



Genome-wide characterization of Nuclear Factor Y (NF-Y) gene family of sorghum [*Sorghum bicolor* (L.) Moench]: a bioinformatics approach

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Abstract Nuclear factor Y (NF-Y) is a heterotrimeric transcription factor (TF) complex with preferential binding to CCAAT elements of promoters, regulating gene expression in most of the higher eukaryotes. The availability of plant genome sequences have revealed multiple number of genes coding for the three subunits, namely NF-YA, NF-YB and NF-YC in contrast to single NF-Y gene for each subunit reported in yeast and animals. A total of 33 NF-YTF comprising of 8 NF-YA, 11 NF-YB and 14 NF-YC subunits were accessed from the sorghum genome. The bioinformatic characterization of NF-Y gene family of sorghum for gene structure, chromosome location, protein motif, phylogeny, gene duplication and *in-silico* expression under abiotic stresses have been attempted in the present study. The identified *SbNF-Y* genes are distributed on all the 10 chromosomes of sorghum with variability in the frequency and 18 out of 33 *SbNF-Ys* were found to be intronless. Segmental duplication event was found to be predominant feature based on gene duplication pattern study. Several orthologs and paralogs groups were disclosed through the comprehensive phylogenetic analysis of *SbNF-Y* proteins along with 36 *Arabidopsis* and 28 rice NF-Y proteins. *In-silico* expression analysis under abiotic stresses using rice transcriptome data revealed several of the sorghum *NF-Y* genes to be associated with salt, drought, cold and heat stresses.

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Introduction

Nuclear Factor Y (NF-Y) is a transcription factor complex with three subunits namely NF-YA, NF-YB and NF-YC that binds to the CCAAT box of eukaryotic promoters and regulate gene expression (Dolfini and Mantovani 2013; Laloum et al. 2013; Petroni et al. 2012). NF-YA also called HAP3 (Heme Activator Protein 3) or CBF-A (CCAAT-Binding Factor A), are characterized having two domains which are highly conserved in all higher eukaryotes (Maity and de Crombrugge 1998). The N-terminal domain is involved in NF-YB: NF-YC heterodimer interaction and the C-terminal domain is concerned with DNA binding site recognition (Xing et al. 1994). At the stage of transcription activation NF-YB forms a heterotrimer, with NF-YA and NF-YC (Kim et al. 1996). Like NF-YC proteins consist of HFM domain, but these are more closely related to the core histone H2A (Dolfini et al. 2012).

These subunits are required for their association and transcriptional regulation in both vertebrates and plants (Sinha et al. 1995). A single gene encodes each NF-Y subunit in animals and yeast, but in plants the three subunits are encoded by multiple genes (Edwards et al. 1998; Gong et al. 2004; Riechmann et al. 2000). Based on the availability of plant genome sequences NF-Y gene families of *Arabidopsis*, rice, wheat, *Brachypodium distachyon*, canola, soybean, common bean have been reported (Cao et al. 2011; Liang et al. 2014; Ripodas et al. 2014; Siefers et al. 2009; Stephenson et al. 2007; Thirumurugan et al. 2008).

Individual NF-Y subunits are identified to be involved in number of important plant processes, yet no complete NF-Y

complex for gene regulation has been studied in the plants. The functional diversity of NF-Y genes corresponding to the three subunits have been elucidated in different plants (Laloum et al. 2013; Petroni et al. 2012). The role of NF-Y in embryogenesis, photoperiod-dependent flowering time regulation, seed oil production, root nodule development, chloroplast biogenesis, seed germination (Petroni et al. 2012), heat stress (Sato et al. 2014), drought tolerance (Zhang et al. 2015), enhanced yield (Yadav et al. 2015) have been reported.

Sorghum bicolor is a grass species belonging to the subtribe Saccharinae, within the grass family Poaceae. Sorghum is a diploid with 10 chromosomes and has a genome of about 730 Mb (Paterson et al. 2009). There are few studies related with transcription factors like MYB, AUX_ARF, bZIP, AP2, WRKY, basic helix loop helix, NAC and Dof in sorghum (Kushwaha et al. 2011; Lu et al. 2013; Sekhwal et al. 2015; Yan et al. 2013) though genome wide analysis of NF-Y gene family of sorghum is still lacking. This manuscript reports genome wide *in-silico* characterization of *SbNF-Y* gene family highlighting putative gene structures, chromosomal localization, motif analysis, gene duplication, ancestral protein sequence analysis, comprehensive phylogenetic analysis with rice and *Arabidopsis* NF-Y gene family. The *in-silico* expression profiling of NF-Y genes of sorghum under stress conditions have also been assessed based on the available rice transcriptome data after deciphering the corresponding orthologs identified in the phylogenetic tree.

Materials and methods

Databases searches for the identification of NF-Y gene family of sorghum and its annotation

The NF-Y A, B and C sequences of *A.thaliana* were retrieved from plant transcription factor database Plant TFDB (<http://plantfdb.cbi.pku.edu.cn/>) version 3.0 (Jin et al. 2014). The nucleotide sequences of NF-Y domain were used to search the potential NF-Y domain homologs hit in the whole genome shotgun (wgs) sequence and nucleotide collection (nr/nt) of sorghum through tblastn at the NCBI database. The 2 kb upstream and 2 kb downstream sequences of NF Y domain homologs were retrieved from whole genome shotgun sequence of sorghum for fishing out the putative NF-Y genes using BioEdit software version 7.2.5 (Hall 1999). The NF-Y gene sequences identified were tentatively designated as *SbNF-YAs*, *SbNF-YBs* and *SbNF-YCs* with corresponding numbers. The annotated sequences were further subjected to bioinformatics server namely FGGENESH (Solovyev et al. 2006) for prediction of full length genes with putative CDS and protein sequences. The putative NF-Y protein sequences of sorghum were subjected to protein functional analysis using PFAM version 27 (Punta et al. 2012), PROSITE version 20.93

(Sigrist et al. 2013), INTERPROSCAN version 4.0 (Quevillon et al. 2005) and MOTIFSCAN databases (Falquet et al. 2002). The physio-chemical feature of NF-Y proteins like isoelectric point (PI) and molecular weight were calculated using ExPASy server (Gasteiger et al. 2003).

Chromosomal distribution and intron/exon gene structure prediction

The annotated *SbNF-Y* genes were positioned on chromosomes through NCBI-BLAST search and manually marked. FGGENESH server was used to analyze the gene structure of predicted *SbNF-Y* genes.

NF-Y protein alignment and phylogenetic analysis

ClustalX 2.0.10 (Thompson et al. 1997) was utilized for Multiple sequence alignment of predicted *SbNF-Y* proteins. Neighbor joining approach (with 500 reiterations) was used to construct phylogenetic tree using MEGA 6.0 software (Tamura et al. 2013).

Inference of duplication time

The Ks and Ka value of paralogous sequences were calculated by DnaSP software version 5.0 (Librado and Rozas 2009). The Ks value was calculated which was further used to calculate the approximate date of the duplication event ($T = Ks/2\lambda$), using the mean value of clock-like rates (λ) of synonymous substitution (6.5×10^{-9}) (Gaut et al. 1996).

Motif prediction and subcellular localization

Motif analysis of *SbNF-YA*, *SbNF-YB* and *SbNF-YC* protein sequences were analyzed through MEME (Multiple EM for Motif Elicitation) program software version 4.4.0 (Bailey and Elkan 1994). For the identification of conserved motifs the maximum number of motifs was set for 10, with a width range between 5 to 55, while other factors were at default selections. The NF-Y protein sequences were passed through the available prediction algorithms and the raw data were used to make preliminary location predictions. ARAMEMNON (<http://aramemnon.botanik.uni-koeln.de>) server (Schwacke et al. 2003) was used for subcellular localization and identification of transmembrane domain.

Prediction of ancestral protein sequences

For the prediction of ancestral sequences, FAST-ML 2.02 (www.tau.ac.il/~talp/supplementary/fastml/fastml.2.02) (Pupko et al. 2002) was used. The unrooted input tree was used, which was automatically rooted by midpoint rooting in FAST-ML Ancestral Sequence Prediction analysis.

In-silico expression analysis

Due to non-availability of transcriptome data for the identified *SbNF-Y* genes of sorghum, attempts were made to analyze the transcriptome data of rice orthologs to generate heatmap as both the crops belong to same family and show high degree of synteny (Wang et al. 2009). The coding sequences of the orthologous pair of sorghum genes were used as queries for BLAST search and the corresponding rice gene sequences were identified based on more than 90 % sequence identities. The Rice eFP browser (<http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi>) (Toufighi et al. 2005) was used to retrieve RMA normalized expression data for abiotic stresses namely drought stress, salt stress, cold stress and heat shock recorded for seedling stages. All transcript data was analyzed with HCE version 2.0 beta web tools (Seo et al. 2004).

