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Pseudomolecule-scale genome assemblies of Drepanocaryum sewerzowii and Marmoritis complanata

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The Nepetoideae, a subfamily of Lamiaceae (mint family), is rich in aromatic plants, many of which are sought after for their use as flavors and fragrances or for their medicinal properties. Here, we present genome assemblies for two species in Nepetiodeae: *Drepanocaruym sewerzowii* and *Marmoritis complanata*. Both assemblies were generated using Oxford Nanopore Q20 + reads with contigs anchored to nine pseudomolecules that resulted in 335 Mb and 305 Mb assemblies, respectively, and BUSCO scores above 95% for both the assembly and annotation. We furthermore provide a species tree for the Lamiaceae using only genome-derived gene models, complementing existing transcriptome and marker-based phylogenies.

Keywords: Lamiaceae; Nepetinae; chromosome-level assembly; nanopore sequencing; Hi-C sequencing

Introduction

The mint family (Lamiaceae) is the sixth largest plant family with a number of species regarded as important for medicinal, aromatic, and ornamental properties (Harley *et al.* 2004; Zhao *et al.* 2021; Rose *et al.* 2022). Within the Lamiaceae, species from the Nepetoideae are renowned for the accumulation of terpenoids, with tissues used for the extraction of essential oils or as traditional herbal medicines (Wink 2003; Frezza *et al.* 2019). The clade includes widely recognized aromatic species such as mint, lavender, lemon balm, and catnip; the volatile terpenoids produced by these plants are responsible for their characteristic fragrances. The ethnobotanical and commercial relevance of this plant family has resulted in considerable scientific interest, including genome assemblies for 36 species at the time of writing (Published Plant Genomes).

Here, we present the genome assemblies for two Nepetoideae species, namely *Drepanocaryum sewerzowii* (Regel) Pojark. and *Marmoritis complanata* (Dunn) A.L.Budantzev. *M. complanata* is endemic to the subnival band of the Himalaya–Hengduan Mountains, a unique arctic–alpine region recognized as a biodiversity hotspot (Myers et al. 2000; Sun et al. 2017). This unique habitat necessitates careful control of seed germination to ensure survival (Peng et al. 2018). *M. complanata* and other species of the genus are also used as traditional herbal medicines to treat a variety of ailments that include digestive, reproductive, musculoskeletal, and skin disorders (Zaman et al. 2022). *D. sewerzowii* is native to

a region that ranges from Iran to Central Asia and Pakistan and is the sole representative of this genus (Serpooshan et al. 2018).

These two species are part of the Nepetinae, a subtribe of the mint family (Lamiaceae, subfamily Nepetoideae, tribe Mentheae) that consists of 375 species and 9-12 genera of which Nepeta L. is considered the type genus encompassing 200–300 species. Other genera in this subfamily include Dracocephalum L., Hymenocrater Fisch. and C.A. Mey., Lophanthus Adans., Agastache Clayton ex Gronov., and Schizonepeta (Benth.) Briq. (Serpooshan et al. 2018; Rose et al. 2023). The phylogenetic relationship of M. complanata and D. sewerzowii relative to Nepeta cataria L., N. racemosa Lam., Agastache rugosa (Fisch. and C.A. Mey.) Kuntze and Schizonepeta tenuifolia (Benth.) Briq. is what prompted our efforts to assemble these genomes. We have been exploring the evolutionary, genomic, and enzymatic innovations of monoterpenoid biosynthesis in these species (Lichman et al. 2019, 2020; Hernández Lozada et al. 2022; Liu et al. 2023). However, the available genomic resources provide limited taxonomic coverage. The genome assemblies presented here will allow us to further explore the evolutionary innovations that have impacted terpenoid biosynthesis in the mint family.

