A Comprehensive Phylogenomic Platform for Exploring the Angiosperm Tree of Life

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Abstract.—The tree of life is the fundamental biological roadmap for navigating the evolution and properties of life on Earth, and yet remains largely unknown. Even angiosperms (flowering plants) are fraught with data gaps, despite their critical role in sustaining terrestrial life. Today, high-throughput sequencing promises to significantly deepen our understanding of evolutionary relationships. Here, we describe a comprehensive phylogenomic platform for exploring the angiosperm tree of life, comprising a set of open tools and data based on the 353 nuclear genes targeted by the universal Angiosperms353 sequence capture probes. The primary goals of this paper are to (i) document our methods, (ii) describe our first data release and (iii) present a novel open data portal, the Kew Tree of Life Explorer (https://treeoflife.kew.org). We aim to generate novel target sequence capture data for all genera of flowering plants, exploiting natural history collections such as herbarium specimens, and augment it with mined public data. Our first data release, described here, is the most extensive nuclear phylogenomic dataset for angiosperms to date, comprising 3,099 samples validated by DNA barcode and phylogenetic tests, representing all 64 orders, 404 families (96%) and 2,333 genera (17%). A “first pass” angiosperm tree of life was inferred from the data, which totalled 824,878 sequences, 489,086,049 base pairs, and 532,260 alignment columns, for interactive presentation in the Kew Tree of Life Explorer. This species tree was generated using methods that were rigorous, yet tractable at our scale of operation. Despite limitations pertaining to taxon and gene sampling, gene recovery, models of sequence evolution and paralogy, the tree strongly supports existing taxonomy, while challenging numerous hypothesized relationships among orders and placing many genera for
the first time. The validated dataset, species tree and all intermediates are openly accessible via the Kew Tree of Life Explorer and will be updated as further data become available. This major milestone towards a complete tree of life for all flowering plant species opens doors to a highly integrated future for angiosperm phylogenomics through the systematic sequencing of standardised nuclear markers. Our approach has the potential to serve as a much-needed bridge between the growing movement to sequence the genomes of all life on Earth and the vast phylogenomic potential of the world’s natural history collections.

**Keywords:** angiosperms, Angiosperms353, genomics, herbariomics, museomics, nuclear phylogenomics, open access, target sequence capture, tree of life.

**INTRODUCTION**

Discovering the tree of life is among the most fundamental of the grand challenges in science today (Hinchliff et al. 2015). The tree of life is the biological roadmap that allows us to discover, identify and classify life on Earth, to explore its properties, to understand its origins and evolution, and to predict how it will respond to future environmental change. Of all eukaryotic lineages, the angiosperms (flowering plants) are among the most pressing priorities for tree of life research. Angiosperms sustain the terrestrial living world, including humanity, as primary producers, ecosystem engineers and earth system regulators. They hold potential solutions to global challenges, such as climate change, biodiversity loss, human health, food security and renewable energy (Antonelli et al. 2020). In light of this, a phylogenetic framework with which to navigate and interpret the species, trait and functional diversity of angiosperms has never been more necessary. However, despite substantial progress, the evolutionary connections among Earth’s ca. 330,000 flowering plant species (WCVP 2020) remain incompletely known.
The angiosperm research community were early and organised adopters of the molecular phylogenetic approach, resulting in numerous benchmark tree of life publications (e.g. Chase et al. 1993; Soltis et al. 2008; Soltis et al. 2011), and a community approach to phylogenetic classification (APG 1998; APG II 2003; APG III 2009; APG IV 2016). Through this distributed effort, a wealth of DNA sequence data is now available in public repositories, covering ca. 107,000 (31%) of the ca. 350,000 species of vascular plants (RBG Kew 2016; WCVP 2020), most of which are angiosperms (see also Cornwell et al. 2019). However, the lack of sequence data for the remaining 69% obstructs their accurate placement in the tree of life. In addition, lack of complementarity in gene sampling across public DNA sequence data impedes phylogenetic synthesis (Hinchliff and Smith 2014). For example, data from either one or both of \textit{rbcL} and \textit{matK}, the two most popular plastid genes for phylogenetics, are available for only 54% of the ca. 107,000 sequenced vascular plant species (RBG Kew 2016).

Comprehensive phylogenetic trees of flowering plants are in high demand (Hinchliff et al. 2015; Eiserhardt et al. 2018), but currently can only be made “complete” using proxies, such as taxonomic classification, to interpolate the unsequenced species (Smith and Brown 2018), which may not accurately reflect relationships. Greater community-wide coordination of both taxon and gene sampling would benefit phylogenetic data integration immensely, creating numerous downstream scientific opportunities.

High-throughput sequencing (HTS) now promises to significantly deepen our understanding of evolutionary relationships among Earth’s species, including angiosperms (Li et al. 2019; Yang et al. 2020). For example, the One Thousand Plant Transcriptomes (1KP) initiative has brought an unprecedented scale of data to bear on the plant tree of life (Wickett et al. 2014; Gitzendanner et al. 2018; Leebens-Mack et al. 2019). Nevertheless, with greatly increased data depth come trade-offs in taxon sampling; the pre-eminent HTS studies cited here account for less than 0.01% of angiosperm species. Undeterred by this sampling
gap, the Earth Biogenome Project (EBP) has launched a “moonshot for biology” by proposing to sequence and characterise the genomes of all of Earth’s eukaryotic species over a 10-year period (Lewin et al. 2018). Projects such as the 10,000 Plant Genomes Project (Cheng et al. 2018) and the Darwin Tree of Life Project (https://www.darwintreeoflife.org/) aim to contribute to this goal by producing numerous chromosome-level genome assemblies across major lineages and regional biotas. However, taxon sampling remains a significant issue, due to the challenges of obtaining the high molecular weight DNA required by these projects (for long-read HTS) from samples that are both authentically identified and compliant with the spirit and letter of the Nagoya Protocol (Secretariat of the Convention on Biological Diversity 2011). Despite its immense potential, the “whole genome” approach to discovering the tree of life remains a future goal that will not be achieved on a large taxonomic scale in the short term. Methodological compromises are required to accelerate progress.

The world’s natural history collections are a goldmine for genomic research (Buerki and Baker 2016), containing tissues of almost all species of life on Earth known to science. However, the condition of these tissues and the DNA therein varies widely, depending on age and preservation techniques, among other factors. In the case of plants, herbarium specimens generally yield degraded DNA, which, though not useful for long-read HTS, is now being intensively exploited for short-read HTS (Bakker et al. 2016; Brewer et al. 2019; Forrest et al. 2019; Alsos et al. 2020). In this context, target sequence capture is growing in popularity as the HTS method most widely applied to herbarium DNA (Dodsworth et al. 2019). This approach (also known as target enrichment, target capture, sequence capture, anchored hybrid enrichment) and its variations (e.g. Hyb-Seq, which combines target sequence capture with genome skimming) use RNA or DNA probes to enrich sequencing libraries for specifically targeted loci (Faircloth et al. 2012; Lemmon et al. 2012; Weitemier et al. 2014). It is proving
to be an increasingly cost-effective means of isolating hundreds of loci for phylogenetic
analysis from even centuries-old specimens (Brewer et al. 2019), bringing comprehensive
taxon sampling from herbarium collections within the reach of any phylogenomic researcher
(Hale et al. 2020).

Numerous target sequence probe sets have been developed for specific angiosperm
groups (e.g. Annonaceae [Couvreur et al. 2019], Asteraceae [Mandel et al. 2014], Dioscorea
[Soto Gomez et al. 2019], Euphorbia [Villaverde et al. 2018]). The design of these probe sets
is informed by available genomic resources, as well as criteria specific to the group of interest
and research questions. As a result, locus overlap between probe sets tends to be minimal.

