

# Allotetraploid origin and divergence in *Eleusine* (Chloridoideae, Poaceae): evidence from low-copy nuclear gene phylogenies and a plastid gene chronogram

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Received: 26 April 2011 Returned for revision: 27 June 2011 Accepted: 13 July 2011 Published electronically: 31 August 2011

- **Background and Aims** *Eleusine* (Poaceae) is a small genus of the subfamily Chloridoideae exhibiting considerable morphological and ecological diversity in East Africa and the Americas. The interspecific phylogenetic relationships of *Eleusine* are investigated in order to identify its allotetraploid origin, and a chronogram is estimated to infer temporal relationships between palaeoenvironment changes and divergence of *Eleusine* in East Africa.
- **Methods** Two low-copy nuclear (LCN) markers, *Pepc4* and *EF-1 $\alpha$* , were analysed using parsimony, likelihood and Bayesian approaches. A chronogram of *Eleusine* was inferred from a combined data set of six plastid DNA markers (*ndhA* intron, *ndhF*, *rps16-trnK*, *rps16* intron, *rps3*, and *rpl32-trnL*) using the Bayesian dating method.
- **Key Results** The monophyly of *Eleusine* is strongly supported by sequence data from two LCN markers. In the cpDNA phylogeny, three tetraploid species (*E. africana*, *E. coracana* and *E. kigeziensis*) share a common ancestor with the *E. indica*–*E. tristachya* clade, which is considered a source of maternal parents for allotetraploids. Two homoeologous loci are isolated from three tetraploid species in the *Pepc4* phylogeny, and the maternal parents receive further support. The A-type *EF-1 $\alpha$*  sequences possess three characters, i.e. a large number of variations of intron 2; clade E-A distantly diverged from clade E-B and other diploid species; and seven deletions in intron 2, implying a possible derivation through a gene duplication event. The crown age of *Eleusine* and the allotetraploid lineage are 3.89 million years ago (mya) and 1.40 mya, respectively.
- **Conclusions** The molecular data support independent allotetraploid origins for *E. kigeziensis* and the *E. africana*–*E. coracana* clade. Both events may have involved diploids *E. indica* and *E. tristachya* as the maternal parents, but the paternal parents remain unidentified. The habitat-specific hypothesis is proposed to explain the divergence of *Eleusine* and its allotetraploid lineage.

**Key words:** Allotetraploid origin, chloroplast markers, East Africa, *Eleusine*, low-copy nuclear markers, phylogeny, Poaceae.

## INTRODUCTION

Hybridization resulting in the formation of allopolyploids is a frequent mode of plant speciation and evolution (Ellstrand *et al.*, 1996; Seehausen, 2004; Paun *et al.*, 2009), and 21.8 % of grass species (Poaceae) are estimated to have arisen as a result of hybridization events (Knobloch, 1968, 1972). Following advances in molecular phylogeny in recent decades, the evolutionary consequences of merging parental genomes into a polyploid nucleus have attracted much attention (Ge *et al.*, 1999; Petersen *et al.*, 2006; Mason-Gamer *et al.*, 2010). Allopolyploid species present substantial challenges to molecular phylogenetic reconstruction. For example, nuclear gene data are more expensive and time-consuming to obtain than plastid DNA (cpDNA) data, and they are difficult to interpret in allopolyploid taxa where genomes have experienced gene recombination, homogenization or copy loss (Wendel, 2000; Petersen *et al.*, 2006). However, low-copy nuclear (LCN) markers have already been successfully used to elucidate the allopolyploid origins

of some plants. Recently, LCN markers have provided useful information (e.g. Fortune *et al.*, 2007; Jakob and Blattner, 2010). Here we apply phylogenetic analyses based on two LCN markers and plastid data of *Eleusine* Gaertn., a genus of the sub-tribe Eleusininae in the C<sub>4</sub> grass subfamily Chloridoideae (Peterson *et al.*, 2010).

*Eleusine* is a genus of six diploid ( $2n = 16, 18$  and  $20$ ) and three tetraploid taxa ( $2n = 36$  or  $38$ ; Phillips, 1995) exhibiting considerable morphological and ecological diversity in East Africa and the Americas, except that the cytologically unknown *E. semisterilis* S. M. Phillips, known only from the holotype specimen from Kenya (Phillips, 1972), is not included in this study (for the chromosome number and the genome composition see Supplementary Data Table S1, available online). The genus is characterized by having digitate or sub-digitate inflorescences with sub-sessile spikelets arranged along the main axis and 1–16 primary long branches, three-nerved awnless lemmas, and caryopses that have simple verrucate or compound reticulate sculpturing (Liu *et al.*, 2005, 2007; Jiang *et al.*, 2011). The genus is distributed in East

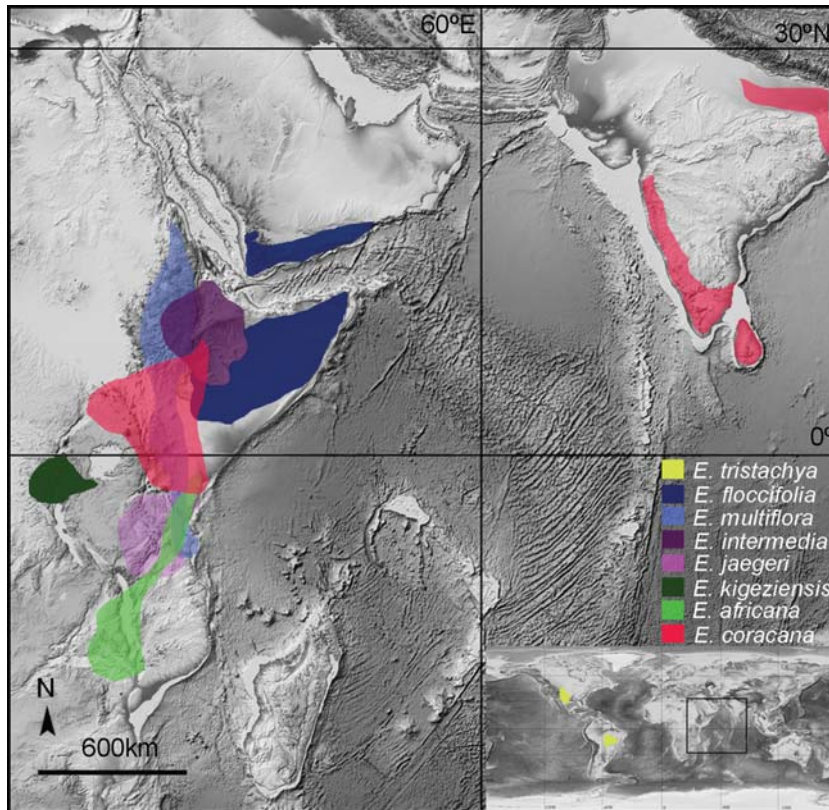


FIG. 1. Natural distribution of *Eleusine* (unmarked *E. indica* widespread on four continents except for Antarctica). The background map was downloaded from <http://www.ngdc.noaa.gov/mgg/global/global.html> (Amante C, Eakins BW. *ETOPO1 1 arc-minute global relief model: procedures, data sources and analysis*. NOAA Technical Memorandum NESDIS NGDC-24, March 2009).

