

# Phylogenetic analysis of the genus *Sorghum* based on combined sequence data from cpDNA regions and ITS generate well-supported trees with two major lineages

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- Background and Aims Wild Sorghum species provide novel traits for both biotic and abiotic stress resistance and yield for the improvement of cultivated sorghum. A better understanding of the phylogeny in the genus Sorghum will enhance use of the valuable agronomic traits found in wild sorghum.
- *Methods* Four regions of chloroplast DNA (cpDNA; *psbZ-trnG*, *trnY-trnD*, *trnY-psbM* and *trnT-trnL*) and the internal transcribed spacer (ITS) of nuclear ribosomal DNA were used to analyse the phylogeny of sorghum based on maximum-parsimony analyses.
- Key Results Parsimony analyses of the ITS and cpDNA regions as separate or combined sequence datasets formed trees with strong bootstrap support with two lineages: the Eu-sorghum species S. laxiflorum and S. macrospermum in one and Stiposorghum and Para-sorghum in the other. Within Eu-sorghum, S. bicolor-3, -11 and -14 originating from southern Africa form a distinct clade. S. bicolor-2, originally from Yemen, is distantly related to other S. bicolor accessions.
- Conclusions Eu-sorghum species are more closely related to S. macrospermum and S. laxiflorum than to any other Australian wild Sorghum species. S. macrospermum and S. laxiflorum are so closely related that it is inappropriate to classify them in separate sections. S. almum is closely associated with S. bicolor, suggesting that the latter is the maternal parent of the former given that cpDNA is maternally inherited in angiosperms. S. bicolor-3, -11 and -14, from southern Africa, are closely related to each other but distantly related to S. bicolor-2.

Key words: Molecular phylogeny, Sorghum, Eu-sorghum, Zea mays, non-coding regions, cpDNA, ITS.

### INTRODUCTION

Sorghum Moench is highly heterogeneous and with Cleistachne Bentham form Sorghastrae (Garber, 1950), one of the 16 subtribes belonging to tribe Andropogoneae. Species of the genus Sorghum have chromosome numbers of 2n = 10, 20, 30 or 40 (Garber, 1950; Lazarides et al., 1991).There are five recognized sections and 25 species within Sorghum. The sections are Eu-sorghum, Chaetosorghum, Heterosorghum, Para-sorghum and Stiposorghum (Garber, 1950; Lazarides et al., 1991). Eu-sorghum includes cultivated sorghums and their closest wild relatives (De Wet and Huckay, 1967). According to De Wet (1978) three species were recognized in section Eu-sorghum, including two perennial species, S. halepense and S. propinguum, and an annual, S. bicolor. However, in the earlier classification by Snowden (1935), Eu-sorghum is considered to comprise two subsections, Arundinacea and Halepensia. The subsection Arundinacea, commonly found in tropical Africa and India, consists of S. bicolor (L.) Moench, S. arundinaceum (Desv.) Stapf and S. drummondii (Steud.) Millsp. S. propinguum (Kunth) Hitchcock, S. halepense (L.) Pers and S. almum Parodi form subsection Halepensia, and are found in the Mediterranean region and Southeast Asia.

The wild Australian *Sorghum* species constitute over two-thirds of the recognized *Sorghum* species, of which one

species each belongs to *Chaetosorghum* and *Heterosorghum*. The section *Para-sorghum* comprises seven species. Of these, five are native to northern monsoonal Australia, Africa and Asia (Garber, 1950; Lazarides *et al.*, 1991). *Stiposorghum* consists of ten species that are endemic to northern Australia (Garber, 1950; Lazarides *et al.*, 1991). The wild and weedy *Sorghum* species present a valuable source of agronomic traits such as pest and disease resistance (Sharma and Franzmann, 2001; Kamala *et al.*, 2002; Komolong *et al.*, 2002) for introgression into *S. bicolor*. Exploitation of these valuable traits requires a thorough understanding of the phylogenetic relationships between cultivated sorghum and the wild sorghum gene pool.

The chloroplast genome is useful in providing information on the inference of the evolutionary patterns and processes in plants (Raubeson and Jansen, 2005). The genome has, either solely or combined with other genomes, been widely used for inferring phylogenetic relationships of different taxa, including *Hordeum*, *Triticum* and *Aegilops* (Gielly and Taberlet, 1994), *Guizotia* (Geleta, 2007), Solanaceae (Melotto-Passarin *et al.*, 2008) and *Sorghum* (Dillon *et al.*, 2007). The non-coding chloroplast regions are phylogenetically more informative than the coding regions at lower taxonomic levels because they are under less functional constraints and evolve rapidly (Gielly and Taberlet, 1994). One of the

chloroplast DNA (cpDNA) regions, *trn*T-*trn*L, used in this study was reported to possess sufficient phylogenetic signal for studies at lower taxonomic levels (Shaw *et al.*, 2005).

The internal transcribed spacer (ITS) region of the 18S–5·8S–26S nuclear ribosomal DNA (nrDNA) has been commonly used for phylogenetic inference at the generic and infrageneric level in plants. The ITS loci properties of biparental inheritance, universality of primers, intragenomic uniformity and intergenomic variability merit their utility for phylogenetic reconstruction (Baldwin *et al.*, 1995). Two ITS regions, ITS1 and ITS2, generally evolve more rapidly than coding regions and have shown to be equally informative, being able to differentiate between closely related species (Baldwin, 1992) and more specifically to resolve phylogenetic relationships of sorghum and related species (Sun *et al.*, 1994; Dillon *et al.*, 2001; Guo *et al.*, 2006).

