Molecular structures and functional exploration of NDA family genes respond tolerant to alkaline stress in Gossypium hirsutum L.

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Abstract

Background: The internal NAD(P)H dehydrogenase (NDA) gene family was a member of the NAD(P)H dehydrogenase (ND) gene family, mainly involved in the non-phosphorylated respiratory pathways in mitochondria and played crucial roles in response to abiotic stress.

Methods: The whole genome identification, structure analysis and expression pattern of NDA gene family were conducted to analyze the NDA gene family.

Results: There were 51, 52, 26, and 24 NDA genes identified in G. hirsutum, G. barbadense, G. arboreum and G. raimondii, respectively. According to the structural characteristics of genes and traits of phylogenetic tree, we divided the NDA gene family into 8 clades. Gene structure analysis showed that the NDA gene family was relatively conservative. The four Gossypium species had good collinearity, and segmental duplication played an important role in the evolution of the NDA gene family. Analysis of cis-elements showed that most GhNDA genes contained cis-elements related to light response and plant hormones (ABA, MeJA and GA). The analysis of the expression patterns of GhNDA genes under different alkaline stress showed that GhNDA genes were actively involved in the response to alkaline stress, possibly through different molecular mechanisms. By analyzing the existing RNA-Seq data after alkaline stress, it was found that an NDA family gene GhNDA32 was expressed, and then the GhNDA32 was silenced by virus-induced gene silencing (VIGS). By observing the phenotype, we found that the wilting degree of silenced plants was much higher than that of the control plant after alkaline treatment, suggesting that GhNDA32 gene was involved in the response to alkaline stress.

Conclusions: In this study, GhNDAs participated in response to alkaline stress, especially NaHCO₃ stress. It was of great significance for the future research on the molecular mechanism of NDA gene family in responding to abiotic stresses.

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Introduction
With the continuous increase in the area of salt-alkaline land worldwide, salt-alkaline stress had become an important abiotic stress that restricted plant growth and development. Salt-alkaline stress was divided into salt stress and alkaline stress. Salt stress was mainly caused by neutral salts such as NaCl and Na$_2$SO$_4$, and alkaline stress was usually caused by Na$_2$CO$_3$ and NaHCO$_3$ [1]. Previous studies found that alkaline stress was more harmful than salt stress [2]. At present, little was known about the mechanism of crop tolerance to alkaline stress. Therefore, it was very important to explore the mechanism of alkaline stress.

Mitochondria were an important place for energy production in plants, previous research found that plant mitochondria contained a complex electron transport chain, in which there was an alternative respiratory pathway (AP), and the difference of the AP was that it was not linked to energy conservation [3]. It was well known that most of the energy produced by plants come from mitochondria, at the meantime, plant mitochondria were also a source of reactive oxygen species (ROS), studies showed that the presence of the AP could prevent over-reduction of the electron transport chain (ETC), thereby minimizing the generation of ROS and protecting plants from oxidative damage [4, 5, 6]. In addition, AP actively participated in the response to abiotic stress, such as temperature, nutrients, heavy metals, high light, drought and oxidative stress [7].

AP contained two important components, the alternative oxidase (AOX) and type II NAD(P)H dehydrogenases (NDs). Expression of the AOX genes was highly responsive to abiotic and biotic stress, as well as dysfunctions in respiratory metabolism [8], and ROS may be an important signal controlling AOX expression [9]. In plants, fungi and some certain bacteria, there were many type II NAD(P)H dehydrogenases (NDs), which were small single proteins attached to either side of the inner membrane [10]. In potato, St-nda1 gene was shown to be tightly bound to the inner surface of the mitochondrial inner membrane, while St-ndb1 was loosely bound to the outer mitochondrial membrane [11, 12]. In Arabidopsis, the NDs could be divided into three subgroups, AtNDA (1, 2), AtNDB (1-4), and AtNDC1. AtNDA1, 2 and AtNDC1 were identified as encoding internal NAD(P)H dehydrogenases, whereas AtNDB1-4 encode external NAD(P)H dehydrogenases [13, 14].

NDAs, belonged to the subclass of NDs, played a vital role in response to abiotic stress. Studies found that they were involved in the response to light, cold and Pi stress. The expression of nda1 in Arabidopsis was dependent on light and circadian regulation, which played an important role in the photosynthesis and other respiratory NADH oxidation [14]. In potato, the immunodetectable NDA1 protein abundance and the internal rotenone insensitive NADH dehydrogenase activity were all affected by light, suggesting that the nda1 gene was involved in photosynthetically associated processes, most likely photorespiration [15]. After cold treatment, potato leaves could induce the production of oxidase and ROS, and the expression of NDA1 gene decreased, which may be related to the redox function of NDA expression [16]. Previous research found that low-level expression of NDAs could lead to the delayed growth and restrict citric acid cycle reactions but apparently had no effect on photosynthesis [17]. In Arabidopsis, a non-phosphorylating mitochondrial electron transport chain consisting of NDA2, NDB2 and AOX was synthesized to maintain respiratory electron flow through the mitochondrial electron transport chain during Pi stress [18]. However, whether NDA family genes responded to alkaline stress in cotton had not been reported, so it was meaningful to study it.