Result and discussion

Genome wide characterization of NF-Y transcription factor genes of sorghum

The nucleotide and amino-acid sequences of the conserved NF-Y domains corresponding to A, B and C subunits subjected to BLAST search resulted into a total of 33 *NF-Y* genes comprising of eight *NF-YA*, 11 *NF-YB* and 14 *NF-YC* from the whole genome of sorghum. The predicted sorghum eight *NF-YA* genes, 11 *NF-YB* and 14 *NF-YC* genes were named as *SbNF-YA1* to *SbNF-YA8*, *SbNF-YB1* to *SbNF-YB11* and *SbNF-YC1* to *SbNF-YC14* respectively. There exists great variability in terms of number of *NF-YA*, *NF-YB* and *NF-YC* subunits in different crops. A total of 10 *NF-YA*, 13 *NF-YB* and 13 *NF-YC* genes were predicted in *Arabidopsis* (Siefers et al. 2009), 14 *NF-YA*, 14 *NF-YB*, five *NF-YC* in *Brassica napus* (Liang et al. 2014) while in *Brachypodium* seven *NF-YA*, 17 *NF-YB* and 12 *NF-YC* genes were annotated (Cao et al. 2011). In case of wheat 10 *NF-YA*, 11 *NF-YB* and 14 *NF-YC* genes have been reported (Stephenson et al. 2007) while in rice, there exists five *NF-YA*, 10 *NF-YB* and 10 *NF-YC* genes (Yang et al. 2005). Recently genome wide analysis of NF-Y genes of two legumes namely soybean (21 *GmNF-YA*, 32 *GmNF-YB* and 15 *GmNF-YC*) and common bean (nine *NF-YA*, 14 *NF-YB* and seven *NF-YC*) has been reported (Quach et al. 2015; Ripodas et al. 2014). The presence of the conserved NF-YA, NF-YB and NF-YC domains in the predicted *SbNF-Y* protein was a typical feature for considering it as a family member of NF-Y transcription factor.

The annotated sequences were further subjected to FGENESH server for prediction of ORFs, CDS and protein sequences. The result of FGENESH for *SbNF-YA*, *SbNF-YB* and *SbNF-YC* gene sequences of sorghum is summarized in Table 1. The identified *SbNF-Y* genes encode peptides

ranging from 90 to 790 aa with the pI value varying from 4.26 to 10.83, and the molecular weight ranging from 10.21 kD to 83.52 kDa as estimated from expasy server (http://web.expasy.org/compute_pi/).

Protein functional analysis and subcellular localization

The protein functional analyses of these 33 putative *SbNF-Y* proteins have been performed using different servers namely PFAM version 27, PROSITE version 20.93, INTERPROSCAN version 4.0, MOTIFSCAN databases and NUCPRED software. The members of *SbNF-YA* proteins subjected to INTERPROSCAN revealed identity with InterPro accession number IPR001289. These proteins were also similar to signature accession namely PS51152 from PROSITE database and Pfam database confirming their identity to NF-YA like proteins. All the *SbNF-YA*, *SbNF-YB* and *SbNF-YC* proteins showed nuclear localization signal (NLS) with RRR amino-acid residue when subjected to NUCPRED software. Further Motifscan integrated with PeroxiBase profiles, PROSITE patterns, PROSITE profiles, HAMAP profiles, Pfam HMMs (local models), Pfam HMMs (global models) databases revealed the presence of alanine, arginine, glutamine and serine rich regions in different *SbNF-YA* proteins.

Similarly, for *SbNF-YB* proteins showed identity with InterPro accession number IPR003957 along with a signature accession namely PF00808 for Pfam database and PS00685 for PROSITE database confirming their identity to NF-YB like proteins. In case of *SbNF-YB* proteins only six out of 11 sequences showed NLS signal with variable amino-acid residues. The NLS signal for *SbNF-YB1*, *SbNF-YB3*, *SbNF-YB5* and *SbNF-YB6* was found to be KRK while for *SbNF-YB7* and *SbNF-YB8*, the NLS signal of KRRK and RRK was observed. Motifscan integrated with PeroxiBase profiles, PROSITE patterns, PROSITE profiles, HAMAP profiles, Pfam HMMs (local models), Pfam HMMs (global models) databases revealed the presence of asparagine, glutamine, glycine, methionine, histidine, alanine and proline rich regions in different *SbNF-YB* proteins.

The *SbNF-YC* sequences showed identity with InterPro accession number IPR027170 along with PF00808 accession of Pfam database while PROSITE database could not identify any putative hits. Only five *SbNF-YC* sequences (*SbNF-YC1*, *SbNF-YC3*, *SbNF-YC5*, *SbNF-YC6*, *SbNF-YC8*) out of 14 have NLS with KRR residue when subjected to NUCPRED software. Motifscan integrated with PeroxiBase profiles, PROSITE patterns, PROSITE profiles, HAMAP profiles, Pfam HMMs (local models), Pfam HMMs (global models) databases revealed the presence of glutamine, proline, alanine, aspartic acid, glycine and leucine in different *SbNF-YC* proteins.

Table 1 List of *NF-Y* genes identified in sorghum with their corresponding proteins, CDS and chromosome positions

<i>Sb NF-Y</i> Gene	Source accession number	Chr. no.	Chromosome location (bp)	No. of introns	mRNA length(bp)	Amino acid length (a.a)	Protein Mol. Wt.(kDa)	Protein PI value
<i>SbNF-YA1</i>	XM 002465725.1	1	68598278–68598532	0	690	229	24.87	8.68
<i>SbNF-YA2</i>	ABXC01000113.1	1	10088592–10088732	1	273	90	10.21	10.83
<i>SbNF-YA3</i>	ABXC01000830.1	2	3779232–3779891	2	636	211	22.83	9.96
<i>SbNF-YA4</i>	ABXC01005669.1	8	52974564–52974779	1	360	119	13.05	10.05
<i>SbNF-YA5</i>	ABXC01001587.1	2	73106344–73106742	5	918	305	33.37	8.49
<i>SbNF-YA6</i>	ABXC01000526.1	1	55640044–55640457	4	783	260	28.35	9.76
<i>SbNF-YA7</i>	ABXC01000138.1	1	12431437–12431853	1	549	182	19.61	9.94
<i>SbNF-YA8</i>	ABXC01005686.1	8	53656544–55656918	3	873	290	30.78	9.88
<i>SbNF-YB1</i>	ABXC01000523.1	1	55505670–55506635	4	2373	790	83.52	5.38
<i>SbNF-YB2</i>	ABXC01002897.1	4	5945582–59456306	0	831	276	29.26	6.45
<i>SbNF-YB3</i>	ABXC01004591.1	7	6276612–6277436	0	828	275	27.67	6.00
<i>SbNF-YB4</i>	XM002459009.1	3	72403131–72403568	0	441	146	16.46	7.14
<i>SbNF-YB5</i>	XM002459011.1	3	72412515–72413060	0	544	182	19.09	6.15
<i>SbNF-YB6</i>	XM002463118.1	2	73025022–73025675	0	657	218	22.73	6.30
<i>SbNF-YB7</i>	002G135100.1	2	20148061–20146210	1	520	174	19.17	4.74
<i>SbNF-YB8</i>	007G117100.1	7	49678386–49682600	0	4214	297	33.44	5.14
<i>SbNF-YB9</i>	010G119200.1	10	2352136–2353131	0	822	273	28.66	7.36
<i>SbNF-YB10</i>	003G346500.1	3	66773363–66774961	5	388	167	18.03	6.11
<i>SbNF-YB11</i>	009G239600.1	9	57918864–57921191	3	419	137	15.04	4.87
<i>SbNF-YC1</i>	ABXC01004605.1	7	6754771–6755478	0	762	253	28.38	5.04
<i>SbNF-YC2</i>	ABXC01000653.1	1	64266744–64267415	0	744	247	26.18	5.31
<i>SbNF-YC3</i>	ABXC01001448.1	2	63138811–63139227	0	609	202	21.42	5.37
<i>SbNF-YC4</i>	ABXC01003563.1	5	51186980–51187096	5	558	185	20.34	8.26
<i>SbNF-YC5</i>	ABXC01004477.1	6	61341870–61342172	0	387	128	13.58	5.74
<i>SbNF-YC6</i>	ABXC01005047.1	7	63565543–63565956	2	726	241	25.81	5.87
<i>SbNF-YC7</i>	ABXC01006233.1	9	53768950–53769063	3	357	118	12.85	5.80
<i>SbNF-YC8</i>	ABXC01006864.1	10	56155673–56156437	0	798	255	28.39	5.11
<i>SbNF-YC9</i>	003G040500.1	3	3812916–3813569	0	654	217	22.95	4.29
<i>SbNF-YC10</i>	005G089100.1	5	12927331–12928204	1	793	263	29.35	4.26
<i>SbNF-YC11</i>	007G054700.1	7	5576465–5577850	0	1386	416	51.82	4.73
<i>SbNF-YC12</i>	008G43000.1	8	4255910–4257304	0	1395	464	51.95	4.72
<i>SbNF-YC13</i>	008G071900.1	8	9705760–9706977	0	1218	405	45.58	5.03
<i>SbNF-YC14</i>	007G070100.1	7	7704760–7707221	0	675	224	25.14	4.45