This study sought to resolve the phylogenetic relationships between species of the genus *Sorghum* based on four regions of the cpDNA: *trnY-trnD*, *psbZ-trnG*, *trnY-psbM* and *trnT-trnL* and the ITS of nrDNA and also to evaluate the usefulness of the five non-coding regions of cpDNA in resolving relationships among the closely related species within section *Eu-sorghum*.

### MATERIALS AND METHODS

### Plant material

Details of twenty-two *Sorghum* species along with GenBank germplasm and GenBank sequence accession numbers used in this study are given in Table 1. The germplasm accessions included wild sorghum and several cultivated sorghum obtained from the Australian Tropical Crops Genetic Resource Centre, Biloela, Queensland, Australia. In addition, five accessions of *S. bicolor* and one of *S. arundinaceum* were obtained from the Zambian National Plant Genetic Resources Centre.

## DNA extraction, PCR and sequencing

Each *Sorghum* species was represented by 1–2 accessions, except for *S. bicolor* for which 11 accessions were used. Genomic DNA was extracted from fresh leaf tissues of seedlings raised in the greenhouse at approx. 2 weeks of age using a modified CTAB extraction method (Doyle and Doyle, 1987). The quality of the DNA was analysed by agarose gel electrophoresis and DNA concentration was determined using a Nanodrop® ND-1000 spectrophotometer (Saveen Werner, Malmö, Sweden).

Primers for amplification and sequencing of the *trnS-trnfM*, *trnY-psbM* and *trnT-trnD* regions were designed for this study while the *trnT-trnL* region was amplified and sequenced using the universal primers designed by Taberlet *et al.* (1991). A primer pair was used for each of the cpDNA regions. However, two primer pairs were designed for amplification of the *trnY-psbM* region. Universal primers ITS4 and ITS5 (White *et al.*, 1990) were used for amplification and sequencing of the ITS region.

The sequences of the primers and information on specific primers supplied by Eurofins MWG GmbH (Ebersberg,

Germany) used in this study are given in Table 2. A GeneAMP PCR system 9700 thermocycler was used for amplification with the following temperature regime: denaturation at 94 °C, followed by 30 cycles of 1 min denaturing at 94 °C. 1 min primer annealing at 51 °C and primer extension for 2 min at 72 °C, and a final 7-min extension at 72 °C. Successfully amplified samples were purified using the QIAquick PCR purification kit (Qiagen GmbH, Hilden, Germany) and microcentrifuge according to the manufacturer's instructions. Nine microlitres of purified PCR products was mixed with 1 µL of sequencing primers and sent to the sequencing facility in the University of Oslo, Norway (http:// www.bio.uio.no/ABI-lab/), where DNA sequencing was done. The quality of the sequences was evaluated using Sequence Scanner version 1.0 (Applied Biosystems, www. appliedbiosystems.com/) and only high-quality sequences were used for the analysis. All regions were sequenced using both forward and reverse primers. The sequences from the forward and reverse primers were aligned for each sample in order to generate a consensus sequence. As the sequences were of high quality, the forward and reverse sequences are identical, except in a few cases. These few discrepancies were resolved by repeated PCR and sequencing.

Sequence alignment and data analyses

The quality of the sequences was visually inspected using Sequence Scanner version 1.0 (Applied Biosystems). Multiple sequence alignment was performed using ClustalX version 2.1.10 (Larkin *et al.*, 2007). The sequences were edited using BioEdit version 7.0.9 (Hall, 1999) and PAUP\* 4.0 Beta 10 was used for phylogenetic analyses. The phylogenetic analyses were approached in three ways. In the first approach, the ITS sequences of the nrDNA were analysed separately. In the second approach, the sequences of the four noncoding regions of the cpDNA were also analysed separately. In the final approach, a combined analysis of the cpDNA regions and the ITS was carried out. In all the cases indel positions were treated as missing data. *Zea mays* (GenBank accession no. U04796) was used as an out-group species.

### **RESULTS**

Sequence characteristics of the Sorghum species

The sequence characteristics and parsimony-based tree statistics of four non-coding regions of cpDNA and the ITS are summarized in Table 3. The aligned sequences derived from all the cpDNA regions and the ITS revealed differences in sequence length between the *Sorghum* species. The longest sequences were obtained from the *trnY-psbM* spacer and ranged from 1028 nt (*S. drummondii*) to 1053 nt (*S. exstans*). The eight *S. bicolor* sequences from this spacer exhibited 2–3 nt differences between them. By contrast, the *psbZ-trnG* spacer provided the shortest sequences, which ranged between 286 nt (*Eu-sorghum* species) and 291 nt (*S. intrans*). The similarity in sequence length between the *Eu-sorghum* species could be attributed to the occurrence of 5-nt indels within the *psbZ-trnG* intergenic spacer. Indels of similar size at corresponding positions were also observed in *S. laxiflorum* and *S. macrospermum*.

