RESOURCE ARTICLE



Plant DNA-barcode library and community phylogeny for a semi-arid East African savanna

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Abstract

Applications of DNA barcoding include identifying species, inferring ecological and evolutionary relationships between species, and DNA metabarcoding. These applications require reference libraries that are not yet available for many taxa and geographic regions. We collected, identified, and vouchered plant specimens from Mpala Research Center in Laikipia, Kenya, to develop an extensive DNA-barcode library for a savanna ecosystem in equatorial East Africa. We amassed up to five DNA barcode markers (rbcL, matK, trnL-F, trnH-psbA, and ITS) for 1,781 specimens representing up to 460 species (~92% of the known flora), increasing the number of plant DNA barcode records for Africa by ~9%. We evaluated the ability of these markers, singly and in combination, to delimit species by calculating intra- and interspecific genetic distances. We further estimated a plant community phylogeny and demonstrated its utility by testing if evolutionary relatedness could predict the tendency of members of the Mpala plant community to have or lack "barcode gaps", defined as disparities between the maximum intra- and minimum interspecific genetic distances. We found barcode gaps for 72%-89% of taxa depending on the marker or markers used. With the exception of the markers rbcL and ITS, we found that evolutionary relatedness was an important predictor of barcode-gap presence or absence for all of the markers in combination and for matK, trnL-F, and trnH-psbA individually. This plant DNA barcode library and community phylogeny will be a valuable resource for future investigations.

KEYWORDS

barcode gap, biodiversity, comparative phylogenetics, East Africa, Forest Global Earth Observatory, Mpala Research Centre

1 | INTRODUCTION

DNA barcoding (Hebert, Cywinska, Ball, & DeWaard, 2003) is a tool to identify species and infer the ecological and evolutionary relationships between them. Determining that a "barcode gap" is present

at a locus enables researchers to reliably differentiate species from each other. Biologists use DNA barcodes to study food webs (Pompanon et al., 2012), biological invasions (Dejean et al., 2012), and for biomonitoring (Baird & Hajibabaei, 2012), wildlife forensics (Dawnay, Ogden, McEwing, Carvalho, & Thorpe, 2007), and natural

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product validation (Filonzi, Chiesa, Vaghi, & Marzano, 2010). Genetic information provided by DNA barcodes can also be used to estimate phylogenies that yield insights into community assembly (Erickson et al., 2014; Swenson, 2012), trait evolution (Gill et al., 2016), and lineage diversification (Polato et al., 2018). To attain this wide range of uses, investigators rely upon the availability of DNA-barcode libraries, which do not yet exist for many taxa and for many of the world's most biodiverse regions.

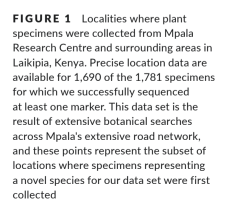
As of January 2019, the Barcode of Life Data System (BOLD) (Ratnasingham & Hebert, 2007) included 20,867 records from vascular plant specimens in Africa. Although Africa comprises ~20% of the global landmass and harbours ~15% of the global plant diversity (Linder, 2014), African plants comprise only ~8% of the vascular plant records available in BOLD. Within Africa, most of the available DNA barcodes for plants are from Western and Southern Africa (13,998 specimens; 67% of African specimens) (Bezeng et al., 2017; Lahaye et al., 2008; Parmentier et al., 2013). These global and continental biases in the availability of genetic resources for plants limit both basic and applied research priorities in Africa (Daru, Berger, & Wyk, 2016).

