In the past decade, the use of high-throughput sequencing in plant systematics, more specifically Illumina-based short-read sequencing, has changed from being a potentially revolutionary technique to a relatively commonplace approach (Delseni et al., 2010; McKain et al., 2018). While earlier studies relied on genome skimming to obtain organellar (i.e., plastid, mitochondrial, ribosomal) and high-copy-number nuclear markers (Godden et al., 2012; Straub et al., 2012), it soon became clear that for plant systematics, sequencing methods that isolate specific sets of low-copy-number nuclear genomic regions would be more powerful. The first reports of the use of array-free probes for multiplexed in-solution capture and sequencing using high-throughput sequencing platforms were Applications in Plant Sciences 2021 9(7): e11422; http://www.wileyonlinelibrary.com/journal/AppsPlantSci © 2021 The Authors. Application in Plant Sciences published by Wiley Periodicals LLC on behalf of Botanical Society of America. This is an open access article under the terms of the Creative Commons Attribution-NoCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.
published in 2011 (Bajgain et al., 2011), followed a few years later by the first family-wide probe set (Mandel et al., 2014).

The Compositae1061 probe set (also known as Compositae COS; Mandel et al., 2014) was one of the first probe sets to be designed specifically for a single family. The sunflower family (Compositae or Asteraeaceae) comprises more than 25,000 species, and many of its lineages have experienced recent and rapid radiations, large-scale gene family expansions, and ancient polyploidization events (Barker et al., 2008, 2016; Semple and Watanabe, 2009; Huang et al., 2016). Prior to the design and use of the Compositae1061 kit, many of the most important evolutionary questions about the family’s diversity were difficult to address, due to the poor resolution of, and lack of support for, the major backbone nodes of the family’s phylogeny. With the affordability and efficiency of high-throughput sequencing making genomic approaches attainable in many systems, and under the direction of Compositae expert Vicki A. Funk, members of the synantherology community sought to develop phylogenomic tools to address long-standing evolutionary questions in the family. In early 2011, the Compositae1061 probe set was developed using a set of expressed sequence tag (EST) loci obtained from three economically important members of the sunflower family, lettuce (Lactuca sativa L.), safflower (Carthamus tinctorius L.), and sunflower (Helianthus annuus L.), and included roughly 10,000 probes targeting the exons of 1061 orthogonal genes (Mandel et al., 2014).

Since 2014, this probe set has been used to study the family-wide phylogeny (Mandel et al., 2015, 2017, 2019); the relationships among different tribes (Watson et al., 2020); the relationships at the tribe level in the Cardueae (Herrando-Moraira et al., 2018, 2019), Vernonieae (Siniscalchi et al., 2019a), and Perityleae (Lichter-Marck et al., 2020); and the infrageneric relationships in Antennaria Gaertn. (Thapa et al., 2020). Data generated using Compositae1061 have also been used as a source to mine for microsatellite markers (Siniscalchi et al., 2019b; Thapa et al., 2019). The utility of the probe set at different evolutionary levels, within the family, and using different starting materials (e.g., herbarium samples vs. samples stored in silica gel) has been extensively explored by Jones et al. (2019). Moreover, the probe set is also able to successfully capture and recover loci for species in families closely related to the Compositae, such as the Calyceraeae and Goodeniaceae (Mandel et al., 2019).

Overall, the probe set has been accepted by researchers working on the family, with several ongoing studies yet to be published. In this sense, it fulfills one of the original goals of its design: the creation of a set of markers that could generate easily shareable data across the Compositae.

The wide and varied use of the Compositae1061 probe set has highlighted some of its limitations. One major issue is paralogy (multiple copies of a specific gene), mostly due to the rampant occurrence of both ancient and recent polyploid events within the family (Jones et al., 2019). A second issue is the low phylogenetic resolution at shallow taxonomic levels, such as in studies of closely related taxa or clades resulting from rapid radiation events (Thapa et al., 2020). This issue arises from the probes being designed exclusively from exonic regions, where there might not be enough sequence variation to accurately distinguish the species. Finally, even though the probe set contains 1061 loci, the mean number of loci recovered across studies has been ~700 (Herrando-Moraira et al., 2018).

The recent development of the universal Angiosperms353 kit opens up new opportunities for systematic studies combining deep and shallow phylogenetic levels (Dodsworth et al., 2019). This probe set was specifically developed to choose the minimum number of target instances needed to successfully recover 353 nuclear orthologs from any flowering plant. Its design included 31 Compositae species and a representative of each of the closely related families Goodeniaceae and Menyanthaceae (Johnson et al., 2019). Johnson et al. (2019) showed an average recovery of ~283 loci per species for the Angiosperms353 probe set, and at least 100 loci for over 600 angiosperm species, but this can be increased further using the pipeline recently described by McLay et al. (2021). It has been successfully implemented with low-quality material such as herbarium specimens (Brewer et al., 2019; Shee et al., 2020) and performed well for resolving shallow-level relationships (e.g., radiations [Larridon et al., 2020; Shee et al., 2020] and even at the intraspecific level [Van Andel et al., 2020; Beck et al., 2021; Slimp et al., 2021]). A few studies have compared the performance of Angiosperms353 with other taxon-specific probe sets (e.g., in Cyperus L. [Larridon et al., 2020], in the subtribe Malinae of the Rosaceae [Ufimov et al., 2021], and in the Ochnaceae [Shah et al., 2021]), but only Larridon et al. (2020) directly tested the mergeability of different data sets. Alternatively, new lineage-specific kits are being designed that incorporate part or all of the Angiosperms353 targets (e.g., for the Melastomataceae [Jantzen et al., 2020] and the Gesneriaceae [Ogutcen et al., 2021]).