Cotton is an important fiber crop and a model crop for studying polyploidy and evolution [19]. Gossypium hirsutum (G. hirsutum) and Gossypium barbadense (G. barbadense) are the two most widely cultivated allotetraploid cotton species in the world, which are formed by inter-genomic hybridization of Gossypium arboreum (G. arboreum) and Gossypium raimondii (G. raimondii) [20]. Cotton is also a pioneer crop in salt-alkaline land, with a certain degree of salt-alkali tolerance, and is a common crop for us to study salt-alkaline stress [21]. Exploring the alkaline stress response genes will help promote the cotton's tolerance to alkaline stress. From the RNA-Seq data, an NDA gene GhNDA32 was found to respond to alkaline stress, so it is necessary to analyze its phylogenetic relationship and study the alkaline stress tolerance of the family. This study aimed to explore gene structure, evolutionary relationship, expression pattern, and cis-acting elements of NDA family. An NDA gene belonged to clade a, GhNDA32, was isolated and characterized, and we performed a preliminary analysis of the expression pattern of GhNDA32. This study provided potential candidate genes for cotton gene functional verification.
Results

Identification of NDA proteins

To identify the NDA gene family in cotton, a local blast on the proteome of the *G. hirsutum*, *G. barbadense*, *G. arboreum*, and *G. raimondii* was performed, and genome with the hidden Markov model (HMM) profile of PF07992 was used as a query condition. Redundant sequences were detected and deleted by manual, 51, 52, 26, 24 genes were identified in *G. hirsutum* (GhNDAs), *G. barbadense* (GbNDAs), *G. arboreum* (GaNDAs), and *G. raimondii* (GrNDAs), respectively (Fig. 1). Statistical analysis of the number of genes in four *Gossypium* species showed that the number of NDA genes in two tetraploid cotton species (*G. hirsutum* and *G. barbadense*) were almost double of diploid cotton species (*G. arboreum* and *G. raimondii*). To understand the genes of NDA family more conveniently, genes were renamed according to the position of genes on chromosomes. *GhNDA1*-*GhNDA51* was assigned for *G. hirsutum*, *GbNDA1*-*GbNDA52* for *G. barbadense*, *GaNDA1*-*GaNDA26* for *G. arboreum* and *GrNDA1*-*GrNDA24* for *G. raimondii*, other species were also renamed according to this rule (Additional file 2: Table S2). Subsequently, the physical properties of the GhNDAs genes were analyzed and predicted, including protein length, protein molecular weight (MWs), isoelectric point (pI), protein hydrophilicity and hydrophobicity analysis and subcellular location (Additional file 3: Table S3). For *G. hirsutum*, all of the 51 genes encoded proteins ranging from 357 (*GhNDA35*) to 2209 (*GhNDA48*) amino acids, with an average of 603.922 amino acids. The MWs varied from 38.742 (*GhNDA35*) kDa to 242.628 (*GhNDA22*) kDa with an average of 66.107 kDa and pI varied from 5.553 (*GhNDA36*) to 10.024 (*GhNDA35*) with a mean of 7.909. The values of the grand average of hydropathy were all negative, which proved that the proteins of NDA family were hydrophilic. The prediction of subcellular localization showed that there were 20 genes located in the mitochondria, 17 in cytoplasmic and 14 in chloroplast.

Phylogenetic analysis of NDA genes

To understand the evolutionary relationship of NDA genes among four *Gossypium* species and other plant species, multiple sequence alignment of 247 protein sequences (including 51 in *G. hirsutum*, 52 in *G. barbadense*, 26 in *G. arboreum*, 24 in *G. raimondii*, 23 in *Arabidopsis thaliana* (*A. thaliana*), 12 in *Solanum tuberosum* (*S. tubersum*), 19 in *Glycine max* (*G. max*), 11 in *Oryza sativa* (*O. sativa*), 8 in *Populus trichocarpa* (*P. trichocarpa*), 5 in *Theobroma cacao* (*T. cacao*), 9 in *Vitisvinifera Genoscope* (*V. vinifera*) and 7 in *Zea mays* (*Z. mays*)) was performed and a phylogenetic tree was constructed using MEGA 7 software based on the neighbor-joining (NJ) method. The phylogenetic tree was subsequently decorated using EvolView (https://www.evolgenius.info/evolvview) (Fig. 2B). According to previous method of dividing the phylogenetic tree [22], we divided NDA genes into 8 clades based on the sequence similarity, tree topology and structural characteristics in each sequence. The clade a had the most extensive genes,
of which 14 were *G. hirsutum*, 14 were *G. barbadense*, 7 were *G. arboreum*, and 6 were *G. raimondii*, 7 were *A. thaliana*, 3 were *S. tubersum*, 2 were *O. sativa*, 2 were *G. max*, 1 was *Z. mays*, 1 was *V. vinifera*, 1 was *T. cacao* and 1 was *P. trichocarpa*, respectively. All species had gene pairs derived from the same node, demonstrating that the NDA genes in all species had experienced gene duplication events that causing the expansion of the NDA gene family. The number of genes in *G. barbadense* and *G. hirsutum* was much higher than that of other species, indicating that the NDA gene family in two allotetraploid species showed a large-scale expansion during the evolution process. These results indicated that gene duplication was the main reason for the expansion of the NDA gene family in cotton.

In addition, to study the relationship between the common ancestors of diploid cotton (*G. arboreum* and *G. raimondii*) and allotetraploid cotton (*G. hirsutum* and *G. barbadense*), NJ tree of four *Gossypium* species was constructed (Fig. 2A). With reference to the classification of subgroups, we found that the NDA genes of the four *Gossypium* species were distributed in each subgroup, and each branch contained proteins from diploid and allotetraploid *Gossypium* species and the NDA genes in tetraploid cotton were almost twice as many as diploid cotton in each subgroup.