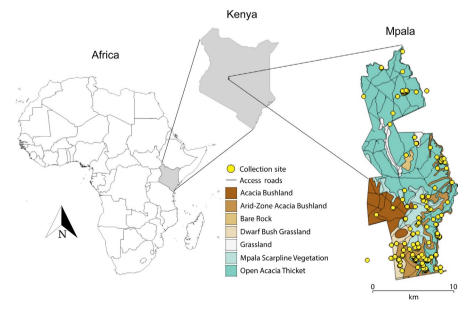
Existing African plant DNA barcodes have helped resolve the systematics of ecologically and economically important taxa, including rosewoods (Hassold et al., 2016), acacias (Boatwright, Maurin, & Bank, 2015; Kyalangalilwa, Boatwright, Daru, Maurin, & Bank, 2013), aloes (Daru et al., 2013; Manning, Boatwright, Daru, Maurin, & Bank, 2014), and the Combretaceae (Gere et al., 2013; Jordaan, Wyk, & Maurin, 2011a, 2011b; Maurin, Chase, Jordaan, & Bank, 2010). Researchers in the field of community phylogenetics (also called phylogenetic community ecology), have used African plant DNA barcodes to understand plant community responses to herbivory (Yessoufou et al., 2013), classify biogeographical regions of Southern Africa (Daru, Bank, et al., 2016), and to assess the evolutionary history of African cycads (Yessoufou, Bamigboye, Daru, & Bank, 2014), underground trees (geoxyles) (Maurin et al., 2014) and thorny savanna plant assemblages (Charles-Dominique et al., 2016).

Increasing the taxonomic and geographic coverage of DNA barcodes for African plants will enrich our understanding of these species, communities, and ecosystems.

In equatorial East Africa, extensive research into conservation biology and the structure and function of savanna ecosystems has taken place at the Mpala Research Centre (MRC) in the Laikipia Highlands of central Kenya (0.293 N, 36.898 E; 1,700-2,000 m a.s.l.: Figure 1). This ~200 km² unfenced conservancy and working ranch sustains diverse wildlife and domestic livestock species, and is the site of several long-term manipulative experiments that aim to elucidate the effects of herbivory and environmental change on semi-arid savanna ecosystems (Goheen et al., 2018). These studies include the Kenya Long-term Exclosure Experiment (KLEE) (Young, Okello, Kinyua, & Palmer, 1998), the Glade Legacies And Defaunation Experiment (Augustine & McNaughton, 2006), and the Ungulate Herbivory Under Rainfall Uncertainty experiment (UHURU) (Goheen et al., 2013; Kartzinel et al., 2014). Additionally, MRC hosts the only savanna site currently participating in the Forest Global Earth Observatory (ForestGEO) network; accordingly, botanical research at MRC contributes to global comparisons of long-term vegetation dynamics.

In 2012, we established a pipeline for the collection of plant vouchers and DNA barcoding at MRC (Kartzinel et al., 2015). We collected both woody and herbaceous plant specimens throughout the property and neighbouring landscapes. For each morphospecies identified in the field, we sequenced up to five plant DNA-barcode markers for one to four specimens per taxon. At the National Museums of Kenya's East Africa Herbarium (EA), expert botanists examined specimens and made taxonomic determinations, revising field-based morphospecific identifications as necessary. Here, we present our DNA barcode reference library for MRC, estimate a community phylogeny, and demonstrate the utility of these resources by testing if relatedness can predict the presence or absence of barcode gaps within this plant community.





2 | MATERIALS AND METHODS

2.1 | Site description, specimen collection, and taxonomic identification

The vegetation at MRC includes semi-arid savanna, acacia bushland, and wooded grasslands with interspersed riparian zones and rocky hills. On average, temperatures range from 11 to 24°C and precipitation totals ~600 mm/year, accumulating mainly during three rainy periods (April-May, July-August, November) (Franz, Caylor, Nordbotten, Rodríguez-Iturbe, & Celia, 2010). Distinct habitat types occur on soil types characterized as red sandy loams (northern and southeastern areas), heavy-clay "black-cotton" vertisols (southwestern), and transitional soils between red-sand and black-cotton habitats (Pringle, Prior, Palmer, Young, & Goheen, 2016). This diversity of habitats types is represented across the ~200 km² MRC land-scape, and similar soil types supporting similar plant communities occur throughout the 9,500 km² Laikipia region and more broadly across East Africa, including Nairobi National Park and parts of the Serengeti-Mara ecosystem.