Unlike the Sanger sequencing era, in which researchers converged on tractable genes such as
*rbcL* and *matK*, the lack of complementarity between probe sets curtails prospects for data
integration across broad taxonomic scales. In addition, development of custom probe sets is
expensive, requiring considerable genomic resources and bioinformatic expertise. A publicly
available, universal probe set for angiosperms targeting a standard set of loci would resolve
these issues (Buddenhagen et al. 2016; Chau et al. 2018). In response to this, we designed the
Angiosperms353 probe set (Johnson et al. 2019), drawing on 1KP transcriptome data from
c. 650 species across the angiosperms (Leebens-Mack et al. 2019). The probe set targets 353
genes from 410 low-copy, protein-coding nuclear orthologs previously selected for
phylogenetic analysis across green plants (Leebens-Mack et al. 2019), enriching up to ca. 260
kbp from any flowering plant. Angiosperms353 probes are an open data resource that can be
used without the expense of design or access to prior genomic data (Baker et al. 2021) and
have already been successfully applied across different taxonomic scales (e.g. Larridon et al.
2019; Murphy et al. 2020; Pérez-Escobar et al. 2020; Shee et al. 2020), including at the
population level (Van Andel et al. 2019; Slimp et al. 2020; Beck et al. 2021).
Here, we describe a large-scale effort to establish a new phylogenomic platform for exploring the angiosperm tree of life, comprising a set of open tools (Angiosperms353 probes, laboratory protocols, analysis pipeline, data portal) and data (sequence data, assembled genes, alignments, gene trees, species tree). This platform, which directly addresses the challenges outlined above, is an outcome of the Plant and Fungal Trees of Life project (PAFTOL; www.paftol.org) at the Royal Botanic Gardens, Kew (RBG Kew 2015).

As a step towards the ultimate goal of a complete species-level tree, we aim to gather DNA sequence data for the Angiosperms353 genes from one species of all 13,862 angiosperm genera (WCVP 2020). This unprecedented dataset of standard loci draws extensively on herbarium collections for comprehensive sampling, especially of genera that have not been sequenced before (Brewer et al. 2019). Extensive new data have been generated, analysed and released into the public domain, along with corresponding phylogenetic inferences. By providing our data in open and accessible ways, including an interactive tree of life, we aim to foster a transparent and collaborative environment for future data re-use and synthesis.

This paper serves as the baseline reference for our platform, (i) documenting our methods, (ii) describing our first data release, comprising 17% of angiosperm genera, including initial insights on phylogenetic performance, and (iii) presenting a novel data portal, the Kew Tree of Life Explorer, through which our data and corresponding tree of life can be interrogated and downloaded. We conclude with reflections on the prospects for our approach, future development requirements and the role of open data for enhancing cross-community collaboration towards a complete tree of life.

**Materials and Methods**
This section describes the workflow (Fig. 1) used by the PAFTOL project to generate our first data release (i.e. Data Release 1.0), which is publicly accessible through our open data portal, the Kew Tree of Life Explorer (https://treeoflife.kew.org), described below. The workflow consists of three main stages: (i) sample processing, encompassing sample selection and laboratory protocols for target sequence capture data generation (Fig. 2), (ii) data analysis, including target gene assembly, data mining, data validation and phylogenetic inference (Figs. 2, 3), and (iii) data publication via the data portal (Fig. 4). The data accessible via the portal comprise raw data (unprocessed sequence reads) and results from “first pass” analyses (gene assemblies, alignments, gene trees, species tree). Though not exhaustive, these first explorations of the data apply methods that are both rigorous and tractable at our scale of operation.

Details of the first data release are also given in the data release notes in the portal via our secure FTP (http://sftp.kew.org/pub/treeoflife/) and are also archived at the Royal Botanic Gardens, Kew (RBGK) Research Repository (https://doi.org/10.34885/paftol). A new release note will be published in the same locations with each future data release and will detail any changes in methods used relative to the first release described here.

**Sampling**

We aimed to generate novel data from across the angiosperms, using a stratified sampling approach of one species per genus. Our sampling was standardised to the complete list of angiosperms within the World Checklist of Vascular Plants (WCVP 2020), which currently recognises 13,862 accepted genera in 418 families, aligned to the 64 orders of the APG IV classification (APG IV 2016). We prioritised genera that were not represented by published transcriptomic or genomic data in public sequence repositories (e.g. GenBank), and avoided genera that had already been sampled in large genomic initiatives such as the 1KP.
The selection of species within genera was made pragmatically, although we prioritised the species of the generic type where possible. Plant material was obtained from a variety of sources (Fig. 2), primarily from the collections of RBGK (herbarium, DNA bank, silica gel-dried tissue collection, living collection and the Millennium Seed Bank, https://www.kew.org/science/collections-and-resources/collections). Additional material (tissue samples, extracted DNA) was generously provided by individuals in our collaborative networks (see Acknowledgements). To be selected, the material must have been (i) legally sourced and made available for use in phylogenomic studies, (ii) identified to species level, preferably by an expert in the group, and (iii) ideally collected in the wild. As far as was practically achievable, we ensured that the identity of each sample was substantiated by a voucher specimen deposited in a publicly accessible herbarium.

All metadata were captured using a relational database that allowed us to track processing of samples from the selection of material, through the library preparation pipeline to the completion of sequencing. Data were recorded in four main tables (Specimen, Sample, Library, Sequencing). The database architecture allowed us to record multiple sequence datasets (fastq files) from one or several libraries, and one or several DNA extracts from a single specimen. Relevant voucher specimen information was also captured in the database (e.g. collector(s), collector number, herbarium acronym (following Index Herbariorum http://sweetgum.nybg.org/science/ih/), country of origin, date of collection, specimen barcodes). Voucher data are available via our data portal (see below). Images of specimens sampled from the RBGK Herbarium are in the process of being captured in RBGK’s online herbarium catalogue (http://apps.kew.org/herbcat/) and, where available, are linked to the appropriate records in the Kew Tree of Life Explorer.
DNA extraction

DNA was extracted from 40 mg of herbarium material, 20 mg of silica gel-dried material (Chase and Hills 1991), or 100 mg of fresh material using a modified CTAB extraction method (Doyle and Doyle 1987; Fig. 2). Plant tissue was pulverized using a Mixer Mill MM400 (Retsch GmbH, Germany). DNA extractions were purified by a magnetic bead clean-up using Agencourt AMPure XP beads (Beckman Coulter, Indianapolis, IN, USA), according to the manufacturer’s protocols. Samples obtained from the RBGK DNA bank (http://dnabank.science.kew.org/homepage.html) had been extracted using a modified CTAB method (Doyle and Doyle 1987) followed by caesium chloride/ethidium bromide density gradient cleaning and dialysis. DNA samples provided by external collaborators had been extracted using a wide variety of extraction methods from living, silica gel-dried and herbarium material.

All DNA samples were quality checked for concentration and degree of fragmentation. DNA concentration was measured using a Quantus (Promega, Madison, WI, USA) or Qubit (Thermo Fisher Scientific, Inchinnan, UK) fluorometer. DNA fragment size range was routinely assessed on a 1% agarose gel using ethidium bromide and visualized with a UVP Gel Studio (AnalytikJena, Jena, Germany). For samples with a low DNA concentration (i.e. not visible on a gel), fragment sizes were assessed on a 4200 TapeStation using Genomic DNA ScreenTape (Agilent Technologies, Cheadle, UK).

Library preparation

Genomic DNA samples were diluted to 4 ng/µl with 10 mM Tris (pH 8.0). Those with an average fragment size greater than 350 bp were sonicated to an average fragment size ca. 400 bp, using a Covaris M220Focused-ultrasonicator (Covaris, Woburn, MA, USA) by adding 50 µl of diluted genomic DNA to a 130 µl Covaris microAFA tube. The sonication time was adjusted for each sample based on its average DNA fragment size (15 to 100 secs,
following the manufacturer’s protocols. Additional parameters used were peak incident
power to 50W, duty factor to 10% and 200 cycles per burst.