The sub-cellular localization of *SbNF-Y* proteins based on consensus sequences showed chloroplast as important target site. Few of the *SbNF-Y* proteins were also destined for mitochondria while few followed secretory pathway (Table 2). Several of the *SbNF-Y* proteins are attached to a membrane by hydrophobic anchors. Alpha helix transmembrane spans with average hydrophobicity were predicted in twelve of the *NF-Y* proteins. Anchoring of protein to GPI (glycosylphosphatidyl inositol) *via* C-terminal attachment was predicted in three of the *NF-YB* proteins namely *SbNFYB1*, *SbNFYB3* and *SbNFYB6*. Through gene fusion experiments it was demonstrated that proteins destined to receive a GPI anchor carry a C-terminal signal sequence for GPI-anchorage. (Caras et al. 1987).

Chromosomal locations and gene structure analysis of *SbNF-Y* genes

The identified 33 *SbNF-Y* genes of sorghum are distributed on all the ten chromosomes of *S. bicolor* as given in GenBank chromosome data. In case of *SbNF-YA* genes, a maximum of four genes *viz.* *SbNF-YA1*, *SbNF-YA6*, *SbNF-YA7* and *SbNF-YA2* were identified on chromosome one while chromosome two and eight possess two genes each. The chromosome location of *SbNF-YB* sequences revealed a maximum of three genes on the chromosome three namely *SbNF-YB5*, *SbNF-YB4* and *SbNF-YB10* while chromosome two and seven each have two genes. For *SbNF-YC*, a maximum of four genes *i.e.* *SbNF-YC6*,

Table 2 Identification of subcellular localization and transmembrane domain of *SbNF-Y* proteins using ARAMEMNON server

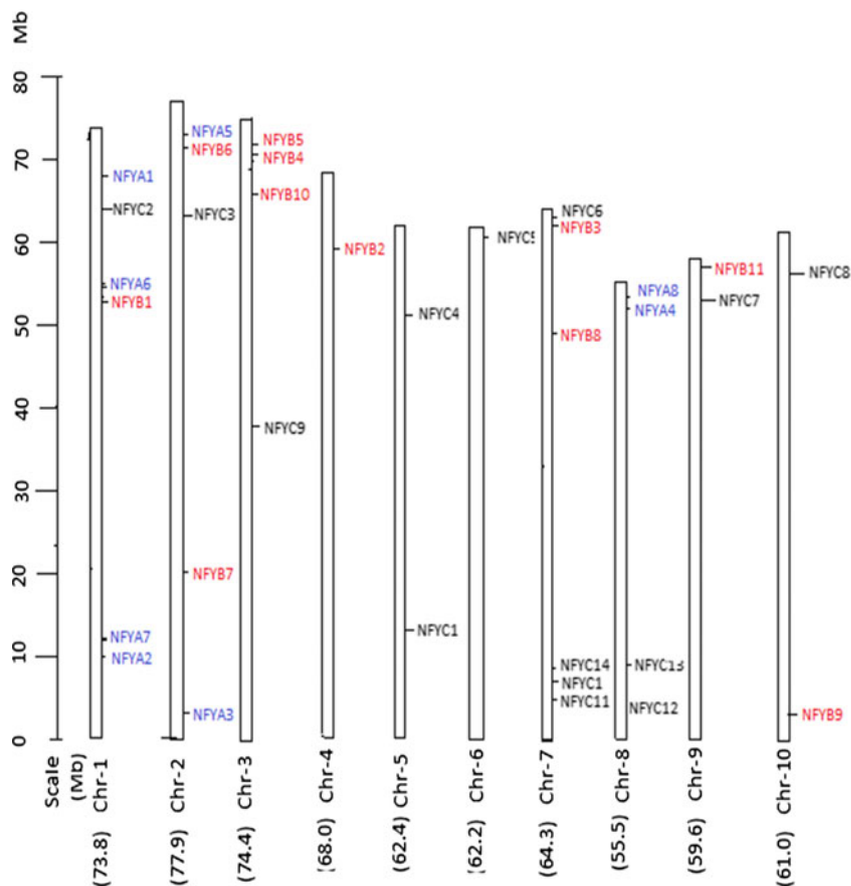
Gene	Transmembrane spans	Subcellular location			Lipid modification
		Chloroplast	Mitochondria	Secretory pathway	
<i>SbNF-YA1</i>	α helix TMspans	✓			
<i>SbNF-YA2</i>		✓			
<i>SbNF-YA3</i>	α helix TMspans				
<i>SbNF-YA4</i>					
<i>SbNF-YA5</i>		✓			
<i>SbNF-YA6</i>	α helix TMspans	✓			
<i>SbNF-YA7</i>		✓			
<i>SbNF-YA8</i>	α helix TMspans	✓	✓		
<i>SbNF-YB1</i>	α helix TMspans				GPI-SOM
<i>SbNF-YB2</i>			✓	✓	
<i>SbNF-YB3</i>	α helix TMspans				GPI-SOM
<i>SbNF-YB4</i>			✓	✓	
<i>SbNF-YB5</i>		✓	✓		
<i>SbNF-YB6</i>	α helix TMspans			✓	GPI-SOM
<i>SbNF-YB7</i>					
<i>SbNF-YB8</i>		✓		✓	
<i>SbNF-YB9</i>		✓		✓	
<i>SbNF-YB10</i>		✓			
<i>SbNF-YB11</i>		✓			
<i>SbNF-YC1</i>					
<i>SbNF-YC2</i>	α helix TMspans	✓			
<i>SbNF-YC3</i>		✓			
<i>SbNF-YC4</i>	α helix TMspans	✓	✓		
<i>SbNF-YC5</i>					
<i>SbNF-YC6</i>					
<i>SbNF-YC7</i>	α helix TMspans	✓	✓		
<i>SbNF-YC8</i>					
<i>SbNF-YC9</i>					
<i>SbNF-YC10</i>	α helix TMspans	✓	✓		
<i>SbNF-YC11</i>					
<i>SbNF-YC12</i>					
<i>SbNF-YC13</i>	α helix TMspans	✓			
<i>SbNF-YC14</i>					

SbNF-YC14, *SbNF-YC1* and *SbNF-YC11* were observed on chromosome seven while chromosomes five and eight have two genes each. The distribution of *SbNF-Y* genes on 10 chromosomes of sorghum is shown in Fig. 1. The variability of *NF-Y* gene density on chromosomes of different crops have been reported like *PvNF-Y* genes were distributed on 10 out of the 11 common bean chromosomes (Ripodas et al. 2014), *BdNF-YA* and C subunits were absent on chromosome five, and no *BdNF-YB* on chromosome four and five (Cao et al. 2011), *OsHAP* gene were dispersed on 11 chromosome out of the 12 rice chromosomes (Thirumurugan et al. 2008).

The exon-intron organization provides an insight into evolutionary relationships among genes or organisms and needs

substantial investigation (Koralewski and Krutovsky 2011). The gene structure analysis could provide possible mechanisms of structural evolution of *NF-Y* genes in sorghum and thus an attempt was made to compare the exon-intron structures of all the 33 annotated *NF-Y* genes. The putative gene structure of these *SbNF-Y* gene families is shown in Fig. 2. Predominance of intronless *SbNF-Y* genes was evident based on the fact that a total of 18 out of 33 *SbNF-Y* genes lacked intron. In case of *SbNF-YA* genes, there exists great variability in terms of introns varying from intronless *SbNF-YA1* to a maximum of five introns for *SbNF-YA5*. For *SbNF-YB* gene family seven out of 11 were found to be intronless while a maximum of five introns was observed for *SbNF-YB10*.

