Psychotrieaeae and Spermacoceae alliances, but the tribes Colletocemateae, Ophiorrhizeae, and Urophyllaeae show inconsistent and/or poorly supported relationships in most analyses (Robbrecht & Manen, 2006; Bremer & Eriksson, 2009; Rydin et al., 2009a; Manns et al., 2012; Wikström et al., 2015). Also, some of the relationships among tribes in the Psychotrieaeae and Spermacoceae alliances are problematic (Rydin et al., 2009b; Razafimandimbison et al., 2017). In the Cinchonoideae, relationships among the four well supported groups that have emerged in recent analyses are not understood, and alternative resolutions have resulted in widely different ideas about the geographic origins and expansions of not only the subfamily, but of the Rubiaceae as a whole (Antonelli et al., 2009; Manns et al., 2012). In the Ixoroideae, some of the relationships in the Vangerieaeae and Coffeeeae alliances (Kainulainen et al., 2013; Moly et al., 2014), and between the early diverging Condamineae, Mussaendeae–Sipaneeae, and Henriquezeea–Posoquerieae–Sipaneeae clades have been problematic to resolve (Kainulainen et al., 2013).

As in most plant groups (Zimmer & Wen, 2012; Davis et al., 2014; Ruhfel et al., 2014; Rothfels et al., 2015), previous work in Rubiaceae have relied almost exclusively on data from the plastid genome. There have been indications of conflicts between analyses that have strictly used data from the plastid genome, and those that have combined their plastid data with data from the nuclear genome (Robbrecht & Manen, 2006; Rydin
et al., 2009a, 2009b), but the reasonably well resolved and primarily plastid-based picture of overall phylogenetic relationships in the family was until recently more or less unchallenged. A recent work by Rydin et al. (2017) changed this and they demonstrated unambiguous and well supported conflicts between this plastid-based consensus and relationships obtained in their analyses of mitochondrial data. Among the supported conflicts they reported, two stood out as particularly striking: the rejection of monophyly of subfamilies Cinchonoideae and Ixoroideae with two tribes from subfamily Cinchonoideae, the “quinine-tribe” Cinchoneae and the Isertieae, nested well inside subfamily Ixoroideae; and the grouping of Jackieae from the Vanguerieae alliance with Airspermeae from the Coffeeae alliance (Rydin et al., 2017). Neither relationship has been reported before, and the authors speculated that the conflicts possibly reflect ancient hybridization events between early members of the subfamilies Cinchonoideae and Ixoroideae and of the Vanguerieae and Coffeeae alliances (Rydin et al., 2017). 

Adopting a “more characters” phylogenomic approach (Rokas et al., 2003; Delsuc et al., 2005; Davis et al., 2014) we here use complete data from the maternally inherited plastid genome with the aim to resolve relationships that have remained problematic in previous plastid-based analyses of the family. We conduct phylogenetic analyses on 59 Rubiaceae and four outgroup taxa. The effect of any compositional heterogeneity in our data, induced by synonymous substitutions, is investigated by comparing results from nucleotide-based analyses of the protein coding data with corresponding analyses of codon degenerated data that uses ambiguity codes to eliminate the distinction between synonymous codons. To further explore the consequences of a biased employment of plastid data, we also conduct analyses on corresponding taxa using data from the biparentally inherited nuclear rDNA cistron.

2 Material and Methods
2.1 Taxon sample

Taxa were targeted with the objective of having all, or as many as possible, tribes currently recognized in the family represented in the analyses. In total, 56 of the 65 commonly recognized tribes of Rubiaceae were included. Aleisanthieae, Condamineae, Crossopterygeae, Cyanoneuroneae, Foonchewieae, Henriquezieae, Peraeae, Schraderieae, and Strumpfieae were the tribes not included. The genus Glionnetia from the Seychelles, currently unclassified at the tribal level, was also included. The phylogenetic position of Glionnetia has been problematic to estimate but the genus is commonly resolved in an unsupported position in the Vanguerieae alliance (Razafimandimbison et al., 2011; Kainulainen et al., 2013; Wikström et al., 2015). Two non-Rubiaceae taxa from Gentianales were included as outgroup taxa in the analyses of ribosomal rDNA data: Mostuea Didr. (Gelsemiaceae), and Asclepias L. In the analyses of plastid data; these two taxa were complemented with two additional outgroup taxa: Rhazya Decne. (Apocynaceae) and Gentiana L. (Gentianaceae). Detailed species and voucher information for all included taxa is given in Table 1.

Table 1. List of taxon names, DNA voucher information, and accession numbers for sequences used in the analyses

| Taxon                                          | DNA voucher/reference | rDNA cistron accession numbers | Plastid accession number |
|------------------------------------------------|-----------------------|--------------------------------|--------------------------|
| Alberta magna E.Mey.                           | J. E. Tonkin 200 (UPS) | MK607891                       | KY348839                 |
| Amphidasya ambigua (Standl.) Standl.           | Clark and Watt 736 (QCNE, MO, UPS) | MK607892 | KY378703 |
| Anthospermum spathulatum Spreng.               | Bremer et al. 4405 (UPS) | MK607893                       | KY378687                 |
| Argostemia yappii King                         | Bremer and Bremer 1609 (S) | MK607894 | KY378693 |
| Asclepias syriaca L.                           | Straub et al. (2013)   | --                             | NC_022432                |
| --                                             | Straub et al. (2011)   | JF312046                       | --                       |
| Augusta australcaledonica (Brogn.) J.H.Kirkbr. | Mousley et al. 237 (P) | MK607890                       | KY492076                 |
| Bertiera longithyrsa Baker                     | Kärhed et al. 256 (UPS) | MK607895                       | KY348833                 |
| Boholalia nematostylis Merr.                   | D. Bicknell 1561A (S)  | MK607896                       | KY348840                 |
| Cephalanthus occidentalis L.                   | NTBG-960457.002        | MK607897                       | KY378678                 |
| Cinchona pubescens Vahl                       | Bremer 2733 (UPS)      | MK607898                       | KY378682                 |
| Coffea arabica L.                              | Samson et al. (2007)   | --                             | NC_008535                |
| --                                             |                       | NW_020849278                   | --                       |
| Colletoeccema dewevei (De Wild.) E.M.A.Petit   | Lisowski 47195 (K)     | MK607899                       | KY378707                 |
| Coptosapelta flavescentis Korth.               | Puff 950720-12 (WU, BKF) | MK607900 | KY378704 |
| Craterispermum sp.                             | Eriksson et al. 999 (S) | MK607901                       | KY378662                 |
| Cubanola domingensis (Britton) Aiello          | Bremer 4500 (S)        | MK607902                       | KY378677                 |
| Danais xanthorrhoea (K.Schum.) Bremek.         | Bremer 3079 (UPS)      | MK607903                       | KY378686                 |
| Deppea grandiflora Schltdl.                   | Bremer 2724 (UPS)      | MK607904                       | KY378675                 |
| Dunnia sinensis Tutcher.                       | Longmen 10 (in Ge et al., 2002) | MK607905 | KY378692 |