From 2012 until 2018, we sampled plant species as extensively and thoroughly as possible from all vegetation zones in this ecosystem. Initial collections occurred within the UHURU (Goheen et al., 2013; Kartzinel et al., 2014) and KLEE (Young et al., 1998) experiments, where plant surveys are conducted at regular intervals. Subsequent collections spanned the extensive road network of MRC and the surrounding landscapes. Collection efforts were led by parataxonomists with >10 years of botanical research experience at MRC, with specialized training directed by botanical experts at EA, and supplemented by input from ecologists at multiple institutions. We collected voucher specimens from three individuals per morphospecies and deposited them in the EA, the Smithsonian Institution herbarium (US), and MRC's research and teaching collection. A tissue sample for genetic analyses was collected from each of these specimens, along with up to one additional individual from the field (not vouchered).

At the time of collection, we provisionally identified the specimens, took photographs, and recorded GPS coordinates. Multiple researchers contributed to this collection effort and not all specimens could reliably be identified to species upon initial collection: consequently, some taxa were collected on multiple occasions and we obtained DNA barcodes from as many as 16 specimens per taxon. Expert botanists at the EA identified the specimens to the finest taxonomic level possible (~96% species-level identifications). We reserve use of the word "species" to refer to the 433 taxa assigned accepted Latin binomials in our data set, and we more inclusively use the word "taxa" with reference to these species and an additional 27 provisionally distinguishable taxonomic entities that are currently only resolved to family- or genus-level (N = 460 taxa in total). We present these data using the taxonomic nomenclature currently recognized by the Angiosperm Phylogeny Group (Chase et al., 2016) and The Plant List (2013). Because this nomenclature follows the controversial splitting of African Acacia

spp. into the genera *Senegalia* and *Vachellia* (Smith & Figueiredo, 2011), these latter names appear in our tables and figures, but we refer to "acacias" inclusively.

2.2 | Generation of DNA barcodes

We used standard protocols to bidirectionally sequence five markers commonly used for plant DNA barcoding: *rbcL*, *matK*, *trnL-F*, *trnH-psbA*, and ITS (protocols in Text S1; primers in Table S1) (Fazekas, Kuzmina, Newmaster, & Hollingsworth, 2012; Ivanova, Fazekas, & Hebert, 2008; Kartzinel et al., 2015; Kuzmina et al., 2017). Sequences (*N* = 4,696) from a subset of specimens that we collected between 2012 and 2015 were initially included in a reference library used to support a DNA-metabarcoding study of herbivore diets (Kartzinel et al., 2015). The current data set includes 1,762 new sequences and improvements in the accuracy of the taxonomic determinations that we completed between 2015 and 2018. To build consensus sequences, we trimmed and assembled forward and reverse reads in GENEIOUS R11 (Kearse et al., 2012). The most current version of these DNA barcode sequences is provided as a publicly accessible data set on BOLD.

2.3 | Sequence alignment

To align DNA sequences for genetic-distance, barcode-gap, and phylogenetic analyses, we used the R (R Core Team, 2017) package DECIPHER (Wright, 2016). These plant DNA barcode data sets included substantial sequence-length variation arising both from insertion-deletion polymorphisms and incomplete Sanger sequence reads. We considered including all sequences in these data sets meeting the minimum acceptable sequence-length thresholds set by Genbank (100 bp for coding genes and 200 bp for noncoding genes), but found that confidently establishing significant sequence homology in alignments required us to reduce the number of short sequences in further analyses. We therefore excluded the shortest ~25% of sequences obtained for each marker from further analyses. To evaluate the identities of specimens that are not identified to the species-level (N = 27 out of 460 taxa), representatives of these taxa were included in the alignment used to estimate the community phylogeny. However, to prevent taxonomic uncertainty from influencing genetic-distance and barcode gap analyses, we included only specimens that had been identified to species-level (N = 433 species).