**Chromosomal location of four *Gossypium* species**

To study the chromosomal distribution of NDA genes in four *Gossypium* species, the physical location of these genes on chromosomes was drew. 149 out of 153 NDA genes were distributed to their specific chromosomes, while the remaining only 4 NDA genes, *GhNDA20*, *GhNDA23*, *GhNDA49* and *GaNDA26*, were not located on any one chromosome because they were located on scaffold (Fig. 3). This result showed that the genetic evolution process of NDA genes was mature and stable. There was no *GhNDA* gene on Chr8 of *G. hirsutum* At sub-genome (GhAt), Chr5 of *G. hirsutum* Dt sub-genome (GhDt), Chr1, 5, 8 of *G. arboreum*, Chr4 of *G. raimondii*, Chr5 of *G. barbadense* (Fig. 3A, B, E, F), which may be related to the chromosome deletion of *G. hirsutum* or the translocation of large fragments during the evolution. The distribution of NDA genes on 13 chromosomes of different cotton species was uneven, and the number of each chromosome did not show a significant positive correlation with its length.

Among 52 identified NDA genes of *G. hirsutum*, 3 genes were located on scaffold (Fig. 3A, B). Chr9 in GhAt had most NDA genes, with a total of 4 genes, and Chr9 in GhDt had most NDA genes, with a total of 5 genes (Fig. 4). In *G. barbadense*, Chr9 in GbAt and GbDt had most NDA genes, with a total of 5 genes, which was
Fig. 3 Chromosomal location of NDA genes from four *Gossypium* species. **A** Chromosomal location of NDA genes on chromosomes in *G. hirsutum* At sub-genome (GhAt), **B** Chromosomal location of NDA genes on chromosomes in *G. hirsutum* Dt sub-genome (GhDt), **C** Chromosomal location of NDA genes on chromosomes in *G. barbadense* At sub-genome (GbAt), **D** Chromosomal location of NDA genes on chromosomes in *G. barbadense* Dt sub-genome (GbDt), **E** Chromosomal location of NDA genes on chromosomes in *G. arboreum* (Ga), **F** Chromosomal location of NDA genes on chromosomes in *G. raimondii* (Gr) genes. The gene ID on the right side of each chromosome correspond to the approximate locations of each NDA gene. The scale of the genome size was given on the left. Red lines represented tandem gene duplications.
Fig. 4 Number of NDA genes in each chromosome
similar to the situation of *G. hirsutum* (Fig. 4). Among 26 identified NDA genes of *G. arboreum*, 25 genes were distributed on 13 chromosomes while one gene was found at scaffold (Fig. 3E). In *G. arboreum*, Chr9 had most NDA genes with a total of 5 genes while 5 genes in Chr6 in *G. raimondii* (Fig. 3E, F).

**Gene structure and motif composition analysis**

To further examine the structural characteristics of the NDA genes in *G. hirsutum*, the conserved motifs of NDA proteins were predicted by using the online website MEME (http://meme-suite.org/). 10 conserved motifs were found (named motif 1 to motif 10) and the results were represented in schematic diagrams (Fig. 5B). The number of motifs was different in each protein, varying from 3 to 9. NDA members within the same clade were found to have a similar motif composition, which was conspicuous in class a, b, c, h, f. All the proteins belonged to the h and f subgroups contained motif 8, but the other clades did not have motif 8. Each clade had motif 1, and only clade a had the motif 4, while the other clade did not have, which may be the reason that clade a had obtained the special motif 4 through evolution or other clade had lacked specific conserved motifs during evolution. The different compositions of the motifs may represent the diversity of function. The clade a, b, c, h, f, h showed the similar motif arrangements, indicating that the protein was conserved within a specific subgroup. However, the functions of these conserved motifs remained to be further verified.

The diversity of gene structure was mainly influenced by the evolution of multigene families [23]. To further explore the diversity of NDA gene's structure, the characteristics of exon-intron structures were analyzed. As can be seen from Fig. 5C, NDA genes of the same subfamilies had similar intron-exon arrangement, and different subfamilies displayed variation in exon-intron structures. The structures of NDA genes can be divided into two types, with intron less and multiple-exon. In *G. hirsutum*, the number of exons of all NDA genes varied from 1 to 22. Clade b had the least exon number (4), while Clade d had the most exon number (22). It was noting that one gene *GhNDA24* belonged to clade h had only one exon, no intron.

**Fig. 5**  Phylogenetic analysis, conserved motifs, and gene structures of *GhNDA* genes. A  Phylogenetic tree of *GhNDA* genes, the 8 major phylogenetic subgroups, designated as a to g, are marked with different colored backgrounds; B  Conserved motifs of *GhNDA* genes; C  Gene structures of *GhNDA* genes. Green boxes indicated exons; black lines indicated introns
Gene duplication and collinearity analysis
In general, plants had a higher rate of gene replication than other eukaryotes. Whole genome duplication, segmental duplication, and tandem duplication were considered to be the main causes of expansion in plant gene family [24]. Segmental duplications in the genome region may result in the expansion of the gene family [25]. To investigate the evolutionary process of NDA genes, the gene duplication pattern of four Gossypium species was performed using MCScanX. In this study, a total of 361 gene pairs were identified as whole genome duplication, 2 gene pairs were the tandem duplication and 102 were the segmental duplication (Fig. 6). Among the 2 tandem repeat gene pairs, one was GaNDA6/7, the other was GrNDA17/18 (Fig. 3E, F). The collinearity gene pairs between G. hirsutum and G. barbadense were the most
among the 10 groups, with 80 gene pairs, while *G. raimondii* and *G. raimondii* had the least, with only 3 gene pairs, which was in line with the comparison of the number of diploid and tetraploid genes. From these results, we presumed that whole genome duplication and segmental duplication played an important role in the evolution of NDA gene family.