The Angiosperms353 probe set is currently being used in the Plant and Fungal Trees of Life (PAFTOL; http://paftol.org) program at the Royal Botanic Gardens, Kew (Richmond, Surrey, United Kingdom), to produce data for one representative of all angiosperm genera, including Compositae genera. It is also being applied across the Australian flora by the Genomics for Australian Plants consortium (https://www.genomicsforaustralianplants.com/). In this context, a comparison between the Compositae1061 and Angiosperms353 probe sets in the Compositae is timely. Furthermore, understanding how the Angiosperms353 probe set performs in a plant lineage known to contain extensive paralogy issues and how it compares with a family-specific probe set, and verifying if data generated with different probe sets can be combined, is essential in a time where data sharing and collaborative projects abound. Here, we compared the data generated using both probe sets in eight genera of the Compositae. We address the following questions: (1) do the Compositae1061 and Angiosperms353 enrichment panels share any loci?; (2) how do issues of paralogy compare between the two probe sets?; (3) how can we best integrate data generated using these two approaches?

**METHODS**

**Identification of shared loci**

The BLAST Command Line Applications were used to examine whether there are any shared loci between the Compositae1061 and Angiosperms353 probe sets. The sequences of the loci contained in the Compositae1061 were used to create a local BLAST database, using the makeblastdb command. As this probe set was based on three EST libraries from sunflower, lettuce, and safflower, some loci are represented by more than one sequence. The sequences of the loci contained in the Angiosperms353 set were obtained from GitHub (https://github.com/mossmatters/Angiosperms353 [accessed 15 April 2020]) and separated into individual FASTA files. The Angiosperms353 probe set contains up to 18 different probe sequences per locus, as it is intended to be applicable across all...
flowing plants (details reported by Johnson et al., 2019). Each locus FASTA file separated from the Angiosperms353 set was then queried against the local Compositae1061 BLAST database using BLASTN.

Taxon selection, plant material, DNA extraction, library preparation, and sequencing

Eight taxa were chosen because they overlapped between Mandel et al. (2019) and those available from the PAFTOL program: *Cota tinctoria* (L.) J. Gay (Asteraceae); *Fallenis maritima* (L.) Greuter (Inuleae); *Calendula arvensis* (Vaill.) L. (Calenduleae); *Cardopatium corymbosum* (L.) Pers. (Cardueae); *Cichorium intybus* L. (Cichorieae); *Deinandra corymbosa* (DC.) B. G. Baldwin from PAFTOL and *D. minthornii* (Jeps.) B. G. Baldwin from Mandel et al. (2019) (Heliantheae); *Helichrysum stoechas* (L.) Moench (Gnaphalieae); and *Roldana gilgii* (Greenm.) H. Rob. & Brettell from Mandel et al. (2019) and *R. petasitis* (Sims) H. Rob. & Brettell from PAFTOL (Senecioneae). The taxa sequenced with the Compositae1061 probe set were previously published by Mandel et al. (2019), and details on sample origin and library preparation can be found in Mandel et al. (2014, 2019) and Jones et al. (2019). Sequence data from this project are also available at the National Center for Biotechnology Information Sequence Read Archive under BioProject PRJNA540287. The data set obtained from the Angiosperms353 probe set was collected as part of the PAFTOL program at the Royal Botanic Gardens, Kew, following the protocol described by Johnson et al. (2019), and is available along with voucher information at https://treeoflife.kew.org/ (accessed 4 April 2021).

Sequence assembly and data analysis

Eight samples were assembled for each probe set, as specified above. All sequences were trimmed using Trimmomatic version 0.39 (Bolger et al., 2014), using SLIDINGWINDOW mode with a five-base window and a quality cutoff of 20; reads shorter than 36 bp were removed. The trimmed and paired files were then assembled using HybPiper version 1.3.1 (Johnson et al., 2016), with the respective probe set target sequences as a reference. The trimmed and paired reads were first mapped against the target loci using BWA version 0.7.17 (Li and Durbin, 2009), and were then assembled de novo into contigs using SPAdes version 3.13.1 (Bankevich et al., 2012). Exonerate version 2.2 (Slater and Birney, 2005) was subsequently used to extract the longest unique contig that mapped to a specific target. The final gene matrices were aligned using MAFFT version 7.407 (Katoh and Standley, 2013) with the “-auto” option.

In a second step, each data set was assembled with the opposite probe set reference file to verify the bycatch of loci contained in the other probe set, which was part of our strategy to determine whether data from different sequencing runs generated with different probe sets could be successfully integrated. Even if the two sets contain some of the same loci, the locus lengths can differ depending on the initial source used for probe development. Basic statistics for all assemblies were obtained using the hybpiper_stats.py script in HybPiper and the software AMAS (Borowiec, 2016). Lists of all loci and those loci flagged as paralogs were also obtained with HybPiper. In cases where data generated with one probe set were assembled using the opposite reference file, the recovered loci were further analyzed to identify whether they were in the pool of loci shared by both probe sets, as obtained in the BLAST step described above. All data obtained from these analyses are summarized in Appendices S1–S7 (see Supporting Information).