For *rbcL*, we aligned all sequences simultaneously using ALIGNSEQS(). For *matK*, we used ALIGNTRANSLATION() to align the amino acid translation of our DNA sequences and back-translated to DNA. Because distantly related species often have highly divergent sequences for *trnL-F*, *trnH-psbA*, and ITS, we split the sequence data to align by family using the R package PHYLOTOOLS (Zhang, 2017) and aligned using ALIGNSEQS(). For analyses considering data from all markers together, we concatenated all alignments to create a supermatrix of all markers, including markers that we had split into separate alignments by plant family (Kress et al., 2009).

2.4 | Calculation of intra- and interspecific genetic distances

To assess levels of genetic variation within and among plant species in our data set, we calculated uncorrected intra- and interspecific genetic distances for each marker separately (rbcL. matK, trnL-F, trnH-psbA, and ITS) and for all markers together (rbcL + matK +trnL-F + trnH-psbA + ITS) using DISTANCEMATRIX() in DECIPHER (Wright, 2016). To limit the influence of missing data, terminal gaps, gap-to-letter matches, and gap-to-gap matches were not included in the calculation of distances. Pairwise comparisons of genetic distances among all species were conducted for the global alignments of rbcL, matK, and the supermatrix of all markers. By-family alignments for trnL-F, trnH-psbA, and ITS allowed for pairwise comparisons at the family level only. We evaluated the presence of "barcode gaps" based on these genetic distances, which can be identified based on the disparities between the maximum intraspecific and minimum interspecific genetic distances (Hebert et al., 2003).

2.5 | Estimation of community phylogeny

We built an alignment that included only a single specimen per taxon, choosing the specimen represented by the most available sequence data across at least three markers. We tested for the best model of nucleotide substitution and partitioning scheme to use in subsequent phylogenetic reconstruction steps using the program PARTITION FINDER 2 (Lanfear, Frandsen, Wright, Senfeld, & Calcott, 2016). Based on AIC, the best model of nucleotide substitution was GTR + I + Γ which we applied to each of the following partitions: (a) rbcL codon position one, (b) rbcL codon position two, (c) rbcL codon position three, (d) matK codon positions one and two, (e) matK codon position three, (f) trnL-F, (g) trnH-psbA, and (h) ITS. To ensure accurate reconstruction of established phylogenetic relationships, we constrained family-level relationships known a priori using the "R20120829" tree, available from PHYLOMATIC (Webb & Donoghue, 2005) in the R package BRRANCH-ING (Chamberlain, 2016). Using the supermatrix, constraint tree, GTR + I + Γ model of nucleotide substitution, and partitioning scheme determined using PARTITION FINDER 2, we ran a maximumlikelihood analysis to estimate a phylogeny in the program RAXML (Stamatakis, 2014) through the CIPRES Science Gateway (Miller, Pfeiffer, & Schwartz, 2010). To maximize the RAXML phylogeny's utility for this and future studies, we needed to rescale its branch lengths to represent absolute time. We applied PHYLOCOM'S BLADJ() function (Webb, Ackerly, & Kembel, 2008) to the RAXML phylogeny based on 36 fossil calibration dates (Bell, Soltis, & Soltis, 2010; Gastauer & Meira-Neto, 2016). We used the R package MONOPHY (Schwery & O'Meara, 2016) to test the monophyly of genera. For this analysis, we maintained supported nodes and collapsed unsupported nodes (bootstrap support <63) to polytomies (Farris, Albert, Kallersjo, Lipscomb, & Kluge, 1996).