**Calculation of non-synonymous (Ka) to synonymous (Ks) substitution rates during evolution**

To study the selection pressure of duplicated gene pairs in the evolutionary process, Ka, Ks, and Ka/Ks of 416 homologous gene pairs from 10 combinations of four *Gossypium* species were determined, including Ga-Ga, Ga-Gb, Ga-Gr, Gb-Gb, Gb-Gr, Gh-Ga, Gh-Gb, Gh-Gh, Gh-Gr and Gr-Gr (Additional file 4: Table S4). Among them, 2 (0.48%) duplicated gene pairs with Ka/Ks ratio > 1, which occurred in Gh-Ga and Gh-Gb, respectively, indicating that these genes may experience relatively rapid evolution and undergo the positive selection pressure. 414 (99.52%) duplicated gene pairs with Ka/Ks ratio < 1, exhibiting pure selection (Fig. 7).

**Gene ontology (GO) annotation analysis of GhNDA**

To further study the functions of GhNDA, three major categories, including biological process, cellular component, and molecular function, were analyzed via CottonFGD (Fig. 8). According to GO analysis, we found that the molecular functions were mainly involved in oxidoreductase activity (51), flavin adenine dinucleotide binding (46) and NADP binding (6). Biological processes were mainly in the following three aspects, oxidation-reduction process (51), cell redox homeostasis (14) and glutathione metabolic process (5). The cellular components were mainly in cytoplasm (16).

**Analysis of promoters and expression profiles of GhNDA genes under alkaline stress**

*Cis*-acting elements played vital roles in the response to abiotic stress [26]. Some plant hormones such as ethylene (ET), abscisic acid (ABA), methyl jasmonate (MeJA) and gibberellin (GA) were necessary for plants to adapt

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**Fig. 7** Analysis of non-synonymous (Ka) to synonymous (Ks) ratio. **A** Prediction of number of duplicated gene pairs involved in different combinations from four *Gossypium* species, **B** Ka and Ks divergence values for (Gh-Gh), (Gb-Gb), (Ga-Ga), (Gr-Gr), (Gh-Gb), (Ga-Gr), (Ga-Gh), (Ga-Gb), (Gr-Gh) and (Gr-Gb) are shown in circular chart. Different colors represent Ka/Ks gene pairs of 10 groups.

**Fig. 8** Gene ontology (GO) analysis of GhNDA genes.
to abiotic stress, and some transcription factors could combine with hormone-related cis-acting elements to regulate the expression of genes [27, 28]. To identify putative cis-acting elements in the NDA genes, the region of 2000 bp upstream of the start codon of the NDA genes was selected, and the promoter region related to stress and plant hormones was extracted. In this study, most GhNDA genes in G. hirsutum contained cis-acting elements related to plant hormones (ABA, MeJAs, GA3) and various stresses (low temperature, light and wound stress) (Fig. 9B), the cis-acting elements of NDA genes in the same subfamily were different despite had similar motifs, and therefore their functions may differ. Most of NDA genes contained cis-acting elements related to plant hormones and various stresses. GhNDA5 had 11 cis-acting elements (3 were light responsiveness, 1 were abscisic acid responsiveness, 3 were MeJA responsiveness, 3 were wound responsiveness), whereas GhNDA33 and GhNDA11 had only one cis-acting element (light responsiveness). Almost all GhNDA genes were involved in MeJAs cis-responsive elements except clade g.

It was found that gene expression levels were highly correlated with gene function [29]. We detected the expression pattern of GhNDA genes under different stresses (AS, SAS) by analyzing RNA-seq data. By analyzing the heat map, we found that only four genes, GhNDA8, GhNDA14, GhNDA32 and GhNDA40, only responded to SAS stress, GhNDA1, GhNDA22, GhNDA26 and GhNDA48 only responded to AS stress, GhNDA11, GhNDA19, GhNDA25, GhNDA35 and GhNDA51 responded to both AS and SAS stress in roots (Fig. 9C). In addition, we found that most of the gene expression levels in leaves were higher than those in roots (Fig. 9C). The expression patterns of NDA genes in roots and leaves were different, which required us to use qRT-PCR to further verify the differences in gene expression.

Expression analysis of GhNDA genes under different alkaline stress

After being exposed to AS and SAS stress, qRT-PCR assays were conducted to explore the expression patterns of GhNDA genes in roots and leaves (Figs. 10, 11). 18 genes were picked up randomly from the NDA genes actively responding to SAS and AS stress for further analysis through qRT-PCR. We found that the GhNDA genes all responded to alkaline stress, although they showed different up-regulation relationships. In addition, GhNDA7, GhNDA8, GhNDA48 showed extremely significant up-regulation under AS stress in roots, GhNDA1, GhNDA22, GhNDA25, GhNDA35, GhNDA40,
Fig. 10 Expression analysis of GhNDA genes in response to different alkaline stress in roots using qRT-PCR assays. Cotton seedlings were treated with 125 mM NaHCO$_3$ for 12 h. The mean values were from three independent biological replicates. Statistical analyses were performed by Student’s t-test (*P < 0.05 and **P < 0.01).