The recovered loci were used in different phylogenetic analyses. Gene trees were obtained using RAxML version 8.2.9 (Stamatakis, 2014) in the rapid bootstrap mode, with 100 searches. The GTR+I+Γ model was used for all loci, as it is the most complex model currently available and has been shown to accurately infer topologies in real-life and simulated conditions (Abadi et al., 2019). The multispecies pseudo-coalescent method implemented in ASTRAL III version 5.6.3 (Zhang et al., 2018) was used to obtain a species tree from each data set. Support values in the form of local posterior probabilities were obtained using the “-q” option, and were considered high if equal to or higher than 0.95 and moderate if between 0.90 and 0.94. Four trees were produced: data generated with Compositae1061 and assembled with Compositae1061 (treatment A), data generated with Angiosperms353 and assembled with Angiosperms353 (treatment B), data generated with Compositae1061 and assembled with Angiosperms353 (treatment C), and data generated with Angiosperms353 and assembled with Compositae1061 (treatment D) (Table 1). In a second step, loci flagged as paralogs during the assembly were removed from all four data sets, as defined above, and new gene trees and species trees were obtained with these “cleaned” data sets.

Two additional unrooted ASTRAL trees containing all 16 samples were also produced: one with all 16 samples assembled with the Angiosperms353 probe set as the reference and another with the samples assembled with the Compositae1061 probe set as the reference. A third tree was produced, containing six taxa sequenced with Compositae1061 and two taxa sequenced with Angiosperms353, all assembled using Compositae1061 as the reference and with the loci flagged as paralogs removed, to confirm whether data set integration is indeed possible. All trees were visualized using FigTree version 1.4.4 (https://github.com/rambaut/figtree).

Gene tree discordance was evaluated using PhyParts (Smith et al., 2015). This program requires rooted trees for its analysis; therefore, all four species trees obtained in the ASTRAL analysis, as well as the respective gene trees used as input to generate them, were rooted using the function pxrr in the package phylx (Brown et al., 2017). Due to the lack of an outgroup taxon belonging to a different family, all species trees were rooted with *Cardopatium*, as the tribe Cardueae emerges as sister to the subfamilies Cichorioideae and Compositae1061.

Table 1. Summary of the four treatments used in the study.

| Treatment | Sequenced with | Assembled using the reference | No. of samples included | Recovered loci, average (range) | Percent of paralogous loci |
|-----------|----------------|-------------------------------|--------------------------|---------------------------------|-----------------------------|
| A         | Compositae1061 | Compositae1061               | 8                        | 721 (3–1012)                    | 0–46%                       |
| B         | Angiosperms353 | Angiosperms353               | 8                        | 287 (242–323)                  | 0.6–13%                     |
| C         | Compositae1061 | Angiosperms353               | 6                        | 25 (29–38)                     | ca. 5%                      |
| D         | Angiosperms353 | Compositae1061               | 8                        | 35 (21–59)                     | 2–25%                       |
and Asteroidae in most phylogenetic analyses (e.g., Mandel et al., 2019). The gene trees were rooted using a hierarchical scheme (as some missing trees might contain missing taxa) in the following order, after the topology from Mandel et al. (2019): Cardopatium, Cichorium, Pallenis, Deinandra, Calendula, Roldana, Cota, and Helichrysum. The phypartsiecharts.py script (https://github.com/mossmasters/phylocscripts/tree/master/phypartsiecharts [accessed 15 April 2020]) was used to plot the results from PhyParts as pie charts in each tree node.

RESULTS

Identification of shared loci

The results from the BLAST search show that 59 target instances from Angiosperms353 had hits when searched against the Compositae1061 database. These 59 target instances represent 30 individual loci from the Angiosperms353 probe set, as it contains multiple sequences for each locus. These loci from Angiosperms353 each matched with only one Compositae1061 locus, although some of them had positive hits for more than one probe in the data set, as presented in Table 2. The identity percentage between query and subject sequences varied from 72% to 98% in the searches. Most of the searches generated partial overlaps between the query and the subject, given that locus length is different between each panel. The difference in size for the same loci in each panel varied from 3 to 1836 bp. The results from the searches are summarized in Appendix S1.

TABLE 2. Loci shared across both probe sets. For cases where more than one species representative is included for the same gene, all loci that had hits were included.