Methods and Materials Plant growth conditions

D. sewerzowii seeds were obtained from the Millennium Seed Bank at the Royal Botanic Gardens, Kew (serial no. 0694027).

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M. complanata seeds were collected from Puyong Pass Shangri-la County, Yunnan Province, SW China (99°55′E, 28°24′N), 4,620 m a.s.l. (Peng et al. 2018). Seeds were germinated on 1% water agar in a growth room set to 16 h day length, temperature of 20 (±2) °C, relative humidity of 60% (±10%), and a NS12 light spectrum at 120 µmol m⁻² s⁻¹ PPFD using Valoya L28 LED lights (Helsinki, Finland). Once a radical emerged, the seedlings were transferred to 7 cm square pots containing Levington Advance Seed and Modular FS2 (ICL Professional Horticulture) seedling soil that was pretreated with Calypso (Bayer). Once established, a single individual was selected and maintained as a clonal population by propagation using cuttings.

Genome size and heterozygosity estimation

Genome size estimations were determined through flow cytometry (FCM) using the method of Dolezel *et al.* (2007). Briefly, the LB01 buffer was used together with *N. cataria* and *N. racemosa* tissues to prepare a reference standard with previously reported genome sizes (Mint Evolutionary Genomics Consortium 2018). A CytoFLEX LX (Beckman Coulter) FCM with a 561 nm excitation laser, 610/20 emission filter, and a flow rate of 30 μ L/min was used. The threshold was set to 488 nm forward scatter to exclude instrument noise and background signal from the buffer. Fluorescence intensities of stained nuclei were used to analyze the nuclear DNA content and determine the genome size relative to the aforementioned *Nepeta* spp. reference samples.

Paired-end 150 Illumina short reads were used to estimate genome size and heterozygosity by determining k-mer frequencies with KMC v3.1.2 (Kokot *et al.* 2017) using a k-mer of 21 and transformed into a histogram using kmc_tools with a count cutoff of 15,000. Genomescope v2.0 (Ranallo-Benavidez *et al.* 2020) with ploidy set to 2 was used to estimate the genome size and level of heterozygosity.

Nucleic acid isolation

High molecular weight DNA isolation and sequencing

High molecular weight (HMW) DNA was extracted in duplicate from ~1 g of young leaf tissue using the Nucleobond HMW DNA Extraction kit (Macherey-Nagel, Germany). HMW DNA purity and concentration were assessed by Nanodrop and Oubit, whereafter the extractions were combined. Small fragment DNA elimination was performed with the Circulomics short-read eliminator kit (PacBio). Briefly, an equal volume of SRE reagent was added to the sample, and this was centrifuged for 1 h at 12,000×g. The pellet was washed with 70% ethanol before resuspending in TE buffer with low EDTA. DNA quality and quantity were assessed with a nanodrop spectrophotometer (Thermo Fischer Scientific), Agilent Tapestation (running genomic DNA screentape), and Qubit fluorimeter (Invitrogen). Sequencing was performed with the ligation sequencing kit SQK-LSK114 (Oxford Nanopore Technologies), as per the manufacturer's guidelines, with limited modifications; namely extending the reaction times for end preparation to 30 min at each temperature, and extending adapter ligation steps to an hour). Sequencing was performed on a single promethION FLO-PRO114 flowcell (Oxford Nanopore Technologies) per species, with nuclease flush and sample reload steps performed every 24 h through the run time. For M. complanata two additional runs using the SQK-LSK112 ligation sequencing kit (Oxford Nanopore Technologies) and FLO-MIN112 minION flowcells (Oxford Nanopore Technologies) were performed.

Base calling of the ONT reads was performed using guppy (Oxford Nanopore Technologies) version 6.1.5 for D. sewerzowii and version 6.3.9 for *M. complanata* with the super accuracy (sup) model. Read length and quality were assessed using Nanoplot (De Coster and Rademakers 2023). *D. sewerzowii* reads were filtered for a 10 kb minimum length using Nanofilt (De Coster *et al.* 2018). For *M. complanata* we combined all reads from the promethION and minION runs and then filtered using Nanofilt (De Coster *et al.* 2018) with a 3 kb length and Q15 quality cutoff.