Continued
| Taxon | DNA voucher/reference | rDNA cistron accession numbers | Plastid accession number |
|-------|-----------------------|--------------------------------|-------------------------|
| Faramea multiflora A.Rich. ex DC. | Bremer et al. 3331 (UPS) | MK607906 | KY378701 |
| Gaertnera phyllostachya Baker | Kårehed et al. 272 (UPS) | MK607907 | KY378695 |
| Gardenia conferta Guillaumin | Barabbé 330 (NOU) | MK607908 | KY348835 |
| Gentiana straminea Maxim. | Ni et al. (2014) | – | NC_027441 |
| Gloineria sericea (Baker) Tirveng. | Bremer et al. 5404 (S) | MK607909 | KY378665 |
| Greenea oblonga Craib. | Larsen et al. 33451 (P) | MK607910 | KY378664 |
| Guettarda scabra (L.) Vent. | Rova 2260 (GB) | MK607911 | KY378680 |
| Gynochthodes officinalis (F.C.How) Razafim. & B.Bremer | Ding et al. (2015) | – | NC_028009 |
| Hillia triflora (Oerst.) C.M.Taylor | Bremer 3101 (UPS) | MK607912 | KY378676 |
| Hymenodictyon parvifolium Oliv. | B. Bremer 3809 (UPS) | MK607913 | KY378679 |
| Isertia laevis (Triana) B.M.Boom | B. Bremer et al. 3364 (UPS) | MK607914 | KY378683 |
| Ixora hookeri (Oudem.) Bremek. | Kainulainen et al. 182 (S) | MK607915 | KY378663 |
| Ixora parviflora Lam. | Gillis 7892 (FTG) | MK607916 | KY378672 |
| Jackiopsis ornata (Wall.) Ridsdale | K. S. Tan s.n. | MK607917 | KY378669 |
| Kohautia caespitosa Schnizl. | Bremer et al. 425668 (UPS) | MK607918 | KY378684 |
| Lasianthus sp. | Kainulainen et al. 17 (S) | MK607919 | KY378708 |
| Luculia grandifolia Ghose | B. Bremer 2713 (S) | MK607920 | KY378705 |
| Mitchella repens L. | Atha and Gonzalez 1443a (MEXU) | MK607921 | KY378710 |
| Mitriostigma axillare Hochst. | Bremer and Rydin 5014b (S) | MK607922 | KY348837 |
| Morinda citrifolia L. | Bremer 3302 (UPS) | MK607923 | KY378694 |
| Mostuea brunosii Didr. | B. Bremer et al. 5077 (S) | MK607924 | KY378706 |
| Mussaenda densiflora H.L.Li | Razafimandimbison et al. 747 (S) | MK607925 | KY348834 |
| Ophiorrhiza mungos L. | Bremer 3301 (UPS) | MK607926 | KY378702 |
| Paederia foetida L. | Wong and Keong s.n. (KLU) | MK607927 | KY378691 |
| Palicourea guianensis Aubl. | B. Bremer et al. 3332 (UPS) | MK607928 | KY378697 |
| Paragepnepa lancifolia (Bojer ex Baker) Tirveng. & Robbr. | Bremer et al. 5413 (S) | MK607929 | KY348838 |
| Pavetta abyssinica Fresen. | De Block 6 (BR) | MK607930 | KY378673 |
| Pentas lanceolata (Forssk.) Deflers | Bremer 2702 (S) | MK607931 | KY378685 |
| Plocama pendula Alton | K. Andreassen 1 (UPS) | MK607932 | KY378690 |
| Posoqueria abyssinica (Forssk.) Roem. & Schult. | Bergius Botanic Garden SU-C-88.10 | MK607933 | KY378662 |
| Prismaticeris fragrans Geddes | Kainulainen et al. 39 (S) | MK607934 | KY378699 |
| Psychotria kirkii Hiern | Bremer 3102 (UPS) | MK607935 | KY378696 |
| Psyradra obovata (Klotzsch ex Eckl. & Zeyh.) Bridson | Bremer 3762 (UPS) | MK607936 | KY378666 |
| Retiniphyllum pilosum (Spruce ex Benth.) Müll.Arg. | Wurdack and Adderley 43270 (S) | MK607937 | KY378670 |
| Rhaya stricta Decne. | Park et al. (2014) | – | KJ485849 |
| Rondeletia odorata Jacq. | Bremer and Andreassen 3504 (UPS) | MK607938 | KY378681 |
| Rubia horrida (Thunb.) Puff | Bremer et al. 4266 (UPS) | MK607939 | KY378689 |
| Sabicea diversifolia Pers. | B. Bremer et al. 5262 (S) | MK607940 | KY378671 |
| Schizocolea linderi (Hutch. & Dalziel.) Bremek. | Adam 2016 (UPS) | MK607941 | KY378700 |
| Scyphiphora hydrophyllacea C.F.Gaertn. | K. Bremer et al. 99 (S) | MK607942 | KY378668 |
| Sipanea sp. | Rova et al. 1981 (GB) | MK607943 | KY378674 |
| Stachyarrhena heterochroa Bridson | Persson et al. 821 (GB) | MK607945 | KY348836 |
| Steenisia pleurocarpa (Airy Shaw) Bakh.f. | Puff BF 990619-1/4 (WU) | MK607944 | KY378709 |
| Theligenon cyanocrambe L. | Reuterswärd and Forsslund 2 (S) | MK607946 | KY378688 |
| Traillioidaoxa gracilis W.W.Sm. & Forrest | Boufford et al. 35041 (MO) | MK607947 | KY378667 |

References are given for sequences taken from Genbank.
2.2 DNA extraction and sequencing
DNA was extracted from herbarium, live, or silica-dried material using a cetyltrimethylammonium bromide (CTAB) protocol (Doyle & Doyle, 1987; Doyle, 1991). The extracted DNA was cleaned using the QiAquick PCR cleaning kit from Qiagen (Hilden, Germany) following the protocol specified by the manufacturer.

High-throughput sequencing was done at the Science for Life Laboratory (ScLifeLab, Stockholm, Sweden) following the manufacturer's instructions for the Illumina HiSeq 2500 platform (Illumina, San Diego, California, USA). Pair-end runs with 300-bp insert size fragments and 2 × 150 bp read lengths were performed. Samples were multiplexed with 14, 17, or 28 other samples in one lane in three consecutive runs. Library preparation at the ScLifeLab was done using the Illumina TruSeq DNA PCR-free library preparation kit (Illumina) for samples with high quantities of DNA, and the Thruplex DNA-seq library preparation kit from Rubicon (Rubicon Genomics, Ann Arbor, Michigan, USA) for samples with lower quantities of DNA. Demultiplexing and conversion was done using CASAVA v1.8.2 (Illumina).

2.3 Plastid sequence assembly
To isolate plastid sequences from the original reads, a BLAT (BLAST-like alignment tool v36; Kent, 2002) search of forward and reverse reads against an initial database of whole plastid genomes of eight Lamiai taxa was performed. The initial database included the plastid genomes of Coffea arabica (Rubiaceae; NC_008535), Gynochthodes officinalis (Rubiaceae; NC_028009), Nicotiana undulata (Solanaceae; NC_016068), Catharanthus roseus (Apocynaceae; NC_021423), Asclepias nivea (Apocynaceae; NC_022431), Rhyza stricta (Apocynaceae; KJ485849), Boea hygrometrica (Gesneriaceae; NC_016468), and Scrophularia takasimensis (Scrophulariaceae; MK590983). Forward and reverse reads were both saved if either showed at least 70% similarity to any of the Lamiai genomes. After the BLAT search, reads were extracted from the original fastq data files using pullseq v1.0.1 (github.com/bthomas/pullseq) into new forward and reverse fastq data files representing a “chloroplast” subset. De novo assembly of the “chloroplast” subset of reads was performed for each taxon using ABYSS v1.5.2 and eight different k-mer lengths (55, 61, 67, 73, 85, 91, 97, 103). All generated contigs were pooled and mapped onto a reference genome using bwa v0.7.5a-r405. Initially, the chloroplast genome of Coffea arabica (NC_008535) was used as reference. This resulted in complete or near complete draft genomes. All original reads were subsequently mapped onto the draft genomes using bwa v0.7.5a-r405 allowing unfinished gaps to be filled and sequencing depths to be evaluated. Assemblies were reviewed and edited using gaps from the Staden Package v2.0.0b10 (Staden, 1996; Staden et al., 2000). After the completion of each chloroplast genome, it was added to the database of chloroplast genomes used in the BLAT search and also considered as a potential reference genome in the final mapping stage in subsequent assemblies. Protein coding (CDS), tRNA, and rRNA genes were annotated using Sequin v15.10 (available by anonymous FTP at https://www.ncbi.nlm.nih.gov/Sequin/; National Center for Biotechnology Information, Bethesda, Maryland, USA) by transferring the annotation of Coffea arabica (NC_008535).