| Angiosperms353 loci | Compositae1061 loci |
|---------------------|---------------------|
| Ambtr-6412, NVSO-6412 | sun-f-A1g23400, saff-At1g23400 |
| Ambtr-6467, IHP-6467 | saff-f-At2g27290, sun-f-At2g27290 |
| Ambtr-6462, EDXZ-6462, LYPZ-6462, NUZN-6462 | sun-f-At2g27450, saff-f-At2g27450, lett-f-At2g27450 |
| AQQ-5614, LVUS-5614, XRCC-5614 | sun-f-At3g03790, lett-f-At3g03790 |
| AQQ-5870 | sun-f-At2g15240 |
| Arath-5477 | lett-f-At1g14620 |
| Arath-5840, EZZT-5840, KDC-5840, NVSO-5840 | lett-f-At4g35250, saff-f-At4g35250, sun-f-At4g35250 |
| Arath-5857, BKF-5857, UYED-5857 | lett-f-At2g43030, sun-f-At2g43030, saff-f-At2g43030 |
| AZBL-5841, HYZL-5841, QJKZ-5841, SSVG-5841, UMUL-5841 | sun-f-At3g05070, lett-f-At3g05070 |
| BEFC-6449 | sun-f-At1g09830 |
| BIDT-5562 | lett-f-At4g35250, saff-f-At4g35250, sun-f-At4g35250 |
| BIDT-5910 | lett-f-At4g35250 |
| BIDT-6733 | lett-f-At4g35250 |
| BIDT-6946, UMUL-6946, VUSY-6946 | sun-f-At3g03790, saff-f-At3g03790, lett-f-At3g03790 |
| BIDT-6954, IDLU-6954 | sun-f-At1g09830 |
| DOV-7371 | lett-f-At4g35250 |
| DUNI-6498, HUG-6498, OXYP-6498, RCJX-6498 | sun-f-At3g05070, lett-f-At3g05070 |
| EMB-5918, JYMN-5918 | sun-f-At3g05070, lett-f-At3g05070 |
| HOKG-6458 | sun-f-At1g09830 |
| JEPE-4527 | lett-f-At3g05070, saff-f-At3g05070, sun-f-At3g05070 |
| JNKW-6705 | sun-f-At3g05070, lett-f-At3g05070, saff-f-At3g05070 |
| JYNN-7141 | sun-f-At3g05070, lett-f-At3g05070, saff-f-At3g05070 |
| LSJW-5933, MDKJ-5933 | sun-f-At3g05070, lett-f-At3g05070, saff-f-At3g05070 |
| NUZN-6139 | sun-f-At3g05070, lett-f-At3g05070, saff-f-At3g05070 |
| NVSO-7194 | sun-f-At1g09830 |
| Oryst-6038 | sun-f-At1g09830 |
| QJIK-7367, WAIL-7367, ZCUA-7367 | sun-f-At2g03420 |
| QTJ-7368, UYED-7368 | sun-f-At3g05070, lett-f-At3g05070, saff-f-At3g05070, sun-f-At3g05070, lett-f-At3g05070, saff-f-At3g05070 |
| WPY-6913, WWG-6913 | sun-f-At1g09830 |
| XRCX-5594 | sun-f-At1g09830 |

Recovered loci and paralogy

The main results from the assembly are summarized in Fig. 1, Table 3, and Appendices S2 and S3. Alignments produced from data sequenced with Angiosperms353 and assembled with the matching reference (treatment B) tended to be longer and have more parsimony-informative (PI) sites (Fig. 1A), while those sequenced and assembled with Compositae1061 (treatment A) tended to have similar lengths. The assemblies with the opposite reference (treatments C and D) tended to produce shorter alignments, and in the case of treatment D, several alignments did not present PI sites. For the samples sequenced with the Compositae1061 probe set and assembled using the same probe set as reference (treatment A), the percentage of reads on target varied from 2.8% in Cichorium to 56% in Helichrysum (Fig. 1B). The number of recovered loci varied from three in Cichorium to 1012 in Deinandra, with an average of 721 loci recovered. Loci flagged as paralogous were recovered for six of the eight species and the percentage of paralogous loci in relation to the recovered loci varied from 9% (Cardopatium) to 47% (Calendula). When the same data set was assembled with the Angiosperms353 probe set (treatment C), it showed percentage of reads on target ranged from 0.1% (Cichorium) to 1.6% (Pallenis). The number of recovered loci varied from 28 (Cardopatium) to 38 (Calendula). Only Calendula and Helichrysum presented paralogs, both with around 5% of the recovered loci being flagged. Cichorium and Roldana had very few loci recovered, being dropped from the final assembly with Angiosperms353, which was probably due to issues during genomic library preparation, well before sequencing.

For the samples sequenced with the Angiosperms353 probe set and assembled using this probe set as reference (treatment B), the percentage of reads on target varied from 9.5% (Cota) to 18.4% (Calendula) (Fig. 1B) and the number of recovered loci was somewhere between 242 (Cota) and 323 (Pallenis), with an average of 287 loci. The percentage of loci flagged as paralogous ranged from 0.3% (Pallenis) to 13% (Calendula). When this data set was assembled with the Compositae1061 probe reference (treatment D), the percentage of reads on target varied from 1.8% (Cota) to 5.2% (Calendula). The number of recovered loci varied from 21 (Cota) to 59 (Calendula), with an average of 32 loci. The percentage of paralogous loci varied from 2% (Pallenis) to 25% (Calendula).

The assembly of data generated in treatment C generated 39 unique loci. From these 39 loci, 29 are contained in the pool of 30 loci that are represented in both probe sets. The opposite scenario, treatment D, generated 71 unique loci, among which all 30 loci shared by both probe sets are represented (Appendix S4).

In treatment A, 640 of the 1061 loci (~60%) that compose the probe set were flagged as paralogous during assembly with HybPiper. Of these 640 loci, 388 were flagged for two or more taxa, while the remaining 252 loci were flagged only in one taxon (Appendix S5).
For treatment B, 16 loci were flagged as paralogous in two or more samples and 43 in only one sample (Appendix S6), totaling 58 loci (16% of the total in the probe set). Paralogous loci recovered from the sequences assembled with the opposite reference (treatments C and D) are summarized in Appendix S7.