Africa and the Americas, and has been introduced to Asia (Fig. 1). Six species are distributed along the eastern branch rift, *E. kigeziensis* S. M. Phillips is endemic to the western branch rift of East Africa (for definition of two branch rifts see Lærdal and Talbot, 2002), *E. tristachya* (Lam.) Lam. is endemic to the Americas, and *E. indica* (L.) Gaertn. has a cosmopolitan distribution (Phillips, 1995; Liu and Peterson, 2010). Over 2 Mt per annum of *E. coracana* (L.) Gaertn. (finger millet) is produced in Africa, and >4.5 Mt per annum is produced worldwide (National Research Council, 1996). In East Africa, finger millet adapts well to highlands of 1000–2000 m elevation and produces high quality caryopses that are able to withstand frequent drought ((Oduori, 2005). All parts of the finger millet plant provide the raw materials for antiphlogistic medicines used to treat leprosy, liver disease, measles, pleurisy, pneumonia, and smallpox (Duke and Wain, 1981).

The taxonomic difficulty of *Eleusine* is reflected in the controversy of the taxonomic status of *E. africana* Kenn.-O'Byrne *sensu stricto* (*s.s.*), *E. coracana s.s.*, and *E. indica*. Phillips (1974) recognized two species, *E. coracana* and *E. indica*, with 'africana' being considered a subspecies of *E. indica* due to its long, easily shattered spikelets. Phillips (1995) subsequently treated the three taxa as separate species due to the presumed genetic isolation between *E. indica* and two tetraploid species. This also is the opinion of several other authors (Hiremath and Salimath, 1991; Bisht and Mukai,

2002). More recently, Neves *et al.* (2005) recognized two species, *E. coracana* and *E. indica*, and relegated 'africana' as a subspecies of *E. coracana*, owing to the presumed wild progenitor status of 'africana'. Therefore, phylogenetic analyses of these taxa are necessary if we are to understand their status and genetic isolation more clearly.

The allotetraploid origin of finger millet has not been satisfactorily explained. Genomic *in situ* hybridization (GISH) studies identified *E. indica* (AA genome) and *E. floccifolia* (Forssk.) Spreng. (BB genome) as candidate progenitors for two tetraploids, *E. africana* and *E. coracana* (Bisht and Mukai, 2001a, b). However, phylogenetic analyses of nuclear ribosomal DNA (nrDNA) internal transcribed spacer (ITS) and plastid *trnT-trnF* sequences contradicted this hypothesis (Neves *et al.*, 2005). In the ITS locus A clade (see Neves *et al.*, 2005), *E. indica* was confirmed to be closely related to three allotetraploids (*E. africana*, *E. coracana* and *E. kigeziensis*), and this group, in turn, was sister to *E. tristachya*. However, no diploid was detected as the second genome donor for the B locus. The GISH results provided valuable information with respect to the genetic similarity of chromosomes, but were not useful in elucidating interspecific phylogenetic relationships. Bisht and Mukai (2002) reported similar hybridization signals between *E. coracana* and two hybrid pairs, *E. indica*–*E. floccifolia* and *E. tristachya*–*E. floccifolia*, but, so far, the degree of relationship between *E. indica* and *E. tristachya* remains unresolved.

Plastid sequence similarity between *E. indica* and *E. tristachya* suggests that *E. tristachya* became separated from its ancestor very recently (Neves *et al.*, 2005). Therefore, phylogenetic analyses using additional DNA markers are necessary in order to determine the maternal parents.

Accurate estimates of phylogeny and lineage ages are of critical importance in describing the causative reasons for divergence of genera and allotetraploid lineages. In East Africa, the eastern branch rift became uplifted in the early Miocene, the western branch rift subsequently uplifted in the late Miocene, and these processes continued for both branches throughout the middle Pleistocene (Pallister, 1971; Lærdal and Talbot, 2002). It was proposed that the island-like habitats formed by geological lifting movements facilitated the diversification of plants in East Africa (Hedberg, 1969), such as the mountain-dwelling giant senecios along the eastern branch rift (Knob and Palmer, 1995). Palaeoclimatic studies of East Africa have shown that glacial–interglacial oscillations were characterized by cooler and drier climates during the Pliocene–Pleistocene interval (DeMenocal, 1995; Cerling *et al.*, 1997), and such extreme variations in palaeoclimates might explain the observed divergence in allopolyploid lineages (Stebbins, 1980). However, the paucity of information relating to plant divergence times has impeded our understanding of whether generic diversification was catalysed by palaeotectonic movements. More importantly, the temporal relationships that exist between variation in the palaeoclimate and the divergence of allopolyploid lineages requires investigation.

In this study, we sampled nine extant species of *Eleusine* within its native distribution range and employed two LCN markers and six plastid markers to explore their evolutionary history. Our study aims to: (1) investigate interspecific phylogenetic relationships within *Eleusine*; (2) identify the maternal parents and suggest reasons for the equivocal paternal parents of three allotetraploids; and (3) estimate divergence times for *Eleusine* and its tetraploid lineages, and determine their temporal relationships with palaeoclimate variation in East Africa.

## MATERIALS AND METHODS

### *Taxon sampling*

Nineteen accessions were sampled to cover the native distribution range of *Eleusine* in East Africa and the Americas. Four species of *Astrebla* F. Muell., *Coelachyrum* Hochst. & Nees and *Vaseyochloa* Hitchc. were chosen as outgroups, based on a recent phylogeny of the Chloridoideae (Peterson *et al.*, 2010). Seed germination and seedling cultivation followed the method of Peterson and Annable (1991). Voucher specimens were deposited at four herbaria: IBSC, K, RSA-POM and US (see Supplementary Data Table S1).