2.3.1 Assembly of nuclear rDNA cistrons
Nuclear rDNA cistrons were assembled in a corresponding way as the plastids (see above). The initial database for the BLAT search comprised the rDNA cistron sequence of Asclepias syriaca (JF312046), which was also used as the initial reference sequence for mapping the de novo generated contigs. Annotation of the sequences was done using Sequin v15.10 and boundaries for the 18S, 5.8S, and 28S rDNA genes were inferred from alignment with the Coffea arabica sequence NW_020849278 REGION: complement (11780..17784).

2.4 Phylogenetic analyses
2.4.1 Plastid data
2.4.1.1 Alignment and pretrimming Protein coding (CDS), intron, tRNA, rDNA, and intergenic spacer (IGS) regions were extracted from the annotated GenBank files using an in-house python script built on biopython v1.63 (Cock et al., 2009). Script is available from the Dryad Digital Repository (https://doi.org/10.5061/dryad.q8t1f4). Stop codons and regions of uncertain homology were identified by eye and removed using a python script developed and made available at the Dryad Digital Repository (https://doi.org/10.5061/dryad.q8t1f4) by Fučíková et al. (2016). Extracted DNA regions were aligned individually using Muscle v3.8.31 (Edgar, 2004), with protein-coding gene regions aligned as amino acid sequences and converted back to nucleotides using Seaview 4 (Gouy et al., 2010).

2.4.1.2 Nucleotide based analyses Individual protein-coding genes were concatenated into a combined CDS data set. Also, non-CDS data were assembled comprising the intron, tRNA, and rDNA regions, and phylogenetic analyses were subsequently conducted on two alternative data sets: one including only the CDS data, and a second with CDS and non-CDS data combined. Intergenic spacer regions were as a rule too variable to produce reliable alignments and were therefore not included in any of the analyses. Bayesian analyses were conducted in MrBayes v3.2.6 (Huelsenbeck & Ronquist, 2001; Ronquist & Huelsenbeck, 2003). The CDS data were partitioned by codon position and the non-CDS data into two partitions, one comprising the tRNA and rDNA regions and a second including the intron data. The GTR+I+I substitution model was used for all partitions based on the corrected Akaike information criterion (AICc) as calculated using the program MrAIC c1.4.6 (Nylander, 2004) and PHYML v3.0 (Guindon et al., 2010). Substitution models were unlinked across all partitions. MrBayes was run for 10 000 000 generations sampling trees and parameter estimates every 1000 generations. Two independent runs, each with four chains and heating parameters set to default values, were conducted for all analyses. Bayesian posterior probability values were calculated after discarding the first 25% of the trees and parameters as burnin, and this was well beyond the burnin phase of the chains based on the PSRF convergence diagnostic (Gelman & Rubin, 1992) reported by MrBayes.

Maximum Likelihood (ML) analyses were conducted in RAxML v8.2.9 (Stamatakis, 2014) with CDS and non-CDS data partitions and nucleotide substitution models set up as in the Bayesian analyses. Branch support was estimated using the
of sequenced fragments varied across samples from $1.5 \times 10^6$ fragments in *Cephalanthus occidentalis* to $16.7 \times 10^6$ fragments in *Prismatomeris fragrans* with an average number of $8.9 \times 10^6$ fragments. Assembled sequences were submitted to the DDBJ/EMBL/GenBank databases and a comprehensive list of accession numbers for all deposited sequences, together with taxon names and DNA voucher information is given in Table 1.

### 3.2 Assembled rDNA cistrons

Nuclear rDNA cistrons were assembled for the corresponding accessions. Assembled sequences comprise incomplete external transcribed spacer (ETS), 18S ribosomal RNA (18S rRNA) gene region, internal transcribed spacer 1 (ITS1), 5.8S ribosomal RNA (5.8S rRNA) gene region, internal transcribed spacer 2 (ITS2), 28S ribosomal RNA (28S rRNA) gene region, and incomplete non-transcribed spacer (NTS). Assembled sequences ranges in length from 5835 bp in *Rubia hordida* to 5980 bp in *Lasianthus sp.* See Table 1 for a comprehensive list of accession numbers for all deposited sequences.

### 3.3 Phylogenetic analyses

#### 3.3.1 Plastid data

#### 3.3.1.1 Nucleotide based analyses

The plastid CDS data set include a total of 77 protein-coding gene regions and 69 828 aligned nucleotide characters for 59 ingroup Rubiaceae (57 newly generated here) and four outgroup (one newly generated here) non-Rubiaceae Gentianales taxa. The combined CDS + non-CDS data set for corresponding taxa include a total of 95 494 nucleotide characters: 69 828 CDS and 25 666 non-CDS (18 899 intron; 2225 tRNA; 4542 rDNA). The complete partitioned and annotated data sets are available from Dryad Digital Repository (S1 and S2 Files; [https://doi.org/10.5061/dryad.q573n5tf4](https://doi.org/10.5061/dryad.q573n5tf4)). Results from the phylogenetic analyses are reported as phylograms from the Bayesian analyses in Fig. 2 (analysis of CDS data) and Fig. S1 (analysis of combined CDS and non-CDS data). Branch support is indicated in all trees by Bayesian posterior probabilities (BPP) and ML bootstrap frequencies (BS).

In the CDS analyses relationships are generally resolved and, with a few exceptions, well supported (Fig. 2). In subfamily Rubioideae the tribes Ophiorrhizeae, Urophylleae, and Colletocereeae are in unsupported positions as early diverging lineages. Lasiantheae followed by Cousssareae are successive sisters to a clade comprising the Psychotrieae and Spermacoceae alliances. In the Psychotrieae alliance Schizocoleeae are sister to remaining tribes followed by Craterispermeae, Psychotrieae + Palicoureeae, Prismatomerieae + Gaertnerieae, and Morindeae + Mitchelleae. In the Spermacoceae alliance Danaideae are grouped with Spermacoceae plus Knoxieae and together they are sister to remaining tribes. Anthospermeae and Dunnieae are successive sisters to an unsupported group (BPP = 0.75; BS < 50%) with Argostemmateae sister to a clade comprising Paederieae, Potorieae, Theligoneae, and Rubieae. Within the latter clade, Rubieae are grouped with Theligoneae, with Potorieae and Paederieae as successive sisters. Coptosapelteae (BPP = 0.79; BS = 75%) and Luculieae (BPP = 0.91; BS = 81%) are resolved in positions at the base of subfamily Rubioideae.

In subfamily Cinchonoideae, Naucelleae together with Hymenodictyeeae are resolved sister to a clade (BPP = 1.00; BS = 77%) comprising remaining tribes in the subfamily. Chioococceae are sister to Hillieae plus Hamelieae, and they
Fig. 2. Continued
are sister to a clade (BPP = 0.98; BS = 50%) where Rondeletieae plus Guettardeae are sister to Cinchoneae plus Ixoroideae.

Relationships in the subfamily Ixoroideae are generally supported by both Bayesian and ML analyses, excepting the relationships of Glionnetia sister to Vanguerieae and the Greeneae–Ixoroideae clade (BPP = 0.93; BS = 77%), the sister group relationship between Trailingeae and Scyphiphoreae (BPP = 0.99; BS = 68%), and some relationships in the Gardenieae complex. Sipaneae with Poqueroieae as sister to remaining Ixoroideae followed by Sabiceae with Mussaenaeae, Steenisiaeae, Retiniphyllaeae, and the Coffeeae and Vanguerieae alliances. In the Vanguerieae alliance Jackieae are sister to remaining tribes followed by Trailingeae with Scyphiphoreae (BPP = 0.99; BS = 68%). Glionnetia, and an unsupported node (BPP = 0.93; BS = 77%) comprising the Vanguerieae and a Greeneae + Ixoroideae clade. In the Coffeeae alliance Airospermeae are sister to remaining tribes followed by Augusteae, Albertaeae, Coffeeae with Bertiereae, Cordiereae with Octotropideae (BPP = 1.00; BS = 69%), and Gardenieae sister to the unsupported Pavetteae + Sherbournieae clade (BPP = 0.99; BS = 68%). In the analyses of combined CDS and non-CDS data the results obtained in the CDS analyses are as a rule corroborated with increased support (Fig. 5). Exceptions are the relationships of Urophyllaeae, Colletocoemateae, Theligoneae, and Octotropideae. Urophyllaeae and Colletocoemateae are resolved as sistergroups (BPP = 0.74; BS = 62%), the support for grouping Theligoneae and Rubieae is substantially decreased (BPP = 0.92; BS < 50%), and Octotropideae are no longer supported as sister to Cordiereae, but placed in an unsupported position as sister to a group with Gardenieae, Sherbournieae, and Pavetteae (BPP = 0.91; BS < 50%).