Table 1. Accession identity and geographical origin of each accession of Sorghum species used in the study

		, , , , , ,					
Species	Section	Germplasm accession no.*	DNA sequence accession no.				
			trnY-trnD	psbZ-trnG	trnY-psbM	trnT-trnL	ITS
S. almum	Eu-sorghum	AusTRCF302386 <sup>A</sup>	GQ121828	GQ121769	GQ121810	GQ121791	GQ121750
S. amplum-1	Stiposorghum	AusTRCF302455 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. amplum-2	Stiposorghum	AusTRCF302623 <sup>A</sup>	GQ121822	GQ121755	GQ121799	GQ121783	GQ121727
S. angustum-1	Stiposorghum	AusTRCF302588 <sup>A</sup>	GQ121824	N/A	GQ121793	GQ121775	GQ121737
S. angustum-2	Stiposorghum	AusTRCF302606 <sup>A</sup>	N/A	GQ121761	N/A	N/A	N/A
S. arundinaceum	Eu-sorghum	ZMB 7203 <sup>Zm</sup>	GQ121832	GQ121766	GQ121806	GQ121790	GQ121746
S. bicolor-1	Eu-sorghum	AusTRCF304111 <sup>TA</sup>	N/A	N/A	N/A	N/A	N/A
S. bicolor-2	Eu-sorghum	AusTRCF304113 <sup>YA</sup>	N/A	N/A	N/A	N/A	GQ121748
S. bicolor-3	Eu-sorghum	AusTRCF304114 <sup>ZwA</sup>	N/A	N/A	N/A	N/A	N/A
S. bicolor-4	Eu-sorghum	AusTRCF304115 <sup>BA</sup>	N/A	N/A	N/A	N/A	GQ121745
S. bicolor-5	Eu-sorghum	AusTRCF312813 <sup>ZmA</sup>	N/A	N/A	N/A	N/A	N/A
S. bicolor-14	Eu-sorghum	ZMB 5395 <sup>Zm</sup>	N/A	N/A	N/A	N/A	N/A
S. bicolor-12	Eu-sorghum	ZMB 5757 <sup>Zm</sup>	GQ121829	GQ121770	GQ121813	GQ121792	GQ121743
S. bicolor-15	Eu-sorghum	ZMB 6665 <sup>Zm</sup>	N/A	N/A	N/A	N/A	N/A
S. bicolor-10	Eu-sorghum	ZMB 7016 <sup>Zm</sup>	N/A	N/A	N/A	N/A	GQ121744
S. bicolor-11	Eu-sorghum	ZMB 7034 <sup>Zm</sup>	N/A	N/A	N/A	N/A	N/A
S. bicolor-13	Eu-sorghum	ZMB 7112 <sup>Zm</sup>	N/A	N/A	N/A	N/A	N/A
S. brachypodum-1	Stiposorghum	AusTRCF302480 <sup>A</sup>	GQ121818	GQ121756	GQ121802	GQ121774	GQ121736
S. brachypodum-2	Stiposorghum	AusTRCF302481 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. bulbosum-1	Stiposorghum	AusTRCF302418 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. bulbosum-2	Stiposorghum	AusTRCF302646 <sup>A</sup>	GQ121823	GQ121758	QG121803	GQ121781	GQ121732
S. drummondii-1	Eu-sorghum	AusTRCF300263 <sup>EA</sup>	N/A	N/A	N/A	N/A	N/A
S. drummondii-2	Eu-sorghum	AusTRCF300264 <sup>KA</sup>	GQ121831	GQ121765	GQ121809	GQ121789	GQ121747
S. ecarinatum-1	Stiposorghum	AusTRCF302450 <sup>A</sup>	GQ121831 GQ121821	GQ121754	GQ121800	GQ121784	GQ121747
S. ecarinatum-1	Stiposorghum	AusTRCF302662 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. exstans-1	Stiposorghum	AusTRCF302401 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. exstans-2	Stiposorghum	AusTRCF302473 <sup>A</sup>	GQ121816	GQ121759	GQ121796	GQ121782	GQ121735
S. halepense-1	Eu-sorghum	AusTRCF302473 AusTRCF300167 <sup>A</sup>	GQ121810 GQ121830	GQ121768	GQ121790 GQ121808	GQ121788	N/A
S. halepense-2	Eu-sorghum Eu-sorghum	AusTRCF300107 AusTRCF300188 <sup>A</sup>	N/A	N/A	N/A	N/A	GQ121749
S. interjectum-1	Stiposorghum	AusTRCF300188 AusTRCF302396 <sup>A</sup>	GO121817	GO121753	GO121797	GO121772	GQ121749 GQ121738
S. interjectum-1 S. interjectum-2	Stiposorghum	AusTRCF302390 AusTRCF302433 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. intrans	Stiposorghum	AusTRCF302453 AusTRCF302390 <sup>A</sup>	GQ121825	GQ121752	GQ121795	GQ121780	GQ121733
S. laxiflorum-1	Heterosorghum	AusTRCF302503 <sup>A</sup>	GQ121823 GQ121833	GQ121771	GQ121793 GQ1218011	GQ121786 GQ121786	GQ121733 GQ121741
S. laxiflorum-1	Heterosorghum	AusTRCF302503 AusTRCF302607 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. leiocladum-1	Para-sorghum	AusTRCF302007 AusTRCF300148 <sup>A</sup>	GQ121814	N/A N/A	GQ121805	N/A	N/A N/A
S. leiocladum-1	Para-sorghum	AusTRCF300148 AusTRCF300170 <sup>A</sup>	N/A	GQ121763	N/A	GQ121778	GQ121739
	Chaetosorghum	AusTRCF300170 AusTRCF302367 <sup>A</sup>	GQ121834	GQ121767	GQ121812	GQ121778 GQ121787	GQ121739 GQ121742
S. macrospermum	0	AusTRCF302521 <sup>A</sup>					
S. matarankense-1	Para-sorghum	AusTRCF302636 <sup>A</sup>	GQ121826	GQ121757 N/A	GQ121804 N/A	GQ121776	GQ121731 N/A
S. matarankense-2	Para-sorghum	AusTRCF302539 <sup>A</sup>	N/A N/A	N/A N/A	N/A N/A	N/A GQ121785	N/A N/A
S. nitidum-1	Para-sorghum						
S. nitidum-2	Para-sorghum	AusTRCF302558 <sup>A</sup>	GQ121815	GQ121764	GQ121807	N/A	GQ121740
S. plumosum-1	Stiposorghum	AusTRCF302399 <sup>A</sup>	GQ121819	GQ121762	GQ121798	N/A	N/A
S. plumosum-2	Stiposorghum	AusTRCF302489 <sup>A</sup>	N/A	N/A	N/A	GQ121773	GQ121729
S. plumosum-3	Stiposorghum	AusTRCF302635 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A
S. stipoideum-1	Stiposorghum	AusTRCF302393 <sup>A</sup>	GQ121827	GQ121751	GQ121794	N/A	GQ121734
S. stipoideum-2	Stiposorghum	AusTRCF302669 <sup>A</sup>	N/A	N/A	N/A	GQ121779	N/A
S. timorense-1	Para-sorghum	AusTRCF302381 <sup>A</sup>	GQ121820	GQ121760	GQ121801	GQ121777	GQ121727
S. timorense-2	Para-sorghum	AusTRCF302459 <sup>A</sup>	N/A	N/A	N/A	N/A	N/A

