Article

The Full-Length Transcriptome Sequencing and Identification of Na+/H+ Antiporter Genes in Halophyte Nitraria tangutorum

Bobrov

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Abstract: Nitraria tangutorum Bobrov is a halophyte that is resistant to salt and alkali and is widely distributed in northwestern China. However, its genome has not been sequenced, thereby limiting studies on this particular species. For species without a reference genome, the full-length transcriptome is a convenient and rapid way to obtain reference gene information. To better study N. tangutorum, we used PacBio single-molecule real-time technology to perform full-length transcriptome analysis of this halophyte. In this study, a total of 21.83Gb of data were obtained, and 198,300 transcripts, 51,875 SSRs (simple sequence repeats), 55,574 CDS (coding sequence), and 74,913 lncRNAs (long non-coding RNA) were identified. In addition, using this full-length transcriptome, we identified the key Na+/H+ antiporter (NHX) genes that maintain ion balance in plants and found that these are induced to express under salt stress. The results indicate that the full-length transcriptome of N. tangutorum can be used as a database and be utilized in elucidating the salt tolerance mechanism of N. tangutorum.

Keywords: full-length transcriptome; SMRT; Nitraria tangutorum; NtNHXs

1. Introduction

Nitraria is a type of typical sand shrub in the Zygophyllaceae [1,2]. Nitraria tangutorum is one of the species endemic to the northwestern region of China [3,4]. N. tangutorum is one of the pioneer species that is resistant to salt, alkali, and sand burial, thereby playing an important role in maintaining the ecological balance of northwestern China [3,5]. In addition, this species has both economic value and benefits. Its leaves can be consumed, and its fruits can be used as food and medicine [6,7].

However, the current research on the omics of N. tangutorum is relatively scarce. In recent years, omics has rapidly developed into a powerful tool for studying various biological processes. Genomics, transcriptomics, and other omics can provide a large amount of genetic information [8], which may be utilized to conduct comprehensive studies on N. tangutorum.

The transcriptome is the sum of all mRNAs in a cell, tissue, or organism in a specific environment [9]. Transcriptome sequencing is helpful in the discovery of potential genes and can obtain a large number of gene sequences for molecular studies. The full-length transcriptome includes full-length mRNA sequences and complete transcripts with complete structural information [10]. Due to the limitation of reading length, second-generation sequencing technology can only obtain transcriptomes by splicing. Single-molecular
real-time technology (SMRT) based on third-generation sequencing has obvious advantages in the reading length of sequencing [11,12]. It does not need the splicing of transcripts to obtain the full-length transcripts directly, which can more truly reflect the sequence information of a species and also help in the accurate analysis of the expression of homologous genes and family genes.

Whole-genome sequencing can provide extremely rich genetic information [13]. However, only 548 plant genomes were sequenced before March 8, 2021 (https://www.plabipd.de/index.ep). Species that do not have completed whole-genome sequencing often have limited genetic information that hinders further research. Full-length transcriptome sequencing can provide extensive genetic information that can be used as a database to provide support for molecular and gene function research [14].

In this study, we used PacBio SMRT sequencing technology to obtain the full-length transcriptome of N. tangutorum, which was used in bioinformatics analysis, including full-length transcriptome correction and analysis, CDS prediction transcription factor analysis, SSR analysis, and IncRNA analysis, followed by functional annotation of the obtained transcripts based on the NR, Swiss-Prot, GO, COG, KOG, and KEGG databases. At the same time, we analyzed the full-length transcriptome, identified Na+/H+ antiporter genes, and verified their expression patterns under salt stress. These studies lay the foundation for elucidating the salt tolerance mechanism of N. tangutorum using molecular and omics technologies.

2. Materials and Methods

2.1. Plant Materials

N. tangutorum seeds were collected from Dengkou County, Inner Mongolia. The seeds were mixed in sand at 4 °C for about two months and then were planted in a mixed soil matrix with peat soil: perlite at a ratio of 4:1 in a greenhouse with a 16 h-light/8 h-dark light cycle and 60% air humidity at 23 °C. When the seedlings were 2 months old, the roots, stems, and leaves of three individual seedlings showing similar growth rates were collected and stored at −80 °C for RNA extraction.

2.2. RNA Extraction, Library Construction and SMRT Sequencing

Total RNA was extracted using an RNA extract kit (RNAiso Plus Co 9108, Takara, Japan), the qualified RNA of three seedlings were mixed together, then used oligo dT as primer for cDNA synthesis using PrimeScript 1st Strand cDNA Synthesis Kit (Takara, Shiga, Japan) and amplified by PCR, then the cDNA fragments were screened by a BluePippin nucleic acid recovery system (Sage Science, APG BIO, USA). Then, the full-length cDNA is nicked and repaired, end-repaired, and SMRT adapters were connected to complete the library construction. Then, Agilent 2100 was employed to detect the insert size of the library, qRT-PCR was performed to accurately quantify the library and then sequenced using SMRT technology.