Fig. 11 Expression analysis of GhNDA genes in response to different alkaline stress in leaves using qRT-PCR assays. Cotton seedlings were treated with 125 mM NaHCO$_3$ for 12 h. The mean values were from three independent biological replicates. Statistical analyses were performed by Student’s t-test (*P < 0.05 and **P < 0.01).
GhNDA50, GhNDA51 showed extremely significant up-regulation under AS stress in leaves. Almost all genes were responsive to SAS stress in roots except GhNDA1 and GhNDA50. At the same time, gene expression in leaves also showed a similar trend to that in roots, except GhNDA1, GhNDA25, GhNDA32, GhNDA40 and GhNDA50. Therefore, we believed that most GhNDA genes responded to both AS and SAS stress.

Cotton plants with GhNDA gene silenced by VIGS were sensitive to NaHCO₃ alkaline stress

To verify whether the GhNDA genes responded to alkaline stress, a high-expressed gene GhD06G1340.1 (GhNDA32) from transcriptome data was selected for further study. First, we tested the tissue specificity of the GhNDA32 gene, and the results showed that the relative expression in roots was the highest, followed by leaves and stems (Fig. 12A). Then we analyzed the relative expression of GhNDA32 gene at different time periods treated with alkaline stress (125 mM NaHCO₃) and it showed a trend of first increasing and then decreasing, while reached the peak at 6 h. After that, a VIGS silencing vector pYL156: GhNDA32 was constructed, when the seedlings carrying pYL156: PDS appeared albino phenotype, qRT-PCR was used to detect the expressions of GhNDA32 gene in the pYL156 plants and pYL156: GhNDA32 plants, the results showed that the expression level of GhNDA32 in the pYL156: GhNDA32 was significantly lower than that of the control pYL156 (Fig. 12D), indicating that gene silencing was successful. After treatment with 125 mM NaHCO₃, the leaves of the seedlings all appear wilting, and we found that the pYL156: GhNDA32 cotton seedlings wilted more severely than the pYL156 (Fig. 12C). Therefore, GhNDA32 was also involved in the response to alkaline stress.

Interaction network of GhNDA32 protein

According to the homologous gene in Arabidopsis, we used STRING database (https://string-db.org/) to construct an interaction network to analyze the function of GhNDA32 protein (Fig. 13). From the Fig. 13, we could conclude that NDA1 protein of Arabidopsis, which was homologous to GhNDA32 protein of G. hirsutum, could interact with AOX1A, AOX1C, AOX2, PUMP1, and CIB22 proteins. By analyzing KEGG pathway of GhNDA32 protein from the transcriptome data, we found that GhNDA32 protein was mainly involved in oxidative phosphorylation (ko00190), so we speculated that GhNDA32 protein interacted with AOX1A, AOX1C, AOX2, PUMP1 and CIB22 proteins, which could remove excess reducing energy and balance the redox poise of...
the cell, thereby improving the resistance of cotton seedlings to oxidative stress.

**Discussion**

Cotton is a major cash crop in the world and is also a powerful model for studying genome polyploidization in plants [30]. With the completion of de-novo-assembled genome of *G. hirsutum*, *G. barbadense* [31] and *G. arboreum* [32], this undoubtedly provides us with an important basis for the whole genome analysis of cotton. In the past several years, studies found that the ND family was mainly involved in the response to abiotic stress, such as drought, elevated light treatments and oxidative stress [10, 33], and NDA family genes were mainly in the response to cold stress and photorespiration [15, 34]. However, no systematic study of NDA genes is conducted in cotton, especially under alkaline stress. In this work, we undertook a comprehensive analysis of cotton NDA family and their involvement in alkaline stress.

In this study, 51, 52, 26 and 24 candidate NDA genes were systematically identified in *G. hirsutum*, *G. barbadense*, *G. arboreum* and *G. raimondii*, respectively. The NDA genes of *G. hirsutum* and *G. barbadense* were almost twice the number of *G. arboreum* and *G. raimondii*, which may be the reason that allotetraploid cotton was formed by inter-genomic hybridization of At-genome diploids and Dt-genome diploids, and followed by chromosome doubling [35]. In *G. hirsutum* and *G. barbadense*, the number of homologous NDA genes in At and Dt subgenome was identical, so we could infer that the translocations and reverse transcript insertion rarely occurred in the generation of cotton NDA gene family. The distribution of NDA varied greatly among selected plant species, such as 5 in *T. cacao*, compared with 51 in *G. hirsutum*, indicating that NDA genes underwent large

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**Fig. 13** Interaction network of GhNDA32 protein. The NDA1 represented the protein AtNDA1 corresponding to the protein in *Arabidopsis* with the highest homology to GhNDA32.
scale expansion in higher plants. By analyzing the physical properties of GhNDAs, we found that the average MWs of GhNDAs were 66.107 kDa, which were consistent with the situation of the potato and Arabidopsis with the molecular weight of 55–57 kDa [11].

According to phylogenetic characteristics and gene structure, the NDA genes can be divided into 8 clades in four Gossypium species (Fig. 2A), and clade a and clade h had the largest number of NDA genes with 59 and 35, respectively. By analyzing the motif composition of NDA genes in G. hirsutum, it was worth noting that clade a and clade h contained the largest number of motifs, clade a had a specific motif 4 and none of the other clades, while clade h had motif 7 but clade a did not. Gene structure analysis showed that exon number of NDA genes varied greatly from 1 to 22, which might be due to the directional evolution in the function and structure of NDA genes in the process of evolution. There was a strong connection between the intron-exon structure, conserved motifs and phylogenetic tree analysis of the NDA family in G. hirsutum. Therefore, we speculated that the differences in gene structure and motifs led to different branches of the NDA family, and conserved motifs could also reflect the relations of different subgroups. For example, both clade a and f had motif 9, while other clades did not, which may be related that they had acquired some important functions during evolution, and this might be used as an important basis for identifying these two subgroups.