N/A, not applicable.

Sequence length variations were also observed between *Sorghum* species in the *trnT-trnL* spacer, ranging from 684 nt (*S. arundinaceum*) to 693 nt (*S. leiocladum* and *S. laxiflorum*). Low sequence length differences of 2 nt in the *trnT-trnL* spacer were observed among the *S. bicolor* accessions. Significant sequence variations arising from transitions and transversions were observed at eight positions, which resulted in the discrimination of *S. bicolor*-12, -13 and -14 from the rest

of the *S. bicolor* accessions. The sequences obtained from the *trnY-trnD* spacer were between 318 nt (*S. amplum*, *S. angustum*) and 329 nt (*S. exstans*). The sequences obtained from the ITS showed narrow length differences of 528–534 nt between the *Sorghum* species; base substitutions in the ITS1 accounted for most of this variation. The *S. bicolor* accessions exhibited sequence length differences arising from a single nucleotide indel in the ITS1 region.

<sup>\*</sup> Superscripts at the end of the accession number denote the country of origin and the donor of that particular accession; if only a single country code is present then that country is both a donor and the origin of the accession. A, Australia; B, Burundi; E, Ethiopia; K, Kenya; T, Tanzania; Y, Yemen; Zm, Zambia; Zw, Zimbabwe.

Table 2. Primers used to amplify and sequence the five non-coding regions of cpDNA and the ITS of nrDNA

Region of cpDNA	Primer name	Primer sequence $(5' \rightarrow 3')$	Source of primer sequence	
psbZ-trnG	$tnSM - fw^{\dagger}$	TGCTTCTCCTGATGGTTGGT	This study	
	$tnSM - rv^{\dagger}$	GCTCGCTACATTGAACTACGC	·	
trnY-psbM	psBD - fw*	CTGTCAAGGCGGAAGCTG	This study	
•	$psBD - rv^{\dagger}$	GGGTCACATAGACATCCCAAT	·	
	$trYB - fw^{\dagger}$	GGTTAATGGGGACGGACT		
	$trYB - rv^{\dagger}$	AGGAAGTTAAGATGAGGGTGG		
trnY-trnD	$trTD - fw^{\dagger}$	TGACGATATGTCTACGCTGGT	This study	
	trTD - rv*	AATCCCTGCGGGGTGTAT	·	
trnT-trnL	$trTL - fw^{\dagger}$	CATTACAAATGCGATGCTCT	Taberlet <i>et al.</i> (1991)	
	$trTL - rv^{\dagger}$	TCTACCGATTTCGCCATATC		
ITS	ITS5 $-fw^{\dagger}$	GGAAGTAAAAGTCGTAACAAGG	White et al. (1990)	
	$ITS4 - rv^{\dagger}$	TCCTCCGCTTATTGATATGC	, ,	

<sup>\*</sup> Primer was used for amplification only.