### 3.3.2 Degenerate codon analyses

The complete partitioned and annotated codon degenerated data set is available in Nexus file format from Dryad Digital Repository (S3 File; https://doi.org/10.5061/dryad.q573n5tf4). Results from the phylogenetic analyses are reported as a phylogram (from the Bayesian analyses) in Fig. 3, with branch support indicated for each node by Bayesian posterior probabilities (BPP) and ML bootstrap frequencies. In the subfamilies Rubioideae and Ixoroidae the analyses show results highly similar to the nucleotide analyses. Both place Ophiurhizae as sister to remaining tribes in subfamily Rubioideae, but the support for this relationship (BPP = 0.99; BS = 76%) increases considerably here in the degenerate codon analyses (Fig. 3). Also the sister group relationship between Argostemmatae and the clade comprising Paederieae, Putorieae, Theligoneae, and Rubieae becomes supported (BPP = 1.00; BS = 85%), whereas the support for grouping Prismatomeridea with Gaertnerieae decreases (BPP = 0.83; BS = 59%) in the degenerate codon analyses. In subfamily Ixoroidae the support for grouping Cordiereae with Octotropideae is lost (BPP = 0.90; BS < 50%), and Pavetteae goes from an unsupported position as sister to Sherbournieae to a position as sister to Gardenieae (BPP = 0.95; BS = 69%).

In the subfamily Cinchonoideae the situation is different. Here the results clearly conflict with those from the nucleotide based analyses (Figs. 2, S1) with Isertieae together with Cinchoneae forming a sistergroup relationship to remaining tribes (BPP = 0.99; BS = 59%), and with Naucleeae together with Hymenodictyaeae now in a more derived but unsupported position (BPP = 0.89; BS = 50%) as sister to a group comprising Rondeletieae and Guettardeae.

### 3.3.2 Nuclear rDNA data

The nuclear rDNA data set included a total of 6045 aligned nucleotide characters for 58 ingroup Rubiaceae (57 newly generated here) and two outgroup (one newly generated here) non-Rubiaceae Gentianales taxa. Included were data from the 18S rDNA gene (1814 bp), 5.8S rDNA gene (156 bp), 28S rDNA gene (3427 bp), ITS1 rDNA spacer (331 bp; 207 excluded), and the ITS2 rDNA spacer (317 bp; 153 excluded). The complete partitioned and annotated rDNA data set is available in Nexus file format from Dryad Digital Repository (S4 File; https://doi.org/10.5061/dryad.q573n5tf4).

Results from the phylogenetic analyses of the rDNA gene data set, including only the 18S, 5.8S, and 28S rDNA gene regions, are reported as a phylogram (from the Bayesian analyses) in Fig. 4. In the subfamily Rubioideae relationships are generally resolved and well supported. Colletocoemateae, Ophiurhizaeae, and Urophyllaeae form a monophyletic sister group (BPP = 1.00; BS = 89%) to remaining tribes (BPP = 0.98; BS = 93%) followed by Lasiantheae. Coussareaeae are resolved sister (BPP = 0.98; BS = 76%) to the Spermacoceae alliance (BPP = 1.00; BS = 82%), Anthospermeae are sister to Knoxieae plus Spermacoceae (BPP = 1.00; BS = 70%), and together they form a monophyletic sister group to a clade comprising remaining tribes (BPP = 0.95; BS < 50%) in the Spermacoceae alliance. Among the remaining tribes Dunnieae followed by Danaeaeae, Argostemmataeae, and Paederieae are resolved as successive sister groups to a clade where Rubieae are sister to Theligoneae plus Putorieae (BPP = 0.93; BS = 92%), but the Paederieae (BPP = 0.93; BS = 73%) and Danaeaeae (BPP = 0.89; BS < 50%) positions are poorly supported. In the Psychotrieae alliance (BPP = 0.99; BS = 83%) Morindeaeae together with Mitcheleeaeae (BPP = 0.98; BS = 76%) are resolved sister to Prismatomeridea (BPP = 1.00; BS = 93%), Psychotrieaeae together with Palicouraeaeae are resolved sister to Gaertnerieaeae, and together the six tribes are resolved as monophyletic with
Fig. 3. Continued
In subfamilies Cinchonoideae and Ixoroidae we see a highly deviating pattern. Here relationships are mainly unresolved or poorly supported, and neither the two subfamilies, nor the two informal Ixoroidae alliances are retrieved by the analyses (Fig. 4). Hillieae are sister to Hamelieae (BPP = 1.00; BS = 78%), and Naucleeae are sister to Hymenodictyeae (BPP = 1.00; BS = 98%), but remaining relationships of the Cinchonoideae tribes are unresolved or poorly supported. Among Ixoroidae, the tribe Ixoreeae is sister to Greeneeae (BPP = 1.00; BS = 81%), with Glionnetia as their sister group (BPP = 1.00; BS = 65%). Octotropideae are sister to Sherbournieae (BPP = 1.00; BS = 81%) and together with Gardenieae and Pavetteae they form a monophyletic group (BPP = 1.00; BS = 80%). Coptosapelteae are grouped with Luculieae and together in an unresolved position at the base of the family, but the grouping of Coptosapelteae and Luculieae is not supported by the ML bootstrap (BS < 50%).

Results from the analyses of rDNA data including also the variable ITS1 and ITS2 spacer regions are reported as supporting information (Fig. S2). Compared to the analyses not including the ITS regions there are only minor differences, and the general pattern with most relationships in subfamily Rubioidae resolved and well supported, and with almost no relationships resolved and well supported in subfamilies Cinchonoideae and Ixoroidae remain. Support and resolution are sometimes lower than those of the analysis without nrITS, in particular within the subfamily Rubioidae. The opposite can, however, also be observed for some nodes; there is for example, increased support for grouping Coussareeae with the Spermacoceae alliance, the Psychotrieae alliance, and the sister relationships between Pavetteae and Gardenieae, and between Isertieae and Cinchoneae.

4 Discussion

It is clear that simply adopting a “more characters” phylogenomic approach does not resolve the phylogenetic relationships of Rubiaceae. The plastid based analyses presented here use 69,828 bp from 77 protein-coding genes, and an additional 25,666 bp of nonprotein coding data, yet some of the relationships that have been difficult to resolve in previous analyses remain problematic and appear to be unaffected by the massive addition of new data contributed in the present analyses. The tribes Luculieae and Coptosapelteae, for example, are still placed in poorly supported positions at the base of the family, although the analyses of combined CDS and non-CDS data support a sister group relationship between Luculieae and subfamily Rubioidae (BPP = 0.98; BS = 89%). Also the relationships of the early diverging tribes Urophylleae and Colletoeceae in subfamily Rubioidae remain difficult to resolve (Figs. 2, 3, S1).

It is also clear that there are conflicting phylogenetic signals in the data analyzed. Some of those conflicts come from the plastid data alone where different analyses of the same protein coding data, treated either as nucleotides (Fig. 2) or as codon degenerated data (Fig. 3), resolve and support conflicting topologies, in particular in the subfamily Cinchonoideae. As these analyses use the same data the conflicting signals cannot be “real” in a biological sense, but are likely resulting from erroneous assumptions in the models used to reconstruct our phylogenies (Philippe et al., 2011; Cooper, 2014).