Fig. 1 Chromosomal localization of 33 *SbNF-Y* genes on ten chromosomes of *Sorghum bicolor*. The chromosome numbers and its size are indicated at the bottom of each bar. Sizes of chromosomes are represented on using vertical scale



Similarly for *SbNF-YC* gene family, 10 out of 14 were intronless and *SbNF-YC4* possessed a maximum of 5 introns. Thus *SbNF-Y* gene family shows great variability in terms of intron/exon distribution pattern. The relevance of the gene structure analysis is based on the fact that introns are believed to be the essential entities of eukaryotic genes as their loss or gain causes structural diversity and complexity, which might contribute to the evolution of multiple gene families like the NF-Y gene family. Evidence now exists that introns have many functions, including regulation and exon shuffling and alternative splicing (Fedorova and Fedorov 2003; Le et al. 2003). Further introns might be associated with enhancement of intragenic recombination and moderating the evolutionary rate of genes (Roy and Gilbert 2006).

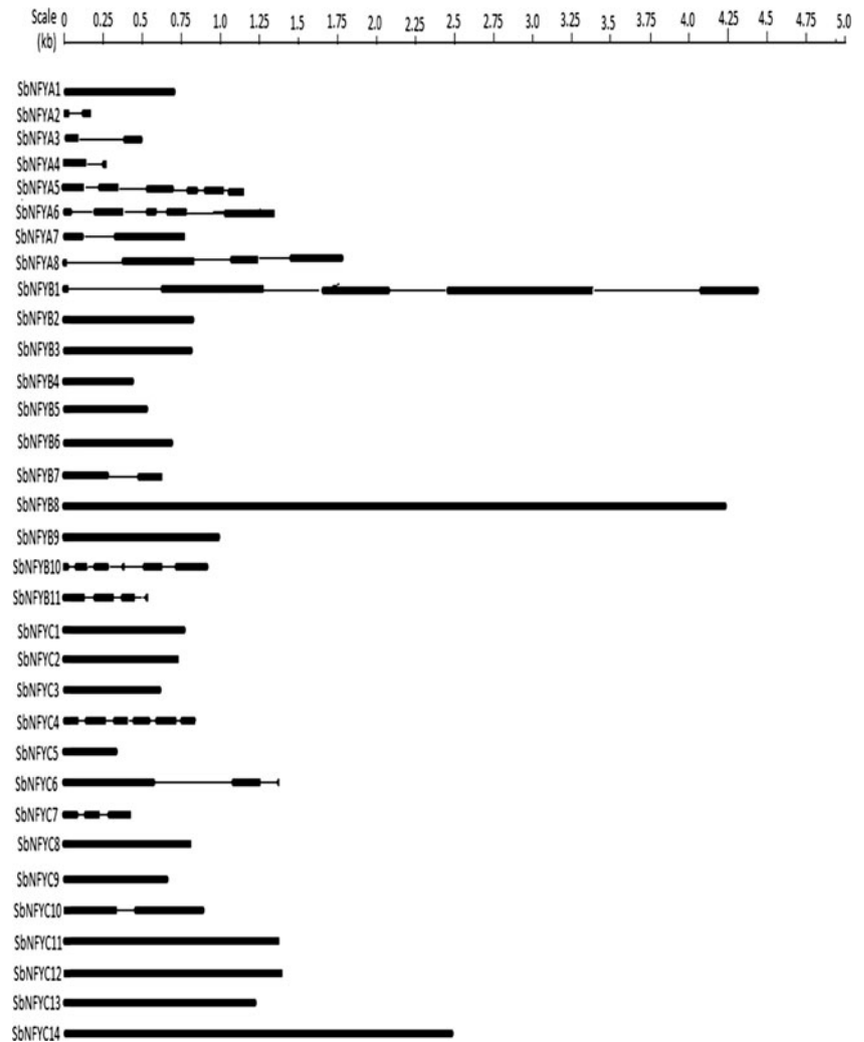
Multiple sequence alignment and phylogenetic assessment of *SbNF-Y* proteins

The 33 *SbNF-Y* proteins were aligned using ClustalX 2.0.10 revealing conserved DNA binding domains for *SbNF-YA*, *SbNF-YB* and *SbNF-YC* proteins as shown in Fig. 3. The *SbNF-Y* proteins have a highly conserved region flanked by largely non-conserved sequences as reported in *Arabidopsis* NF-Y members (Siefers et al. 2009). These conserved regions, in mammals and yeast, are found to be necessary and

sufficient for heterodimerization, heterotrimerization and DNA interactions at CCAAT sites (Kim et al. 1996; McNabb et al. 1997; Romier et al. 2003; Testa et al. 2005; Xing et al. 1993, 1994). The core region sequences of the *SbNF-YA* peptides are highly conserved in comparison to *SbNF-YB* and *SbNF-YC* (Fig. 3). Structural and functional studies suggests that NF-YA proteins are more evolutionarily constrained as compared to NF-YB or NF-YC proteins and further mutations in this pentamer region gives complete or near complete loss of NF-Y binding (Romier et al. 2003).

The NF-YA proteins, consists of two domains that are highly conserved in all higher eukaryotes studied till date (Laloum et al. 2013). The first conserved domain of 20 amino-acid as revealed by studies on yeast and mammalian system forms a alpha helix and found to be essential for the interaction with NF-YB and NF-YC (Mantovani 1999; Romier et al. 2003; Xing et al. 1993). The next neighboring domain of 21 amino-acids, separated from first domain by a conserved linker sequence, is required for specific binding to CCAAT boxes. Other than the conserved domains, plant NF-YA proteins are variable in size and structure as evident for sorghum NF-YA proteins. These conserved domains are placed in the C-terminus of the protein in mammals whereas they are present more centrally in plant NF-YAs. The three histidine (H) and three arginine (R) residues known to be essentially required for

Fig. 2 Intron/exon structure of *SbNF-Y* genes, *black bars* represents the exon while the *lines* represent the intron. The size of exons and introns can be estimated using the horizontal scale



DNA binding as in the case of mammalian NF-YA (Xing et al. 1993) was also observed to be highly conserved in all the eight SbNF-YA proteins (Fig. 3a). Similarly in *Arabidopsis*, rice, *Brachypodium* and wheat these conserved residues were observed (Cao et al. 2011; Siefers et al. 2009; Stephenson et al. 2007; Thirumurugan et al. 2008). The NF-YB and NF-YC subunits in contrast are considered to make DNA contact with the neighboring region of CCAAT sequences and are loosely conserved in comparison to NF-YA (Romier et al. 2003).