Table 3. Sequence characteristics and tree statistics of the cpDNA and ITS regions from maximum-parsimony (MP) analysis

	cpDNA regions				Combined or DNA	Combined anDMA	
	psbZ-trnG	trnY-trnD	trnY-psbM	trnT-trnL	ITS	Combined cpDNA regions	Combined cpDNA regions and ITS
LAS	286-291	318-329	1028-1053	684-693	528-534	2316-2366	2844-3111
PICs*	8 (2.7 %)	12 (3.6%)	32 (3.9 %)	19 (2.7 %)	69 (12.8 %)	71 (3.0 %)	140 (4.5 %)
TL	16	48	101	57	190	536	743
CI	0.9375	0.8958	0.6931	0.8947	0.8737	0.6250	0.6743
HI	0.0625	0.1048	0.31	0.1053	0.1263	0.3750	0.3257
RI	0.9846	0.9734	0.93	0.9757	0.9764	0.8463	0.8938
RC	0.9231	0.8720	0.6489	0.8730	0.8531	0.5252	0.6027

<sup>\*</sup> Inclusive of the out-group.

### Parsimony analysis of the ITS sequences

The aligned sequences of the ITS of the nrDNA provided the highest number of parsimony-informative characters (69: 12.8 %) of the regions used in this study, which could be attributed to an overall faster rate of base substitutions in the ITS than in the non-coding regions of the cpDNA. The ITS provided consistency and retention indices of 0.87 and 0.97, respectively (Table 3). The 50 % majority rule consensus tree from the phylogenetic analysis of DNA sequences of the ITS of 21 Sorghum species and Zea mays as an out-group species is shown in Fig. 1. Two lineages, A and E, were resolved. Lineage A was resolved with strong bootstrap support (100 %) and contained the Eu-sorghum species (clade B, 100 % bootstrap) and clade C with similar bootstrap support containing S. laxiflorum and S. macrospermum. The moderately supported internal clade D (61 %) contains unresolved relationships of S. bicolor accessions with other Eu-sorghum species but excludes S. bicolor-2 originally from Yemen. The other lineage, E, with 92 % bootstrap support contained the remaining native Australian Sorghum species which, except for S. nitidum, are contained in clade F with moderate bootstrap support (88 %; Fig. 1).

Analysis of the non-coding regions of cpDNA sequence data

The cpDNA regions, psbZ-trnG, trnY-psbM, trnY-trnD and trnT-trnL, revealed differences in the number of

parsimony-informative characters, consistency and retention indices (Table 3). The cpDNA data show less homoplasy than the ITS data (Table 3), resulting in more fully resolved 50 % majority rule consensus trees and generally greater bootstrap values for various nodes. The trnY-psbM spacer provided the highest number of parsimony-informative characters (32; 3.9 %). The psbZ-trnG region provided the lowest number of parsimony-informative characters (eight; 2.7%). The trnT-trnL and trnY-trnD intergenic spacers generated sequences that had 19 (2.7%) and 12 (3.6%) parsimonyinformative characters, respectively. As measures of accuracy for the topologies obtained, consistency and retention indices were highest (0.94 and 0.98, respectively) for psbZ-trnG among the cpDNA regions used. The trnY-psbM spacer had the lowest consistency index (0.69) and retention index (0.93). The 50 % majority rule consensus of 100 mostparsimonious trees is shown in Fig. 2. Lineage A is resolved and includes all the Eu-sorghum species, clade B with strong support (100%), and S. laxiforum and S. macrospermum (clade C) with equal bootstrap support. The strongly supported (94 %) clade D includes all *Eu-sorghum* species but excludes S. arundinaceum. The strongly supported (96%) internal clade H containing S. almum and S. bicolor-2 from Yemen excludes S. drummondii-2. All wild Sorghum species from Australian except S. laxiflorum and S. macrospermum form the second lineage (lineage J), which has very strong bootstrap

<sup>†</sup> Primer used for both PCR amplification and sequencing.

LAS, length of aligned sequences; PICs, parsimony-informative characters (number and per cent); TL, tree length; CI, consistency index; HI, homoplasy index; RI, retention index; RC, rescaling consistency index.

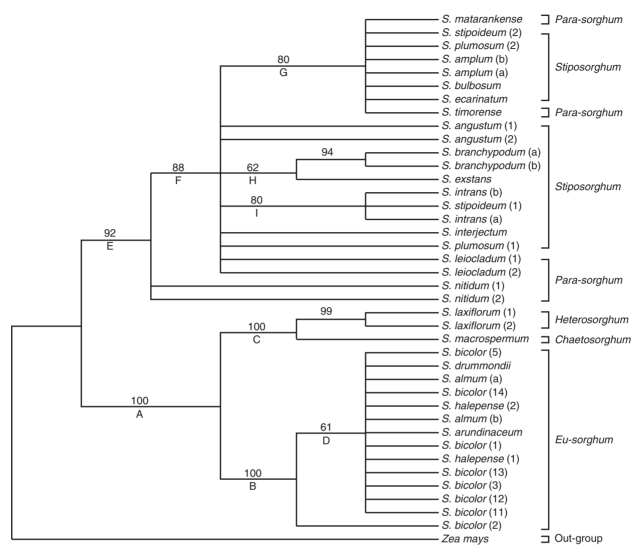


Fig. 1. Maximum-parsimony 50 % majority rule consensus tree (1000 bootstrap replicates with 100 random additions; MaxTrees = 100) generated from a phylogenetic analysis of DNA sequence data from the internal transcribed spacers of the nrDNA of 21 *Sorghum* species and *Zea mays* as an out-group species. Indels are treated as missing data. Clades are indicated by letters below the branch. Bootstrap values of >50 % are indicated above the branches.

support (100%; Fig. 2). Clade K, with moderate bootstrap support (71%), includes all *Stiposorghum* species and some *Parasorghum* species except *S. leiocladum* and *S. nitidum*. The internal relationships within clade K are either moderately to strongly supported by the bootstrap data (76–95%) or remain unresolved (Fig. 2).