Libraries were prepared using the NEBNext Ultra II DNA Library Prep Kit (New
England Biolabs, Ipswich, MA, USA; Fig. 2). Size selection was not employed for samples
with highly degraded DNA. In the early stages of the project, libraries were prepared
following the manufacturer’s protocols exactly, but the majority were prepared using half of
the recommended volumes throughout to reduce costs. All DNA fragments were indexed
using NEBNext Multiplex Oligos for Illumina (Dual Index Primer sets 1 and 2, New England
Biolabs, Ipswich, MA, USA).

The distribution of fragment sizes in each library was assessed with a 4200
TapeStation using standard D1000 tapes. Library concentration was measured using a
Quantus fluorometer. If the library concentration was less than 10 nM, up to eight additional
PCR cycles were performed, following the NEBNext Ultra II Library Prep Kit protocol with
IS5_reamp.P5 and IS6_reamp.P7 primers (Meyer and Kircher 2010). Library quality
assessment was then repeated.

Pooling and hybridisation

Prior to hybridisation (Fig. 2), all libraries were normalised to 10 nM, using 10 mM
Tris (pH 8.0) and then combined into pools of 20 to 24 libraries, each containing 10 µl (0.1
pmol) of each normalized library (i.e. a total of ca. 600-700 ng DNA in each pool, assuming
an average fragment size of ca. 450 bp). To ensure even sequencing across all samples in a
pool, species for pooling were selected to minimize the range of DNA fragment sizes and
ensure a narrow taxonomic breadth. The latter criterion was needed because samples that are
more closely related to the taxa used to construct the probe set tend to preferentially
hybridise. This can lead to an over-representation of their sequences in the DNA data if
appropriate care is not taken when selecting species for the sequencing pool. In rare cases,
such as smaller pools (ca. 10 libraries) of short fragment (i.e. <300 bp) libraries, it was necessary to recalculate the standard volume of normalized libraries to be added to ensure that the final pool contained ca. 500 ng of DNA.

The pooled libraries were dried in a SpinVac (Eppendorf, Dusseldorf, Germany), resuspended in 8 µl of 10 mM Tris (pH 8.0) and enriched by hybridising with the Angiosperms353 probe kit (Johnson et al. 2019; Arbor Biosciences myBaits Target Sequence Capture Kit, ‘Angiosperms 353 v1’, Catalogue #308196) following the manufacturer’s protocol, version 4.0. Hybridisation was typically performed at 65°C for 24 h, with reactions topped with 30 µl of red Chill-out Liquid Wax (Bio-Rad, Hercules, CA, USA) to prevent evaporation. However, for short libraries (i.e. <350 bp) the temperature was reduced to 60°C, following the recommendations of Arbor Biosciences.

The target-enriched pools were amplified using the KAPA HiFi 2X HotStart ReadyMix PCR Kit (Roche, Basel, Switzerland) or NEBNext Q5 HotStart HiFi PCR Master Mix (New England BioLabs, Ipswich, MA, USA) for eight to 14 cycles. Amplified pools were then purified using Agencourt AMPure XP Beads (at 0.9X the sample volume) and eluted in 15 µl of 10 mM Tris (pH 8.0).

Products were quantified with a Quantus fluorometer and re-amplified if the concentration was below 6 nM, with three to six PCR cycles (see above). Final products were assessed using the TapeStation to determine the distribution of fragment sizes. The target-enriched pools were normalized to 6 nM (using 10 nM Tris, pH 8.0) and multiplexed for sequencing, with the number of target-enriched pools combined in each sequencing pool varying from two to 20 (comprising a total of 48-384 samples) depending on the sequencing platform and service provider requirements.
DNA sequencing

Initially, DNA sequencing was performed on an Illumina MiSeq at RBGK with version 3 chemistry (Illumina, San Diego, CA, USA) and ran for 600 cycles to generate 2 × 300 bp paired-end reads. Subsequently, DNA sequencing was outsourced (Macrogen, Seoul, South Korea, or Genewiz, Takeley, UK) and performed on an Illumina HiSeq producing 2 × 150 bp paired-end reads. Raw reads were deposited in the European Nucleotide Archive under an umbrella project (accession number PRJEB35285) and can be accessed from the individual sample records in the Kew Tree of Life Explorer.

Sequence assembly

Coding sequences were recovered from target-enriched sequence data using our pipeline recoverSeqs (accessible from our GitHub repository https://github.com/RBGKew/KewTreeOfLife, pypaftol ‘paftools’ submodule) to retrieve sequences orthologous to the Angiosperms353 target gene set (Johnson et al. 2019; https://github.com/mossmatters/Angiosperms353). This target set contained multiple reference sequences per gene, thereby covering a large phylogenetic breadth to facilitate read recovery across angiosperms.

The process comprised four main stages (Fig. 2), applied to each sample: (i) sequence reads were trimmed using Trimmomatic (Bolger et al. 2014) with the following parameters: ILLUMINACLIP: <AdapterFastaFile>: 2:30:10:2:true, LEADING: 10, TRAILING: 10, SLIDINGWINDOW: 4:20, MINLEN: 40, with the adaptor fasta file formatted for palindrome trimming, (ii) trimmed read pairs were mapped to the Angiosperms353 target genes with TBLASTN. A representative reference sequence for each gene was then selected by identifying the sequence with the largest number of mapped reads. (iii) This representative gene was used as the reference for assembling the gene-specific reads using an overlap-based
assembly algorithm (--assembler overlapSerial option). OverlapSerial was developed specifically for this project (see our GitHub repository) with the aim of improving gene recovery, in terms of gene length and number, relative to the widely used HybPiper (Johnson et al. 2016) and was used as follows. First, the reads were aligned to and ordered along the reference sequence based on a minimum alignment size of 50 bases (--windowSizeReference option) with a minimum sequence identity of 70% (--relIdentityThresholdReference option). Consecutive reads ordered along the reference sequence were aligned in a pair-wise manner to find read overlaps. If an overlap of at least 30 bases (--windowSizeReadOverlap option) and 90% sequence identity (--relIdentityThresholdReadOverlap option) was found, the aligned reads were used to construct a consensus contig with ambiguous bases represented by ‘N’. This last parameter resulted in one or more sets of aligned reads with ≥90% sequence identity, each set being merged into a single contig. In the final stage, the exonerate protein2genome program was used to identify the exon-intron structure within each contig. One or more contigs were chosen that best represented the structure of the exon(s) in the reference gene chosen in step (ii). If the exons existed in multiple contigs, those contigs were joined together to form the recovered gene coding sequence.

Target gene recovery success was assessed for each sample by calculating the number of genes recovered and the sum of the recovered gene lengths. Samples were removed from downstream analyses if the sum of the recovered gene lengths fell below 20% of the median value across all samples.

Public data mining

In addition to newly generated target sequence capture data, the Angiosperms genes were mined from publicly available genomic data (Fig. 2). For Data Release 1.0, we focused on mining data from the 1KP Initiative (Carpenter et al. 2019; Leebens-Mack et al.
Baker et al.

2019) and published genomes with gene annotations (https://plants.ensembl.org/), although other data sources (e.g. the Sequence Read Archive) will be data-mined for future releases. The genes were retrieved from assembled transcript sequences (1KP) or coding sequences (CDS; genomes) using paftools retrievetargets from our pipeline, which uses BLASTN to identify and extract the genomic or transcriptomic sequences corresponding to the 353 genes. BLASTN relies on sequence identity (>70%) and the transcript or CDS with the highest identity is considered to be the ortholog of a given target. Because initial recovery of genes from 1KP transcripts using the standard Angiosperms353 target gene set (Johnson et al. 2019) was unsatisfactory, we used an expanded Angiosperms353 target set to improve matching and retrieval of genes. The expanded dataset is a reduced version of the 1KP alignments (Leebens-Mack et al. 2019) produced by Johnson et al. (2019) for the design of the Angiosperms353 probe set from which non-angiosperm sequences had been removed and gap-only sites trimmed. The expanded target set is available from https://datadryad.org/stash/dataset/doi:10.5061/dryad.s3h9r6j and a reformatted version from our GitHub. As with the novel target sequence capture assemblies, data were removed from downstream analyses if the sum of the gene lengths fell below 20% of the median value across all samples.