Genomic DNA isolation and Illumina sequencing

Genomic DNA (gDNA) was extracted from 100 mg of young leaf tissue, in duplicate, using a CTAB extraction method (Doyle and Doyle 1990) and treated with RNAse A. Removal of RNA was confirmed through gel electrophoresis followed by gDNA quality and quantity assessment with a nanodrop spectrophotometer and a Qubit fluorometer (Invitrogen). A total of 508 ng and 752 ng of gDNA for *D. sewerzowii* and *M. complanata*, respectively, was sent for library preparation and paired-end Illumina sequencing with Novogene (Cambridge, UK).

RNA isolation and sequencing

RNA was extracted from 80 to 100 mg of tissue with the Direct-Zol RNA extraction kit (Zymo Research, CA, USA) as per the manufacturer guidelines. For *D. sewerzowii*, young and mature leaves, closed and open flowers, and stems were used. For *M. complanata* root, young and mature leaf and stem tissues were used. RNA quality was assessed with an Agilent bioanalyzer. Library preparation and paired-end Illumina sequencing were performed by Novogene (Cambridge, UK).

Hi-C sequencing

Freshly harvested young leaf tissue was fixed in 1% formaldehyde and washed as per the Phase Genomics (Seattle, WA, USA) sample preparation protocol. Following fixation, the tissue was flash frozen in liquid nitrogen and homogenized using a tissue lyser. The Hi-C libraries were prepared and sequenced by Phase Genomics.

Genome assembly

Filtered nanopore reads for the respective genomes were used for assembly and error correction. Both species were first assembled using Flye v2.9 (Lin *et al.* 2016; Kolmogorov *et al.* 2019; --iterations 0 and --nano-hq flags). *M. complanata* was also assembled with NECAT v0.0.1 (Chen *et al.* 2021) using the default configuration file settings. Our error correction pipeline entailed polishing with long reads by two rounds of RACON v1.5 (Vaser *et al.* 2017), with reads mapped using minimap2 v2.24 (Li 2018), followed by two rounds of MEDAKA v1.6 (Medaka: Sequence Correction Provided by ONT Research 2018) polishing. Short reads were mapped using bwa-mem v0.7.17 (Li 2013) and duplicate reads were marked using Picard v2.25.5 ("Picard Toolkit" 2019) prior to two iterative rounds of polishing with Pilon v1.23 (Walker *et al.* 2014). Short-read alignment and pairing rates were determined using flagstat command from samtools v.1.17 (Danecek *et al.* 2021).

The M. complanata Flye and NECAT assemblies were merged with Quickmerge v0.3 (Chakraborty et al. 2016; Solares et al. 2018) due to the low N50 scores. The overlap cutoff (-c flag) was 5 and the length cutoff (-l) was 100,000 with the NECAT assembly used as the query. The NECAT-Flye merged assembly underwent another two rounds of short read error correction using Pilon. For M. complanata, we purged the merged assembly of haplotigs prior to HiC scaffolding while D. sewerzowii was purged after HiC scaffolding. Haplotig purging was performed using the purge haplotigs pipeline (Roach et al. 2018). Contigs were scaffolded into pseudomolecules by Phase Genomics using the Proximo Genome Scaffolding Platform. Contiguity and completeness were assessed throughout the assembly pipeline using BUSCO (Benchmarking for University Single Copy Orthologs) v5.4.2 with the embryophy-ta_odb10 dataset (Manni *et al.* 2021).

Genome annotation

Repeats and transposable elements were annotated using the Earl Grey v3.2 (Baril et al. 2023, 2024) pipeline with default settings followed by softmasking of the repeats using the maskfasta function of bedtools. The BRAKER3 pipeline (v3.0.6; Stanke et al. 2006, 2008; Gotoh 2008; Iwata and Gotoh 2012; Buchfink et al. 2015; Hoff et al. 2016, 2019; Kovaka et al. 2019; Pertea and Pertea 2020; Brůna et al. 2021; Bruna et al. 2024) was used to predict gene models using mRNA and protein evidence. For protein evidence, we generated a representative database from 52 Mint species (48 Lamiaceae and 4 from Lamiales families) using the transcriptomes from (Mint Evolutionary Genomics Consortium 2018). MMseqs2 (Steinegger and Söding 2017) was used to remove identical sequences from the database. For mRNA evidence we aligned RNAseq reads from the different tissues using STAR v2.7 (Dobin et al. 2013) with default settings and the "--outSAMstrandField intronMotif" flag. The respective bam outputs were merged using samtools (Danecek et al. 2021) and used as input for BRAKER3. The BRAKER annotation output was reformatted to GFF3 using AGAT v1.1 (Dainat et al. 2020) followed by extraction and translation of the longest open-reading for each predicted coding sequence. Annotation completeness was assessed using BUSCO v5.4.2 (Manni et al. 2021) in protein mode with the embryophyta_odb10 dataset.