### Combined analysis of cpDNA and ITS sequence data

The combined cpDNA and ITS sequences generated a total of 3096 characters, 140 of which (4.5%) were parsimony-informative (Table 3). The maximum-parsimony (MP) analysis involving the combined data from the cpDNA regions and the ITS sequence data, with gaps either considered as missing values (Fig. 3) or scored as presence or absence characters (data not shown), produced two main lineages. Lineage A contains all the *Eu-sorghum* species (clade B), which includes all *S. bicolor* and their immediate wild relatives,  $S. \times almum$ ,

S. halepense, S. drummondii and S. arundinaceum with 100 % bootstrap support. The other lineage, lineage J, consists of all Australian wild Sorghum species except S. laxiflorum and S. macrospermum with high bootstrap support (Fig. 3). S. laxiflorum and S. macrospermum not only form a single clade (C) with strong bootstrap support but are also more closely related to the Eu-sorghum species with 100 % bootstrap support than to other Australian wild Sorghum species. Within the Eu-sorghum section, clade D excludes S. arundinaceum from the rest of the species, but a subgroup comprising S. halepense-1, S. drummondii, S. almum, and S. bicolor-1, -2, -5 and -13 is formed as clade F with 99 % bootstrap support (Fig. 3). The strongly supported (94%) clade E consists of three accessions of S. bicolor (-3, -11 and -14). The S. bicolor accessions in this clade originated from southern Africa, one from Zimbabwe (S. bicolor-3) and the other two from Zambia. S. bicolor-2, an accession from Yemen, seems to be distantly related to S. bicolor accessions from southern Africa but has a

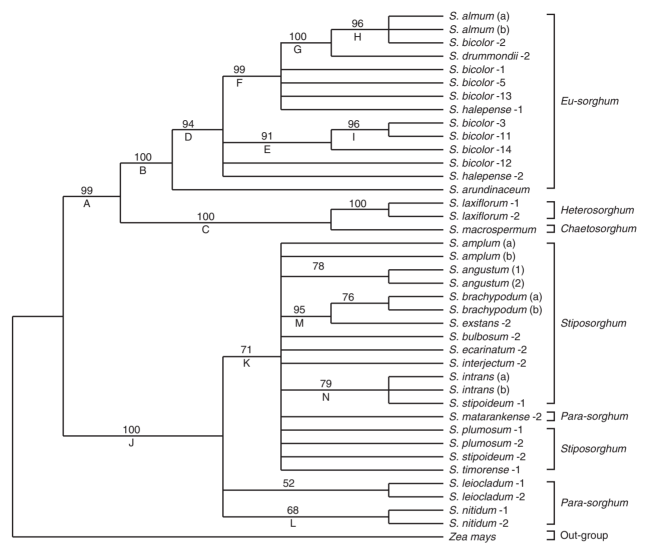


Fig. 2. Maximum-parsimony 50% majority rule consensus tree (1000 bootstrap replicates with 100 random additions; MaxTrees = 100) from a phylogenetic analysis of DNA sequence data from the four regions of cpDNA of 21 *Sorghum* species and *Zea mays* as an out-group species. Indels are treated as missing data.

Clades are indicated by letters below the branch. Bootstrap values of >50% are indicated above the branches.

stronger association (clade H) with *S. almum* with strong bootstrap support (Fig. 3).

Stiposorghum and Para-sorghum form clade J with 100 % bootstrap support (Fig. 3). The internal nodes of this particular clade, however, lack strong bootstrap support. Most of the Para-sorghum and all of the Stiposorghum species form clade K with moderate bootstrap support and the two accessions of S. nitidum form a single clade (L) with equally moderate bootstrap support (Fig. 3). Clade M consists of S. brachypodum and S. exstans with 95 % bootstrap support. S. intrans and S. stipoideum-1 form clade N, and S. amplum and S. ecarinatum form clade O but with only moderate bootstrap support (78 %; Fig. 3).

### DISCUSSION

Comparative DNA sequencing has become a widespread tool for inferring phylogenetic relationships and in systematic studies as it is relatively fast and convenient. Phylogenetic inference and elucidation of the evolutionary processes that generate biological diversity have been accomplished even at lower taxonomic levels using non-coding regions of the chloroplast genome and the ITSs of the nrDNA (Mort et al., 2007; Kårehed et al., 2008). In the present study, all the five cpDNA primers used successfully amplified the target regions in the Sorghum species. Mort et al. (2007) assessed the phylogenetic utility of the ITS and nine rapidly evolving cpDNA loci (including trnS-trnfM, trnD-trnT, psbM-trnD and trnT-trnL) involving six taxa sets of 13-23 taxa using published primer sequences (Shaw et al., 2005). Failure of PCR amplification was reported in Tolpis (Asteraceae) and Chrysosplenium (Saxifragaceae) with the primer pair trnD-trnT. Attempts to amplify the trnT-trnL region was not successful in all the taxa used. This implies that successful amplification using published primers for some cpDNA regions of one taxon may not have universal application across taxa. In this study, trnY-psbM provided the highest number of parsimonyfollowed informative characters by trnT-trnL