### *DNA extraction, amplification and sequencing*

Two LCN markers, phosphoenolpyruvate carboxylase 4 (*Pepc4*) and eukaryotic elongation factor 1-alpha (*EF-1α*), were chosen for this study. The *Pepc4* gene product catalyses carboxylation of phosphoenolpyruvate to form oxaloacetate and inorganic phosphate during the C<sub>4</sub> photosynthetic

pathway (Lepiniec *et al.*, 1994), whereas the *EF-1α* gene product plays a crucial role in the channelling and compartmentalization of protein synthesis in the translational apparatus of higher eukaryotic cells (Negrutskii and El'skaya, 1998). Both of these loci are considered good markers for phylogeny reconstruction. *Pepc4* has been successfully used to reconstruct the phylogeny of Poaceae (Christin *et al.*, 2008), and a preliminary screen of both regions indicated no gene homogenization (Q. Liu *et al.*, unpubl. res.). The two LCN markers also appear to represent independent phylogenetic estimates due to their locations on different chromosomes. The *Pepc4* gene is mapped on *Oryza* group 1 homologous chromosomes, whereas the *EF-1α* gene is mapped on *Oryza* group 3 homologous chromosomes (Sasaki *et al.*, 2002; The Rice Chromosome 3 Sequencing Consortium, 2005).

The primer combination *Pepc4*-8F (5'-ACAACCCTGGC TGCCATC-3') and *Pepc4*-10R (5'-GGAAGTTCTTGATGT CCTTGTCG-3') was designed for *Pepc4* based on sequences derived from *Cynodon dactylon* (L.) Pers. and *Tragus racemosus* (L.) All. Primers *EF-1α*-2F (5'-CATTGACTCCACCA CTGGT-3') and *EF-1α*-3R (5'-TAACGGGCCTTGAG TACTT-3') were designed for *EF-1α* based on sequences from *Hordeum vulgare* L., *Oryza sativa* L. and *Saccharum officinarum* L.

All procedures were performed in the Laboratory of Analytical Biology (LAB) at the Smithsonian Institution. Total genomic DNA was extracted from silica-dried leaves using DNeasy Plant Mini Kits (Qiagen). PCR amplifications were performed using 2 μL of genomic DNA (at 10 ng μL<sup>-1</sup>), 0.6 μL of each primer (at 10 pmol), 2.0 μL of dNTPs (at 10 mM), 1 μL of dimethylsulfoxide, 1 μL of MgCl<sub>2</sub> (at 2.5 mM), 1 U of *Taq* polymerase (Bioline), 2.5 μL of 10× Bioline *Taq* polymerase buffer and 0.5 μL of bovine serum albumin (at 10 mg mL<sup>-1</sup>) in a volume of 25 μL under the following conditions: 95 °C/3 min, 35 × (94 °C/30 s, 53 °C/30 s, 72 °C/80 s), 72 °C/10 min, ending with 4 °C holding.

Amplified PCR products were purified using the PEG method (Hiraishi *et al.*, 1995). The cycle sequencing reactions were conducted in 10 μL volumes containing 0.25 μL of BigDye 3.1, 0.5 μL of primers, 2.0 μL of purified PCR products and 1.75 μL of sequencing buffer. The sequencing reactions were run on an ABI PRISM 3730XL DNA Analyzer (Applied Biosystems, Foster City, CA, USA). Direct sequencing revealed two homoeologues for four accessions of the three tetraploids. Therefore, their purified PCR products were cloned into pCR<sup>®</sup>4-TOPO vectors and transformed into *Escherichia coli* TOP10 competent cells in accordance with the protocol for the TOPO TA cloning kit (Invitrogen). Fragments obtained from white clones were amplified using the original PCR primers. Accessions with unique substitutions were repeatedly sequenced to avoid amplification error. GenBank accession numbers are presented in Table S1.

### *Data analyses*

Sequencher 4.5 (Gene Codes Corporation, 2005) was used to evaluate chromatograms for base confirmation and to edit contiguous sequences. Sequences were initially aligned with Muscle (Edgar, 2004), followed by manual adjustments using Se-Al v. 2.0a11 (Rambaut, 2007). Several chimeric



sequences, identified by the repeated sequencing of the same individual and closer inspection of alignments prior to analysis, were excluded from phylogenetic analyses (Cronn *et al.*, 2003).

Maximum parsimony (MP) searches were performed with 1000 random addition sequence replicates, followed by tree bisection–reconnection (TBR) branch swapping, MultTrees option in effect and character state changes weighted equally in PAUP\* v. 4.0b10 (Swofford, 2003). The gaps of the sequence matrix were treated as missing data. The bootstrap percentages for support (PB) of internal nodes were obtained with 1000 replicates. In each replicate, we performed ten random sequence addition replicates followed by the TBR swapping algorithm and saved all trees in each replicate (Felsenstein, 1985). Maximum likelihood (ML) analysis was implemented in GARLI v. 0.951 (Zwickl, 2006), starting from random trees and using 10 000 000 generations per search. The ML bootstrap (LB) support was estimated from 1000 bootstrap replicates in GARLI. The output file containing the best trees for bootstrap reweighted data was then read into PAUP\*, where the majority-rule consensus tree was constructed and bootstrap support values were calculated. For ML analysis, the substitution model was identified under the Akaike Information Criterion (AIC) implemented in Modeltest v. 3.7 (Posada and Crandall, 1998). Bootstrap values of 90–100 were interpreted as strong support, 70–89 as moderate and 50–69 as weak (Mason-Gamer and Kellogg, 1996). Bayesian posterior probabilities (PP) were estimated using MrBayes v. 3.1.2 (Ronquist and Huelsenbeck, 2003). Two independent runs of Bayesian Markov Chain Monte Carlo (MCMC) analyses were conducted simultaneously, each run having one cold chain and three incrementally heated chains, all starting from a random tree and sampling one out of 1000 generations. Tracer v. 1.5 was used to check the chain stationarity (Rambaut and Drummond, 2007). The first 10 % of trees were discarded as burn-in, and the remaining trees were used to calculate the PP. The sequence and tree statistics of two LCN markers are presented in Table 1.

We rooted the *Pepc4* tree using species of *Astrebla* and *Coelachyrum* as outgroups and rooted the *EF-1α* tree using species of *Vaseyochloa* as outgroups because we could not get clean *EF-1α* sequences for *Astrebla* and *Coelachyrum* in the laboratory. The appropriate choice of outgroups was confirmed by phylogenetic proximity (the monophyletic ingroup being supported), genetic proximity (short branch length being observed) and base compositional similarity (ingroup-like GC%, Table 1) (Rota-Stabelli and Telford, 2008).