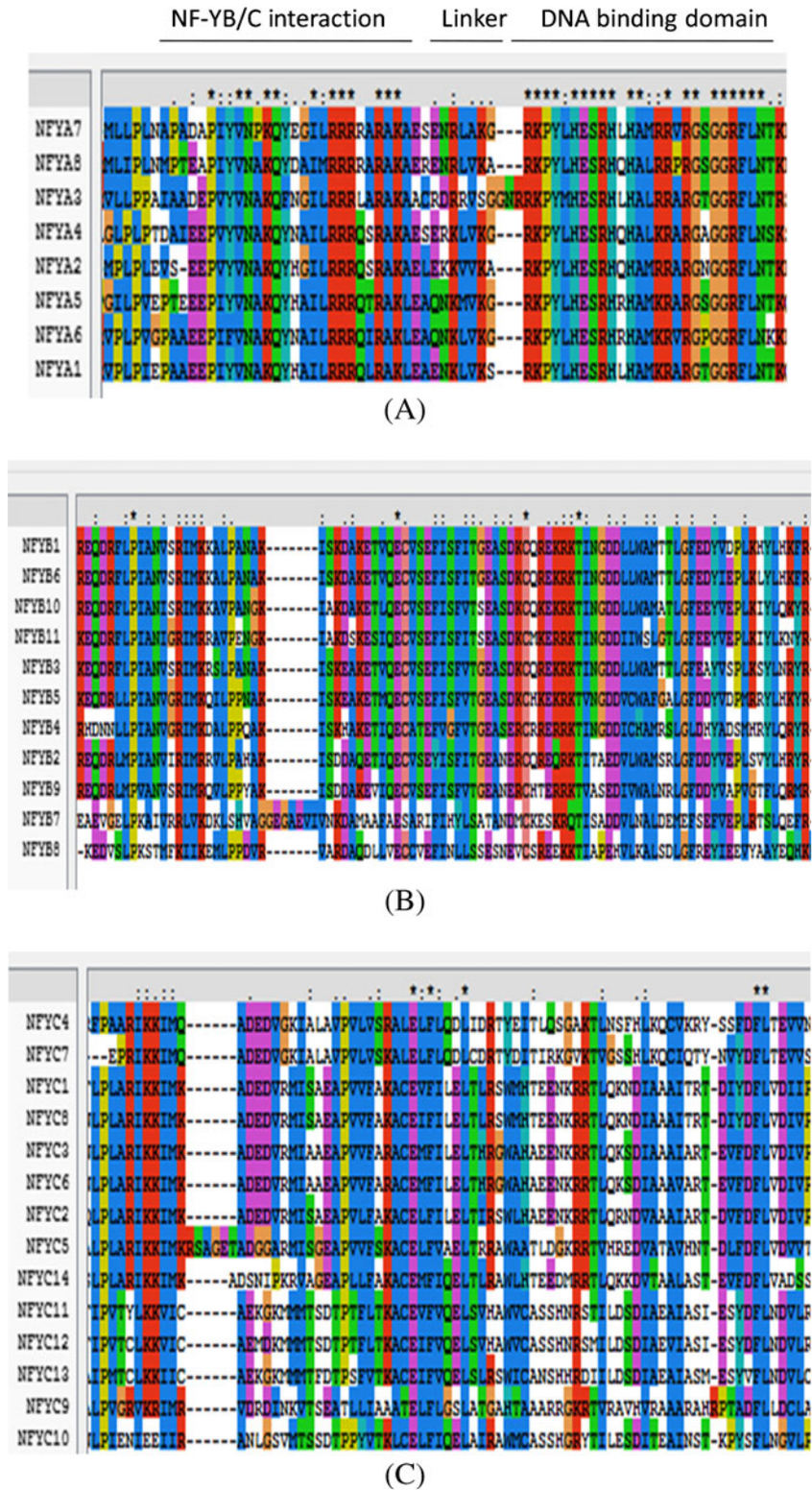
The evolutionary relationship among different SbNF-Y proteins was assessed by subjecting the deduced amino-acid sequences of 33 SbNF-Y proteins for multiple sequence alignment followed by phylogenetic tree construction. The phylogenetic tree revealed three major clusters: cluster I, II and III for SbNF-YA, SbNF-YB and SbNF-YC respectively (Fig. 4). The closely related members within the clusters showed almost similar amino-acid sequence lengths, isoelectric point and molecular weights as listed in Table 1. The result revealed that SbNF-YB8 was much closer to SbNF-YA proteins, while other SbNF-YB proteins showed close relationship with

SbNF-YC proteins. From this phylogenetic tree we could presume that SbNF-YB has been evolved from SbNF-YA gene and are closely related to SbNF-YC genes.

Assessment of gene duplication time for SbNF-Y proteins

The phylogenetic tree constructed for SbNF-YA, SbNF-YB and SbNF-YC genes identified a total of eight pairs of paralogous genes in the terminal nodes as supported by strong bootstrap values. The presence of these paralogous genes on different chromosomes (Fig. 1) clearly suggests the possibility of segmental duplication event, which might be associated with the expansion of the NF-Y gene family in sorghum. Further, study has been done to predict the immediate ancestral protein sequences of these eight pairs of paralogous genes and the identified protein motif structure of the duplicated genes (Fig. 5 and Supplementary Table 1). The ratio of non-synonymous (Ka) to synonymous (Ks) nucleotide substitution rates denotes the selective pressures on genes. This can be used to identify combinations of genes in the phylogenetic

Fig. 3 *S. bicolor* NF-Y protein sequence alignments. Sequence alignments were obtained using Clustal X 2.0.10 program. Sequences were identified and in Genomic databases and renamed as SbNF-YA 1–8 (a), SbNF-YB 1–11 (b) and SbNF-YC 1–14 (c), respectively. Dashes indicate gap in the sequence. Strongly conserved residues are indicated above the alignment with asterisks, colon indicate the residues variation occurring within the strongly conserved groups and dots indicates the residues variation occurring within weaker conserved residue groups

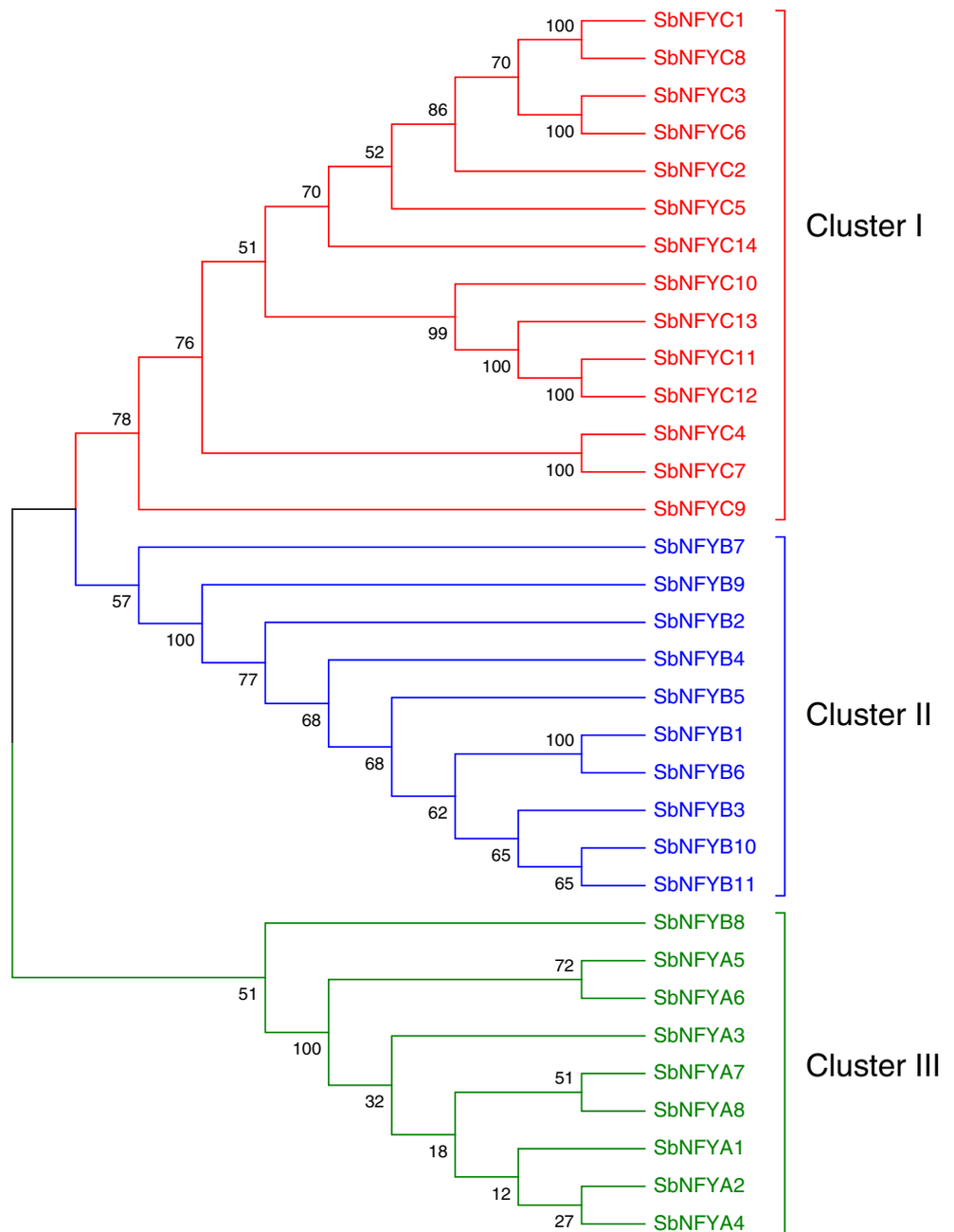


trees, where encoded proteins might be subjected to functional alteration. This ratio when greater than 1 indicates positive selective pressure, whereas when it is around 1, it indicates either neutral evolution at the protein level or an averaging of sites under positive and negative selective pressures. A ratio

less than 1 indicates the possibility of selective pressures associated with conserved protein sequences (Hurst 2002; Yang and Bielawski 2000).