Conflicting signals of a different kind are seen in the analyses of plastid data on the one hand, and of nuclear rDNA data on the other. The analyses of plastid data behave as one would expect, with results that are more or less congruent with the coherent picture of phylogenetic relationships in the family built during the last 20–30 years. The analyses of nuclear rDNA data, however, yield less congruent results. In the subfamily Rubioidae the analyses both resolve and support a large number of relationships (Figs. 4, S2), many also congruent with those indicated by the analyses of plastid data, but other relationships are not only well supported, but also clearly in conflict with the relationships supported by the analyses of plastid data. In the subfamilies Cinchonoideae and Ixoroidae we see a very different general pattern in the results of the phylogenetic analyses. Here the analyses of nuclear rDNA data almost completely fail to resolve and support any phylogenetic relationships, yet also here there are clear indications of conflicting signals in the analyses of plastid versus nuclear data. Plants typically have a maternal inheritance of the plastids (Corriveau & Coleman, 1988; Birky, 1995, 2001), whereas nuclear rDNA have a biparental inheritance, although it may appear uniparental as a result of concerted evolution and differential elimination of one of the parental rDNA (Volkov et al., 2007). Hence, these conflicting signals possibly represent “real” conflicts, in the sense that the different relationships actually reflect alternative evolutionary histories of the two data types.

4.1 Conflicting signals in data of the same genomic compartment within Cinchonoideae

In the subfamily Cinchonoideae our analyses of the same protein-coding gene sequence data treated either as nucleotides (Fig. 2), or as codon degenerated data, where nucleotide ambiguity codes are used to eliminate the
Fig. 4. Continued
distinction between synonymous codons (Fig. 3), provide unequivocal support for conflicting relationships. The subfamily has a primary distribution in the Neotropics but two of the tribes, the Naucleeae and the Hymenodictyeae, have predominantly paleotropical distributions (Razafimandimbison & Bremer, 2001; Antonelli et al., 2009; Manns & Bremer, 2010). Our nucleotide based analyses of the CDS data set (Fig. 2) support that the paleotropical tribes Hymenodictyeae and Naucleeae are sisters to remaining tribes, whereas the codon degenerated version of the same data indicate corresponding support for a conflicting topology with the neotropical tribes Cinchoneae and Isertieae in that same position (Fig. 3). As both types of data are derived from the same set of observations, they must have evolved under the same species tree and the supported conflict must result from erroneous assumptions in the models used to reconstruct our phylogenies. Both topologies have, however, been obtained and reported in previous studies (Andersson & Antonelli, 2005; Robbrecht & Manen, 2006; Antonelli et al., 2009; Manns & Bremer, 2010; Manns et al., 2012; Wikström et al., 2015; Yang et al., 2016), and the alternative positions for the paleotropical tribes have contributed to widely different and conflicting ideas about the geographic origins, not only of the subfamily Cinchonioideae, but of Rubiaceae as a whole. Antonelli et al. (2009) resolved the paleotropical Hymenodictyeae and Naucleeae as sisters to remaining tribes with the neotropical tribes Isertieae and Cinchoneae nested well inside a clade of mainly Central American and Antillean tribes. They considered the pattern to corroborate a boreotropical origin of the family, and suggested Rubiaceae as a whole to have originated in the Paleotropics, and to have dispersed into North America via the North Atlantic Land Bridge (NALB) in the Late Paleocene (Antonelli et al., 2009). The neotropical tribes Isertieae and Cinchoneae were suggested to have dispersed into South America more recently, sometime in the Early Eocene (Antonelli et al., 2009). Rydin et al. (2009a) reported an alternative topology with an initial split in subfamily Cinchonioideae separating the neotropical tribes Rondeletiaceae and Guettardeae, but most recent studies have reported a topology with the neotropical Cinchoneae and Isertieae as sister to remaining tribes (Manns & Bremer, 2010; Manns et al., 2012; Wikström et al., 2015). The alternative positions for the paleotropical tribes have, however, remained poorly supported in most of these analyses, but following ancestral-area and divergence-time reconstructions Manns et al. (2012) rejected entirely the hypothesis of Rubiaceae having used the NALB to reach the Neotropics in the Late Paleocene. They suggested a reversed scenario with the ancestor of both subfamilies Cinchonioideae and Ixoroiodeae present in South America already during the Late Cretaceous, and with the ancestor of Hymenodictyeae and Naucleeae having dispersed out of America, from the Neotropics to the Paleotropics, using the NALB in the Late Paleocene or Eocene (Manns et al., 2012). We consider this out of America scenario as best supported by the analyses presented here with paleotropical tribes Hymenodictyeae and Naucleeae nested well inside the subfamily Cinchonioideae (Fig. 3).

The conflicting positions of the paleotropical Hymenodictyeae and Naucleeae in the nucleotide based analyses, as sisters to remaining tribes (Fig. 2), appear to be caused by a composition bias resulting from synonymous substitutions. Synonymous substitutions primarily result from changes in the third codon position, whereas substitutions in first and second codon positions are mostly nonsynonymous, except for codon variants of Leucine, Arginine, and Serine. Before the era of phylogenomics, analyses of one or a few genes often found “greater phylogenetic signal” in rapidly evolving third codon positions relative to their corresponding first and second codon positions (Källersjö et al., 1998; Björklund, 1999; Sennblad & Bremer, 2000; Simmons et al., 2002, 2006). However, recent analyses of both plants (Zhong et al., 2011; Cox et al., 2014; Li et al., 2014) and animals (van den Bussche et al., 1998; Tarrio et al., 2001; Philippe et al., 2011; Regier & Zwick, 2011; Breinholt & Kawahara, 2013; Morgan et al., 2013) clearly indicate that composition and dependent codon–usage biases, resulting from synonymous substitutions, are potential causes behind conflicting and erroneous phylogenetic signals.

4.2 Conflicting signals in data from different genomic compartments

Our analyses of plastid data on the one hand, and nuclear rDNA data on the other reveal significant conflicts in their phylogenetic signals. Relationships, well supported by the analyses of plastid data, are clearly in conflict with relationships equally well supported in the analyses of nuclear rDNA data, and these conflicting signals we interpret to represent “real” conflicts, in the sense that the different relationships actually reflect alternative evolutionary histories of the two data types. In the subfamily Rubiioideae there are several such conflicts: the close relationship between the tribe Coussareaeae and tribes traditionally united in the Spermacoceae alliance; the sister group relationship between the tribe Anthospermeae and the Knoxiaeae–Spermacoceae clade; the relationships of Danaideae to the Argostemmataeae–Putoriaeae clade, not close to the Knoxiaeae–Spermacoceae clade; and the sister group relationship between Gaertneraeae and the Psychorieaeae–Paliouriaeae clade. These relationships are supported in the analyses of nuclear rDNA data (Figs. 4, S2), but have to our knowledge never been supported by any previous analyses. Two of the conflicts concern tribes in the Spermacoceae alliance, and resolving the relationships in this group have been problematic.
with relationships either unresolved or poorly supported in previous analyses (Robbrecht & Manen, 2006; Rydin et al., 2009b; Wikström et al., 2015). From a data and taxon sample point of view Rydin et al. (2009b) is the most comprehensive analysis of the group, and they considered conflicting signals in different loci to cause some of the problems they encountered in their analyses. They used five different plastid loci in their analyses, but they also included data from the nuclear ITS rDNA spacer regions. The limited amount of non-plastid characters did, however, prevent them from further evaluating conflicting signals between plastid and nuclear data. Our results clearly indicate that there are such conflicting signals, not only in the Spermacoceae alliance but also in the Psychotrieae alliance, and instead of combining data from the plastid genome, that has an almost strict maternal inheritance pattern (Corriveau & Coleman, 1988; Birky, 1995, 2001), with biparentally inherited data from the nucleus, we need to start teasing apart the different signals and investigate the underlying causes behind their conflicts.