#### Divergence time estimates

Because divergence time estimates using cpDNA data are able to avoid potential errors caused by heterogeneity of substitution rates or the existence of paralogous copies of nuclear markers for allopolyploid lineages (Renner, 2005), six plastid markers (*ndhA* intron, *ndhF*, *rps16-trnK*, *rps16* intron, *rps3*, and *rpl32-trnL*) were amplified using published primers (Peterson *et al.*, 2010) for eight accessions representing eight species, except for *E. indica* (downloaded from GenBank). The six amplified plastid sequences were then added to an

TABLE 1. Sequence and tree statistics of two LCN markers used in the study

LCN marker	Aligned full sequence length (bp)	GC%	Coding region length (bp)	Proportion of coding region (%)	No. of variable sites	No. of informative characters	Proportion of informative characters (%)	Ti/Tv	Tree length	CI	RI	Model selected by AIC
<i>Pepc4</i>	1047	54.3	793	75.7	271	171	16.3	1.37	350	0.85	0.96	TVM + G
<i>EF-1α</i>	1143 (remainder 954)	37.0	101	10.6	561	534	56.0	0.97	1189	0.75	0.92	HKY + I + G

alignment matrix derived from 268 species (Peterson *et al.*, 2010) and five species of Centropodieae in the Chloridoideae (Peterson *et al.*, 2011) in order to estimate the divergence times of *Eleusine* and its tetraploid lineages.

In order to evaluate the molecular clock assumption for our data, likelihood scores for clock and non-clock models were compared using a likelihood ratio (LR) test (Felsenstein, 1981). The LR test was calculated as  $2 \times (\ln L_{\text{clock}} - \ln L_{\text{nonclock}})$  and assumed to follow a  $\chi^2$  distribution, with the number of degrees of freedom ( $n$ ) equal to the number of terminals minus two. The assumption of rate constancy was rejected for this study because the constrained and unconstrained analyses differed significantly (LR = 1647.2395, d.f. = 279,  $P = 0$ ). A Bayesian method, which allows a relaxed evolutionary model to estimate divergence times, was then employed.

The Bayesian analysis was conducted in BEAST v. 1.5.3 (Drummond and Rambaut, 2007), which employed a Bayesian MCMC to co-estimate topology, substitution rates and node ages. BEAUTi was used to set criteria for the analysis. The AIC estimated by Modeltest 3.7 (Posada and Crandall, 1998) was used to determine which nucleotide substitution model best fit our data. The oldest  $C_4$  lineage in Chloridoideae was dated to be 32.0 million years ago (mya) (Christin *et al.*, 2008; Vicentini *et al.*, 2008), and thus the crown age of the subfamily was set at 32.0 mya since isotopic surveys provide no evidence for an older date of  $C_4$  grass origin so far (Osborne and Beerling, 2006). In addition, an uncorrelated lognormal model of rate variation among branches in the tree was assumed and a Yule prior on the birth rate of new lineages employed (Drummond *et al.*, 2006). Posterior distributions of parameters were calculated by two independent MCMC analyses of 20 000 000 generations with 10 % burn-in. Results were checked in Tracer to ensure that plots of both analyses converged on the same area and then were combined using TreeAnnotator v. 1.5.2 (part of the BEAST package). Final trees were checked and edited in FigTree v. 1.3.1 (Rambaut, 2009), and the divergence times were shown as the mean and the 95 % highest posterior density (HPD) in millions of years.

## RESULTS

### Analysis of *Pepc4* sequences

The matrix of *Pepc4* comprises 1047 characters, including partial exon 8, complete exon 9, partial exon 10, intron 8 and intron 9 at a length of 71, 531, 181, 131 and 133 bp, respectively. Among the 1047 characters, 171 are parsimony-informative (16.3 %) (Table 1). The log likelihood scores of 56 substitution models range from 3520.6231 to 3599.8337, and Modeltest indicates that the best fit model under AIC is TVM + G. The ML tree has the same topology as the MP strict consensus tree and the Bayesian majority-rule consensus tree. The ML tree is presented in Fig. 2, along with bootstrap support values from MP and ML analyses and posterior probabilities from the Bayesian analysis. The monophyly of *Eleusine* was strongly supported by the three support indices (PB = 100 %, LB = 100 %, PP = 1.00).

A single sequence type for the *Pepc4* locus was identified for all diploid species of *Eleusine* (Fig. 1). These sequences formed three monophyletic groups: the *E. indica*–*E. tristachya* clade (PB = 85 %, LB = 94 %, PP = 1.00); the *E. multiflora* clade (PB = 100 %, LB = 100 %, PP = 1.00); and the strongly supported *E. floccifolia*–*E. jaegeri* clade (PB = 100 %, LB = 100 %, PP = 1.00). Two *Pepc4* sequence types (A- and B-) were identified in one accession of *E. africana* and *E. coracana*, and in two accessions of *E. kigeziensis*, consistent with tetraploidy. These sequences fell into three distinct groups. Clade P-A (PB = 100 %, LB = 89 %, PP = 1.00) contains a sub-clade with *E. intermedia* and the A-type sequences of *E. africana* and *E. coracana* (PB = 63 %, LB = 65 %, PP = 1.00), the A-type sequences of *E. kigeziensis* (PB = 65 %, LB = 64 %, PP = 1.00) and the *E. indica*–*E. tristachya* clade (PB = 85 %, LB = 94 %, PP = 1.00). Clade P-B1 contains B-type sequences of *E. africana* and *E. coracana* (PB = 90 %, LB = 100 %, PP = 1.00). Clade P-B2 contains a monophyletic clade and contains B-type sequences of *E. kigeziensis* (PB = 100 %, LB = 100 %, PP = 1.00). Clade P-A is sister to *E. multiflora*, and this group, in turn, is sister to clade P-B1 (PB = 76 %, LB = 85 %, PP = 0.70). Clade P-A + *E. multiflora* + clade P-B1 is sister to the *E. jaegeri*–*E. floccifolia* clade, while clade P-B2 (*E. kigeziensis*) is sister to the remaining species of *Eleusine* (Fig. 2).

### Analysis of *EF-1 $\alpha$* sequences

The matrix of *EF-1 $\alpha$*  comprises 1143 characters, including partial exon 2, partial exon 3 and complete intron 2 at a length of 78, 23 and 1042 bp, respectively. An ambiguous region (positions 235–423) in intron 2 is excluded from phylogenetic analyses. Among the remaining 954 characters, 534 are parsimony-informative (56.0 %) (Table 1). The log likelihood scores of 56 substitution models range from 5882.0400 to 6096.7363, and Modeltest indicates that the best fit model under AIC is HKY + I + G. The ML tree has the same topology as the MP strict consensus tree and the Bayesian majority-rule consensus tree. The ML tree is presented in Fig. 3, along with bootstrap support values from MP and ML analyses and posterior probabilities from Bayesian analysis. The monophyly of *Eleusine* is strongly supported by the three support indices (PB = 100 %, LB = 100 %, PP = 1.00).