Family identification validation

To verify the family identification of our processed samples, we implemented two validation steps, which were run in parallel (Fig. 3). The two steps consisted of (i) DNA barcode validation, which utilised nuclear ribosomal and plastid barcodes for DNA-based identification, and (ii) phylogenetic validation, which checked the placement of each sample in a preliminary tree relative to its expected position based on its initial family assignment. Identification checks below the family level were not conducted due to the incompleteness of
adequate reference resources for DNA barcode validation and sparseness of sampling for phylogenetic validation at the genus or species level.

For barcode validation of target sequence capture data (Fig. 3), plastomes and ribosomal DNA were recovered from raw reads using GetOrganelle (Jin et al. 2020) and subsequently queried against databases of reference plant barcodes using BLASTN (Camacho et al. 2009). For 1KP samples, transcriptome assemblies were directly used as queries in BLASTN. Note that we considered the family identity of annotated genomes to be correct and hence a barcode validation was unnecessary. Six individual barcode reference databases were built from the NCBI nucleotide and BOLD databases (https://www.ncbi.nlm.nih.gov/nuccore; https://www.boldsystems.org/, accessed on 29/10/2020), one for the whole plastome, and the remaining five for specific loci (nuclear ribosomal 18S, as well as plastid rbcL, matK, trnL, and trnH-psbA). As for samples, the taxonomy of reference sequences was standardized to WCVP (WCVP 2020). BLAST results were further filtered with a minimum identity >95% and a minimum coverage of reference locus ≥90% (except for whole plastomes, for which only a filtering based on minimum length was applied).

Tests could only be completed if a sample’s given family was present in the barcode databases and if at least one BLAST match remained after filtering. Thus, zero to six barcode tests were conducted per sample. A sample passed an individual test if the first ranked BLAST match (ranked by percentage of identity) confirmed its original family identification and failed otherwise. The final result of the barcode validation following the six individual barcode tests were determined as follows: (i) Confirmed, if one or more barcode tests matched the family identification of a sample; (ii) Rejected, if more than half of the barcode tests gave the same incorrect family identification (requires at least two barcode tests); (iii) Inconclusive (otherwise). Further details of the barcode validation methods can be found in
Supplementary Material available on Dryad. The scripts and lists of NCBI and BOLD accessions used in barcode databases are available on our GitHub repository.

To conduct phylogenetic validation (Fig. 3), a preliminary phylogenetic tree was built using the complete, unvalidated dataset, following the phylogenetic methods described below. We then assessed which nodes best represented each order and family in the tree. For every node in the tree, two metrics were calculated for all families and orders: (i) the proportion of samples belonging to a given order/family that are descendants of the node, and (ii) the proportion of samples descending from the node that belong to the order/family. The two metrics were then multiplied to produce an overall taxon concordance score. For each family and order, the highest scoring node was subsequently considered to best represent the taxon in the tree (allowing the identification of outlying samples). A node with a score of 1 for a given order/family is the crown node (most recent common ancestral node) of that taxon, which is monophyletic in the tree. See Supplementary Figure S1 for an illustration.

The family identification of each sample was determined as (i) Confirmed: if identified as belonging to a family whose best scoring node had a taxon concordance score >0.5 and found as a descendant of this node in the tree, (ii) Rejected: if identified as belonging to a family whose best scoring node had a taxon concordance score >0.5 but not found as a descendant of this node, or (iii) Inconclusive: if identified as belonging to a family whose best scoring node had a taxon concordance score ≤0.5. Note that for families represented in the tree by a single sample, the validation was performed with respect to their orders. If the order was represented by a single sample, the validation result was coded as inconclusive.

The outputs of the phylogenetic and DNA barcode validation were combined to identify samples for automatic inclusion and exclusion from the final dataset, and samples for which a decision on inclusion/exclusion was subject to expert review (Fig. 3). Exclusions
after expert review were made based on implausible tree placement (e.g. wrong higher clade) or sample misidentification (e.g. match to another family in the barcode validation).

All assembled Angiosperms353 gene data from all samples validated for inclusion form the basis of Data Release 1.0. These were made publicly available via the Kew Tree of Life Explorer.

Phylogeny estimation

We inferred a phylogenetic tree from all validated data (Data Release 1.0) for presentation in an interactive format in the Kew Tree of Life Explorer. This species tree was estimated from gene trees using the multi-species coalescent summary method implemented in ASTRAL-III (Zhang et al. 2018). In addition to the angiosperm samples, ten samples representing seven gymnosperm families from the 1KP initiative were mined for Angiosperms353 orthologs (using retrievetargets, as described above) and included in all analyses as outgroup taxa. Our phylogenomic pipeline, available from our GitHub repository, is summarised below.

For each gene, DNA sequences were aligned with UPP 4.3.12 (Nguyen et al. 2015). At the start of the alignment process a set of 1,000 sequences were selected for an initial backbone tree. Option -M was set to ‘-1’ so that sequences could be selected within 25% of the median full-length sequence. Filtering and trimming of the alignment were performed with AMAS (Borowiec 2016) as follows. Sequences with insufficient coverage (<60%) across well occupied columns of each gene alignment were removed. Well occupied columns were defined as those with more than 70% of positions occupied. Then, alignment columns with <0.3% occupancy were removed to remove very rare or unique insertions. Finally, sequences with a total length of less than 80 bases were removed, and genes with <30 overlapping bases (at the 70% threshold mentioned above) were excluded.
Gene trees were estimated with IQ-TREE 2.0.5 (Minh et al. 2020) inferring branch support using the ultrafast bootstrap method (option -B; Hoang et al. 2017) with the maximum number of iterations set to 1,000 (option -nm) and using a single model of evolution (option -m GTR+F+R). The use of a single model without testing many models of evolution was a pragmatic choice, following Abadi et al. (2019). TreeShrink 1.3.4 (Mai and Mirarab 2018) was used to remove abnormally long branches from gene trees using default settings, except option -b, which was set to 20. The alignment and gene tree estimation steps were then repeated on the samples retained by TreeShrink. Before reconstructing the species tree using ASTRAL-III, nodes in the gene trees with bootstrap support values less than 30% were collapsed using nw_ed from Newick Utilities 1.6.0 (Junier and Zdobnov 2010). This value was deduced from interpreting Figure 1 in Hoang et al. (2017), adjusting the standard bootstrap threshold of 10% (recommended for ASTRAL-III), to 30% for the ultrafast bootstrap.

All gene alignments, gene trees and the ASTRAL-III species tree are available for download from secure FTP and the Kew Tree of Life Explorer. In addition, the species tree is available to browse through an interactive tree viewer implemented within the Kew Tree of Life Explorer (see also Supplementary Fig. S2).

Data portal implementation

To disseminate results, a data portal (the Kew Tree of Life Explorer; https://treeoflife.kew.org) was designed and implemented (Fig. 4) with a layered architecture that comprised: (i) a MariaDB running on a Galera multi-master cluster as a database management system; (ii) an API written in Python using the Flask framework and the SQLAlchemy library; (iii) a front-end written using the Vue.js framework and Nuxt.js for the tabular data (used to provide access to gene and specimen data) and content pages; (iv) a tree
visualisation module developed from the open source application PhyD3 (Kreft et al. 2017) using D3.js (Bostock 2012) for data visualisation; and (v) deployment on a Linux (CentOS 7) server using Nginx as web server and load balancer.

The data, with appropriate metadata and documentation, are available for public download over secure FTP (http://sftp.kew.org/pub/treeoflife/) and the Kew Tree of Life Explorer under a Creative Commons Attribution 4.0 International (CC BY 4.0) license. When superseded by new releases, archived earlier releases will remain accessible via secure FTP.