Functional annotations were assigned by searching predicted proteins against the Arabidopsis TAIR10 proteome (Lamesch et al. 2012) and Magnoliopsida (taxon ID 3398) Swiss-Prot database (2024_03 Release; UniProt Consortium 2023). Searches were performed using DIAMOND v2.1.0 (Buchfink et al. 2021) with an *e*-value of 0.001 and retaining the top hit only. Functional domains were assigned with hmmsearch through HMMER v3.4 (http://hmmer.org/) against the PFam v37.0 database (Mistry et al. 2021; UniProt Consortium 2023) with an *e*-value cutoff of 0.001 with the highest scoring domain hit retained.

Species tree and macrosynteny analysis

Markerminer v1.0 (Chamala et al. 2015) was used to identify singlecopy genes using predicted coding genes from representative Lamiaceae genomes (Supplementary Table 1) and Paulownia fortunei (Seem.) Hemsl. as an outgroup. Genes present in 26 of the 27 species were included. The MAFFT alignments generated as part of the Markerminer pipeline were trimmed for gaps using the gappyout algorithm of trimAl v1.4.1 (Capella-Gutiérrez et al. 2009) and concatenated into a supermatrix with partitions using the catfasta2phyml script (https://github.com/nylander/ catfasta2phyml). A species-tree was inferred by maximum likelihood with partition models (Chernomor et al. 2016) using IQ-TREE 2 (Minh et al. 2020) with ModelFinder (Kalyaanamoorthy et al. 2017), ultrafast bootstraps (UFBoot2, ×1000; Hoang et al. 2018), and SH-aLRT supports (x1000; Guindon et al. 2010). In addition, a species tree using protein sequences was inferred using the STAG (Species Tree inference from All Genes) method of Orthofinder v2.5.4 (Emms and Kelly 2015, 2017, 2018, 2019). Pairwise macrosynteny analyses were performed against A. rugosa (Park et al. 2023) and S. tenuifolia (Liu et al. 2023) using the JCVI v.1.2.7 (Tang et al. 2024) implementation of MCScan (Tang et al. 2008). MCScan orthologs were identified in full mode with predicted protein sequences and default settings.

Expression analysis

RNAseq read alignments were evaluated with STAR v2.7 (Dobin et al. 2013) and assed with qualimap v2.2.1 (García-Alcalde et al. 2012; Okonechnikov et al. 2016). Qualimap reports were aggregated with MultiQC v1.13 (Ewels et al. 2016). Expression counts as transcripts per million (TPM) were generated using Salmon v1.10.0 (Patro et al. 2017). The transcript index for Salmon was generated using the full set of predicted coding sequences from BRAKER3.

Results and discussion

Genome size and heterozygosity estimation

The genome size was estimated by fluorescence of stained nuclei (Dolezel *et al.* 2007) relative to that of *N. cataria* and its previously reported genome size (Mint Evolutionary Genomics Consortium 2018). *D. sewerzowii* was estimated to be 330 Mb and *M. complanata* to be 337 Mb in size. The k-mer estimation of genome size (Supplementary Fig. 1) was similar to that of FCM with a genome size of 334 Mb and 328 Mb for *D. sewerzowii* and *M. complanata*, respectively. K-mer analysis estimated the heterozygosity at 0.08% and 0.6% for *D. sewerzowii* and *M. complanata*, respectively.