A single sequence type for the *EF-1 $\alpha$*  locus was identified for all diploid species of *Eleusine*. These sequences formed three monophyletic groups: the *E. multiflora* and *E. indica*–*E. tristachya* clade (PB = 100 %, LB = 97 %, PP = 1.00); the strongly supported *E. floccifolia*–*E. jaegeri* clade (PB = 90 %, LB = 92 %, PP = 1.00); and the *E. intermedia* clade (PB = 100 %, LB = 100 %, PP = 1.00). The *EF-1 $\alpha$*  sequences of the three allotetraploids diverged from diploid lineages. As with *Pepc4*, two sequence types (A- and B-) were recovered from one accession of *E. africana* and *E. coracana*, and two accessions of *E. kigeziensis*. These sequences fell into two distinct groups (Fig. 3). Clade E-A contains the A-type sequences of *E. coracana* and sister *E. africana* and *E. kigeziensis* (PB = 100 %, LB = 100 %, PP = 1.00), while clade E-B contains the B-type sequences of *E. coracana*, *E. africana* and *E. kigeziensis* (PB = 100 %, LB = 100 %, PP = 1.00).

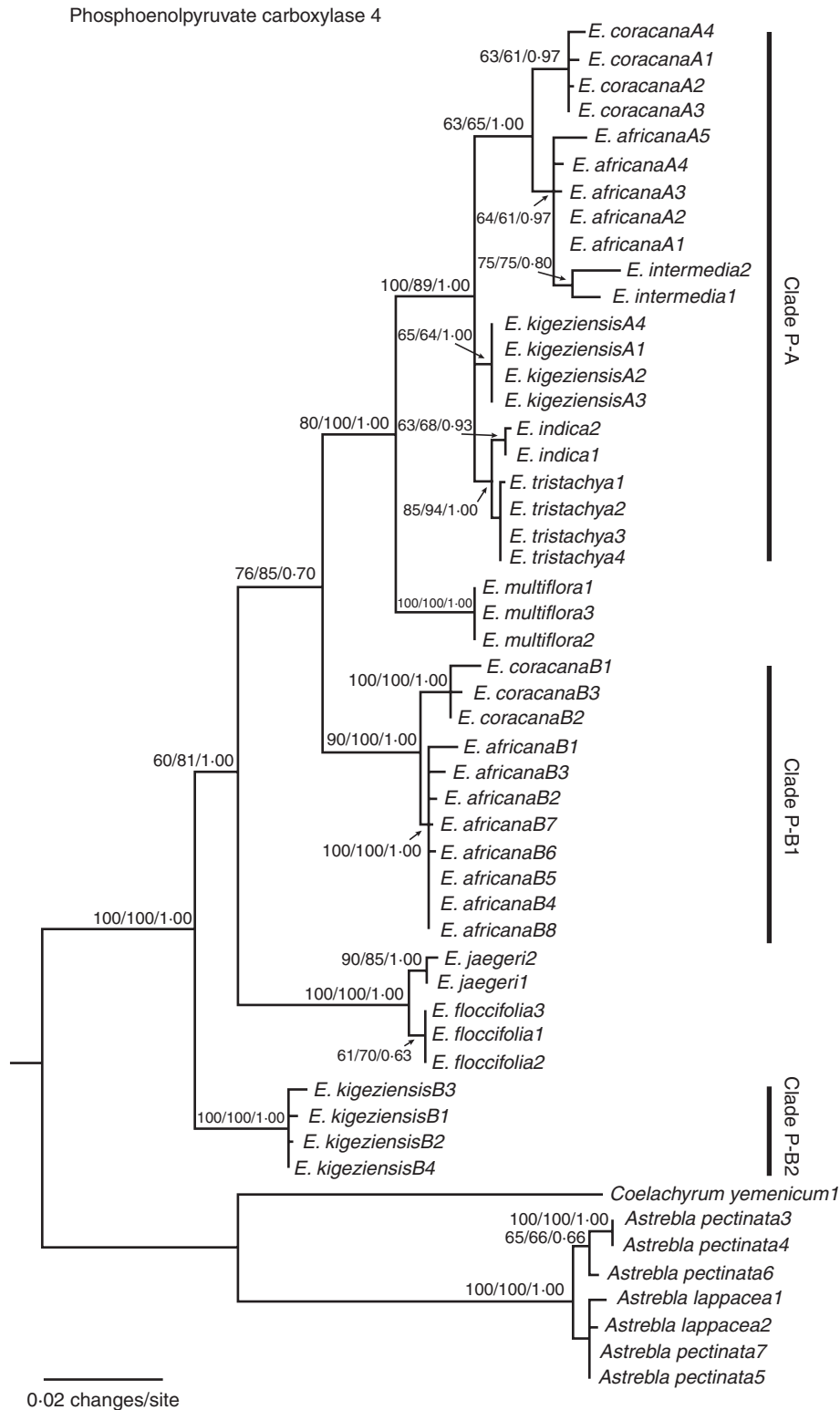


FIG. 2. Maximum likelihood phylogeny of *Eleusine* inferred from the nuclear *Pcp4* data. Numbers above nodes indicate bootstrap values obtained from parsimony and maximum likelihood analyses, and Bayesian posterior probabilities.

LB = 90 %, PP = 1.00). Clade E-A is sister to clade E-B (PB = 86 %, LB = 100 %, PP = 1.00), and together they are sister to a clade containing *E. multiflora* + the *E. indica*–

*E. tristachya* clade + the *E. floccifolia*–*E. jaegeri* clade (PB = 97 %, LB = 100 %, PP = 1.00). *Eleusine intermedia* is sister to all remaining species of *Eleusine*.