2.3. SSR Analysis

For SSR analysis, we screened unigene sequences of >1000 bp in length. The MISA software (http://pgrc.ipk-gatersleben.de/misa/) was used to identify six types of SSRs, namely, mononucleotides, dinucleotides, trinucleotides, tetranucleotides, pentanucleotides, and hexanucleotides.

2.4. CDS, TF, and IncRNA Analysis

TransDecoder software was used to identify CDSs, iTAK software was used to predict TFs (transcription factors), and CNCl (Coding-Non-Coding Index) tools (https://github.com/www-bioinfo-org/CNCl), CPC (Coding Potential Calculator) software(http://cpc2.gao-lab.org/), and Pfam (protein family) were employed to predict the coding potential of sequencing data.
2.5. Gene Functional Annotation

BLAST was used to compare unigene sequence with those in the NR (Non-Redundant protein sequence database, ftp://ftp.ncbi.nih.gov/blast/db), Swiss-Prot (Swiss-Prot protein sequence database, http://www.uniprot.org), GO (Gene Ontology Consortium, http://www.geneontology.org), COG (Cluster of Orthologous Groups of proteins, http://www.ncbi.nlm.nih.gov/COG), KOG (eukaryotic Orthologous Groups, http://www.ncbi.nlm.nih.gov/KOG), and KEGG (Kyoto Encyclopedia of Genes and Genomes, http://www.genome.jp/kegg/) databases to obtain unigene annotation information.

2.6. Identification and Multi-Segment Alignments of Na+/H+ Antipporter Genes

Using the spliced sequence as the library, the BLAST program and Hmmer (v.3.0.1b) were used to search for genes containing the Na+/H+ antiporter domain (PF00999) with an E-value of < 1e-5, and then the SMAT online tool (http://smart.embl-heidelberg.de) was employed to identify genes containing the domain. Multiple alignments were performed using DNAMAN v9.0 (Lynnon Corporation, San Ramon, CA, USA) with the deduced amino acid sequences.

2.7. Phylogenetic and Motif Analyses

Phylogenetic reconstruction was performed using MEGA-X v10.1 (Temple, Philadelphia, PA, USA) using the neighbor-joining method, and bootstrap values were calculated according to 1000 repetitions. For motif analysis, the online analysis software MEME (https://meme-suite.org/meme/tools/) was used to analyze protein conserved motifs, and the Tbtools [15] were used for drawing.

2.8. Gene Cloning

Total RNA was extracted from N. tangutorum seedlings using RNA extraction kit (Estep super total RNA extraction kit Co LS1040, Promega, China), and the first strand of the cDNA was synthesized using HiScript III 1st Strand cDNA Synthesis Kit (Vazyme, China). The specific primers used to amplify the target gene fragments were designed with Oligo 7.60 (Cascade, Co 80809, USA) and are shown in Supplementary Table 7.

2.9. qRT-PCR Analysis

RNA was extracted from N. tangutorum seedlings exposed to 500 mM NaCl for 1, 6, 12, 24, and 48 h and then reverse transcribed for quantitative real-time-PCR (qRT-PCR). qRT-PCR was performed using a SYBR mix (AceQ qPCR SYBR Green Master Mix Co Q121-02, Vazyme, China). The PCR conditions were 94 °C for 15 s, 60 °C for 30 s, and 72 °C for 30 s. The primers used are shown in Supplementary Table 8, and each reaction was performed in triplicate. Relative expression values were calculated using the relative quantization method (2-ΔΔCT).

2.10. Subcellular Localization

ProtComp 9.0 (http://linux1.softberry.com/berry.phtml) was employed for subcellular location prediction, and TMHMM (http://www.cbs.dtu.dk/services/TMHMM/) was utilized for protein transmembrane analysis. For gene subcellular localization, pJIT-166 was used as the target vector, and XbaI and BamHI were employed to generate restriction sites to construct a fusion expression vector. The primers used are shown in Supplementary Table 9. The expression vector was transfected into onion epidermal cells by micro-projectile bombardment and incubated for 24 h in the dark [16]. The cells were subsequently observed using a fluorescence microscope (X-cite 120Q, Carl Zeiss).
3. Results

3.1. Sequencing Data Statistics

Total RNA was extracted from the roots, stems, and leaves of *N. tangutorum* to construct a sequencing library, and SMRT was employed to sequence the qualified library, removing errors and redundant consensus sequences in the original sequence and the obtained data statistics are shown in Table 1. A total of 21.83 Gb raw data were obtained, representing a total of 17,951,056 subreads. Most of the lengths were within the size range of 1 to 2000 bp (Figure 1), the average length was 1,216 bp, and the assembled N50 reached 1427 bp.