Chromosome level assemblies

We sequenced the genomes for D. sewerzowii and M. complanata using Oxford Nanopore long reads and Proximo HiC scaffolding (Phase Genomics) resulting in two chromosome-level assemblies (Fig. 1). A total of 99.24 Gb of super accurate nanopore reads were generated for D. sewerzowii with 80 Gb of reads being greater than 10 kb at a mean read quality (Q-score) of 16.6. The size filtered reads provided 242 × coverage when using the genome size determined through FCM (Supplementary Fig. 1). The initial Flye assembly resulted in 472 contigs, an N50 of 17 Mb, a total assembly length of 333.75 Mb and a BUSCO score of 98.7%. Polishing with long and short reads reduced the number of contigs to 134 and the assembly size to 332.85 Mb while maintaining an N50 of 17 Mb. The BUSCO score increased slightly to 98.8% after polishing. HiC scaffolding orientated the assembly to 9 pseudomolecules (Supplementary Fig. 2a), which is in agreement with the chromosome counts reported by Bordbar (2023). The 9 pseudomolecules contained 97.6% of the contigs, representing 324.87 Mb of the total assembly at an N50 of 35.2 Mb and L50 of 5 (Table 1).

We obtained 59.3 Gb of reads after length and quality filtering, providing 176 × coverage when using the 337 Mb FCM genome size estimation with a mean Q-score of 18. We tried various different read filtering cutoffs for both length and quality with all attempts using Flye failing to reach an N50 greater than ~335 kb. After polishing the best Flye assembly was 420 Mb in size with an N50 of 335 kb, 3,001 contigs and a BUSCO score of 98.5%, of which 17.1% were duplicated. NECAT resulted in a more contiguous genome assembly of 457 Mb with an N50 of 1 Mb, 869 contigs, and 98.6% BUSCO, of which 36.7% were duplicated. The inflated genome size and high number of duplicate BUSCO genes suggested that the fragmented assemblies contained a high number of haplotigs (contigs of a single haplotype), that would artificially inflate genome size.

In an attempt to increase the continuity of the assembly (N50 score) we merged the Flye and NECAT assemblies. The NECAT assembly had fewer contigs and greater N50 and was, therefore, selected to be the query genome with the Flye assembly used to improve the query genome. We evaluated the impact of haplotig



Fig. 1. Circos plots for the genome assemblies of *D. sewerzowii* and *M. complanata* depicting density (1 Mb bins) of genes, total repeats, gypsy, and copia elements along the 9 pseudomolecules.

Table 1. Assembly and annotation metrics.

	D. sewerzowii	M. complanata
Assembly statistics		
Assembly size (Mb)	332.85	305.55
Number of pseudomolecules	9	9
N50 (Mb)	35.13	27.69
L50	5	5
L90	9	28
GC%	38.56	37.45
Number of Ns	2,600	18,800
Mapped short-reads	95%	92.2%
Properly paired short-reads	93.8%	84.1%
Annotation statistics		
Assembly BUSCO ^a	C: 99.0%	C: 95.7%
n = 1,440	S: 96.3%	S: 85.7%
	D: 2.7%	D: 10.0%
Annotation BUSCO ^a	C: 95.0%	C: 95.3%
n = 1,440	S: 92.1%	S: 86.1%
	D: 2.9%	D: 9.2%
Predicted coding genes	24,221	25,080
Predicted proteins	26,989	28,384
Percentage repeats	Total: 62%	Total: 53%
	DNA: 2.38%	DNA: 4.43%
	LINE: 0.53%	LINE: 2.92%
	LTR: 40.61%	LTR: 25.83%

^a Complete (C), single (S), duplicated (D).

purging before and after merging. Each assembly was purged of haplotigs prior to merging and compared to a merged assembly that was purged as the final step. In each iteration, we polished twice with short reads (18 Gb of PE150 reads) after merging. The merging increased the N50 to 3 Mb regardless of when we purged the haplotigs. The timing of the purging step had a large impact on the number of contigs together with a minor impact on the duplicated BUSCOs. Merging, polishing, and then purging the haplotigs resulted in the most contiguous assembly (305.6 Mb) with the fewest number of contigs (338) and a BUSCO score of 95.6%. HiC scaffolding assembled the contigs into 9 pseudomolecules (Supplementary Fig. 2b), which is in agreement with karyotype information (Sun 2016), totaling 258 Mb (85% of the total assembly). The pseudomolecules had a BUSCO score of 91.3% with the total assembly having a BUSCO of 95.7% (Table 1).