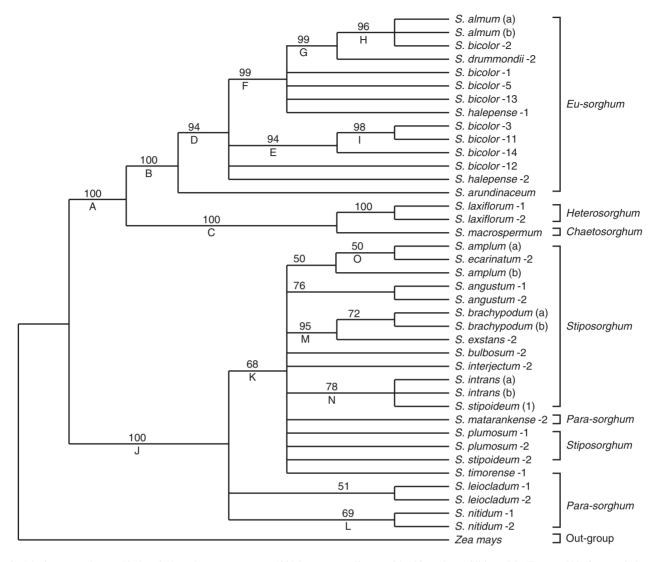


Fig. 3. Maximum-parsimony 50% majority rule consensus tree (1000 bootstrap replicates with 100 random additions; MaxTrees = 100) from a phylogenetic analysis of DNA sequence data from the four regions of cpDNA and the internal transcribed spacers of the nrDNA of 21 Sorghum species and Zea mays as an out-group species. Indels are treated as missing data. Clades are indicated by letters below the branch. Bootstrap values of >50% are indicated above the branches.

trnY-trnD. Based on the potentially informative characters generated, trnT-trnL and psbM-trnD were identified as suitable for low taxonomic level phylogenetic studies (Shaw et al., 2005). Of the cpDNA regions used in this study, trnY-psbM, trnT-trnL and trnY-trnD intergenic spacers were useful in the inference of phylogenetics at low taxonomic level in general and in the genus Sorghum in particular.

In the ITS analysis, all *Stiposorghum* and *Para-sorghum* species were resolved into a lineage separate from *Eu-sorghum*, *Heterosorghum* and *Chaetosorghum* species with strong bootstrap support (92%). These results are consistent with findings based on the analysis of the ITS sequences (Sun *et al.*, 1994; Dillon *et al.*, 2001). However, in general the internal relationships between species within section are unresolved (Fig. 1). As implied and based on its utility in numerous studies, the ITS is a useful marker for resolving phylogenetic relationships at various taxonomic levels, in particular the infrageneric level. However, caution needs to be

taken when analysing ITS sequence data to avoid problems resulting from concerted evolution on the rDNA arrays. Concerted evolution may homogenize different paralogous gene copies in a genome leading to the loss of all but one of the copies, i.e. different copies may be present in different organisms by chance and consequently this will create disagreement between the gene trees and species trees (Álvarez and Wendel, 2003). A fundamental requirement for historical inference based on nucleic acid or protein sequences is that the genes compared are orthologous as opposed to paralogous. However, there are inherent risks in relying exclusively on rDNA sequences for phylogenetic inferences given the 'nomadic' nature of the rDNA loci between inclusion of paralogous genes and exclusion of orthologous comparisons (Álvarez and Wendel, 2003).

The combined analysis of the cpDNA and ribosomal ITS sequence data, as when only the combined cpDNA dataset was used, resolved two major lineages (Figs 2 and 3). In

one lineage, A, the Eu-sorghum species form a clade B with 100 % bootstrap support. These results indicate a close association between species within the section Eu-sorghum. The present results are in agreement with the findings from an assessment of phylogenetic relationships among Sorghum taxa based on 30 allozyme loci (Morden et al., 1990), which could not show clear delimitation between the Eu-sorghum taxa. Weedy forms of sorghum (e.g. S. drummondii) occur wherever cultivated sorghum and S. arundinaceum grow sympatrically (De Wet, 1978). Sympatric speciation, one of the theoretical models for the phenomenon of speciation, is the genetic divergence of various populations from a single parent species inhabiting the same geographical region, such that these populations become different species. However, the present study has shown emergence of two subgroups within Eu-sorghum with strong bootstrap support (Fig. 2). A strong phylogenetic affinity was obtained between S. bicolor-3, an accession from Zimbabwe, three other S. bicolor accessions (-11, -12 and -14) from Zambia and S. halepense-1, as shown in clade E. The other subgroup, clade F, contains all other S. bicolor accessions (-1, -2, -5 and -13; Fig. 2). Within this clade, S. almum is closely associated with S. bicolor-2, an accession from Yemen. S. almum is believed to be a recent fertile hybrid between S. halepense and S. bicolor (Doggett, 1970). As the chloroplast genomes are believed to display maternal inheritance in the majority of angiosperms (Mogensen, 1996; Keeling, 2004; Udall and Wendel, 2006), the present phylogenetic results suggest that S. bicolor could be the maternal parent of S. almum.