Table 1. Full-length transcriptome sequencing data.

| Sample       | Subreads Base (G) | Subreads Number | Average Length of Subreads | N30   | N50   | N90   |
|--------------|-------------------|-----------------|-----------------------------|-------|-------|-------|
| *N. tangutorum* | 21.83             | 17,951,056      | 1216                        | 1788  | 1427  | 744   |

Figure 1. Distribution of subread length. The abscissa is the length of the subreads, and the ordinate is the number of subreads of a particular length.

3.2. Data Correction

CCS sequencing analysis can effectively reduce the error rate of sequencing to obtain more accurate data [17]. To further correct the accurate full-length transcriptome data, we analyzed the statistical results of the consensus sequence distribution in Table 2. At the same time, we also calculated the consensus length and constructed a consensus length distribution map and found that the main length of the consensus sequence was 1–2000 bp (Figure 2).

Table 2. CCS quantity statistics data.

| Sample       | CCS   | Nfl-Reads | Flnc-Reads | Mean-Flnc | Consensus Reads |
|--------------|-------|-----------|------------|-----------|-----------------|
| *N. tangutorum* | 21.83 | 179,510,56 | 1216       | 1788      | 1427            |

CCS (circular consensus sequence), flnc-reads (full-length non-chimeric reads), Mean-flnc (average length of full-length non-chimeric reads), Consensus-reads (reads of consensus sequence obtained after clustering).
Figure 2. Consensus length distribution. The abscissa is the length of consensus, and the ordinate is the number of the consensus length.

3.3. CDS, TFs, IncRNA Analysis

Using TransDecoder software sequencing data to perform coding sequence prediction, a total of 55,574 sequences were obtained. The length distribution of the sequences is shown in Figure 3, and the distribution frequency is shown in Table S1. Using iTAK software, a total of 3,745 transcription factors were predicted that belonged to 75 gene families (Figure 4), including WRKY (157), MYB (236), NAC, AP2/ERF (175), bZIP (155), and other transcription factors. CNCI, CPC software, and Pfam database were used to predict the coding potential of the sequencing data. CNCI analysis identified 74,913 IncRNAs (Table S2). The CPC software was used to classify and identify IncRNA. A total of 139,217 IncRNAs were identified (Table S3). Using the Pfam database to perform multiple sequence alignments and HMMs searches on PacBio sequencing data, a total of 90,061 IncRNAs were identified (Table S4). Finally, a Venn diagram was constructed to depict the IncRNAs identified by the three methods to identify shared sequences (Figure 5).

Figure 3. Coding sequence length distribution. The abscissa is the predicted length of CDSs, the left ordinate is the number of transcripts of the CDSs, and the right ordinate is the percentage of transcripts.
3.4. SSR Analysis

Using MISA software to perform SSR analysis on unigenes of >1.000 bp in length, a total of 51,875 SSRs were identified, most of which were mononucleotides, dinucleotides, and trinucleotides, accounting for 71.68%, 15.04%, and 12.13% of the total number of SSRs, respectively (Table 3), whereas tetranucleotides, pentanucleotides, and hexanucleotides collectively accounted for 11.47% of the total number of SSRs. At the same time, SSR density analysis also generated similar results (Figure 6).

Table 3. Results of SSR analysis.

| Feature                          | Number  |
|---------------------------------|---------|
| Total number of sequences examined | 71,089  |
| Total size of examined sequences (bp) | 111,431,918 |
| Identified SSRs                  | 51,875  |
| SSR containing sequences         | 29,249  |
| Sequences containing more than 1 SSR | 11,082  |
SSRs present in compound formation

| Type of SSRs           | Number   |
|------------------------|----------|
| Mononucleotides        | 37,182   |
| Dinucleotides          | 7804     |
| Trinucleotides         | 6294     |
| Tetranucleotides       | 330      |
| Pentanucleotides       | 89       |
| Hexanucleotides        | 176      |

Figure 6. SSR density distribution. The abscissa is the SSR type, and the ordinate is the number of SSRs of the corresponding type in each Mb of sequence.

3.5. Functional Annotation

The 65,361 unigenes were functionally annotated using the COG, GO, KEGG, NR, Swiss-Prot, and KOG databases (Table 4), with annotation coverage rates of 34.46%, 69.19%, 31.43%, 99.53%, 78.99%, and 61.89%, respectively.