2.6 | Testing for phylogenetic signal in barcode gaps

Phylogenetic signal is the tendency of closely related species to resemble each other because of shared evolutionary history (Felsenstein, 1985). While phylogenetic signal is ubiquitous, estimates of weak or no phylogenetic signal can result from some evolutionary processes (Hansen & Martins, 1996) and properties of data sets such as incomplete taxonomic sampling (Blomberg, Garland, & Ives, 2003; Cavender-Bares, Keen, & Miles, 2006). Thus, the presence of phylogenetic signal in a particular local plant assemblage cannot be assumed. To demonstrate the utility of this phylogeny for studies of phylogenetic community ecology, we tested for phylogenetic signal in the presence or absence of barcode gaps for each marker and for all markers combined using continuous-time Markov models of discrete-trait evolution using the R package GEIGER (Harmon, Weir, Brock, Glor, & Challenger, 2008). Specifically, we compared model fits when the tree-transformation parameter λ (Pagel, 1999) was set to zero (no influence of phylogeny) against those obtained when λ was determined by maximum likelihood (potential influence of phylogeny). Significantly better model fits when the tree transformation parameter λ was determined by maximum likelihood indicate significant phylogenetic signal in barcode gap presence or absence. Significant phylogenetic signal in barcode gap presence or absence in the flora of MRC would allow us to identify clades in which DNAbased species identifications would be straightforward or problematic using these markers.

3 | RESULTS

3.1 | Collection and sequencing success

Of the approximately 500 plant species known or thought to occur at the MRC, to date we have obtained 1,843 specimens from at least 433 species and an additional 27 provisionally distinguished taxa currently resolved to family- or genus-level (*N* = 460 taxa in total; ~92% of the known flora). These specimens belong to two phyla, three classes, 29 orders, 66 families, and 245 genera. Collectively, this barcode library includes data for 1,781 specimens, including 1,523 *rbcL*, 1,197 *matK*, 1,595 *trnL-F*, 1,260 *trnH-psbA*, and 883 ITS sequences. All sequence data have been published on BOLD as "DS-UHURUR2" (dx.doi.org/10.5883/DS-UHURUR2) and on Genbank (accessions in Data set S1).

3.2 | Genetic distances and barcode gaps

The information provided by all markers combined in the supermatrix revealed barcode gaps for 311 of the 429 well-identified species that we assessed (72%). Considering each of the five markers separately, 73%–89% of species exhibited a barcode gap (rbcL = 231 of 316 (73%), matK = 243 of 315 (77%), trnL-F = 257 of 324 (79%), trnH-psbA = 227 of 289 (79%) and ITS = 159 of 178 (89%); Figure 2; Data set S2).

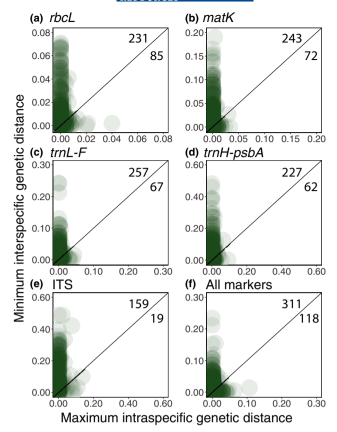


FIGURE 2 Barcode gap plots for species from Mpala Research Centre, showing the relationships between the maximum intraspecific genetic distance (horizontal axes) and minimum interspecific distance (vertical axes) for each DNA-barcode marker and all markers combined. A barcode gap is indicated by points that fall above the solid line. Points have been made translucent such that high densities of points result in darker colours. Numbers on upper and lower halves of plots represent the number of species with (top) or without (bottom) barcode gaps

3.3 | Barcode phylogeny and phylogenetic signal in barcode gaps

Our final phylogeny includes 324 taxa (70%) for which we obtained sufficiently long sequence data (Figure 3). We visualized the results in detail using subtrees for the monocots (Figure S1), superastrids (Figure S2), and superrosids (Figure S3). Of the 186 genera included in the tree, 40 were monophyletic, 20 were not monophyletic, and 126 were monotypic (precluding tests of monophyly). Many of the nonmonophyletic genera were members of the orders Poales, Asterales, Lamiales, and Malvales (Figures S4–S6).