Pseudomolecule termini were manually inspected for the presence of the TTTAGGG telomeric repeat. Seven of the *D. sewerzowii* pseudomolecules contained this repeat on at least one end with Chr. 4 and 7 having it on both ends. For *M. complanata* we found this repeat on 6 pseudomolecules with Chr. 7 and 8 having it on both ends. The presence of this repeat on both ends indicates a telomere-to-telomere assembly for these chromosomes.

A limited number of genome assemblies are available for species within the Nepetinae with the nearest sequenced species, N. cataria and N. racemosa, having highly fragmented assemblies. We, therefore, compared our assemblies to the closest relatives with pseudomolecule assemblies, namely A. rugosa (Park et al. 2023) and S. tenuifolia (Liu et al. 2023). A. rugosa has a 9 chromosome assembly with macrosynteny revealing that Chr. 3, 4, and 6 have remarkably similar genomic structures for both assemblies reported here (Fig. 2). Pairwise comparisons (Supplementary Fig. 3) highlight the extent of genomic rearrangements within the Nepetinae, for example, fusion events resulted in the 6 chromosome structure of S. tenuifolia. D. sewerzowii, and M. complanata have very similar chromosome structures to each other (Supplementary Fig. 3); however, large relative inversions are present on M. complanata Chr. 3 and D. sewerzowii Chr. 2, with a translocation on Chr. 7. The high proportion of properly mapped and paired short-reads (Table 1) suggests that D. sewerzowii is a nearly complete assembly. The mapping and pairing rates for M. complanata short reads (Table 1) indicate a less accurate assembly, likely due to the higher level of heterozygosity (0.6%). Misassembly due to the merging of M. complanata NECAT and Flye assemblies may also be a reason for the lower read pairings. Higher accuracy reads and phasing will be required to fully resolve the problematic regions. Nevertheless, the M. complanata and D. sewerzowii genome assemblies provide a valuable genomic resource for intergeneric analyses.



Fig. 2. Pairwise macrosynteny analysis of the assembled genomes relative to the chromosome level assembly of A. *rugosa*. Conserved collinear blocks are linked by the gray lines.

Repeat and genome annotations

Repeat annotation revealed that 62% of the *D. sewerzowii* genome and 53% of the *M. complanata* genome are repeats (Table 1). The largest portion of the repeats were long terminal repeats (LTRs), occupying 40.6% and 25.8% of the respective genomes. Subsequent to repeat masking our gene annotation, using ab initio, protein and mRNA predictions, resulted in 24,221 and 25,080 gene regions that encode for 26,989 and 28,384 proteins for the respective genomes. BUSCO analysis of the primary isoforms was 95% for both genomes. Gene and repeat density showed an inverse relationship along the chromosomes (Fig. 1).

The RNAseq data we produced found evidence for the expression of the majority of genes. RNAseq reads mapped to gene models showed that 85% (22,794/26,815) of the genes were expressed in at least 1 tissue type for *D. sewerzowii* and 88% (24,919/28,384) of the genes in *M. complanata*. Expression matrices as TPM are available in Supplementary Tables 2 and 3 with functional annotations available in Supplementary Tables 4 and 5.

Phylogenies using genome-derived gene models

The phylogenetic relationships of the Lamiaceae have been reported using plastid, nuclear, and transcriptome approaches. We used genome-derived gene models to construct a species tree via phylogenomic inference, complementing existing species trees (Fig. 3; Mint Evolutionary Genomics Consortium 2018; Serpooshan *et al.* 2018; Rose *et al.* 2022, 2023). The STAG species-tree used multicopy gene families (i.e. orthogroups) predicted by Orthofinder using protein sequences. The consensus tree (Fig. 3a) shows internal bipartition support for 5,296 orthogroups in which all species are present. The ML tree (Fig. 3b) was inferred from a single-copy gene supermatrix totaling 340,706 nucleotide sites with all but two branches showing above 98% support for both ultrafast bootstraps and SH-aLRT. The two branches indicated by the asterisk were not well supported, bootstrap and SH-aLRT <85%.