The *Ks* and *Ka* value of paralogous sequences calculated using DnaSP software is shown in the Table 3. The

Fig. 4 The combined phylogenetic tree of 8 Sb NF-YA, 11 Sb NF-YB and 14 Sb NF-YC proteins constructed by neighbor-joining methods using MEGA 6.0 with a bootstrap of 500 reiterations



paralogous genes namely SbNF-YA5/SbNF-YA6 and SbNF-YB11/SbNF-YB12 reveals the possibility of neutral mutation owing to almost equal K_a and K_s values while paralogous genes *SbNF-YC7/SbNF-YC8*; *SbNF-YB10/SbNF-YB11*; *SbNF-YC1/SbNF-YC8*; *SbNF-YC3/SbNF-YC6*; *SbNF-YB10/SbNF-YB11*; *SbNF-YC3/SbNF-YC6* and *SbNF-YC4/SbNF-YC7* showed positive mutation. One of the paralogous genes *SbNF-YB1/SbNF-YB6* showed negative (purifying) selection as K_a value is greater than K_s . Therefore, statistics of the two variables evaluated for different paralogous genes from different evolutionary lineages could provide a powerful tool for quantifying molecular evolution. The date of duplication

events were also estimated approximately using the K_s value. The segmental duplications of the *SbNF-Y* genes is found to be originated from 10.1 Mya (million years ago, $K_s = 0.1324$) to 155.3 Mya ($K_s = 2.0192$), with an average mean of 59.7 Mya. This data suggests that the sorghum *NF-Y* family gene expansion is due to segmental duplication events, and further these *SbNF-Y* genes retained their function after duplication.

Motif analysis

The distributions of conserved motifs were assessed for SbNF-Y proteins by means of MEME software (Fig. 6). The



Fig. 5 Distribution of motifs highlighting gene duplicates and their ancestors

Table 3 The gene duplication time identified in different paralogous pairs of *NF-Y* gene families. The K_a represents the number of non-synonymous substitution per non-synonymous site while K_s is the

number of synonymous substitution per synonymous site and K_a/K_s represents the ratios of non-synonymous (K_a) versus synonymous (K_s) mutations

Paralogous genes	Location on chromosome	Duplication event	K_s value	K_a value	K_a/K_s	Date (Millions years ago)
<i>SbNF-YA5</i> / <i>SbNF-YA6</i>	Chr2/Chr1	Segmental	0.5934	0.4467	0.7527	45.6
<i>SbNF-YA7</i> / <i>SbNF-YA8</i>	Chr1/Chr8	Segmental	0.9666	0.3352	0.3467	74.3
<i>SbNF-YB1</i> / <i>SbNF-YB6</i>	Chr1/Chr2	Segmental	0.1731	0.3741	2.1611	13.3
<i>SbNF-YB10</i> / <i>SbNF-YB11</i>	Chr3/Chr9	Segmental	1.0555	0.2092	0.1981	81.1
<i>SbNF-YC1</i> / <i>SbNF-YC8</i>	Chr5/Chr10	Segmental	0.9015	0.1409	0.1562	69.3
<i>SbNF-YC3</i> / <i>SbNF-YC6</i>	Chr2/Chr7	Segmental	0.3720	0.1103	0.2965	28.6
<i>SbNF-YC4</i> / <i>SbNF-YC7</i>	Chr5/Chr9	Segmental	2.0192	0.3094	0.1532	155.3
<i>SbNF-YC11</i> / <i>SbNF-YC12</i>	Chr7/Chr8	Segmental	0.1324	0.1169	0.8829	10.1

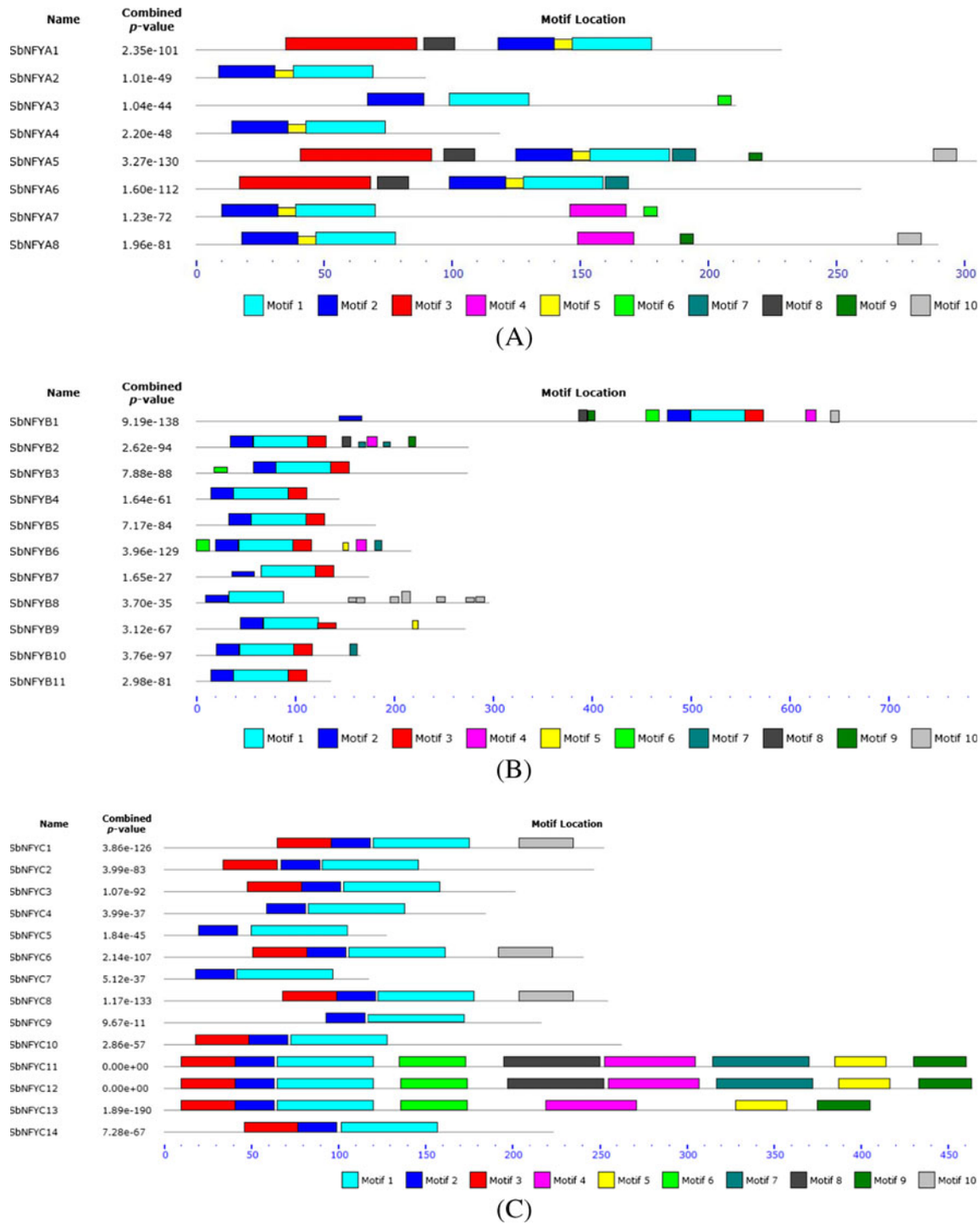


Fig. 6 Motif distribution among SbNF-YA (a); SbNF-YB (b) and SbNF-YC (c) proteins respectively using MEME ver. 4.4.0

variation in motif distributions among NF-YA, NF-YB and NF-YC proteins of sorghum is listed in the Supplementary Table 2. In case of SbNF-YA sequences, motif 1 and 2 were ubiquitously present while rest of the 8 motifs showed variability in terms of distribution among the SbNF-YA proteins. Similarly Sb NF-YB proteins revealed uniform presence of highly conserved motif 1 among all the 11 SbNF-

YA proteins. A motif of 22 amino-acid and 18 amino-acid sequence was observed outside the conserved core region of SbNF-YB. The motif analysis of NF-YC proteins of sorghum also revealed a highly conserved motif 2 observed in 13 out of 14 SbNF-YC proteins. These conserved motifs reflects typical diagnostic features for different subunits of NF-Y proteins in general and hence provides confirmatory

identification of SbNF-Y proteins from the sorghum genome.