We detected significant phylogenetic signal in the presence or absence of barcode gaps for all markers together (χ^2 = 14.976, df = 1, p < 0.001), and singly for matK (χ^2 = 21.085, df = 1, p < 0.001) and trnH-psbA (χ^2 = 6.746, df = 1, p = 0.009). Phylogenetic signal for trnL-F (χ^2 = 3.122, df = 1, p = 0.077) was marginally significant. We did not detect significant phylogenetic signal for rbcL (χ^2 = 2.357, df = 1, p = 0.125) or ITS (χ^2 = 1.875, df = 1, p = 0.171). Thus, for four of the six markers or marker combinations tested, relatedness was

an important predictor of whether a particular species exhibits a barcode gap in this community. Orders with many species lacking barcode gaps included Poales, Malvales, Lamiales, and Fabales (Table S2; Figures S7–S12). These classifications corresponded to many ecologically important savanna species including grasses (Poaceae), sedges (Cyperaceae), acacias (Fabaceae), and mallows (Malvaceae), and represent many of the same clades that included nonmonophyletic genera in this community.

4 | DISCUSSION

Our DNA barcode data set provides a publicly accessible record of an ongoing and long-term botanical inventory of MRC and the surrounding region. The concordance between morphological species identifications and DNA barcodes indicates that this reference library is generally reliable for species identification (72%–89% marker resolution, depending on the marker used). Furthermore, the robust community phylogeny presented here will enable more detailed analyses of this plant community and local species interactions in an evolutionary context. Over time, we expect to increase our barcode coverage of the MRC flora and intend to publish updated versions of record for the data set.

The development of a comprehensive plant DNA-barcode library for MRC has proven challenging and will require continued collection and taxonomic efforts. Some plant species rarely produce fertile specimens, are only distinguishable by their reproductive structures, or are dioecious, undescribed, or cryptic (Dick & Kress, 2009). Furthermore, even if specimens of all local species can ultimately be identified with complete accuracy, DNA barcoding rarely provides perfect discriminatory ability. Here, consistent with the limitations reported in other plant DNA-barcode studies (Braukmann, Kuzmina, Sills, Zakharov, & Hebert, 2017; Chase et al., 2005; CBOL Plant Working Group et al., 2009; Hollingsworth, Graham, & Little, 2011; Kress & Erickson, 2007; Kress et al., 2009; Kress, Wurdack, Zimmer, Weigt, & Janzen, 2005; Lahaye et al., 2008; Li et al., 2015; Newmaster, Fazekas, Steeves, & Janovec, 2008), this data set achieves a high degree of taxonomic resolution that is nevertheless imperfect.

Analyses of intra- and interspecific genetic distances and the presence or absence of barcode gaps reveal the relative utility of different DNA barcode markers and can help identify future research priorities. Phylogenetic signal in barcode gaps indicates that the efficacy of DNA barcoding for species identification in this plant community can be predicted based on knowledge of the evolutionary relationships among species. Species representing diverse and abundant savanna plants are particularly prone to the absence of a barcode gap, and thus sequencing efforts in the future can be scaled up or down depending on the expected marker resolution for the species groups of interest. Studies involving easily resolved clades might require only a subset of the markers utilized here to achieve perfect discrimination capabilities, while other clades might require more detailed genomic analyses (Li et al., 2015). Indeed, prime