**RESULTS**

**Initial dataset**

The initial dataset prior to processing and analysis comprised data from 3,272 angiosperm samples, representing 413 families of angiosperms (99%) and 2,428 genera (18%; Table 1). We generated novel target sequence capture data for 2,522 of these samples, which included 104 angiosperm genera that have never been sequenced before. Data for the remainder were mined from public sources (689 1KP transcriptomes, 61 annotated genomes). The majority of target sequence capture data were generated from the RBGK collections as follows: DNA Bank (43%), herbarium (28%), silica gel-dried tissue collection (8%), living collection (2%), and Millennium Seed Bank (0.3%). The remaining 19% of samples included in this study were provided by various collaborators of the PAFTOL project, either as DNA samples or as dried tissue (see Acknowledgements).

Sequence recovery from all 2,522 target sequence capture samples (prior to any quality controls) is visualised in Figure 5. Eighty-four target sequence capture samples and eleven 1KP transcriptomes were removed from downstream analyses because the sum of gene lengths did not meet the quality threshold of 20% of the median value across all samples.
Family identification validation

The remaining 3,177 samples (Table 1) were processed through our sample family identification validation pipeline (Fig. 3, Supplementary Tables S1 and S2). Of these, 3,064 (97%) were automatically cleared for inclusion and 67 were automatically excluded (Supplementary Table S1). The remaining 46 samples were held for expert review, after which 35 were cleared for inclusion and 11 were excluded due to implausible tree placements. The majority of excluded samples (64 out of 78) were from the novel target sequence capture data, although 14 were 1KP transcriptomes, highlighting the risk of sample misidentification in even the most highly curated datasets. Further details regarding the results obtained during the family identification validation by DNA barcoding can be found in Supplementary Material available on Dryad.

The final validated dataset for Data Release 1.0 consisted of 3,099 angiosperm samples (Table 1), only 5% fewer than were present in the initial dataset. These samples represent all 64 orders, 404 families (96%; 212 represented by >1 sample), 2,333 genera (17%) and 2,956 species (0.01%).

Data Release 1.0: sequence quality and gene recovery

Nine statistics were used to assess the sequence quality across the 3,099 samples of Data Release 1.0 (Table 2). For the 2,374 target sequence capture samples, the mean percentage of on-target reads was 8%, the mean read depth per sample across all recovered genes was 90x with a median value of 38x and the mean percentage length of recovered genes per sample was 62%. The number of genes and the sum length of gene sequence recovered per sample were tightly associated as expected, varying continuously across the dataset up to the full set of Angiosperms353 genes and a total gene length of 256.9 kbp, close to the maximum expected length of 260 kbp for recovering genes with this target gene set (Fig. 5). The total length of sequence recovered from target sequence capture data was shorter
than for samples mined for Angiosperms353 genes from 1KP transcriptomes or annotated genomes data (Table 2). The reason for the shorter length of the recovered genes is that some exons were omitted during the process of refining 1KP alignments to select gene instances for the design of the Angiosperms353 probes (Johnson et al. 2019). These missing exons were however present in the expanded Angiosperms353 target set and were therefore retrieved during data mining from 1KP transcriptomes and annotated genomes. The variation in performance of target enrichment across different samples, illustrated by the measures of variability shown in Table 2, likely reflects the variation in structure and metabolite composition of the starting tissue, which is known to impede DNA extraction from various species and its downstream manipulation. This variation is one of the challenges in dealing with samples from a broad taxonomic range such as across the evolutionary diversity of angiosperms. Variation in gene recovery across orders is visualised in Supplementary Figure S3.

**Phylogenetic results**

The final phylogenetic tree as inferred from Data Release 1.0 is publicly available in interactive form via the Kew Tree of Life Explorer. In the current release, the tree is annotated with local posterior probabilities (LPP, as given by ASTRAL-III) as indicators of branch support. Other measures of support (e.g. quartet scores) can be found within tree files accessible via the RBGK secure FTP. For completeness, the tree is also available in various formats, including Newick (Supplementary Fig. S2).

As a result of filtering and trimming steps during alignment, six genes in Data Release 1.0 were excluded from downstream phylogenetic analysis due to insufficient overlap between sequences. All statistics provided below refer to the remaining dataset. Thus, the species tree is based on 347 gene alignments totalling 824,878 sequences, 489,086,049 base pairs and 532,260 alignment columns. Of these, 509,987 columns (96%) are variable and
475,181 columns (89%) are parsimony informative. The proportion of gaps across all alignments is 61.6% and the median number of genes per sample is 284 (mean: 265.3, standard deviation (SD): 64.3, min: 22, max: 347; Supplementary Table S3). The median number of samples per gene alignment is 2,421 (mean: 2,377.2, SD: 359) and median alignment length is 1,259 (mean: 1,533.9, SD: 985.7; Table 3). The resulting gene trees are highly resolved, with a median support across all nodes (ultrafast bootstrap) of 98% (mean: 87.8%, standard deviation (SD): 18.560) across all nodes in all gene trees (Supplementary Fig. S4). Only 1.3% of all nodes in all gene trees are very poorly supported (ultrafast bootstrap <30%; Supplementary Fig. S4) and thus collapsed prior to species tree inference.

Further statistics for individual gene alignments and gene trees are reported in Table 3 and Supplementary Table S3.

The species tree accommodates 82% of the quartet relationships in the gene trees (ASTRAL normalized quartet score of 0.82). The majority (76.8%) of nodes in the species tree were well-supported (LPP ≥95%, cf. Sayyari and Mirarab 2016), and only seven nodes were informed by too few gene trees (i.e. <20) to evaluate support. Comparing node support in the species tree at different taxonomic levels (Supplementary Fig. S5), median quartet support is progressively higher towards shallower taxonomic levels (Supplementary Fig. S5c), while the effective number of gene trees informing nodes shows the opposite trend (Supplementary Fig. S5e). Local posterior probabilities show a tendency to be lower (1st quartile) at the deepest taxonomic level (Supplementary Fig. S5a). Major groups (i.e. monocots, asterids and rosids) show similar distributions of both local posterior probabilities (Supplementary Fig. S5b) and quartet support values (Supplementary Fig. S5d), despite the fact that the effective number of gene trees supporting nodes is more variable in monocots (Supplementary Fig. S5f), which is the result of the lower recovery rates for some orders in this group such as Alismatales, Commelinales and Liliales (Supplementary Fig. S3).
Discounting taxa represented by a single sample (193 families, one order), 96% of testable families and 83% of testable orders were resolved as monophyletic in the species tree. Most of the samples of non-monophyletic families and orders could be assigned to a clade that represents the family or order well, despite lacking some samples and/or containing some outlier samples from other taxa (“concordant taxa” where taxon concordance score >0.5, see Materials and Methods for details). Only five families (Francoaceae, Hernandiaceae, Phyllanthaceae, Pontederiaceae and Schlegeliaceae, represented by 11 samples) and two orders (Bruniales and Icacinales, represented by six samples) were so dispersed that this was not possible (“discordant taxa” where taxon concordance score ≤ 0.5).

At the family level, 2,893 samples were resolved in the expected family, two samples were resolved in an unexpected position, and 204 samples were not testable because they belonged to a discordant family or a family represented by a single sample. At the order level, 3,060 samples were resolved in the expected order, 32 samples were resolved in an unexpected position, and seven samples were not testable (see Supplementary Tables S4-S6 for lists of specimens from singly represented taxa, poorly resolved taxa, and outliers to well-resolved taxa, respectively). Placements of all but five genera and seven families were consistent with the WCVP/APG IV taxonomic hierarchy of genera, families and orders. Concordance with existing taxonomy was lower at the genus level, with only 74% of testable genera resolving as monophyletic and 47 genera (represented by 130 samples) being discordant; these numbers partly reflect the deliberate inclusion of multiple samples from genera suspected a priori to be potentially non-monophyletic.