In both the ML and STAG topologies, *D. sewerzowii* is recovered as a sister to a clade that includes *M. complanata* and *Nepeta*, which are sister to each other. While our phylogenomic results corroborate existing hypotheses regarding the close relationships among these genera, our trees are incongruent with previously reported topologies (Supplementary Fig. 4). For example, nuclear phylogenetic results by Rose et al. (2023) report *D. sewerzowii* as sister to *Nepeta*, which together are sister to the sister lineages *Hymenocrater* and (*Lophanthus* + *Marmoritis*). This contrasts with plastid-based phylogenetic results reported in the same study, which recover *Nepeta* as sister to a clade comprising the sister taxa, *Drepanocaryum* and *Hymenocrater*, and their sister, (*Lophanthus* + *Marmoritis*), and with results by Serpooshan *et al.* (2018), which recover *D. sewerzowii* as sister to a mixed and partially unresolved clade of *Hymenocrater*, *Lophanthus*, *Marmoritis*, and *Nepeta*. Topological discordances among trees reported in this and previous studies likely reflect differences in taxonomic and molecular sampling, but they also highlight the complexity of resolving intergeneric relationships within Nepetinae.

Understanding these intergeneric relationships is especially relevant in the context of specialized metabolism. Evolutionary innovations in the Mentha longifolia and S. tenuifolia lineages, for example, have resulted in the convergent evolution of biosynthetic genes for monoterpenoid biosynthesis (e.g. pulegone and menthol; Srividya et al. 2020, 2022; Vining et al. 2022; Liu et al. 2023). Similarly, iridoid biosynthesis in N. cataria and N. racemosa evolved through the concomitant emergence of iridoid biosynthetic genes (ISY, NEPS, and MLPL) that also colocalize to form a biosynthetic gene cluster (Sherden et al. 2018; Lichman et al. 2019, 2020). Comparative genomics in these species has suggested that genomic rearrangements and gene movement through ectopic recombination or transposition are likely involved in organizing pathway genes into BGCs (Smit and Lichman 2022). The speciestree presented here (Fig. 3) provide necessary context for further comparative genomics, although interpretations should be considered alongside available transcriptome- and marker-based phylogenies until additional Nepetinae genomes and phylogenomic results become available. Nevertheless, the genomes presented here provide a valuable resource to explore the evolutionary trajectories underpinning the remarkable innovations in specialized metabolism within the Lamiaceae.

Conclusion

Plant genome assemblies are being generated at a remarkable rate, with two-thirds of available plant genome assemblies generated within the last three years (Xie *et al.* 2024). Here, we present the chromosome-level genome assemblies of *D. sewerzowii* and *M. complanata*, representing the first assemblies from these genera. The gene and repeat annotations, along with expression matrices, present a comprehensive resource for comparative genomics. The species-tree using gene models from available Lamiaceae genome assemblies provides a reference point that celebrates the number of sequenced species. These genome assemblies will allow us to decipher the evolutionary innovations that resulted in the



Fig. 3. Species-trees inferred with Lamiaceae genome-derived gene models. a) STAG species-tree inferred with Orthofinder protein orthogroups. Support values show the proportion of trees at which the internal bipartitions occur for all species. b) Maximum-likelihood species tree using single copy nucleotide sequences. Branches with circles are fully supported (>98%) as judged by ultrafast bootstraps and SH-aLRT. Branches indicated by the asterisk are less well supported (<85%).

remarkably diverse number of specialized metabolites found in the Lamiaceae.

Data availability

Genome assemblies are available through Genbank under accession numbers JBCJKZ00000000 and JBCLUX00000000. The raw reads for whole genome and transcriptome sequencing are available in the National Center for Biotechnology Information Sequence Read Archive BioProject PRJNA1097548 and PRJNA1095452. The genome assembly and annotation files are available through figshare as Supplementary data: https://doi.org/10.25387/g3.25671948.

Supplemental material available at G3 online.

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Conflicts of interest

The author(s) declare no conflicts of interest.

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