Comprehensive phylogenetic analysis of NF-Y proteins of sorghum, rice and *Arabidopsis*

A comprehensive phylogenetic tree comprising of 33 NF-Y proteins of sorghum along with 36 *Arabidopsis* and 28 Rice NF-Y proteins was constructed using software MEGA 6.0 by NJ method with bootstrap (500 reiterations) analysis (Fig. 7). The phylogenetic tree gives an insight into the functional

attributes of identified NF-Y members of sorghum based on the availability of functionally characterized NF-Y members of *Arabidopsis* and rice. The phylogenetic tree of NF-YA members of sorghum, rice and *Arabidopsis* revealed few orthologous groups like SbNF-YA8 was placed closely with two NF-YA proteins of *Arabidopsis* namely At5g12840 (AtNF-YA1) (Wenkel et al. 2006) and At3g20910 (AtNF-YA9) (Levesque-Lemay et al. 2003). Regulation of At5g12840 has been implicated in late flowering and thus SbNF-YA8 might represent the most likely candidates for similar function in sorghum. Similarly SbNF-YB2, SbNF-

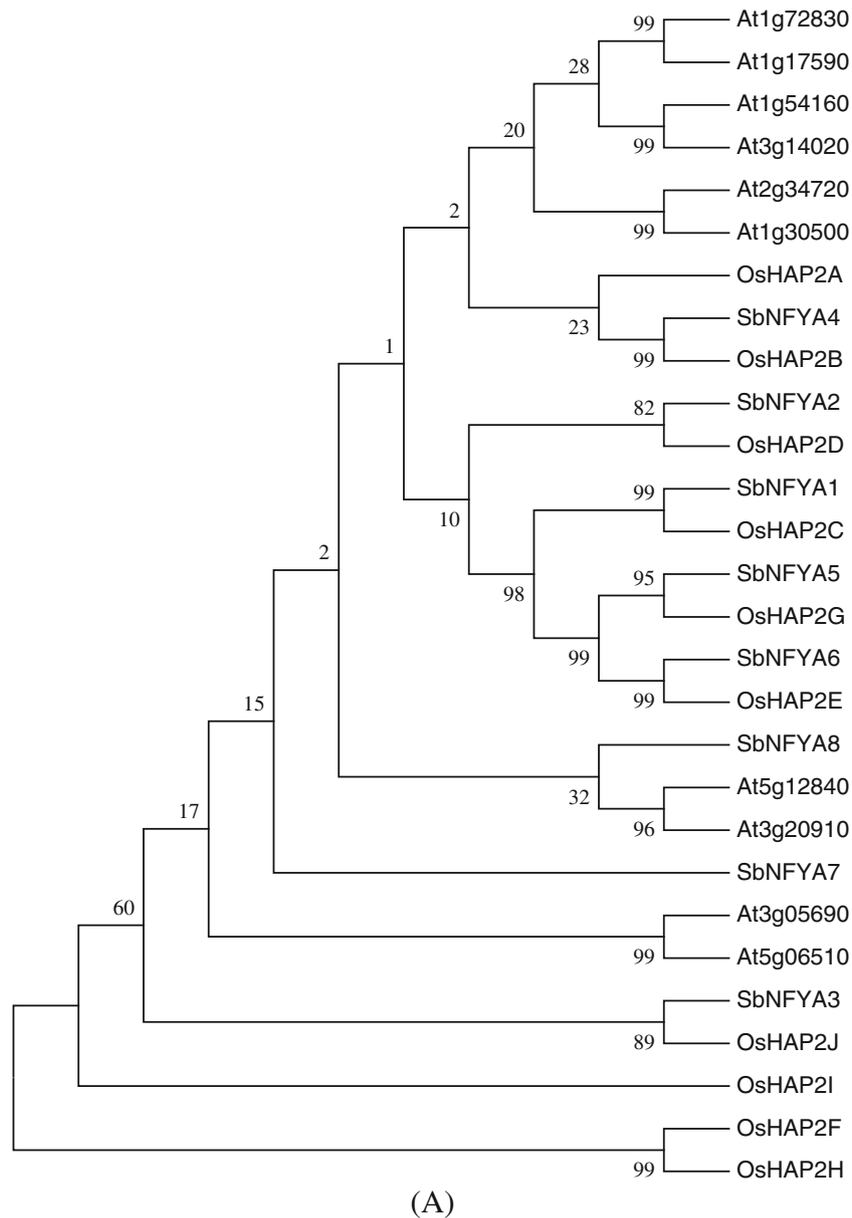
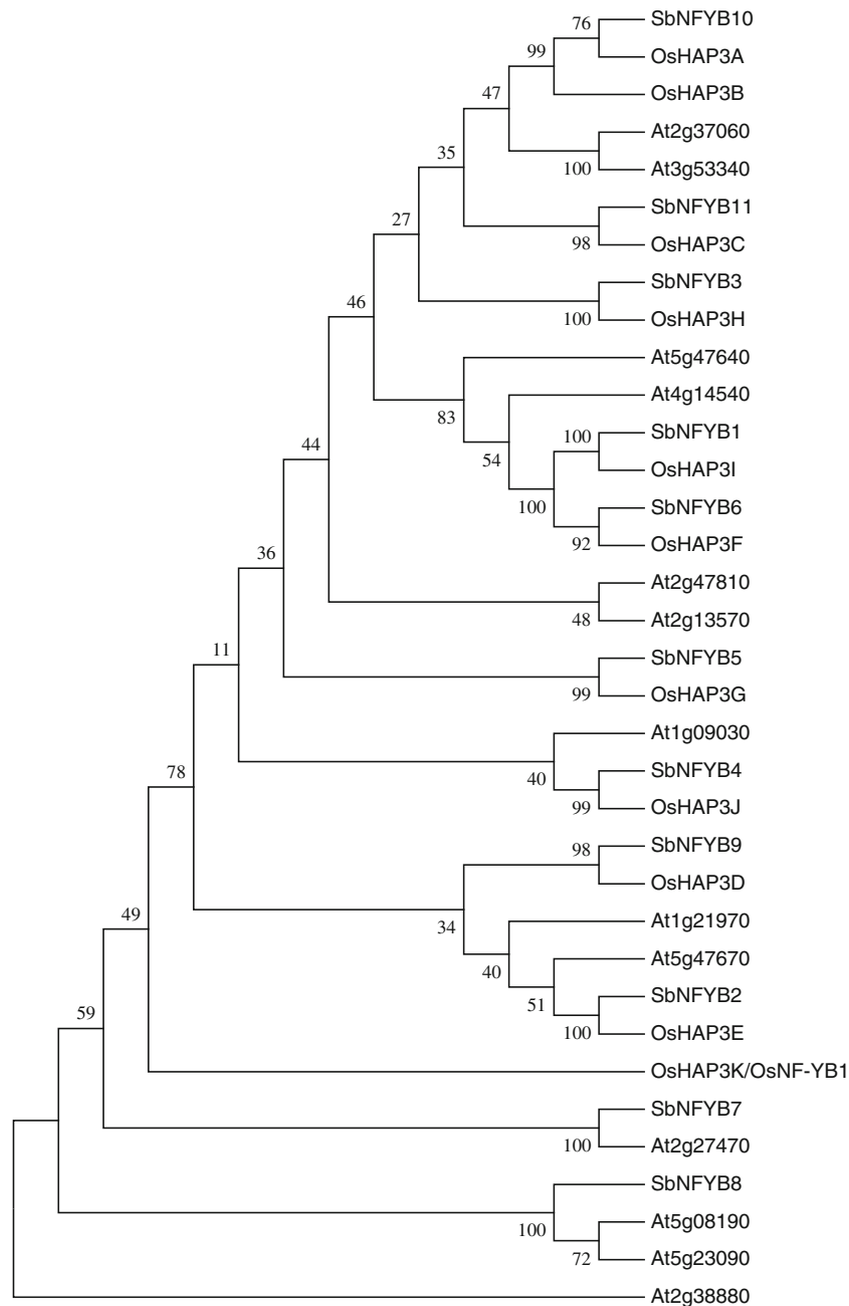


Fig. 7 Phylogenetic tree constructed for (a) a total of 28 NF-Y sequences comprising 10 *Arabidopsis thaliana*, 10 *Oryza sativa* and 8 *Sorghum bicolor* NF-YA proteins; (b) a total of 35 NF-Y sequences comprising 13 *Arabidopsis thaliana*, 11 *Oryza sativa* and 11 *Sorghum bicolor* NF-

YB proteins and (c) a total of 34 NF-Y sequences comprising 13 *Arabidopsis thaliana*, 14 *Oryza sativa* and 14 *Sorghum bicolor* NF-YC proteins using MEGA 6.0 by NJ method with 500 bootstrap reiterations



(B)

Fig. 7 (continued)

YB3, SbNF-YB10 and SbNF-YB11 showed similarity with OsHAP3E, OsHAP3H, OsHAP3A and OsHAP3C respectively. OsHAP3A and OsHAP3C are known to be important in regulation of chloroplast biogenesis (Miyoshi et al. 2003), OsHAP3H are important regulator of photoperiodic flowering (Wei et al. 2010) and OsHAP3E are associated with floral meristem development (Ito et al. 2011). Several HAP5 genes of rice showed similarity with SbNF-YC sequences. The SbNF-YC2 are closely placed with At3g48590 (AtNF-YC1) and At5g63470 (AtNF-YC4) known to associated with

regulation of flowering (Hackenberg et al. 2012) and germination (Liu and Howell 2010; Warpeha et al. 2007). Thus based on phylogenetic tree, attempts can be made to predict the putative functions of SbNF-Y members prior to subjecting it for validation by wet lab experimentations.