In addition to resolving most genera, families and orders as monophyletic, our tree supports more than half (58%) of the relationships among orders presented by the Angiosperm Phylogeny Group (APG IV 2016; Supplementary Fig. S6). Congruence with APG IV varies among major clades, being notably high in magnoliids (100% of APG IV...
relationships supported) and monocots (80%), while being substantially lower in eudicots (47%), especially in rosids (33%). Nodes in our tree that are congruent with APG IV ordinal relationships are slightly better supported on average (mean LPP 0.98, median 1) than nodes that are incongruent with APG IV (mean LPP 0.75, median 0.94).

**Tree of Life Explorer**

The Kew Tree of Life Explorer ([https://treeoflife.kew.org](https://treeoflife.kew.org)) provides open access to taxon, specimen, sequence, alignment and tree data, with associated metadata for the current data release in accordance with the Toronto guidelines on pre-publication data sharing ([Toronto International Data Release Workshop Authors 2009](https://treeoflife.kew.org)). Users can browse by species, gene or interactive phylogenetic tree. The species interface permits searches by order, family, genus or species, and provides voucher specimen metadata (including links to online specimen images, where available), simple sequence metrics, access to assembled genes and raw data. The gene interface documents all Angiosperms353 genes and associated metrics, links to gene identities in UniProt ([https://www.uniprot.org/](https://www.uniprot.org/)) and provides access to assembled genes across taxa. The tree of life interface enables browsing and taxon searching of the species tree inferred from the current release dataset, as well as tree downloads (as PNG or Newick) and zooming into user-defined subtrees. All processed data (assembled genes, alignments, gene trees, species trees) and archived releases are available from RBGK’s secure FTP site ([http://sftp.kew.org/pub/treeoflife/](http://sftp.kew.org/pub/treeoflife/)), whereas raw sequence reads are deposited within the European Nucleotide Archive (project number PRJEB35285) for integration within the Sequence Read Archive.

**DISCUSSION**


The new phylogenomic platform described here is a major milestone towards a comprehensive tree of life for all flowering plant species. The sequencing of a standardised nuclear marker set of this scale for so many taxa is unprecedented, opening doors to a highly integrated future for angiosperm phylogenetics in the genomic era. Much like a “next generation” $rbcL$, which underpinned so many Sanger sequencing-based plant phylogenetic studies, the Angiosperms353 genes offer opportunities for continuous synthesis of HTS data across angiosperms. The foundational dataset presented here can be re-used or extended for tree of life research at almost any taxonomic scale (Johnson et al. 2019; Larridon et al. 2019; Van Andel et al. 2019; Murphy et al. 2020; Pérez-Escobar et al. 2020; Shee et al. 2020; Slimp et al. 2020; Beck et al. 2021). This is the first phylogenetic project to gather novel HTS data across angiosperms with a stratified taxon sampling at the genus level. Our sampling strategy systematically and comprehensively represents both the diversity of angiosperms and their deep-time diversification. As genus-level sampling becomes increasingly complete—a target that is well within reach—this backbone will substantially increase our ability to study the dynamics of plant diversity over time and revisit long-standing questions in systematics (Magallón et al. 2018; Sauquet and Magallón 2018; Soltis et al. 2019). Importantly, it will also sharpen the focus on truly intractable phylogenetic problems (Yang et al. 2020; Zhao et al. 2020), encouraging the exploration of the biological drivers of these phenomena.

Our approach has already led to a burst of community engagement. More than a dozen studies utilising Angiosperms353 probes are already published (e.g. Larridon et al. 2019; Howard et al. 2020; Murphy et al. 2020; Pérez-Escobar et al. 2020; Shee et al. 2020; Slimp et al. 2020; McLay et al. in press), and two journal special issues focused on the probe set are in preparation (Baker et al. in press) arising from a recent symposium (Lagomarsino and Jabaily 2020). The probe set has also been adopted by the Genomics for Australian Plants consortium (https://www.genomicsforaustralianplants.com/), which aims to sequence all angiosperms.
Australian angiosperm genera, coordinating with the PAFTOL project to optimise collective
taxonomic coverage. A subset of the Angiosperms353 genes is now accessible for non-
angiosperm land plants thanks to a probe set developed in parallel (Breinholt et al. 2021),
inviting the prospect of data integration across all land plants. Angiosperms353 genes (as
distinct from the Angiosperms353 probes) are also being leveraged as components of custom-
designed probe sets (e.g. Jantzen et al. 2020; Ogutcen et al. 2021). This approach gives all the
integrative benefits of Angiosperms353, while permitting (i) the tailoring of Angiosperms353
probes to a taxonomic group by using more specific target data to increase gene recovery, and
(ii) the inclusion of additional loci pertinent to the research in question. Angiosperms353
probes have also been directly combined with an existing custom probe set (Nikolov et al.
2019) as a “probe cocktail” in a single hybridisation, capturing both sets of targets
simultaneously with remarkable efficiency (Hendriks et al. in press). These possibilities
render the invidious choice between specific and universal probe sets increasingly irrelevant
(Kadlec et al. 2017).

Although target sequence capture is the most cost-effective way to retrieve the
Angiosperms353 genes at the current time, the opportunity to mine the genes from other
kinds of HTS data (e.g. shotgun sequence data, RNA sequence data) should not be
overlooked. This represents a further opportunity for community engagement, both via
mining of public data in the Sequence Read Archive, for example, and by adding value to
new data being generated with these methods. A stronger understanding of the sequencing
requirements (e.g. coverage) for gene recovery from such data could guide new data
generation so that Angiosperms353 genes can be retrieved routinely as a by-product of other
research. We took several open data measures to encourage community uptake, in both the
design of our tools and the sharing of our data. The Angiosperms353 probe set itself was
designed to be a transparent, “off-the-shelf” toolkit that is open, inexpensive and accessible to
all, especially researchers discouraged by the complexity and cost of custom probe design (Johnson et al. 2019). Our sequence data for Angiosperms353 genes are openly available via the Kew Tree of Life Explorer and the Sequence Read Archive, as a public foundation dataset shared according to pre-publication best practice (Toronto International Data Release Workshop Authors 2009). The Explorer offers enhanced transparency and accessibility by allowing users to navigate the data via a phylogenetic snapshot of the current release, along with metadata (e.g. specimen data) and intermediate data (e.g. gene assemblies, alignments, gene trees). Thanks to these resources, cross-community collaboration via Angiosperms353 is gaining momentum.

Our tree, which is based on the most extensive nuclear phylogenomic dataset in flowering plants to date, is strongly supported, credible and highly congruent with existing taxonomy and many hypothesized relationships among orders (APG IV 2016; Supplementary Fig. S6). The data confirm both the effectiveness of Angiosperms353 probes across all major angiosperm clades and the ability of the genes to resolve relationships across taxonomic scales (Supplementary Fig. S5). Variable sequence recovery notwithstanding (Table 2, Supplementary Fig. S3), most nodes in our tree are underpinned by large numbers of gene trees (Supplementary Fig. S5e), allowing the species tree to be inferred with confidence (Supplementary Fig. S5a) despite gene tree conflict (Supplementary Fig. S5c). However, even the most strongly supported phylogenetic hypotheses must be viewed with caution as they may be biased by model misspecification and wrong assumptions. Moreover, our “first pass” analyses based on a set of standard methods may not suit this dataset perfectly (see below). Nevertheless, our findings are rendered credible by their high concordance with taxonomy, an independent point of reference that has been extensively ground-truthed by pre-phyllogenomic DNA data, especially plastid loci. Agreement with existing family circumscriptions is particularly striking. In contrast, congruence with previously
hypothesized relationships among orders (APG IV 2016) is much lower (Supplementary Fig. S6). Some of these earlier hypothesized ordinal relationships derive from relatively weak evidence (bootstrap/jackknife >50%; APG IV 2016), which may partly explain this disagreement. However, it may also be due to phylogenetic conflict between nuclear and plastid genomes, as the established ordinal relationships rest primarily on evidence from plastid loci, substantiated more recently by plastid genomes (Li et al. 2019). It is hardly surprising, then, that a large-scale nuclear analysis presents strongly supported, alternative relationships (Supplementary Fig. S6). The conundrum remains that these incongruences are visible at the ordinal backbone, but not the family level. A more comprehensive exploration of these relationships, the underlying phylogenetic signal and their systematic implications is currently underway.