Recent work by Rydin et al. (2017) documented conflicting phylogenetic signals very much similar to those seen here, but in their case the conflicts were between plastid data on the one hand, and mitochondrial data on the other. Their analyses of mitochondrial data rejected monophyly of two out of three subfamilies of Rubiaceae and placed the quinine-tribe Cinchoneae and the Isertieae from subfamily Cinchoideae well inside subfamily Ixoroideae. In the subfamily Ixoroideae they also found strong support for grouping the tribe Jackieae, traditionally placed in the Vanguerieae, with Airospermeae from the Coffeeeae alliance, and they considered ancient hybridization events between early occurring members of the subfamilies Cinchoideae and Ixoroideae, and of the Vanguerieae and Coffeeeae alliances as possible reasons behind the conflicting signals (Rydin et al., 2017). Although the analyses of nuclear rDNA data presented here fail to resolve almost any relationships in the subfamilies Cinchoideae and Ixoroideae and of the Vanguerieae and Coffeeeae alliances as possible reasons behind the conflicting signals (Rydin et al., 2017). Although the analyses of nuclear rDNA data presented here fail to resolve almost any relationships in the subfamilies Cinchoideae and Ixoroideae, they do support one of the conflicting relationships seen in the mitochondrial analyses, the close relationship between the tribes Jackieae and Airospermeae. Together they form a well supported group with the tribe Albertaeae (Figs. 4, S2). Boholia (Airospermeae) deviate from most other taxa by having a very long branch in the phylograms, and it is possible that the close relationship to Jackieae is caused by long branch attraction artifacts (Felsenstein, 1978). However, a close relationship between Airospermeae and Albertaeae is not in conflict with the analyses using plastid data (Figs. 2, 3), the conflict concerns their close relationship to Jackieae, and neither Albertaeae, nor Jackieopsis (Jackieae) are associated with long branches in the rDNA phylograms (Figs. 4, S2).

4.3 Usefulness of nuclear DNA

The almost complete failure to resolve the relationships in the subfamilies Cinchoideae and Ixoroideae using nuclear rDNA data is surprising (Figs. 4, S2). One possible explanation could be that the 18S, 5.8S, and 26S nuclear rDNA gene regions analyzed are not variable enough for resolving the relationships in these groups. A simple evaluation of individual phylograms from the posterior distribution of trees of our Bayesian analyses show that the rDNA gene regions, without doubt, are less variable in these groups than in subfamily Rubioideae. However, adding data from the highly variable ITS spacer regions to our analyses have no significant effect on the resolution in the Cinchoideae and Ixoroideae (Fig. S2), which indicate that lack of information is inadequate as explanation for the poorly resolved results based on nuclear data. If anything, the ITS regions appear variable to the extent that they cannot be reliably aligned across the family. Two previous studies (Rydin et al., 2009a; Manns & Bremer, 2010) in fact report conflicting topologies likely caused by their respective alignment of the ITS regions. While data from the 18S, 5.8S, and 26S rDNA gene regions thus may be too conserved to be useful for resolving the relationships in the Cinchoideae and Ixoroideae, the ITS spacer regions appear too variable.

To expand phylogenetic investigations to also allow for detection of potential anastomosing evolutionary patterns caused by for example hybridization will require other approaches than basing studies of yet larger sets of data. Although it has long been known that inheritance of the organellar genomes is not always strictly maternal (see e.g., Rydin et al., 2017 and references therein), the true inheritance patterns for individual taxa of interest have often been left unknown or unconsidered in studies based on organellar DNA. Further, data from biparentally inherited DNA will often be needed in order to dig deeper into “difficult” phylogenetic questions, for example involving taxa of hybrid and/or allopolyploid origin, and nuclear data is therefore an important source of information. As in many phylogenetic studies, we selected the ribosomal DNA for our nuclear data set, DNA regions that are well documented and conveniently present in many copies in the genome, thus easy to sequence. The ribosomal DNA is, however, not useful for addressing questions involving reticulate evolutionary patterns because the many existing copies undergo a process of homogenization, or concerted evolution (Elder & Turner, 1995), that with time will mask the original genetic information in taxa of hybrid origin. It will therefore be necessary to utilize other parts of the nuclear DNA in order to further explore the nature and causes of the conflicting patterns we observe.

4.4 More on Rubiaceae relationships

4.4.1 The unplaced tribes Luculieae and Coptosapelteae

Two South East Asian tribes, the Luculieae and the Coptosapelteae, have been notoriously difficult to associate with any other taxa (Robbrecht & Manen, 2006; Bremer, 2009; Rydin et al., 2009a). Both tribes have commonly been placed in an unresolved “basal” position outside of the three recognized subfamilies (Fig. 1), sometimes together as sister groups (Bremer, 2009; Rydin et al., 2009a; Manns et al., 2012), but more commonly not (Bremer et al., 1999; Robbrecht & Manen, 2006; Bremer & Eriksson, 2009; Manns et al., 2012; Wikström et al., 2015; Yang et al., 2016). Robbrecht & Manen (2006) considered their analyses of nuclear versus plastid data to show conflicting results. Their nuclear data placed the tribes together, inside Rubiaceae and as sister to a clade comprising subfamilies Cinchoideae and Ixoroideae. In contrast, their analyses of plastid data placed them separately, outside of Rubiaceae and sisters to the entire family, or separately with Coptosapelteae outside and sister to the entire family and
Luculieae as sister to subfamily Rubioideae (Robbrecht & Manen, 2006). Rydin et al. (2009a) also included nuclear data in their analyses of deep divergences in the family, in their case data from the ITS rDNA spacer regions, and they reported similar patterns. Their combined analyses of plastid and nuclear ITS data resolved Luculieae and Coptosapelteae together, and in an unresolved “basal” position in the family, whereas their analyses excluding the nuclear ITS data resolved the two tribes separately (Rydin et al., 2009a). Again, their limited amount of non-plastid data prevented them to further evaluate this potential conflict. One clear indication as to the relationships of at least one of these tribes came recently from an analyses of mitochondrial data by Rydin et al. (2017). The relationships of Luculieae remained poorly supported in their analyses, but Coptosapelteae was unequivocally supported as sister to a clade comprising the two subfamilies Cinchonoideae and Ixoroidae. There were, however, several supported conflicts in their analyses of mitochondrial data compared to corresponding analyses of plastid data, specifically with respect to tribal relationships in the subfamilies Cinchonoideae and Ixoroidae. As argued by Rydin et al. (2017), these conflicts potentially result from one, or several, ancient hybridization events between early occurring members of the two subfamilies and, if correct, could make strict comparisons with the plastid-based results presented here invalid.

Despite the substantial increase in the amount of data analyzed, uncertainties surrounding the relationships of these two tribes remain also in the analyses presented here. Both the nucleotide based (Fig. 2), and the degenerate codon (Fig. 3) analyses of plastid CDS data place the tribes separately, and in poorly supported “basal” positions together with the subfamily Rubioideae. Nucleotide analyses of combined CDS and non-CDS data (Fig. S1) also have poor support for placing Coptosapelteae, but resolve Luculieae as sister to subfamily Rubioideae with good support (BPP = 0.98; BS = 89%). Although consistent with most previous analyses of plastid data (Bremer et al., 1999; Robbrecht & Manen, 2006; Bremer & Eriksson, 2009; Manns et al., 2012; Wikström et al., 2015), the continued lack of support for their relationships in most analyses is problematic. Also the nuclear rDNA analyses produce results consistent with those from previous analyses including nuclear data, with Luculieae and Coptosapelteae placed together in a monophyletic group, and in an unresolved “basal” position in the family (Robbrecht & Manen, 2006; Rydin et al., 2009a). The analyses that only include the conserved and unambiguously aligned rDNA gene regions provide unambiguous support for grouping the two tribes in the Bayesian analyses (Fig. 4), but the grouping is not supported by the ML bootstrap, and the Bayesian support is significantly reduced when also the variable and less easily aligned ITS spacer regions are included in the analyses (Fig. S2). This last result is somewhat surprising as the ITS regions were the only nuclear regions included in the analyses by Rydin et al. (2009a), and the ones supporting their grouping of Luculieae and Coptosapelteae.

An alternative to adopting a “more characters” phylogenomic approach for improving our phylogenetic inferences, is to expand on the taxon sample (Heath et al., 2008), as done by Rydin et al. (2009a) in their analyses of deep divergences in the family. Luculieae comprise a single genus with no more than four species, and adding further species to our analyses would likely be of limited value. Coptosapelteae, on the other hand, comprise 2 genera and altogether 56 recognized species, and Acranthera Arnott, the genus not represented in our analyses, was only recently included in the tribe (Rydin et al., 2009a). Morphologically, the genus also deviates significantly from Coptosapelteae, and expanding the taxon sample to also include Acranthera should be done in future attempts at resolving the relationships of this problematic group.