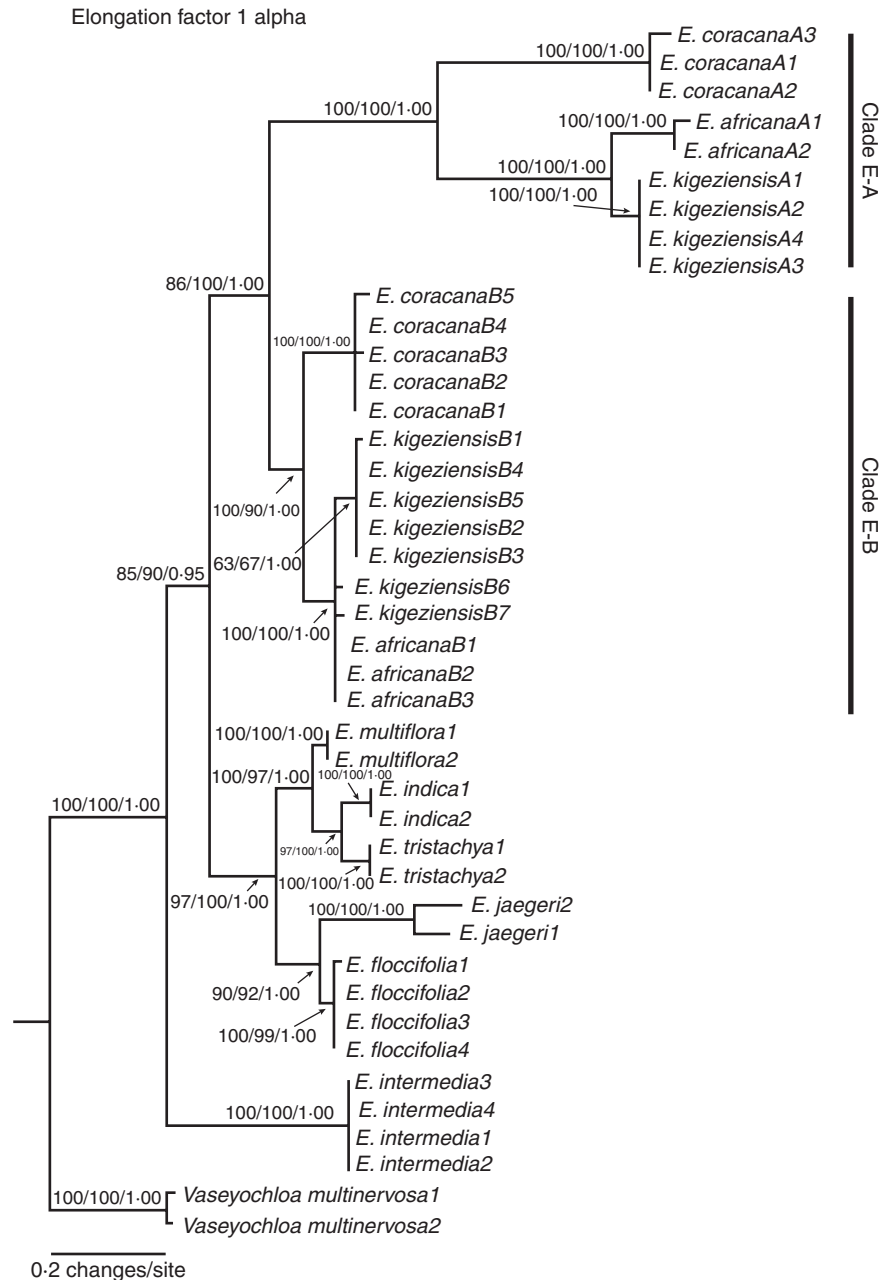


FIG. 3. Maximum likelihood phylogeny of *Eleusine* inferred from the nuclear *EF-1 $\alpha$*  data. Numbers above nodes indicate bootstrap values obtained from parsimony and maximum likelihood analyses, and Bayesian posterior probabilities.

The A-type sequences of the three allopolyploids possess three characters: a large number of variations regularly distributed along intron 2 (e.g. notice the long branch length in Fig. 3); clade E-A distantly diverged from clade E-B and other diploid species (Fig. 3); and seven deletions (6–12 bp in length) in intron 2, implying the likelihood of gene divergence after speciation.

#### Divergence times

A combined matrix of six plastid markers from 281 accessions comprises 6737 characters, of which 2129 are

parsimony-informative (31.6%). Modeltest indicates that the best fit model under AIC is GTR + G. Divergence times for nodes in the phylogeny are shown in Fig. 4.

Using the combination of six plastid DNA markers calibrated with the constrained calibration point of the oldest C<sub>4</sub> lineage in Chloridoideae, the crown age of *Eleusine* was determined to be 3.89 (95% HPD: 2.06–6.20) mya in the Miocene–early Pliocene interval. The crown age of allotetraploid lineage was 1.40 (95% HPD: 0.50–2.71) mya in the middle Pleistocene. The divergence of *E. coracana* was estimated to have occurred 0.67 (95% HPD: 0.15–1.52) mya in the late Pleistocene (Fig. 4). In the BEAST topology, three

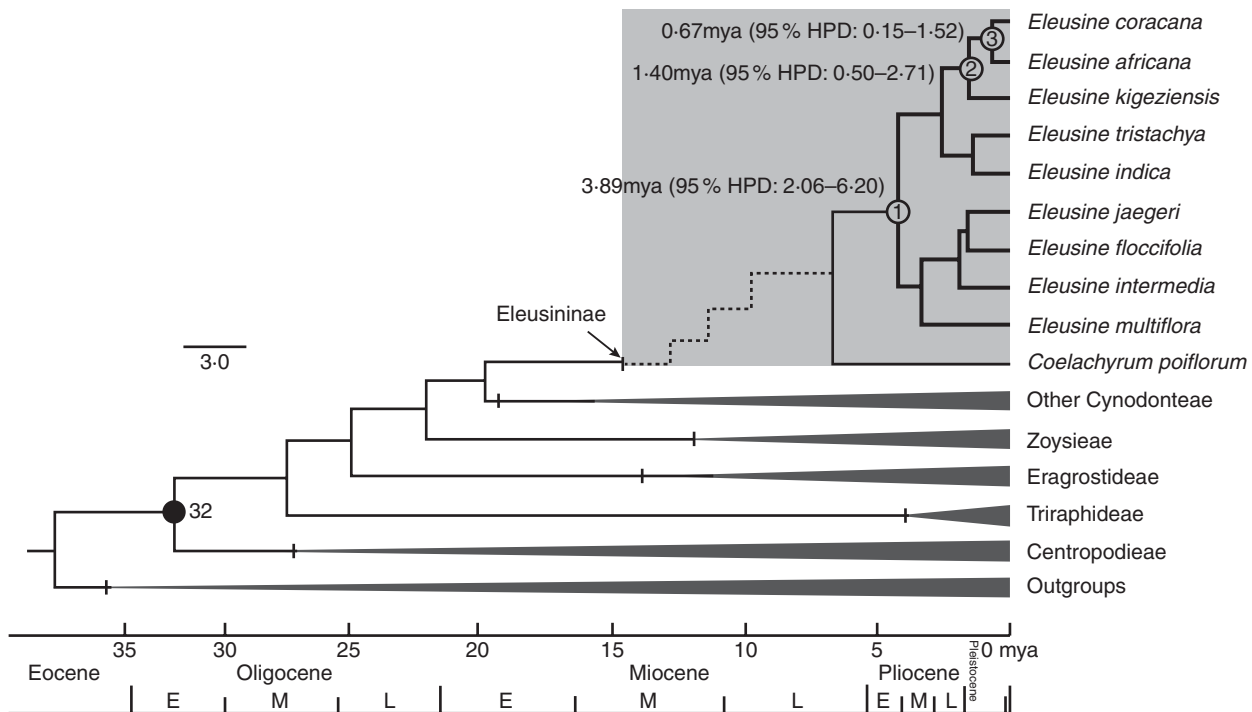


FIG. 4. Bayesian divergence time estimates of *Eleusine* (shaded) based on the six combined plastid gene markers (*ndhA* intron, *ndhF*, *rps16-trnK*, *rps16* intron, *rps3*, and *rpl32-trnL*) of Chloridoideae. Clade constraint is marked by the black circle. An interrupted line indicates the origin of *Eleusine* from sub-tribe Eleusininae. Node numbers 1, 2 and 3 indicate the crown age of *Eleusine*, allotetraploid clade and *E. coracana*, respectively. Numbers in parentheses indicate 95% highest posterior density (HPD) intervals.

tetraploids and the diploid *E. indica*–*E. tristachya* clade comprised a monophyletic lineage, and then the lineage was sister to a clade that included *E. floccifolia*, *E. jaegeri*, *E. intermedia* and *E. multiflora*.