When analyzing the chromosomal location of G. hirsutum and G. barbadense, we found that there were 25 NDA genes in GhAt and 27 genes in GbAt. Interestingly, there was no gene distribution on chromosome 8 of GhAt, which proved that chromosome changes may occur a translocation or the gene on the chromosome 8 was lost in the process of evolution. In addition, there were two genes on chromosome 11 of GhAt, but only one gene on chromosome 11 of GbAt, which proved that G. hirsutum had lost a gene or G. barbadense gained a new gene through evolution. Similarly, there were 26 NDA genes in GhDt and 25 genes in GbDt, and the only difference of chromosomal location of GhDt and GbDt was that the GbDt lacked a gene on chromosome 11 than the GhDt, and the same was that they all did not have the chromosome 5. Our results showed the uneven distribution of NDA genes among all the four Gossypium species, which might be due to the addition or gene loss of NDA genes through tandem, segmental, or whole genome duplication events during evolution.

Studies had found that gene duplication was the most commonly evaluated mechanism for the divergence of gene family [36]. To reveal the expansion mechanism of the NDA gene family, duplicated gene pairs of four Gossypium species were identified by MCScanX, we found that only two tandem duplication events existed in both G. arboreum and G. raimondii, GaNDA6/7 and GrNDA17/18, while the rest of duplicated gene pairs was identified as segmental duplication and whole genome duplication. This results revealed that low-tandem and high-segmental duplications were existed in the NDA gene family and segmental duplication and whole genome duplication played a predominant driving force in the evolution of NDA gene family, and it was consistent with the previous research [37].

To elucidate the differences after gene duplication, the ratio of non-synonymous (Ka) to synonymous (Ks) of four Gossypium species was calculated. In general, Ka/Ks < 1 was considered as a purification selection, indicating that natural selection eliminated harmful mutations and kept the protein unchanged, Ka/Ks > 1 was considered as positive selection, indicating natural selection changed the protein, the mutation site was quickly fixed in the population, and the evolution of the gene was accelerated, Ka/Ks = 1 was considered as neutral selection, indicating that natural selection did not affect mutation [38]. We found the Ka/Ks of most duplicated gene pairs in the four Gossypium species was less than 1, indicating that the four Gossypium species had undergone strong purification selection that occurred after tandem, segmental, and whole genome duplication. This was consistent with recent research that most cotton GRX gene family evolved through purifying selection pressure [39]. However, the selection pressure of most gene pairs among Gb-Gr, Gh-Gr, Gh-Ga and Gb-Ga were generally at 0–0.49, indicating that NDA genes tended to be conservative in evolution. Only the Ka/Ks value of Gh-Ga was greater than 1, suggesting that a certain member of GaNDA gene family members were evolved into some GhNDA gene family members through the environmental selection pressure, but whether they brought harmful traits or beneficial traits remains to be further studied.

Usually, cis-acting elements played an important role when plants were subjected to abiotic stress, and transcription factors (TFs) could bind to cis-acting elements of the promoter to regulate the transcription process and ultimately lead to the expression of genes [26]. In this study, most of the NDA family genes responded to light, which was consistent with the previous light-induced expression of NDA1 [15]. Light was a vital regulator of gene expression in plants, altering the transcription of thousands of genes [40]. However, the vast majority of genes related to light response focused on photosynthesis-associated nuclear genes, little was known about the effects of light on mitochondria and the respiratory chain [41, 42]. In this study, most of the NDA family genes responded to light, therefore, we inferred that
NDA family genes in upland cotton actively responded to light and the light could induce gene expression in mitochondria. Electron transport chains are known to produce ROS, which expose cells to oxidative stress. According to our GO annotation analysis (Fig. 8), most biological processes were concentrated in the oxidation-reduction process, which indicated that the NDA family genes were involved in the oxidation-reduction balance in cells to cope with alkaline stress. As cotton NDA genes are predicted to be localized in multiple subcellular compartments including the cytosol, chloroplasts, and mitochondria (Additional file 3: Table S3), it seems logical that its oxidation-reduction system should also be present in these compartments. In addition, a large number of hormone responsive elements (MeJA, ABA) in the promoters of GhNDAs, indicating that plant hormones probably involved in the regulation of GhNDAs in the upstream.

The expression pattern of genes was often used as an indicator of their functions. Therefore, RNA-Seq data was used to detect the possible function of genes in the NDA gene family by expression levels. Heat map analysis showed that most genes could be affected by SAS and AS stress (Fig. 9C), suggesting that NDA family genes play important roles in the response to alkaline stresses. The expression pattern of genes in different clades was different, but all the genes in the clade b were involved in the response of SAS and AS stress, and the cis-acting elements in the clade b were all involved in light-responsiveness, which demonstrated that GhNDAs in clade b had the function in response to light stress. In addition, some genes actively responded to SAS stress, such as GhNDA40, GhNDA14, GhNDA20, GhNDA45, GhNDA11, GhNDA35, GhNDA51, GhNDA25, GhNDA32 and GhNDA5, while GhNDA1, GhNDA14, GhNDA22 and GhNDA48 actively responded to AS stress. Usually, NAD(P)H dehydrogenases generally may be up-regulated under stress, and the results of qRT-PCR analysis showed that most GhNDAs genes are up-regulated under AS and SAS stress (Figs. 10, 11). GhNDA25 and GhNDA32 had the highest relative expression levels in roots while not in leaves, showing that these two genes may perform specific functions or participate in important signal pathways in the roots. In summary, the differential expression of several GhNDA genes indicated that they played an important role in alkaline stress response, which provided us with important candidate genes for studying alkaline stress in cotton.