In-silico expression analysis under abiotic stresses

To get an insight into the functional attributes of NF-Y genes of sorghum, attempt was made to analyze in-silico expression

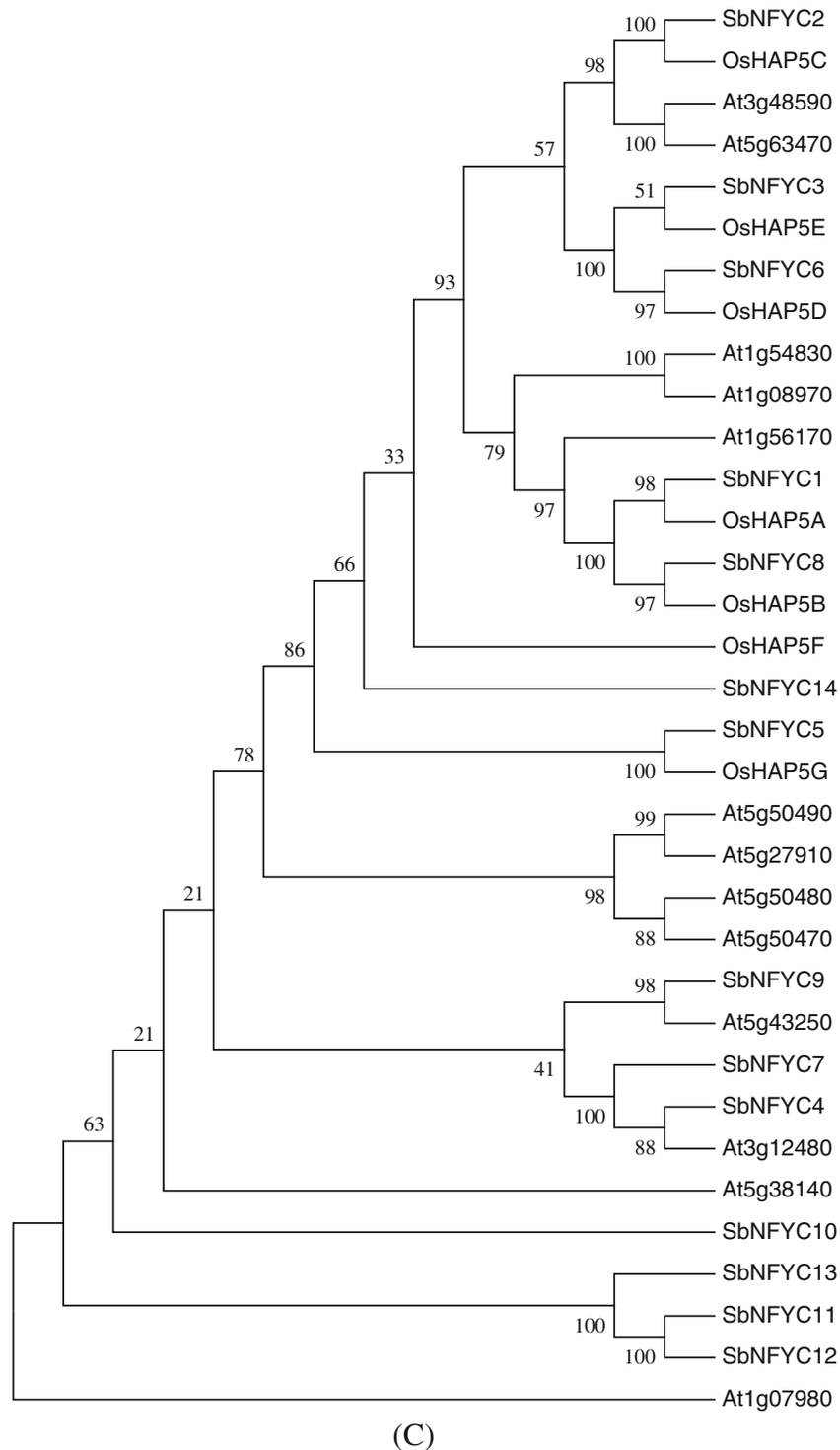


Fig. 7 (continued)

using transcriptome data of rice for abiotic stresses due to non-availability of sorghum transcriptome data. Publicly available gene expression values for abiotic stresses of rice seedling for 21 *NF-Y* genes orthologs to sorghum *NF-Y* genes were retrieved (Fig. 7) and a hierarchical clustering based heat map was constructed (Supplementary Figure). The data reveals

differential expression of *SbNF-Y* genes based on its identity with the corresponding *OsNF-Y* genes identified in the phylogenetic tree constructed. In case of *SbNF-YA* group, *SbNF-YA2*, *SbNF-YA3* and *SbNF-YA5* seems to be expressed at higher level under drought and salt stress condition based on the enhanced transcript level of the rice orthologs *OsHAP2D*,

OsHAP2J and *OsHAP2G* respectively. The *OsHAP2J* has been associated with drought stress (Jiao et al. 2009). Similarly *SbNF-YA1* and *SbNF-YA6* showed relatively higher expression value under cold stress condition as evident from the higher transcript level of the corresponding rice orthologs *OsHAP2C* and *OsHAP2E* respectively. In case of *SbNF-YB* group, *SbNF-YB3*, *SbNF-YB4* and *SbNF-YB10* seems to be potential candidate owing to comparatively higher transcript level under salt, drought and cold stress condition respectively as extrapolated from the evident higher transcript level of rice orthologs namely *OsHAP3H*, *OsHAP3J* and *OsHAP3A* respectively. The functional role of *OsHAP3A* in chloroplast biogenesis (Miyoshi et al. 2003) and *OsHAP3H* in response to photoperiodic flowering, grain yield and plant height (Wei et al. 2010) has been reported. The NF-YB genes of sorghum namely *SbNF-YB1*, *SbNF-YB2*, *SbNF-YB5*, *SbNF-YB6*, *SbNF-YB9* and *SbNF-YB11* corresponding to rice orthologs viz. *OsHAP3I*, *OsHAP3E*, *OsHAP3G*, *OsHAP3F*, *OsHAP3D* and *OsHAP3C* respectively revealed comparatively higher transcript level under heat shock condition. The functional attributes of some of the *OsHAP3* genes have been reported. The *OsHAP3C* is associated with flowering and grain yield (Wei et al. 2010), *OsHAP3E* to floral meristem development, *OsHAP3G* and *OsHAP3I* are related to embryogenesis (Ito et al. 2011). In case of *SbNF-YC* group, *SbNF-YC1* corresponding to rice ortholog *OsHAP5A* seems to be promising based on higher transcript level under drought stresses. Similarly two of the *SbNF-YC* genes namely *SbNF-YC5* and *SbNF-YC8* showed higher transcript level under cold stress condition and could be important for further validation by transgenic technology by making relevant constructs. The higher expression level of *SbNF-YC1* as evident from its corresponding rice ortholog *OsHAP5A* under drought stress could be another potential stress related NF-Y gene of sorghum. The *in-silico* expression profiling attempted in the present study provides an opportunity to target some of the identified potential *SbNF-Y* genes showing comparatively higher expression level under abiotic stresses for its confirmation by appropriate wet lab experimentations.

Conclusion

The availability of genome sequences provides an opportunity for genome wide identification and characterization of transcription factor gene family as they account for more than 5 % of genome and are known to play a significant role in gene regulation. The genome wide identification and characterization of NF-Y gene family of rice, wheat, *Arabidopsis*, *Brachypodium*, *Brassica napus*, soybean and common bean have led to functional validation of several NF-Y subunits associated with important agronomic traits. The multiple members of NF-Y subunits in plants reflect the redundancy

and differentiated functions of these proteins which need to be explored by expression profiling. Using bioinformatics tools attempts have been made in the present study to identify the putative members of NF-Y subunits of sorghum and subject it to extensive *in-silico* characterization for gene structures, motif analysis, chromosomal distribution, conserved motifs, duplication status, ancestral protein sequences and phylogenetic tree construction. The *in-silico* expression profiling based on the available rice transcriptome data for different abiotic stress conditions revealed several potential *SbNF-Y*s showing higher expression level under drought, salt, cold and heat stresses.

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