**Phylogenetic relationships and gene tree discordance**

The recovered phylogenetic relationships varied depending on the data set used to generate them (Fig. 2). The four trees were rooted using *Cardopatium* (Cardueae). All eight samples were recovered in three of the trees, except *Roldana* and *Cichorium* in the tree based on treatment C. In all three completely sampled trees, *Cichorium* (Chiorieae) was sister to the larger subfamily Asteroidae clade. Within this clade, *Deinandra* (Heliantheae) and *Pallenis* (Inuleae) were always sister taxa.

The topologies of the trees obtained from the Compositae1061 data were similar regardless of whether the data were assembled with Compositae1061 (Fig. 2A) or Angiosperms353 (Fig. 2C) as a reference. In root-to-tip order, *Calendula* was in a grade leading to a *Cota–Helichrysum* clade. *Roldana* was sister to this grade in treatment C. In all three completely sampled trees, *Cichorium* (Chiorieae) was sister to the larger subfamily Asteroidae clade. Within this clade, *Deinandra* (Heliantheae) and *Pallenis* (Inuleae) were always sister taxa.

The results of the gene tree discordance analyses (Fig. 6) show a panorama of wide disagreement, increasing toward the tips of the trees. In the tree for treatment A (Fig. 6A), *Cichorium* was dropped and *Roldana* emerged in a clade with *Deinandra* and *Pallenis* instead of grouping with the other three species. *Calendula* emerged as sister to *Cota*, a relationship not seen in other trees. In the tree obtained from treatment D (Fig. 6D), *Cichorium* emerged within the Asteroideae clade, although with very low support. In the two trees that did not present topological changes (Fig. 6B, C), slight changes in support were observed, with the treatment B tree presenting decreased support in the backbone but not in the internal nodes and treatment C presenting the opposite pattern.

In the two trees obtained with the complete 16-species set assembled either with Compositae1061 or Angiosperms353 as the reference (Fig. 4), all species pairs form individual clades, except for *Cichorium* and *Roldana* in the Compositae1061 tree (Fig. 4B); these two species were not recovered from the data generated with Angiosperms353. In the tree obtained from the mix of both data sets (Fig. 5), the two samples sequenced with Angiosperms353 emerged in expected positions, with *Cichorium* as sister to the Asteroideae clade, and *Roldana* in the clade with *Cota, Helichrysum*, and *Calendula*, although the relationships in this four-species clade are different from those seen in other topologies.
In the present study, we sought to compare two different enrichment panels: Compositae1061, developed based on genomic resources available only for the sunflower family (EST libraries), and Angiosperms353, designed from angiosperm-wide genomic resources (transcriptomes and genomes). One of the goals of the present study was to verify the presence of shared loci in both sets. We identified 30 loci that are included in both probe sets, which facilitate complementary analyses with data combined from different studies. These loci appear to be among those that are consistently recovered across the family, given that assembling data generated with one of the probe sets with the opposite reference resulted in the recovery of similar numbers of genes across the samples, as seen in Table 3. Additionally, as seen in Fig. 4, the variation present in the limited number of shared loci is sufficient to allow for samples of the same taxa, sequenced with different probe sets, to group together in phylogenetic analysis. Mixing samples obtained with different probe sets, but assembled with the same reference, also proved possible (Fig. 5), with similar topology and support values seen in the trees generated using the data assembled with their own references.

The varying phylogenetic relationships we recovered are representative of the issues routinely found during phylogenetic studies in the sunflower family. From the eight sampled species, six belong to the subfamily Asteroideae, which includes >60% of the species diversity of the family (Susanna et al., 2020). Relationships among groups of tribes in this subfamily have been notoriously difficult to resolve, such as the tribes within the Heliantheae alliance and the group informally named Fab5 (Anthemideae, Astereae, Calenduleae, Gnaphalieae, and Senecioneae), as seen in multiple Compositae phylogenetic studies (Pelser and Watson, 2009; Huang et al., 2016; Panero and Crozier, 2016; Mandel et al., 2019; Watson et al., 2020). In the trees presented in Figs. 2 and 3, the conflict among species belonging to the Fab5 (Cota, Calendula, Helichrysum, and Roldana) is clear, with their relationships changing in each tree and with the removal of paralogs. It is noteworthy that relationships recovered with the Compositae1061 data set do not reflect those shown in Mandel et al. (2019), although they are based on the same data, suggesting that the reduced sampling used in this study significantly impacted the resolution of the relationships. The low number of loci recovered for Cichorium (3) and Roldana (23) in this data set, likely a result of sequencing issues, could be an additional source of phylogenetic noise and a factor leading to topological incongruences.

With the increasing abundance of large-scale genomic data sets composed mainly of nuclear genes, the issue of paralogy (which results from small-scale gene family expansions to whole-genome duplications) has become more widely discussed among plant systematists. Multiple copies of specific genes or whole gene families are most likely a consequence of the ancient polyploid origin at the base of all flowering plants and the occurrence of further additional ancient polyploid events leading to the base of the Compositae