## DISCUSSION

### *Interspecific phylogenetic relationships in Eleusine*

A sister relationship between *E. indica* and *E. tristachya* was strongly supported by *Pepc4*, *EF-1 $\alpha$*  and cpDNA data sets (Figs 2–4). Biochemical and genetic evidence also supports this relationship: the two species have similar karyotypes (Hiremath and Chennaveeraiah, 1982), sub-equal 2C DNA amounts of 2.9 pg (Hiremath and Salimath, 1991), high Rogers' similarity value of isozyme S = 0.794 (Werth *et al.*, 1994) and a shared intergenic spacer region (IGS) phenotype in ribosomal DNA (rDNA) (Hilu and Johnson, 1992). The close relationship between *E. indica* and *E. tristachya* in the chronogram suggests that the arrival of *E. tristachya* in the Americas was recent (Fig. 4). Therefore, the disjunctive distribution of *Eleusine* is best explained in terms of long-distance dispersal. The hypothesis that the American *E. tristachya* had separated from its African ancestor in ancient times is not supported by our data since it appears to be of recent origin (Hilu and Johnson, 1997).

A sister relationship between *E. floccifolia* and *E. jaegeri* also received strong support from our data. Similarly, biochemical and genetic characters have been useful for

interpreting phylogenetic relationships between the two species, including the possession of isozyme gene *Idh-2* duplicated loci (Werth *et al.*, 1993), a sub-equal 2C DNA amount of 3.3 pg (Hiremath and Salimath, 1991) and highly variable IGS phenotypes in rDNA (Hilu and Johnson, 1992). Apparent morphological differences between the two species could be due to elevation-related phenotypic plasticity. *Eleusine floccifolia* is unique in forming less robust tufts with slender culms and leaf blades with scattered tufts of soft white hairs; the species is distributed in seasonal climate highlands at moderate elevations (about 1800 m) in Ethiopia, Kenya, Somalia and Yemen (Phillips, 1972; Lovett, 1993). *Eleusine jaegeri* is easily recognized since it forms a very robust tussock with fasciculate-branched culms and has glabrous leaf blades (Phillips, 1972, 1995); the species is restricted to extremely arid regions at moderate to high elevations (1800–3200 m) in Tanzania and Kenya (Fig. 1). The phylogenetic lineage thus exemplifies the modern synthetic evolution theory: change is based on a combination of chance and natural selection that affects phenotype and genetic characters unequally (Stebbins, 1986).

The taxonomic separation of *E. africana* s.s., *E. coracana* s.s. and *E. indica* is supported by our study. *Eleusine indica* was reported to be a diploid species (Moinuddin *et al.*, 1994), and only one type of *Pepc4* and *EF-1 $\alpha$*  sequence was detected from the sampled accessions. Thus, it appears that *E. indica* is genetically distant from the two allotetraploids (Chennaveeraiah and Hiremath, 1974). Two types of *Pepc4* and *EF-1 $\alpha$*  sequences were identified from both *E. africana*



and *E. coracana* accessions. In the *Pepc4* phylogeny, A-type sequences of *E. coracana* grouped with A-type sequences of *E. africana* and *E. kigeziensis*, *E. intermedia*, *E. indica* and *E. tristachya* (i.e. clade P-A), while B-type sequences of *E. coracana* grouped with B-type sequences of *E. africana* in clade P-B1 (Fig. 2). In the *EF-1 $\alpha$*  phylogeny, both A- and B-type sequences of *E. coracana* grouped with the A- and B-type sequences of *E. africana* and *E. kigeziensis*, respectively (Fig. 3). The relationships among allotetraploids were incongruent between two nuclear gene trees. A possible interpretation of the conflicting pattern was that a hybridization event was followed by speciation at the polyploid level, producing two species of the allopolyploid clade (*E. africana* and *E. coracana*). Furthermore, the recurrent gene flow might have occurred between *E. kigeziensis* and the lineage of *E. africana* and *E. coracana* (Kellogg et al., 1996; Emshwiller and Doyle, 1998). Contrasting phylogenetic signals in the IGS region (rDNA) occur between *E. coracana* and *E. africana*, and it is evident that finger millet is domesticated from a limited number of wild populations due to a less variable IGS region (Hilu and Johnson, 1992). This finding indicates that finger millet had a more narrow genetic base than *E. africana*, while the unique simple verrucate sculpturing of finger millet caryopses highlighted a rapid process aided by domestication (Paterson et al., 2004; Jiang et al., 2011). Therefore, recognition of these three taxa at the species level as opposed to placing *E. africana* as a subspecies either of *E. indica* or of *E. coracana* is compatible with our results (Phillips, 1995).

#### Allotetraploid origin

As inferred from the results of two sets of molecular markers (plastid and *Pepc4*), three tetraploids are of allopolyploid origin. In the plastid phylogeny, all three allotetraploids share a common ancestor with the *E. indica*–*E. tristachya* clade, which represents a source for the maternal parents (Fig. 4). In the *Pepc4* phylogeny, two homoeologous loci were isolated from each of the allotetraploids, providing strong evidence for the presence of two divergent genomes in each allotetraploid. The maternal lineage identified by plastid phylogeny is also confirmed by the A homoeologues of three allotetraploids in Fig. 2 (clade P-A). However, the *EF-1 $\alpha$*  tree shows that the collected gene copies may be derived through a gene duplication event, but the sets of gene copies being derived from the allopolyploidization event (suggested by the *Pepc4* tree) may be degraded or not collected at all. With the exception of *E. intermedia*, the diploid species formed a single lineage separating from the remaining species in the *EF-1 $\alpha$*  phylogeny, thus providing indirect evidence that hardly supports the species as a potential source for the maternal parents.

The *Pepc4* data support independent allotetraploid origins for *E. kigeziensis* and the *E. africana* and *E. coracana* clade. Based on cpDNA evidence, both events may have involved species from the *E. indica*–*E. tristachya* clade as the maternal parents, but the paternal parents involved in the original hybridization events remain unknown. We propose three hypotheses to explain why the paternal parents remain unidentified: (1) the paternal parents of *E. kigeziensis* may come from

outside of *Eleusine*, and thus they remain unidentified due to restricted sampling at the intergeneric level; (2) genomes derived from paternal parents may have undergone a rapid evolutionary divergence (or degradation) after allotetraploid speciation and are therefore untraceable; or (3) the paternal parents are extinct in the wild.