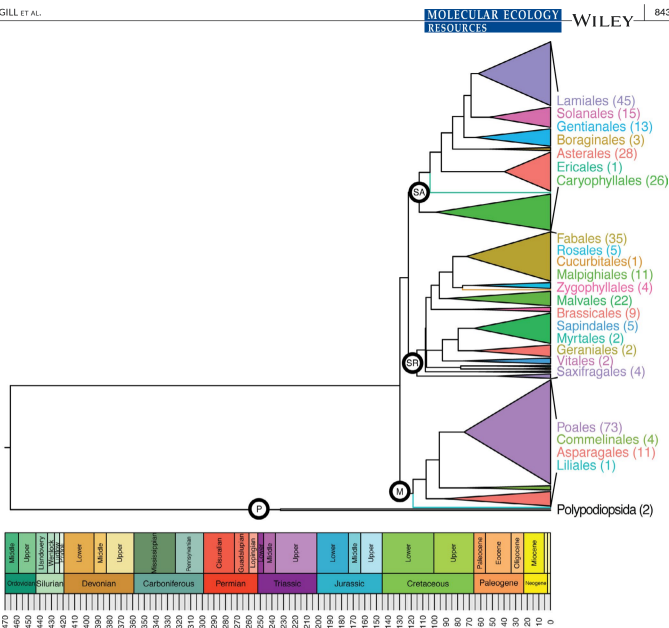


FIGURE 3 Community phylogeny for the plants of Mpala Research Center, Kenya. The size of wedges represents the number of species within orders. Four major lineages are shown, including Polypodiopsida ("P"), monocots ("M"), superrosids ("SR"), and superasterids ("SA"). Detailed subtrees for 89 monocots, 102 superrosids, and 131 superasterids are shown in Figures S1-S3. Numbers after order names indicate the number of species represented

Age (Ma)

candidates for whole-chloroplast sequencing or "genome skimming" (Coissac, Hollingsworth, Lavergne, & Taberlet, 2016) include many grasses (Poaceae), sedges (Cyperaceae), acacias (Fabaceae), and mallows (Malvaceae) for which these standard DNA barcoding protocols appear insufficient.

This DNA barcode release increases the number of plant DNA barcodes from Africa in the BOLD database by 9%, providing a valuable resource for research in the region and filling a recognized gap in the availability of genetic resources for East Africa, and for drylands and savannas and in general (Daru, Berger, et al., 2016). These ecosystems cover more than half of the African continent (Werner, 1991) and traverse a variety of temperature,

rainfall, and soil conditions (House, Archer, Breshears, Scholes, & Tree-Grass Interactions Participants, 2003). The plants that occur within these ecosystems are shaped by herbivory, fire, and land use (Charles-Dominique et al., 2016), and exist at an unstable equilibrium that is prone to phase shifts toward a steady state as either forest or grassland (Staver, Archibald, & Levin, 2011). Worldwide, the study of savanna floras is critical because although savannas vary in vegetation composition and environmental conditions, their characteristic coexistence of trees and grasses provides important habitat for biodiversity and underpins human livelihoods (House et al., 2003; Scholes & Archer, 1997).

We expect that this plant DNA-barcode library will support ongoing and future research by providing a well curated taxonomy for the local flora and improving opportunities to compare and coordinate among the many research programs hosted by MRC. It will further support phytogeographical research across Africa and worldwide through MRC's participation in the global ForestGEO network. Ongoing investigations enabled by the construction of this library include the evaluation of putative new species, community and comparative phylogenetic analyses, and forensic ecological investigations through dietary DNA metabarcoding. This research will contribute to our collective understanding of savanna biodiversity and provide much-needed genetic resources for the region's flora.

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AUTHOR CONTRIBUTIONS

R.M.P. and T.R.K. conceived and designed the research; P.M.M., J.R.G., S.K., and A.A.H. conducted field collections and taxonomic identifications; T.R.K., M.K., and W.J.K. developed the barcoding strategy and performed laboratory analyses; T.R.K. and B.A.G. analyzed the data; B.A.G. drafted the manuscript with subsequent contributions from all authors.

DATA ACCESSIBILITY

R code, alignments, and phylogeny: Dryad doi.org/10.5061/dryad.qk85bp8. Specimen data and DNA barcodes: BOLD dx.doi. org/10.5883/DS-UHURUR2 and Genbank (accessions listed by record in Supplementary Data set S1).

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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