**TABLE 3.** Summary of assembly statistics.

| Treatment             | Sample    | Genes recovered | Genes flagged as paralogs | Genes not flagged as paralogs | Percentage of genes flagged as paralogs |
|-----------------------|-----------|-----------------|---------------------------|--------------------------------|----------------------------------------|
| A (data generated with Compositae1061 and assembled using Compositae1061 as the reference) | Calendula | 1008            | 470                       | 538                            | 47%                                    |
|                       | Cardopatium | 893             | 82                        | 811                            | 9%                                     |
|                       | Cichorium   | 3               | 0                         | 3                              | 0                                      |
|                       | Cota        | 903             | 211                       | 692                            | 23%                                    |
|                       | Deinandra    | 1012            | 301                       | 711                            | 29%                                    |
|                       | Helichrysum  | 951             | 248                       | 703                            | 26%                                    |
|                       | Pallenis     | 977             | 181                       | 796                            | 18%                                    |
|                       | Roldana      | 23              | 0                         | 23                             | 0                                      |
| B (data generated with Angiosperms353 and assembled using Angiosperms353 as the reference) | Calendula | 315             | 41                        | 274                            | 13%                                    |
|                       | Cardopatium  | 314             | 2                         | 312                            | 0.6%                                   |
|                       | Cichorium    | 296             | 3                         | 293                            | 1%                                     |
|                       | Cota         | 242             | 5                         | 237                            | 2%                                     |
|                       | Deinandra     | 272             | 5                         | 267                            | 2%                                     |
|                       | Helichrysum   | 275             | 6                         | 269                            | 2%                                     |
|                       | Pallenis      | 323             | 1                         | 322                            | 0.3%                                   |
|                       | Roldana       | 261             | 11                        | 250                            | 4%                                     |
| C (data generated with Compositae1061 and assembled using Angiosperms353 as the reference) | Calendula | 38              | 2                         | 36                             | 5%                                     |
|                       | Cardopatium  | 29              | 0                         | 29                             | 0                                      |
|                       | Cichorium    | 0               | 0                         | 0                              | NA                                     |
|                       | Cota         | 31              | 0                         | 31                             | 0                                      |
|                       | Deinandra     | 37              | 0                         | 37                             | 0                                      |
|                       | Helichrysum   | 35              | 2                         | 33                             | 5%                                     |
|                       | Pallenis      | 35              | 0                         | 35                             | 0                                      |
|                       | Roldana       | 0               | 0                         | 0                              | NA                                     |
| D (data generated with Angiosperms353 and assembled using Compositae1061 as the reference) | Calendula | 59              | 15                        | 44                             | 25%                                    |
|                       | Cardopatium  | 34              | 0                         | 34                             | 0                                      |
|                       | Cichorium    | 31              | 0                         | 31                             | 0                                      |
|                       | Cota         | 21              | 0                         | 21                             | 0                                      |
|                       | Deinandra     | 31              | 0                         | 31                             | 0                                      |
|                       | Helichrysum   | 30              | 2                         | 28                             | 6%                                     |
|                       | Pallenis      | 48              | 1                         | 47                             | 2%                                     |
|                       | Roldana       | 28              | 1                         | 27                             | 3%                                     |
Indeed, a hexaploid ancestor was proposed for most of the lineages within the family (Barker et al., 2016; Li and Barker, 2020), and paralogy has been a frequent issue in phylogenomic studies ever since their inception.

The design of the Compositae1061 probe set focused on genes considered to be conserved orthologs at the time, thus aiming to reduce the number of possibly paralogous genes in the set (Mandel et al., 2014); however, with its use across different lineages of the family, it became clear that large numbers of loci are present in multiple copies after sequencing. As previously demonstrated by Jones et al. (2019), lineages within the family present different degrees of paralogy, probably associated with the hypothesized presence of further whole genome duplication events in several lineages (Huang et al., 2016; Li and Barker, 2020). Corroborating the results from Herrando-Moraira et al. (2018) and Jones et al. (2019), for the data generated using the Compositae1061 probe set, ~20% of the loci are putatively paralogous in most species, while Cardopatium (Cardueae) presented the lowest levels of paralogy (~9%). There is currently no evidence for ancient polyploidy in the tribe Cardueae, while there are multiple events proposed for the other tribes present in our analysis, such as the Calenduleae (Calendula), the Gnaphalieae (Helichrysum), and the Heliantheae alliance (Deinandra) (Huang et al., 2016; Li and Barker, 2020).

As expected, the removal of paralogs from the analysis caused topological changes (Fig. 2A, 3A), with more marked effects in the position of taxa with higher paralogy, such as Calendula and Helichrysum.

The number of paralogous loci recovered from the Angiosperms353 data assembled with itself as the reference reflect these phylogenetic trends, with Calendula presenting the highest proportion of paralogs (~13%) and Cardopatium (Inuleae) the smallest (0.3%). Pallenis (Inuleae) is the sample with the smallest number of paralogs (0.3%) in this set. The very high paralogy observed in Calendula arvensis in both data sets likely arises from the fact that the species has not only experienced multiple ancient polyploid events, but is also an allotetraploid (Nora et al., 2013; Plume, 2015). The overall lower levels of paralogy seen in the Angiosperms353 data set appear not to interfere with the topology as there are no changes when paralogs are removed, although this generates a slight improvement in support values (Figs. 2B, 3B).

Regarding the assemblies carried out using the other probe set as the reference, in the data generated for treatment C (Fig. 2C), the number of paralogous loci in each species decreased in relation to the number generated for treatment A, while in treatment D (Fig. 2D), the number of paralogous loci increased for the four species (Appendix S7). The treatment D assembly recovered 20 unique loci flagged as paralogs, of which 11 were from the pool of 30 loci.
shared by both probe sets. The recovery of fewer paralogous loci when assembling Compositae1061 data with the Angiosperms353 reference might be explained by an overall lower rate of potentially paralogous loci in the Angiosperms353 kit (see further discussion of this below). The removal of paralogs from these two assemblies caused different effects: the topology remained the same in the first tree, with small changes in support (Fig. 3C), but was altered in the second case (Fig. 3D). The position of Cichorium in this last topology, within the subfamily Asteroideae, is dubious and goes against all previous phylogenetic work and the historical classifications of the Compositae. Calendula was the taxon with the highest number of removed loci, emerging as sister to Cichorium, and is likely the reason for the stark topological changes.