For the *E. africana* and *E. coracana* clade, the *Pepc4* and nrDNA ITS data supported its location nested within *Eleusine*, suggesting that their paternal parents should be members of the genus, but the earliest diverging clade P-B2 in *Pepc4* phylogeny indicated the possibility of B homoeologues of *E. kigeziensis* derived from intergeneric hybridization (hypothesis 1). It would be worthwhile to test the hypothesis with targeted sampling among allied genera (Peterson et al., 2010). The second proposal provides evolutionary scenarios to reconcile conflicting patterns between two LCN trees. The clade P-B1 is sister to clade P-A + *E. multiflora* in the *Pepc4* tree; the similar branching pattern is shared by the *Waxy* tree (Q. Liu et al., unpubl. res.) for the lineage of *E. africana* and *E. coracana*, and so its paternal parents must be members of the genus. However, the long branch of clade E-B in the *EF-1 $\alpha$*  tree implies its derivation from a gene duplication event. The paternal gene copies derived from the allopolyploidization event (suggested by the *Pepc4* tree) are diverged so greatly that they are too rare to find even after extensive cloning of the *EF-1 $\alpha$*  gene (Wendel, 2000; Petersen et al., 2006). In contrast to *EF-1 $\alpha$*  phylogeny, sequence variation of clade P-B2 reflects complex evolutionary scenarios for the genome origin following the allopolyploidization. Furthermore, the third hypothesis is probably reasonable because the B homoeologues of *E. kigeziensis*, *E. africana* and *E. coracana* do not correspond to any diploid taxon sampled for chloroplast, LCN or a previous nrDNA ITS phylogeny (Neves et al., 2005). Since the genus has been thoroughly studied throughout its natural range in East Africa and the Americas, the only unsampled *E. semisterilis* in this study seems an unlikely candidate for paternal parents due to its unusual morphology of laxly arranged spikelets that disarticulate as a unit (vs. between the florets for all other species in the genus; Phillips, 1972). The natural distribution of *E. coracana* also excludes the possibility of a genetic contribution by extra-African diploid taxa. Therefore, the extinction hypothesis is reasonable to explain the intriguing parental origins for the two hybridization events.

#### Palaeoenvironmental hypothesis for divergence of genus and allotetraploid lineage

The habitat-specific hypothesis might apply to the current species distribution pattern within *Eleusine*. This premise could account for the divergence of species within *Eleusine* since phenotypic traits, such as racemes terminating in a fertile spikelet and hypervariable sculpturing of the caryopsis surface, are apparently adaptations to seasonal rainfall in East Africa (Stebbins, 1986; Liu and Peterson, 2010; Jiang et al., 2011). East African summer rainfall has been highly variable due to rain shadow effects, which were shaped by the topographic barrier of the eastern branch rift to moist, maritime air from the Indian Ocean (Sepulchre et al., 2006). The Kenyan and Ethiopian highlands capture moisture, and

summer monsoonal run-off favours the rise and prosperity of  $C_4$  plants (e.g. *Eleusine*) in habitat-specific mountain savannas. The same premise (habitat-specific hypothesis) might account for the divergence of the allotetraploid lineages (e.g. *E. africana*, *E. coracana* and *E. kigeziensis*). The geological records support the claim that East African climates had changed from warmer and wetter at the Miocene–early Pliocene interval to cooler and drier during the glacial–interglacial oscillations of the Pliocene–Pleistocene interval (DeMenocal, 1995; Cerling *et al.*, 1997). The divergence time of the allotetraploid lineage at 1.40 (95 % HPD: 0.50–2.70) mya falls into the interval of 1.6–1.8 mya, when increasing amplitudes of palaeoclimate variability induced plant speciation, adaptation and evolution (Vicentini *et al.*, 2008; Edwards *et al.*, 2010). Widespread interspecific gene exchange might have occurred during hybridization during the allopolyploidization process (Wakeley and Hey, 1997; Emshwiller and Doyle, 1998). The periodic increases in drier conditions in East Africa during the periods of 1.6–1.8 and 0.8–1.2 mya might have established discrete opportunities for ecological fragmentation and genetic isolation, leading to rapid diversification of allotetraploids adapted to mountain savannas at elevations <3200 m (DeMenocal, 1995).

### Conclusions

Our molecular results based on plastid and LCN markers support the premise that *E. africana*, *E. coracana* and *E. kigeziensis* (tetraploid species) are of allopolyploid origin. Our results support two separate allopolyploidization origins for *E. kigeziensis* and for the *E. africana*–*E. coracana* clade. Both events may have involved the diploid *E. indica*–*E. tristachya* clade as the maternal parent, but the paternal parents are unknown. The most likely explanation for the difficulty in determining the paternal parents of three tetraploids is that they no longer exist. Our study has identified two promising sister lineages for further study, i.e. the *E. indica*–*E. tristachya* clade and the *E. floccifolia*–*E. jaegeri* clade. The habitat-specific hypothesis proposes that the divergences of the allotetraploid lineage within *Eleusine* in East Africa may be associated with habitat-specific mountain savannas formed by topographic uplift and increasing amplitudes of palaeoclimate variability at crucial geological periods. In the future, it may be informative to sample extra-generic relationships, and this can be achieved with a broader sampling of chloridoid genera.

### SUPPLEMENTARY DATA

Supplementary data are available online at [www.aob.oxfordjournals.org](http://www.aob.oxfordjournals.org) and consist of Table S1: samples, vouchers and GenBank accessions of *Eleusine* and outgroups for phylogeny reconstruction and dating analysis.

### ACKNOWLEDGEMENTS

We thank USDA-Beltsville National Germplasm System, ILRI-Addis Ababa and J. Travis Columbus for help with carpology and leaf collection; Lee Weigt, Jeffery Hunt, Rong Li and Xinwei Xu for help in the laboratory; Robert J. Soreng,

Dianxiang Zhang, Gabriel Johnson and Tiejiao Tu for helpful discussions; and six anonymous reviewers for their constructive comments that improved the manuscript. This work was supported by the National Geographic Society Committee for Research and Exploration (grant no. 8087-06), the Smithsonian Institution's Small Grants Program, National Natural Science Foundation of China (no. 30700043), the China Scholarship Council Awards (no. 2008491004), the Key Laboratory of Plant Resources Conservation and Sustainable Utilization of Chinese Academy of Sciences (no. 200922) and the Knowledge Innovation Program of the Chinese Academy of Sciences (no. KSCX2-EW-J-28).

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