The analyses presented here are primarily intended as a window onto the information content of our current data release and are not a complete exploration of the data. Thus, downstream application of the current species tree comes with caveats. We used current, widely accepted methods in a pipeline that can be re-run in a semi-automated fashion whenever we release new data. As a consequence, not all possible analysis options and effects have been explored. We anticipate that users of our data will probe it more rigorously and will tailor both sampling and phylogenomic analyses to their specific questions. For example, users may leverage our data by enriching a subset with denser sampling of their own to address more focused evolutionary questions. A further exemplar use case could be deeper re-analysis of our data from raw sequence reads to deepen understanding of gene history and conflict.

Important limitations in our analysis relate to (i) taxon sampling, (ii) gene selection (ii) gene recovery, (iii) models of sequence evolution and (iv) paralogy. Taxon sampling for intermediate data releases is biased by the current state of progress towards our systematic
sampling strategy. This will be addressed in future data releases and can be adjusted by users of our data. In addition, potential phylogenetic biases attributable to the function or other properties of the Angiosperms353 genes remain poorly understood and require further investigation. Gene recovery relied upon the standard Angiosperms353 target file (Johnson et al. 2019), which, by its universal nature, can yield patchy results. However, it has recently been reported that tailoring target sequences to specific taxonomic groups can improve recovery (McLay et al. in press); this will be tested in future data releases. Moreover, we are yet to exploit intronic data captured in the “splash zone” adjacent to our target exons. By necessity, our “first pass” phylogenetic analysis does not explore the fast-evolving spectrum of methodological options available for phylogenomic analysis. For example, we rely on a simple standard model of sequence evolution, but more sophisticated models accounting for codon positions or amino acids may improve phylogenetic inference. Potential paralogy is not addressed by our current pipeline. The genes underpinning our analysis were carefully chosen to represent single-copy genes across flowering plants (Johnson et al. 2019; Leebens-Mack et al. 2019). The very low proportion of ambiguous bases across all gene alignments (0.01%; Table S3) suggests that gene assembly was not strongly impacted by divergent gene copies, such as paralogs. However, some paralogy may have gone unnoticed due to the pervasiveness of gene and genome duplication in plants (Li and Barker 2020). Overall, we expect that the occasional presence of paralogs in our current analysis would more likely lead to inflated estimates of gene tree incongruence, and thus result in reduced support values, than significant topological biases (Yan et al. 2020). Thus, we consider our tree relatively conservative while acknowledging that we are not yet exploiting the full potential of our data. Although a rigorous analysis of paralogy in Angiosperms353 genes was not tractable for this data release, we look forward to deeper insights emerging as community-wide engagement with Angiosperms353 grows.
In the immediate future, we will deliver a further data release through which we expect to reach the milestone of sampling 50% of all angiosperm genera. This target will be achieved through substantial novel data production by PAFTOL and collaborators, augmented by data mined from public sources. In-depth phylogenetic analyses of our data and their evolutionary implications are also underway.

Beyond this point, we see three priority areas in which future platform developments might be concentrated, resources permitting. Firstly, taxon sampling to the genus level must be completed. Our original target of sampling all angiosperm genera remains, but the mode of reaching this is likely to evolve. We anticipate an acceleration in production of Angiosperms353 data by the broader community. The completion of generic-level sampling will require both the integration of community data in the broader angiosperm tree of life as well as strategic investment in filling inevitable data gaps for orphan groups. Secondly, numerous opportunities for refinement exist across our methods. For example, insights from our data might permit the optimisation of the Angiosperms353 probes to improve gene capture. Efficiency of gene assembly from sequence data can also be improved bioinformatically (McLay et al. in press). However, as costs of sequencing decline, target sequence capture in vitro may no longer be necessary, the target genes simply being mined from sufficiently deeply sequenced genomes. Thirdly, for the full integrative potential of Angiosperms353 genes to be achieved, infrastructure for aggregating and sharing this coherent body of data must be improved. While the Kew Tree of Life Explorer provides a proof-of-concept, it is the public data repositories (e.g. NCBI, ENA) that offer the greatest prospects of a mechanism to achieve this. To fully parallel the earlier success of public repositories for facilitating single-gene phylogenetic trees (e.g. *rbcL*, *matK*), new tools are
needed to assist with efficient upload and annotation of target capture loci and associated 
metadata.

Even with a completed genus-level angiosperm tree of life well within reach, the 
monumental task of sampling all species remains. The scale of this challenge is 24-fold 
greater than the genus-level tree towards which we are currently working. However, with 
sufficient investment, increased efficiencies and community engagement, such an ambition 
could potentially be realised. Collections-based institutions are poised to play a critical role in 
this endeavour through increasingly routine molecular characterisation of their specimens, 
perhaps as part of digitisation programmes, and are already facilitating the growing trend 
towards species-complete sampling in phylogenomic studies (e.g. Loiseau et al. 2019; 
Murphy et al. 2020; Kuhnhäuser et al. 2021). Our platform demonstrates how large-scale 
phylogenomic projects can capitalise on natural history collections to achieve a much more 
complete sampling than hitherto possible.

The growing movement to sequence the genomes of all life on Earth, inspired by the 
Earth Biogenome Project (Lewin et al. 2018), significantly boosts the prospects for 
completing the tree of life for all species, but is hampered by the focus on “gold standard” 
whole genomes requiring the highest quality input DNA. Our platform offers the opportunity 
to bridge the gap between the ambition of these projects and the vast phylogenomic potential 
of natural history collections. However, as life on Earth becomes increasingly imperilled, we 
cannot afford to wait. To meet the urgent demand for best estimates of the tree of life, we 
must dynamically integrate phylogenetic information as it is generated, providing synthetic 
trees of life to the broadest community of potential users (Eiserhardt et al. 2018). Our 
platform facilitates this crucial synthesis by providing a cross-cutting dataset and directing 
the community towards universal markers that seem set to play a central role in completing 
an integrated angiosperm tree of life.
DATA AVAILABILITY AND SUPPLEMENTARY MATERIAL

All data generated in this study are publicly released under a Creative Commons Attribution 4.0 International (CC BY 4.0) license and the Toronto guidelines on pre-publication data sharing (Toronto International Data Release Workshop Authors 2009). The data are accessible via the Kew Tree of Life Explorer (https://treeoflife.kew.org) and our secure FTP (http://sftp.kew.org/pub/treeoflife/). Raw sequence reads are deposited in the European Nucleotide Archive (https://www.ebi.ac.uk/ena/browser/home) under umbrella project PRJEB35285. Scripts and other files relating to our phylogenomic pipeline are available at our GitHub (https://github.com/RBGKew/KewTreeOfLife). Supplementary materials cited in this paper plus Data Release 1.0 datasets duplicated from our secure FTP (assembled genes, gene alignments, gene trees, species tree, examples of scripts) are available from the Dryad Digital Repository (https://doi.org/10.5061/dryad.ns1rm8ps7).

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**Figure Legends**

**Figure 1.** Summary workflow. Overview of steps taken by the PAFTOL project to generate Data Release 1.0 of the Kew Tree of Life Explorer ([https://treeoflife.kew.org](https://treeoflife.kew.org)). The stages of the workflow are further elaborated in Figs. 2–4.

**Figure 2.** Sample processing and data analysis workflows. Sample processing (left): processes are indicated by bold headings with reagents and machines used given below; quality control (QC) checkpoints are indicated in dark grey boxes. Data analysis (right): pipeline products are shown in blue-green circles (available to download via the Kew Tree of Life Explorer, [https://treeoflife.kew.org](https://treeoflife.kew.org)); processes are indicated by bold headings with programs used given below.