Samples sequenced with Angiosperms353 present a much smaller proportion of paralogs than those sequenced with Compositae1061, being below 10% in seven of the eight species. This is probably a reflection of the original data used to develop each probe set. While the Compositae1061 set was based exclusively on three EST libraries, the only genetic resources available for the Compositae at the time, the development of the Angiosperms353 probe set relied on a set of 410 alignments of orthologous loci across 1100 green plants, singled out in the context of the 1000 Plants (1KP) project (details in Johnson et al., 2019; Leebens-Mack et al., 2019). This comparison across a wide group of genomic references, including 31 Compositae species, allows for a more refined selection of target regions that will truly present as single-copy loci in most plant species. This data set also seems to be more resilient to paralogy, as there are no changes in topology with the removal of paralogous loci. It is worth noting, however, that even if the loci flagged as paralogous were removed from the analysis, the Compositae1061 set still generates several hundred more loci than Angiosperms353 (538–811 vs. 237–322, respectively) and presents higher proportions of on-target reads (Fig. 1B), which could be decisive when dealing with rapid radiations or very recent divergences. Nevertheless, removing the paralogs created changes in topology in the present study, which could also be a consequence of the very sparse sampling.

Many phylogenomic studies using the Compositae1061 probe set assembled their data using phyluce (Faircloth et al., 2015), which takes a more restrictive approach than HybPiper with regard to paralogs, as a way of dealing with resulting conflicting relationships (Mandel et al., 2019; Siniscalchi et al., 2019a; Thapa et al., 2019). HybPiper flags possible paralogs but keeps the locus in the final alignments by choosing either the copy with the greater sequencing depth or the one with the greatest percentage identity to the reference (Johnson et al., 2016). On the other hand, phyluce removes any locus for which the assembled contigs match multiple loci or

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**FIGURE 3.** Phylogenies obtained using the different data sets and assembly strategies after the removal of loci flagged as paralogs. Values on the nodes are local posterior probabilities obtained using ASTRAL-III. (A) Data generated with Compositae1061 and assembled using Compositae1061 as the reference. (B) Data generated with Angiosperms353 and assembled using Angiosperms353 as the reference. (C) Data generated with Compositae1061 and assembled using Angiosperms353 as the reference. (D) Data generated with Angiosperms353 and assembled using Compositae1061 as the reference.
different contigs match the same locus from the final alignments. One additional difference between pipelines is also the extent of the assembled loci. The first step in HybPiper is mapping against reference loci (target sequences), followed by a de novo assembly (of mapped reads per target locus); thus, the loci ultimately recovered usually span the length of the reference loci and include flanking regions, which are later automatically removed and can be recovered using the “--supercontig” option in the “retrieve_sequences.py” script. Phyluce begins with a de novo assembly of the data, with the contigs being posteriorly matched against the references, which also allows for the assembly of off-target flanking regions or introns, which are not removed. Herrando-Moraira et al. (2018) compared the effects of different assembly methods in the relationships of the tribe Cardueae and noted that phyluce introduces more phylogenetic noise, but without deeply affecting the recovered relationships. This is likely an effect of the unequal recovery of flanking regions; as they are not targeted, the recovery is different across loci and taxa, introducing more missing data in the final matrices.

In the present study, we decided to assemble the data with HybPiper, as it is more widely used by the plant systematics community. We analyzed data sets with and without paralogs. Given the widespread genomic duplications in the family, it is prudent to remove potentially paralogous loci from the final assemblies, or at least investigate the phylogenetic history of possible paralogous copies, for example, using the “paralog_investigator.py” script in HybPiper. Unfortunately, HybPiper does not include an option to easily remove specific paralogs from samples, in which case phyluce tends to be a better option, as this is done automatically in its pipeline. Finally, as HybPiper assemblies tend to be similar in length to the original locus sequences, the sequences and assemblies generated using Angiosperms353 in this study are longer overall than those from Compositae1061 (Fig. 1A), which is probably due to the original size of the loci used as basis to design the probes contained in each kit.

Another issue arising from large phylogenomic data sets becoming more widely available is gene tree discordance, which is usually explained by whole-genome duplication or polyploidy, hybridization, incomplete lineage sorting, or some combination of these processes. Gene tree discordance has been widely documented in plants (as summarized by Smith et al., 2020) and more specifically in the Compositae (Herrando-Moraira et al., 2019; Jones et al., 2019; Siniscalchi et al., 2019a; Watson et al., 2020). Most gene tree discordance analyses in Compositae studies show high levels of disagreement, increasing toward the tips of the trees. This is likely due to the fact that gene recovery is variable in samples sequenced with the Compositae1061 probe set, an issue that might have several origins, such as low probe hybridization efficiency, large and repetitive genomes impairing probe binding, and a high divergence of the probes in relation to the target sequence. However, one study in the Cardueae (Herrando-Moraira et al., 2019) showed the opposite, with the backbone presenting more discordance than the tips. Siniscalchi et al. (2019a) demonstrated that reduced data sets, which eliminated gene trees lacking several taxa, improved the overall discordance by decreasing the number of uninformative gene trees and increasing the proportion of gene trees that agree with the species tree. A similar effect is observed in the discordance analysis presented here, where the two trees obtained with data assembled with the opposite probe set as the reference showed less overall discordance, probably due to the lower number of missing taxa in these gene trees, as presented in Appendix S2.