Unigene annotation using the NR database revealed that these exhibited a high degree of homology with Citrus sinensis and C. clementina (Figure 7). We compared the annotated genes to the GO database and divided them into three independent classifications: cellular component, molecular function, and biological process, respectively (Figure 8).

COG analysis classified these unigenes into 25 categories. Amino acid transport and metabolism was the most frequent functional category, followed by cell motility (Figure 9). KEGG annotation annotated 205,48 unigenes to 150 pathways (Table S5), which can be divided into 22 categories (Figure 10).

Figure 7. Homologous species distribution by Nr database. The best hits with an E-value = 1.0E-5 for each query were grouped according to species.
Table 4. Unigene annotation statistics.

| Annotated Databases | Number of Unigenes | 300–1000 bp | ≥ 1000 bp |
|---------------------|--------------------|-------------|-----------|
| COG                 | 26,526             | 3396        | 22,903    |
| GO                  | 45,222             | 6726        | 37,501    |
| KEGG                | 20,548             | 3528        | 16,970    |
| KOG                 | 40,450             | 6479        | 33,887    |
| Pfam                | 47,073             | 6746        | 40,303    |
| Swiss-Prot          | 51,635             | 8760        | 42,752    |
| Nr                  | 65,055             | 12,596      | 52,211    |
| All                 | 65,361             | 12,728      | 52,376    |

The column header 300–1000 bp indicates the number of unigenes annotated to the database whose length is ≥ 300 bases and ≤ 1000 bases. The column header ≥ 1000 bp indicates the number of unigenes annotated to the database unigene number with lengths ≥ 1000 bases.

Figure 8. GO function classification.

Figure 9. COG annotation classification statistics of expressed genes.
3.6. Identification of Na⁺/H⁺ Antiporter Genes in the Full-Length Transcriptome

The full-length transcriptome has a resolution accuracy of a single nucleotide, thus allowing the discovery of gene transcripts. N. tangutorum is a typical halophyte, but its molecular mechanism for salt tolerance remains unclear. The NHX gene family plays an important role in stress resistance in plants. However, its distribution in N. tangutorum is unknown. Using the full-length transcriptome as a database, we searched for sequences containing the domain according to the Na⁺/H⁺ antiporter (PF00999) Pfam domain and manually compared it to SMAT online analysis for confirmation. Using these steps, we have identified a total of nine NtNHXs, and then we named these based on their homology to AtNHXs. Multiple sequence alignment shows that these all contain a Na⁺/H⁺ antiporter domain (Supplemental Figure 2), then phylogenetic reconstruction of the AtNHXs was performed, which showed that these genes could be divided into three groups (Figure 11).

Figure 10. KEGG metabolic categories in the transcriptome of Nitraria tangutorum. The ordinate is the annotated biological function, and the abscissa is the percentage of these genes in the total number of annotated genes.
3.7. Gene Conserved Motifs Analysis of NHXs

To identify the conserved motifs of these NtNHXs proteins, using MEME software, we determined that NtNHXs had 14 mainly conserved motifs (Figure 12). Almost all identified NtNHXs harbored similar motif sequences, indicating that these motifs were highly conserved.

3.8. Real-Time Quantitative PCR Analysis

NHXs play an important role in maintaining the balance of Na⁺/H⁺ and salt resistance in plants [18,19]. To verify whether these NtNHX genes play a similar role in the salt-tolerance process of N. tangutorum, the seedlings of N. tangutorum were subjected to stress using 500 mM NaCl solution and plant materials were collected at 1, 6, 12, 24, and 48 h to assess NtNHX expression. Figure 13 shows that the NtNHX genes were expressed under salt stress, indicating that NtNHXs may play an important role in the salt resistance of N. tangutorum.
3.9. Clone and Subcellular Localization of NtNHX7

In *Arabidopsis*, AtNHX7 is also known as AtSOS1, which encodes a Na⁺/H⁺ antiporter that is located in the plasma membrane of the cell that promotes Na⁺ efflux and maintains the balance of ions inside and outside the cell [20]. To explore whether NtNHX7 has a similar effect and also to verify the accuracy of the predicted gene, we designed specific primers and cloned NtNHX7. We compared the sequence difference between the cloned gene and the reference sequence from this transcriptome, and we found that their similarity is 99.8% (Figure S1), indicating that the NtNHX7 predicted by the full-length transcriptome has higher accuracy.