4.4.2 Subfamily Rubioideae

The early diverging tribes in the subfamily Rubioideae, the Ophiirrhizheae, Urophylleae, and Colletoecemateae, also remain problematic. Earlier analyses of plastid data have sometimes placed and supported Colletoecemateae as sister to remaining tribes in the subfamily (Robbrecht & Manen, 2006; Rydin et al., 2008, 2009a), but there is no support for this early split in the subfamily when using much larger sets of genomic data, neither based on plastid and nuclear data (the present study), nor based on mitochondrial data (Rydin et al., 2017). Relaxed-clock analyses of plastid data have instead resolved the Colletoecemateae as sister to the Lasiantheae (Manns et al., 2012; Wikström et al., 2015), but also this relationship is rejected here (Figs. 2–4). Here, the degenerate codon analyses of plastid CDS data, as well as the nucleotide based analyses of combined CDS and non-CDS data (Fig. S1), support Ophiorrhizheae as sister to remaining tribes of the subfamily, but the positions of Urophylleae and Colletoecemateae remain unsupported. In contrast to our plastid-based analyses, our analyses of nuclear rDNA data resolve and support the three tribes as a monophyletic group, sister to remaining tribes in the subfamily (Figs. 4, S2). However, as no support is obtained for the relationships of the tribes Urophylleae and Colletoecemateae in the analyses of plastid data, the different relationships indicated for the three tribes cannot be unambiguously attributed to conflicting phylogenetic signals in the two data types. Rydin et al. (2017), in their analyses of mitochondrial data, found strong support for a third alternative with the Colletoecemateae as sister to the Urophylleae, and with this Colletoecemateae–Urophylleae clade sister to remaining tribes of the subfamily. The grouping of the three tribes in a monophyletic group, as supported by our analyses of nuclear rDNA data (Figs. 4, S2), clearly conflicts with this mitochondrial based result reported by Rydin et al. (2017).

In the Spermacoceae alliance the Danaideae were for some time considered problematic. Some analyses placed the Danaideae sister to remaining tribes of the Spermacoceae alliance (Bremer & Manen, 2000; Bremer, 2009; Bremer & Eriksson, 2009; Rydin et al., 2009b; Yang et al., 2016), or sister to a clade comprising the Anthospermeae, Argostemmateae, Paederieae, Rubieae, and the Theligoneae (Robbrecht & Manen, 2006), but more recent analyses of plastid data and mitochondrial data respectively, have more or less firmly placed the tribe as sister to a Knoxieae–Spermacoceae clade (Rydin et al., 2009a; Krüger et al., 2012; Krüger, 2014; Wikström et al., 2015; Rydin et al., 2017). Here however, the placement of Danaideae is once again challenged since our analysis of nuclear data support Danaideae as member of a clade also comprising Durnnieae, Argostemmateae, Paederieae, Rubieae, Theligoneae
and Putorieae. Among these remaining tribes of the alliance the Rubieae, Theligoneae, and the Putorieae have consistently formed a monophyletic group (Robbrecht & Manen, 2006; Backlund et al., 2007; Rydin et al., 2009b; Wikström et al., 2015; Yang et al., 2016; Rydin et al., 2017), and together they have commonly been placed in a sister group relationships to the Paederieae (Robbrecht & Manen, 2006; Backlund et al., 2007; Rydin et al., 2009a, 2009b; Yang et al., 2016). All these relationships in the Spermacoceae alliance are corroborated and well supported also in the present analyses of plastid data (Figs. 2, 3, S1). The relationships of the Anthospermeae, Argostemmataceae, and Dunnieae have been more problematic. Rydin et al. (2009b) specifically addressed tribal relationships in the alliance and included a comprehensive sample of taxa from all tribes in their analyses. They resolved the Anthospermeae, Argostemmataceae, and Dunnieae as successive sister groups to the Rubieae–Theligoneae–Putorieae–Paederieae clade (Rydin et al., 2009b). Their position of the Anthospermeae as the first diverging group was well supported, and this position has been supported also in other analyses (Rydin et al., 2009a; Wikström et al., 2015). The relative positions of the Argostemmataceae and Dunnieae in their analyses were, however, unsupported (Rydin et al., 2009b). Our analyses of plastid data support the Anthospermeae, Dunnieae, and Argostemmateae as successive sister groups to the Rubieae–Theligoneae–Putorieae–Paederieae clade (Figs. 2, 3, S1), and these relationships are entirely consistent with those supported by Rydin et al. (2009b).

Among conflicting tribal relationships in the Spermacoceae alliance supported in our analyses of nuclear rDNA data, the position of Anthospermeae as sister to the Knoxieae–Spermacoceae clade is perhaps the most apparent and striking example. The tribe Anthospermeae comprise 12 genera with around 200 species (Puff, 1982; Thureborn et al., 2019), and although previous analyses of a single or a few genes from the plastid genome have been unable to find support for their precise relationships to other tribes in the alliance, a sister group relationship to the Knoxieae–Spermacoceae clade have never before been reported. A second example is the position of the Coussareae, a morphologically heterogeneous group of Neotropical plants with 10 different genera and a possibly 400 species (Andersson & Rova, 1999; Bremer & Manen, 2000; Löfstrand et al., 2019). Previous analyses of plastid data (Bremer & Manen, 2000; Rydin et al., 2008, 2009a, 2009b; Wikström et al., 2015), the analyses of plastid data presented here (Figs. 2, 3, S1), as well as analyses of mitochondrial data (Rydin et al., 2017) all support a position of the Coussareae as sister to a clade comprising the Spermacoceae and the Psychotrieae alliances. In contrast however, the analyses of nuclear rDNA data presented here support a position of the Coussareae nested inside the Spermacoceae alliance (Figs. 4, S2).

In the Psychotrieae alliance our analyses of plastid data more or less corroborate patterns reported in previous analyses of the group. One exception is Prismatomerideae, here resolved sister to the Gaertnerieae. This sister relationship is, however, only supported in our nucleotide based analysis of plastid CDS and combined CDS and non-CDS data (Figs. 2, S1), not in the degenerate codon analyses of CDS data, where silent substitutions are ignored (Fig. 3), and it has not been seen in previous analyses of the group (Robbrecht & Manen, 2006; Razafimandimbison et al., 2008, 2017; Wikström et al., 2015). Based on our analysis of nuclear data, previously reported relationships in the Psychotrieae alliance are also corroborated with one exception, the position of the tribe Gaertnerieae, which is supported as sister to the Psychotrieae–Palioureae clade in the analysis of rDNA data (Figs. 4, S2), but more closely related to Mitchelleae, Morindeae, and Prismatomerideae based on plastid data (Figs. 2, 3, S1). Mitochondrial data support the sister group relationship of Gaertnerieae to a Psychotrieae–Palicourae clade (Rydin et al., 2017), and yield entirely congruent results with nuclear ribosomal data (the present study) in the Psychotrieae alliance.

Notwithstanding these conflicting signals in plastid and nuclear data, in subfamily Rubieae the analyses of nuclear rDNA data also show considerable congruence to the results of the plastid-based analyses, and thus to the coherent picture of phylogenetic relationships in the family developed during the last 20–30 years. The Rubieae, Theligoneae, and Putorieae, for example, are resolved as monophyletic, which is consistent with most previous analyses (see Bremer, 2009; Rydin et al., 2009b). This Rubieae–Theligoneae–Putorieae clade are grouped with the Paederieae, Argostemmataceae, and Dunnieae (Rydin et al., 2009b; Wikström et al., 2015), although the nuclear rDNA analyses also include the Danaideae in this group. The Spermacoceae alliance are supported as monophyletic (Andersson & Rova, 1999; Bremer & Manen, 2000; Robbrecht & Manen, 2006; Rydin et al., 2009b; Wikström et al., 2015), even though the nuclear rDNA data also place the Coussareae in this group. The Knoxieae and Spermacoceae are supported as sister groups (Bremer, 1996; Robbrecht & Manen, 2006; Kårehed & Bremer, 2007; Rydin et al., 2009b; Wikström et al., 2015). The Psychotrieae alliance are supported as monophyletic (Bremer & Manen, 2000; Robbrecht & Manen, 2006; Rydin et al., 2008; Razafimandimbison et al., 2008, 2017), and the sister group relationships of the Morindeae and Mitchelleae, and the Psychotrieae and Palicourae are both supported (Razafimandimbison et al., 2008; Rydin et al., 2008; Wikström et al., 2015).