**Figure 3.** Family identification validation workflow. Processes are indicated by bold headings. Embedded table (bottom right) indicates decisions made for each sample based on the two validation steps.

**Figure 4.** Data publication workflow. Implementation of Kew Tree of Life Explorer data portal is illustrated. Arrows indicate data flow from internal repository to public interface. Infrastructural components are shown in purple; publicly available information is shown in green. External links available from the portal are listed in the lower left.

**Figure 5.** Density plots of target sequence recovery from our raw data. Data are presented prior to any filtering, illustrating relationships of sum of gene lengths (bp) to (a) the number of mapped reads and (b) the number of recovered genes. Colours indicate density of data.
points. Black dotted lines indicate medians of variables and red dotted lines indicate the
threshold used to remove samples from downstream analyses, set as 20% of the median value
across all samples.
Sample processing
1. Sampling
2. DNA extraction
3. Library preparation
4. Pooling and hybridisation
5. DNA sequencing

Data analysis
6. Sequence assembly
7. Public data mining
8. Family identification validation
9. Phylogeny estimation

Data publication
10. Data portal implementation
**Tables**

**Table 1.** Total number of angiosperm samples included at three stages of data release preparation. The first column represents all samples available in the initial dataset. The second column indicates samples included in our preliminary tree, prior to family identification validation, but after removal of samples for which the sum of the gene lengths fell below 20% of the median value across all samples. The third column provides numbers for the samples made public in the Kew Tree of Life Explorer, Data Release 1.0, and included in our final phylogenetic tree. Numbers of angiosperm families, genera and species in each data subset are provided in brackets (as families/genera/species).

| Data source                  | Initial dataset | Preliminary tree pre-validation | Final tree and Data Release 1.0 |
|------------------------------|----------------|---------------------------------|---------------------------------|
| Target sequence capture data | 2,522 (304/1988/2397) | 2,438 (297/1947/2340) | 2,374 (292/1903/2280) |
| 1KP transcriptomes           | 689 (254/544/682) | 678 (250/530/677) | 664 (245/517/663) |
| Annotated genomes            | 61 (23/43/59)    | 61 (23/43/59)     | 61 (23/43/59)     |
| Total                        | 3,272 (413/2428/3079) | 3,177 (410/2388/3028) | 3,099 (404/2333/2956) |
Table 2. Target sequence capture and gene recovery statistics by sample or gene for Data Release 1.0, including the results of mining of genes from the 1KP and annotated genome datasets. The upper five rows apply to target sequence capture data only. (SD = standard deviation).

|                                | Median   | Mean     | SD    | Minimum | Maximum |
|--------------------------------|----------|----------|-------|---------|---------|
| **Raw reads per sample**       |          |          |       |         |         |
|                                | 1.757 × 10⁶ | 2.822 × 10⁶ | 3.076 × 10⁶ | 1.676 × 10⁴ | 4.054 × 10⁷ |
| **Trimmed reads per sample**   |          |          |       |         |         |
|                                | 1.585 × 10⁶ | 2.549 × 10⁶ | 2.791 × 10⁶ | 1.391 × 10⁴ | 3.605 × 10⁷ |
| **Percentage of reads on-target** |          |          |       |         |         |
| per sample⁠               | 5.676     | 8.020    | 7.704 | 0.005   | 50.953  |
| **Read depth per sample**     | 38        | 90       | 105   | 5       | 2,243   |
| **Read depth per gene**       | 38        | 97       | 37    | 27      | 226     |
| **Recovered genes per sample:** |          |          |       |         |         |
| Target sequence capture data  | 338       | 330      | 24    | 148     | 353     |
| 1KP transcriptomes             | 341       | 328      | 44    | 30      | 353     |
| Annotated genomes              | 346       | 341      | 13    | 287     | 353     |
| **Recovered genes lengths**   |          |          |       |         |         |
| across all samples⁠              | 387       | 477      | 347   | 48      | 3,564   |
| Target sequence capture data  | 717       | 803      | 466   | 50      | 4,689   |
| 1KP transcriptomes             | 341       | 328      | 44    | 30      | 353     |
| Annotated genomes              | 346       | 341      | 13    | 287     | 353     |
### A PHYLOGENOMIC PLATFORM FOR ANGIOSPERMS

| Annotated genomes | 972 | 1,136 | 642 | 45 | 8,601 |
|-------------------|-----|-------|-----|----|-------|

| Sum of recovered gene lengths per sample (bp): |
|-----------------------------------------------|
| **Target sequence capture data**              |
| - 1.613 × 10^5                               |
| - 1.576 × 10^5                               |
| - 4.355 × 10^5                               |
| - 3.433 × 10^5                               |
| - 2.569 × 10^5                               |
| **1KP transcriptomes**                        |
| - 2.753 × 10^5                               |
| - 2.627 × 10^5                               |
| - 6.659 × 10^5                               |
| - 6.498 × 10^5                               |
| - 3.674 × 10^5                               |
| **Annotated genomes**                         |
| - 3.901 × 10^5                               |
| - 3.876 × 10^5                               |
| - 1.868 × 10^5                               |
| - 3.217 × 10^5                               |
| - 4.273 × 10^5                               |

| Percentage length per recovered gene across all samples: |
|---------------------------------------------------------|
| **Target sequence capture data**                        |
| - 63: 88%                                               |
| - 62: 85%                                               |
| - 16: 10%                                               |
| - 27: 44%                                               |
| - 96: 95%                                               |
| **1KP transcriptomes**                                  |
| - 88: 90%                                               |
| - 85: 95%                                               |
| - 10: 16%                                               |
| - 44: 95%                                               |
| - 100: 100%                                             |

| Percentage length of recovered genes per sample:        |
|---------------------------------------------------------|
| **Target sequence capture data**                        |
| - 63: 88%                                               |
| - 62: 85%                                               |
| - 14: 13%                                               |
| - 20: 16%                                               |
| - 95: 100%                                              |
| **1KP transcriptomes**                                  |
| - 88: 90%                                               |
| - 84: 95%                                               |
| - 13: 16%                                               |
| - 16: 100%                                              |

*across all recovered genes

*at bases with ≥4x depth across all recovered genes, calculated by Samtools depth program

*at bases with ≥4x depth across all samples, calculated by Samtools depth program

*see Supplementary Figure S7
percentage length calculated against each representative target gene
Table 3. Properties of the 347 gene alignments and gene trees underpinning the species tree included in the Kew Tree of Life Explorer Data Release 1.0. (SD = standard deviation).

|                          | Median | Mean  | SD   | Minimum | Maximum |
|--------------------------|--------|-------|------|---------|---------|
| Number of samples        | 2,421  | 2,377.2 | 358.8 | 491     | 3,014   |
| % of total samples\(a\) | 77.9   | 76.5  | 11.5 | 15.8    | 96.9    |
| Alignment length         | 1,259.0 | 1,533.9 | 985.7 | 250     | 8,119   |
| % gaps\(b\)             | 58.9   | 57.9  | 11.3 | 14.4    | 85.8    |
| Variable sites           | 1,224  | 1,469.7 | 940.6 | 240     | 7,873   |
| % variable sites         | 96.6   | 96.0  | 2.5  | 81.5    | 100     |
| Parsimony informative sites | 1,137  | 1,369.4 | 859.3 | 233     | 6,792   |
| % parsimony informative sites | 90.7   | 90.0  | 4.20 | 69.1    | 98.9    |
| % nodes in gene trees above 30% UFBS\(c\) | 98.9 | 98.5 | 1.3 | 90.7    | 99.9    |
| Mean support\(c\) of all nodes | 88.1 | 87.8 | 2.7 | 78.9    | 94.3    |
| Median support\(c\) of all nodes | 98.0 | 97.6 | 1.8 | 90.0    | 100     |

\(a\)percentage of samples in species tree present in alignment/gene tree

\(b\)percentage of empty cells in each alignment

\(c\)UFBS: ultrafast bootstrap