S. drummondii, commonly known as Sudan grass, is believed to be a segregate from a natural hybrid between S. bicolor and S. arundinaceum and is thought to have originated in the region from southern Egypt to the Sudan (Hacker, 1992). The cultivated species, S. bicolor, is allied to S. arundinaceum, its assumed wild progenitor (Lazarides et al., 1991). This is consistent with the present results, which place S. arundinaceum in close relationship with S. bicolor with 100 % support (Fig. 3).

Various models of the origin of *S. halepense* have been suggested. Generally, the species is believed to have arisen as a segmental allotetraploid derived from the cross of two diploids (n=10) species. Doggett (1970) suggested that *S. halepense* was derived from the rhizomatous perennial *S. propinquum* and the annual *S. arundinaceum*. In the allozyme variation study involving *Eu-sorghum*, *S. halepense* could not be differentiated from *S. bicolor*, suggesting that the latter was one of the parental species of *S. halepense* (Morden *et al.*, 1990). The present results (Figs 1 and 2) support the suggestion that *S. bicolor* is one of the parents of *S. halepense*.

Eu-sorghum species are closely related to S. macrospermum and S. laxiflorum with strong bootstrap support (Fig. 3), consistent with previous reports based on combined ITS1/ndhF/adh1 (Dillon et al., 2007) and ITS sequence data (Sun et al., 1994). This study has also revealed a very close relationship between S. macrospermum and S. laxiflorum with 100% support (Figs 2 and 3), which suggests these species should not be classified under different sections. The close association between these two species has prompted the suggestion that Chaetosorghum and Heterosorghum be combined in a single

section (Sun et al., 1994; Dillon et al., 2004), which is strongly supported by the present data. The ancestry of cultivated sorghum is not well resolved. Based on the ease of formation of crosses (Doggett, 1970) and chromosome morphological similarities (Gu et al., 1984) within Eu-sorghum, it has been assumed that no other sections except Eu-sorghum provided the ancestral material for cultivated sorghum (van Oosterhout, 1992). However, the close association of S. macrospermum and S. laxiflorum with section Eu-sorghum indicates that there is strong sequence homology among them, suggesting that these species are phylogenetically closely related.

The phylogenetic relationships among the Australian wild *Sorghum* species have been described in detail (Sun *et al.*, 1994; Spangler *et al.*, 1999; Dillon *et al.*, 2001, 2004, 2007; Spangler, 2003; Price *et al.*, 2005). The internal relationships among the Australian wild sorghums are moderately well supported. *S. intrans* and *S. stipoideum* belonging to section *Stiposorghum* form a clade N with moderate support (Figs 2 and 3). These species have also been reported to be comparable in morphology and distribution (Lazarides *et al.*, 1991).

The analysis of the combined data set involving ITS and cpDNA resulted in a tree that is identical to that inferred from cpDNA alone. Similar results were obtained using the two loci on Crassula (Mort et al., 2007). In contrast to a cpDNA-based approach, phylogenetic studies using nuclear DNA sequences have traditionally been hampered by difficulties in distinguishing between orthologous and paralogous sequences (Small et al., 2004). The practice of obtaining sequence data from two or more loci that can reasonably provide independent tests of phylogeny is a proven means of avoiding well-supported but incorrect phylogenies that do not track organismal phylogeny (Mort et al., 2007). Chloroplast DNA loci, which are often assumed to be uniparentally inherited and non-recombining, have been extensively used for systematics and phylogenetics. However, the rate of evolution of the cpDNA genome is slower than that of the nuclear genome. Correspondingly, the cpDNA regions that have been used for phylogenetic studies are less variable than the most extensively used nuclear loci, internal transcribed spacers of nrDNA (ITS) (Small et al., 2004; Mort et al., 2007). It is often difficult to obtain adequate resolution of any phylogeny of closely related taxa using few cpDNA loci due to the low number of phylogenetically informative characters (Rokas et al., 2003). Hence, the practice of acquiring sequence data from several loci is a proven means of acquiring a better resolved phylogeny (Rokas and Carroll, 2005; Mort et al., 2007). In the present study, the phylogeny of the genus Sorghum is well resolved when the combined data from ITS and four cpDNA regions were used.

### Conclusions

The cpDNA regions used in this study have provided phylogenetic relationships even at low taxonomic level. The trnY-psbM, trnT-trnL and trnY-trnD intergenic spacers have specifically been identified to be more useful in inferring phylogenetics even at infraspecies level. The close relationship between S. macrospermum and S. laxiflorum suggest that they should not be classified under different sections and

support the proposal that sections *Chaetosorghum* and *Heterosorghum* be merged. The results also indicated that the *Eu-sorghum* species are more closely related to *S. macrospermum* and *S. laxiflorum* than to any other Australian wild *Sorghum* species. *S. almum* is more closely associated with *S. bicolor* than with *S. halepense*, its known parents. As the chloroplast genome is maternally inherited, the results suggest that *S. bicolor* is the most probable maternal parent of *S. almum*. The *S. bicolor* accessions (-3, -11 and -14) from southern Africa form a distinct and well-supported clade. *S. bicolor*-2, originally from Yemen, is distantly related to other *S. bicolor* accessions used in this study. These results may provide opportunities to use sorghum gene pools outside section *Eu-sorghum* for development and improvement of cultivated sorghum.

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