Under alkaline stress, GhNDA32 gene expression was induced in cotton. However, alkaline stress can also destroy the oxidation-reduction balance of plants, resulting in cotton leaves wilted with water loss. In our study, the wilting degree of VIGS plants was higher than that of non-VIGS plants, indicating that the alkaline tolerance of cotton decreased after GhNDA32 gene was silenced. According to GO annotation, the biological process GhNDA32 gene participated in is oxidation-reduction process, with molecular function as oxidoreductase activity. Combining the results of GO annotation, VIGS and the interaction network diagram, we could infer that GhNDA32 gene might interact with AOX1A, AOX1C, AOX2, PUMP1 and CIB22 protein to regulate the activity of NADH dehydrogenase to regulate the redox reaction in the cotton, thereby eliminating the ROS produced in the body and maintaining cotton from the harm of oxidative stress.

Overall, this study will contribute to further understanding of the biological and molecular functions of cotton NDA and their antioxidant effects under alkaline stress. Our analysis provides a framework for further research into the function of this important gene family. The results of this study provide useful information for studying the effect of NDA genes in upland cotton under alkaline stress. These findings not only provide useful information for studying the effects of the NDA genes in G. hirsutum under alkaline stress but also provide valuable information for potential candidate genes related to plant growth and development and adversity stress.

Conclusions

In this study, phylogenetic relationship, gene structure, chromosomal distribution, collinearity analysis as well as cis-acting elements were conducted, which largely enriched our knowledge of the cotton NDA gene family. In addition, most GhNDAs contained cis-acting elements like light responsiveness. Expression patterns and functional characterization indicated GhNDAs participated in response to alkaline stress, especially NaHCO₃ stress. Taken together, all these results were of great significance for the future research on the molecular mechanism of NDA gene family in responding to abiotic stresses.

Materials and methods

Databases

Genome sequences of four Gossypium species (G. hirsutum, NAU; G. barbadense, HAU; G. arboreum, CRI; and G. raimondii, JGI) were used to identify the gene family. Genomic sequences, coding sequences of all of the four species were downloaded from Cotton Functional Genomic Database (CottonFGD) (http://www.cottonfgd.org/) [43]. Protein sequences of other species like A. thaliana (TAIR 10), T. cacao (version 10), S. tuberosum (version 4.03), O. sativa (version 7), P. trichocarpa (version 3.0), V. vinifera Genoscope (version 12), G. max (version 10), Z. mays (version 10) were obtained from Phytozome v12.1 (https://phytozome.jgi.doe.gov/pz/portal.html).
Identification of NDA family members
To identify the members of the GhNDA gene family, the protein sequence and genome annotation were downloaded from CottonFGD. The Hidden Markov Model (HMM) of PF07992 was downloaded from the Pfam website (https://pfam.xfam.org/). Local blast was used to obtain the protein sequence of PF07992, NCBI Batch Web CD-Search Tool (https://www.ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi) was used to further screen the genes [44]. CottonFGD was used to obtain some other features of GhNDA genes like protein length, molecular weights (MWs), isoelectric points (pIs), grand average of hydropathy. Subcellular location of GhNDA proteins were predicted using several websites, such as WOLF-PSORT (https://wolfpsort.hgc.jp/) and CELLO v2.5 (http://cello.life.nctu.edu.tw/) [45].

Phylogenetic analysis and sequences alignments
The full-length amino acid sequences of NDA genes for twelve plant species including G. hirsutum, G. barbadense, G. arboreum, and G. raimondii (downloaded from CottonFGD), A. thaliana (TAIR 10), T. cacao (version 10), S. tuberosum (version 4.03), O. sativa (version 7), P. trichocarpa (version 3.0), V. vinifera (version 12), G. max (version 10), Z. mays (version 10) (downloaded from Phytozome v12.1) were provided in MEGA7 software using ClustalW program for multiple sequence alignment. Subsequently, MEGA 7 software [46] was used to construct phylogenetic tree using neighbor joining (NJ) method with default parameters.

Chromosomal locations of NDA genes from four Gossypium species
The diagrams of the chromosomal locations from four Gossypium species including G. hirsutum, G. barbadense, G. arboreum and G. raimondii were drawn with the help of TB Tools software by using fragments per kilo base of exon per million fragments mapped (FPKM).

Analysis of the conserved protein motifs and gene structure
Multiple Em for Motif Elicitation (MEME) website (http://meme-suite.org/) was used to identify the conserved protein motifs [48]. TB Tools software was used to map the evolutionary relationship, gene structure, and conserved motifs of GhNDA proteins with mast file from MEME website, nwk file from phylogenetic tree analysis, and gff3 genome file of G. hirsutum.

Collinearity analysis of NDA genes in four Gossypium species
The complete genome sequences of four Gossypium species along with genome annotation files were subjected to MCScanX tool [49] to investigate the collinearity and analyze the syntenic relationship among NDA genes of four Gossypium species. The collinear and homologous chromosomal regions among four Gossypium species were visualized using Advance Circos tool in TB Tools software.