4.4.3 Subfamilies Cinchonoideae and Ixoroidae

The “more characters” approach adopted here also provides some additional resolution to relationships that have been problematic in subfamilies Cinchonoideae and Ixoroidae, at least with respect to the relationships of the maternally inherited plastid. Disregarding the conflicting signal in subfamily Cinchonoideae, caused by synonymous substitutions (see discussion above), relationships among the four previously identified and well supported groups in the Cinchonoideae are with one exception supported in our analyses of plastid data. Still problematic is the position of the Hymenodictyeae–Naucleeae clade, here resolved in a poorly supported position as sister to the Rondeletieae–Guettardae clade (Fig. 3). This position as sister to the Rondeletieae–Guettardae clade is, however, unambiguously supported by analyses of mitochondrial data (Rydin et al., 2017).

In the subfamily Ixoroidae our plastid data analyses mainly corroborate relationships obtained in recent
analyses of the group where the alternative “more taxa” approach was used (Kainulainen et al., 2013, 2017; Mouly et al., 2014). Relationships among the early diverging Condamineae, Mussaendaeae–Sabiceaeae, and Henriquezaeae–Possuquerieae–Sipaneae clades have been difficult to support (Kainulainen et al., 2013; Wikström et al., 2015), but the main problem has been the position of the tribe Condamineae, which we were unable to include in the present analyses. In the Vanguerieae alliance the small tribes Scyphiphoreae and Trailliaedoxeae, and the genus Glionnetia from the Seychelles, currently unclassified at tribal level, have been problematic (Razafimandimbison et al., 2011; Kainulainen et al., 2013; Wikström et al., 2015). Glionnetia have been supported in a group also including the tribes Vanguerieae, Greeneeeae, Ixoreeae, and Aleisantheia (Kainulainen et al., 2013; Wikström et al., 2015), but the relationships between Glionnetia, the Vanguerieae, and a Greeneeeae–Ixoreeae–Aleisantheia clade have been unsupported. The tribes Scyphiphoreae and Trailliaedoxeae have commonly been placed in an unresolved sister group positions to this group of four tribes and Glionnetia (Razafimandimbison et al., 2011; Kainulainen et al., 2013; Wikström et al., 2015). Our analyses of plastid data provide some support for the relationships of Glionnetia. In the nucleotide based and degenerate codon analyses of CDS data Glionnetia remain resolved in a corresponding polytomy as in previous analyses, together with the Vanguerieae and a Greeneeeae–Ixoreeae clade, but a sister group relationship to these tribes obtains support in our nucleotide-based analyses of combined CDS and non-CDS data (Fig. S1). In contrast, there is good support for the relationships of the Scyphiphoreae and Trailliaedoxeae, which are here supported as sister groups to each other, and together supported as sister to the Glionnetia–Vanguerieae–Greeneeeae–Ixoreeae clade (Fig. 3). Also in the Coffeaeae alliance we see only minor differences in the results from our analyses of plastid data compared to analyses that have adopted a “more taxa” approach (Kainulainen et al., 2013, 2017; Mouly et al., 2014). They all supported monophyly of a group comprising four tribes, the Gardenieae, Pavetteae, Sherbourieae, and Cordierae, and with Octotropidea sister to this group. Here, the tribe Octotropidea is instead resolved sister to the Cordierae and they together are sister to a Gardenieae–Pavetteae–Sherbourieae clade (Figs. 2,3). Octotropidea as sister to Cordierae is, however, not supported by the nucleotide analyses of combined CDS and non-CDS data (Fig. S1).

5 Conclusions

It is important to note that most phylogenetic results in the large coffee family are consistently retrieved, regardless of kind of data and analytical approach. However, our study here together with our analyses of mitochondrial data (Rydin et al., 2017) reveals that there are (not so few) clades that are well supported by data from one or two of the genomic compartments, but not from all three, and some of these conflicts may be “real”, reflecting biological processes of the past. A striking example concerns the deepest splits in the family and the previously unresolved position of the small tribe Coptospapelteae, which we show is sister to subfamily Rubioideae plus Luculieae based on plastid data (the present study), sister to Luculieae based on nuclear data (the present study), and sister to Cinchonoideae plus Ixoroideae based on mitochondrial data (Rydin et al., 2017). We also show that phylogenetic questions that have been considered answered are in fact not. For example, the neotropical tribe Coussareae have been consistently resolved as sister to a large clade comprising the Psychotrieae and the Spermacoceae alliances (see e.g., Löfstrand et al., 2019), a result seemingly supported by data from all three genomic compartments including the mitochondrion (Rydin et al., 2017). Here we show, analyzing the entire cistron, that the nuclear ribosomal DNA with strong support places the Coussareae in the Spermacoceae alliance, thus strongly conflicting with our results based on data from the plastome and the mitochondrion. Adding genomic data for additional taxa may be relevant in order to address some still unresolved parts of the Rubiaceae phylogeny, but for the future it will most of all be important to take the possibility of reticulate evolutionary patterns into consideration. A straightforward way will be to utilize biparentally inherited low-copy genes from the nuclear genome, an approach that should be adopted in future analyses trying to resolve these relationships.

Acknowledgements

The authors would like to acknowledge support from Science for Life Laboratory, the National Genomics Infrastructure, NGI, and Uppmax for providing assistance in massive parallel sequencing and computational infrastructure. The project was funded by grants from the Royal Swedish Academy of Sciences, Stockholm University, and the Swedish Research Council (A0529602 to B.B. and 2017-03985 to C.R.).

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Supplementary Material

The following supplementary material is available online for this article at http://onlinelibrary.wiley.com/doi/10.1111/jse.12566/supinfo:

Fig. S1. Phylogram resulting from nucleotide based bayesian analysis of the combined plastid CDS and non-CDS data sets. Nodes indicated by a black dot are well supported and have a bayesian posterior probability (BPP) of 0.95 or more (Alfaro et al., 2003; Erixon et al., 2003). Maximum Likelihood non-parametric bootstrap support (BS) is also indicated for each node in the tree. Support values are 1.00 (BPP) and 100% (BS) unless indicated in the figure. Subfamilies Rubioideae, Cinchonoideae, and Ixoroideae, and the Psychotrieae, Spermacoceae (Rubiodeae), and Vanguerieae, Coffeeae (Ixoroideae) alliances as traditionally recognized are indicated.
on the tree. Taxa in blue show conflicting relationships in different analyses of data from the same (plastid) genomic compartment (see Fig. 3). Taxa in red show conflicting relationships in analyses of data from different genomic compartments (see Figs. 4, S2).

**Fig. S2.** Phylogram resulting from the MrBayes analyses of nuclear rDNA data including also the variable ITS1 and ITS2 spacer regions. Ambiguous ITS1 and ITS2 alignment positions were filtered out prior to the analyses using transitive consistency scores (TCS; Notredame et al., 2000; Chang et al., 2014). Alignment positions with a TCS score of 3 or less were considered ambiguous and 360 of the total 648 ITS spacer characters were excluded from the analyses. Nodes indicated by a black dot are well supported and have a bayesian posterior probability (BPP) of 0.95 or more (Alfaro et al., 2003; Erixon et al., 2003). Maximum Likelihood non-parametric bootstrap support (BS) is also indicated for each node in the tree. Support values are 1.00 (BPP) and 100% (BS) unless indicated in the figure and nodes with BPP < 0.50 are collapsed. Subfamilies Rubioideae, and Cinchonoideae plus Ixoroideae as traditionally recognized are indicated on the tree. Taxa in red show conflicting relationships in analyses of data from different genomic compartments (see Figs. 2, 3).