Overall, our results show that the Compositae data obtained using two different probe sets can be combined due to the presence of 30 shared loci between them, enabling mixed analyses (Figs. 4, 5). One interesting result is that both data sets assembled using the other as a reference recovered more than 30 loci each. Hybridization reactions vary in precision and efficiency, with the occurrence of bycatch being well known (e.g., Jones et al., 2019). One explanation could be the presence of other loci contained in the probe set being part of the bycatch of the hybridization, which then match to targets
Few studies have used Compositae1061 to investigate infrageneric relationships in different Compositae tribes, while there are currently no published reports of the use of Angiosperms353 in the Compositae. Lichter-Marck et al. (2020) and Thapa et al. (2020) investigated infrageneric relationships within Perityle Benth. (Perityleae) and Antennaria (Gnaphalieae), respectively, and compared the effects of concatenated vs. gene tree analyses. Both studies found high levels of paralogy and topological incongruence between the phylogenies generated using different inference methods. Jones et al. (2019) investigated the levels of paralogy and conflict within a species complex in Picris L. (Cichorieae), showing high levels of gene tree conflict but good overall resolution and support. The probe set has been proven useful at lower phylogenetic levels but presented the same issues seen at higher levels, with incongruences between different assembly and phylogenetic inference methods (Herrando-Moraira et al., 2018; Siniscalchi et al., 2019a), possibly indicating issues with the actual loci chosen as targets or a complicated history of genomic evolution within the family.

The possibility of integrating data from different origins opens up opportunities for new collaborations and integrative projects using data that can be universally shared. Given the high amount of paralogy in the loci contained in the Compositae1061 set and taking into account the new genetic resources available for the family, such as three complete genomes and more than 30 transcriptomes, a redesign of this specific probe set could be beneficial. A deeper study of
paralogy across the family could indicate loci that are problematic in several lineages, and these could then be replaced by newly selected ones. Alternatively, the loci contained in the Angiosperms353 set could be included with the Compositae1061 to create a more inclusive set of targeted loci, which has already been done for the Melastomataceae (Jantzen et al., 2020) and Gesneriaceae (Ogutcen et al., 2021). Finally, it is worth noting that Mandel et al. (2019) successfully integrated transcriptomic data from the 1KP project with the Compositae1061 loci, demonstrating how different sources of data can be combined for phylogenetic reconstruction.

This data integration will be useful at higher levels of phylogenetic analyses, such as for adding outgroups to an analysis or in tribe or subfamily phylogenies, as the small number of shared loci between Compositae1061 and Angiosperms353 will probably not be sufficient to resolve relationships in shallower nodes or in cases of rapid radiations. When choosing a probe set to start a new project, it will be important to decide upfront whether integration with previous data sets is an important factor and to choose whichever probe set was used before. Both probe kits are manufactured by the same company and have identical laboratory protocols, although Compositae1061 is slightly cheaper due to the lower number of probes per reaction. Hendriks et al. (2021) present the possibility of integrating Angiosperms353 and a custom probe set in the same hybridization reaction, which has not yet been tested in the Compositae, but is surely an exciting possibility.

We conclude that the Compositae1061 kit provides more loci, even with higher levels of paralogy, than Angiosperms353, which can be useful when working on shallower phylogenetic levels. The Angiosperms353 set yields a more even number of loci across samples that are less affected by paralogy, which can be useful when working across several lineages in the Compositae family. The outlook for phylogenomic studies in the Compositae is promising, especially if researchers across the globe are able to combine genomic data to address the evolutionary history of this large and complex group of flowering plants.

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AUTHOR CONTRIBUTIONS

C.M.S. and J.R.M. had the initial idea for this work. C.M.S., J.R.M., O.H., L. Palazzesi, and J.P. planned this work. J.R.M., O.M., I.J.L., F.F., and W.J.B. oversaw data production and provided the sequences used here. C.M.S. and J.R.M. conducted data analyses. C.M.S. wrote the initial draft of the manuscript, and J.R.M., O.H., L. Palazzesi, J.P., and L. Pokorny provided additional text. All authors read, commented on, and approved the manuscript and the subsequent review.

DATA AVAILABILITY

All gene trees, species trees, reference files, and code used in this article are available from https://github.com/carol-siniscalchi/Comp1061-Angio353. Raw sequence files are available at the National Center for Biotechnology Information Sequence Read Archive under BioProject PRJNA540287 and from http://paftol.org.

SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

APPENDIX S1. Summary of hits of individual Angiosperms353 loci sequences BLAST-searched against the Compositae1061 database.

APPENDIX S2. Basic statistics from each assembly obtained with the hybpiper_stats.py in HybPiper.

APPENDIX S3. Basic assembly statistics for each treatment and locus.

APPENDIX S4. Loci recovered with inverted assemblies.

APPENDIX S5. Paralogous loci.

APPENDIX S6. Loci flagged as paralogous for each sample.

APPENDIX S7. Loci assembled with the opposite reference and corresponding paralogous loci.

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