Calculation of selection pressure
Duplicated gene pairs from four Gossypium species including G. hirsutum, G. barbadense, G. arboreum, and G. raimondii were identified by using MCScanX tool. The nonsynonymous (Ka), the synonymous (Ks), and Ka/Ks were calculated to investigate the selection pressure by using Ka/Ks calculator in TB Tools software.

Gene ontology (GO) annotation analysis of GhNDAs
CottonFGD (https://cottonfgd.org) was used to determine the GO annotation analysis of GhNDAs including biological process, molecular functions and cellular component.

Analysis of cis-acting element in promoters of GhNDAs
The upstream sequences (2000 bp) of the NDA sequences were retrieved from the genome sequence in G. hirsutum. Then, the retrieved promoter sequences were submitted to PlantCARE (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/) [50]. Schematic diagram of phylogenetic tree and cis-acting elements was drawn with the help of TB Tools.

Expression patterns under different alkaline stress
RNA-Seq data (GSE165472) was used to analyze the expression level of differentially expressed genes (DEGs) under different alkaline stress (pH = 8.5 NaOH, 125 mM NaHCO3). The heat map along with phylogenetic tree and cis-acting elements was generated through TB Tools software by using fragments per kilo base of exon per million fragments mapped (FPKM).

Plant materials and alkaline treatments
Gossypium hirsutum cv. Zhong 9807 which is obtained from the Institute of Cotton Research of CAAS was used. Cotton seeds were sown in pots containing the soil under a 16-h light/8-h dark cycle at 25 °C for approximately 30 days. Seedlings with three true leaves and one heart-shaped leaf were washed carefully and transplanted into conical flasks containing 125 mM
NaHCO₃ solution for 0 and 12 h. Leaves and root samples were collected. Each sample was replicated three times. All these samples were immediately frozen in liquid nitrogen and kept at −80 °C for subsequent analysis.

**RNA extraction and quantitative real-time (qRT-PCR) analysis**

The total RNA of cotton roots and leaves were extracted using EASYspin Plus Plant RNA Kit (Aidlab, Beijing, China). The quantity and quality were determined by a NanoDrop 2000 Spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). The cDNA was reverse using the EasyScript® All-in-One First-Strand cDNA Synthesis SuperMix for qPCR (One-Step gDNA Removal). The qRT-PCR was performed using the Applied Biosystems@7500 Fast instrument and TransStart Top Green qPCR SuperMix. The Actin gene was used as a control. All the operational procedures followed the manufacturer’s protocols. Statistical analysis was conducted with biological replicates with mean values of three technique replicates. \(2^{-\Delta \Delta CT}\) method was used to calculate the relative FC for each sample [51]. The primers were designed using NCBI Primer-Blast tool, listed in Additional file 1: Table S1.

**Gene interaction network of the GhNDA32 protein**

The GhNDA32 protein interaction network was analyzed by the STRING database (https://string-db.org/) based on the homologous gene in Arabidopsis thaliana [52]. Cytoscape software (version 3.7.1) was used to construct a co-expression regulation network of the GhNDA32 protein [53].

**Virus-induced gene silencing (VIGS) experiment**

To verify the function of the NDA genes, we selected a highly expressed gene GhNDA32, pYLI156: GhNDA32 vector was constructed with the restriction enzyme cutting site BamHI and SacI. The sequences of the primer pair were detailed in Additional file 1: Table S1. The GV3101 strains carrying pYLI156: GhNDA32, pYL156: PDS, and pYL192 were cultured to OD600 = 1.2–1.5. Each mixture was injected into the underside of cotyledons of upland cotton material Zhong 9807. After injection, the seedlings were placed in the dark overnight, and a 16-h light / 8-h dark cycle was performed at 25 °C. When the plants injected with pYL156: PDS appeared an albino phenotype, it proved that the VIGs experiment was successful.

**Abbreviations**

NDA: The internal NAD(P)H dehydrogenase; ND: NAD(P)H dehydrogenase; VIGS: Virus-induced gene silencing; AP: Alternative respiratory pathway; ROS: Reactive oxygen species; ETC: electron transport chain; AOX: Alternative oxidase; CottonFGD: Cotton Functional Genomic Database; Ga: Gossypium arboreum; Gb: Gossypium barbadense; Gh: Gossypium hirsutum; Gr: Gossypium raimondii; HMM: Hidden Markov Model; MWs: Molecular weights; pI: Isoelectric points; MEME: Multiple Em for Motif Elicitation; Ks: The synonymous; Kt: The nonsynonymous; GO: Gene ontology; DEGs: Differentially expressed genes; FPKM: Fragments per kilo base of exon per million fragments mapped; qRT-PCR: quantitative real-time PCR; NJ: Neighbor-joining; ABA: Abscisic acid; MeJA: Methyl jasmonate; GA: Gibberellin; CK: No infection; SAS: 125 mM NaHCO₃; AS: pH 8.5 NaOH.

**Supplementary Information**

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**Authors’ contributions**

YPF, WWY: Conception and design; YPF, YXZ, CR, WWY: methodology; YPF, YXZ, CR, HZ, NX, JW, MGH, XXL, XGC, DLW, SW, LZX, LI2, HH, JJW, LQS, CC: data analysis; YPF: manuscript writing; YPF, WWY: reviewing and editing. All authors read and approved the final manuscript.

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**Availability of data and materials**

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**Declarations**

Ethics approval and consent to participate

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Consent for publication

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Competing interests

The authors declare no competing interests.

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