Real-time quantitative PCR analysis revealed that NtNHX7 is expressed by salt stress (Figure 13). Subsequently, we used the sequencing results to analyze its expression pattern. First, ProtComp 9.0 and TMHMM were used to predict the subcellular location pattern of NtNHX7, which showed that NtNHX7 has a higher probability of localizing to the plasma membrane of the cell (Table S6, Figure 14 A). To further determine the subcellular localization of NtNHX7, we constructed the NtNHX7-GFP vector and performed transient transformation experiments to confirm our findings. As shown in Figure 14 B, NtNHX7 is mainly expressed on the cell membrane, which also verifies our previous bioinformatics prediction.

**Figure 13.** Relative expression level of NtNHXs in *Nitraria tangutorum* under 500 mM NaCl salt stress.

**Figure 14.** Subcellular localization of NtNHX7. (A) NtNHX7 transmembrane helix domain prediction by TMHMM. (B) NtNHX7 subcellular localization by microprojectile bombardment, 35S: GFP plants as control. Scale bar =100 μm.
4. Discussion

The SMRT technology is one of the relatively fast-developing bioinformatics technologies in recent years [21]. It can be used to improve the gene annotation status of sequenced species and as a reference for unsequenced species to facilitate scientific research. For example, the annotation of the sorghum genome has been improved using the SMRT technology, which will help to further study sorghum [22].

In sugar beet, the SMRT technology has been used to predict new genes and has laid the foundation for further research on this species [23].

Currently, transcriptome sequencing is mainly performed through NGS sequencing [24]. However, due to the limitation of sequencing read length in traditional next-generation sequencing, transcript information is mostly obtained by splicing sequences, and the full-length transcript sequence cannot be directly obtained. This hampers the analysis of transcriptome structure and may be accompanied by a high frequency of false-positive sequences.

Compared with second-generation sequencing, third-generation sequencing has a significantly higher read length, reaching an average of > 10 kb, and the maximum can reach 60 kb [25]. The SMRT technology can obtain a complete transcript without splicing, which is of great significance for transcriptome structure analysis, prediction of new genes [26], and supplementary genome annotation [27].

At present, various species have completed whole-genome sequencing, but there are still numerous organisms that have not completed the construction of a whole-genome map due to limited attention or sequencing cost, which also hinders further research on these species. For species that have not completed whole-genome sequencing, the full-length transcriptome provides a wealth of genetic information. Transcriptome data can be used to clone genes, develop SSR primers [28], and analyze homologous genes.

*N. tangutorum* is a typical drought-tolerant and salt-tolerant plant that is of important ecological significance. However, whole-genome sequencing of this species has not been completed to date, and limited genetic information hinders related molecular research. Although some genes of *N. tangutorum* have been cloned and their functions have been verified, this is only the tip of the iceberg in terms of studying this species from a molecular perspective. Using its full-length transcriptome sequencing data, it is possible to analyze its CDS, transcription factors, and SSRs and at the same time perform functional annotations on its genes and develop SSRs primers for plant population classification [29].

Full-length transcriptomes can also be used as a database to identify target genes [30]. *Nitritaria tangutorum* is a typical halophyte, and we would like to study the expression of salt-resistance-related genes in it. In the study of other plants such as *Arabidopsis* [31] and rice [18], *NHXs* have been shown to be involved in the process of salt stress tolerance. Therefore, in this study, we used the full-length transcriptome data as a library to obtain sequence information on the *NtNHXs* using BLAST. Then we investigated whether these genes responded to salt stress, and we found they can be induced by salt stress, although they had different expression characteristics in different tissues after salt treatment. In general, their expression reached its maximum within 12 h and then decreased (Figure 13), which was similar to the study of *Gossypium barbadense* [32] and *Pyrus betulifolia* [33], but there were also some differences, we suspect that the reason for this difference might be related to differences in salt stress concentration and species. In summary, the gene expression results indicated that the *NtNHX* gene might be involved in the process of resisting salt stress in *Nitritaria tangutorum*, but further gene functional verification studies are needed.

In addition, we selected *NtNHX7* to be cloned and sequenced and found that the cloned gene sequence was basically consistent with the sequence provided by the full-length transcriptome (Figure S1) through sequencing. Further, we constructed the *NtNHX7*-GFP gene fusion expression vector and observed its expression pattern through microprojectile bombardment and found that the signal was mainly expressed on the
plasma membrane and distributed inhomogeneously (Figure 14), which was similar to the expression pattern of the AtNHX7 (also known as AtSOS1) in Arabidopsis [20].

According to the above results, we thought that the full-length transcriptome could be used as a gene database, although it may not be as complete when compared with the full-genome sequence. However, for unsequenced species, it can enrich the scarce omics information and also lay the foundation for further molecular research.

5. Conclusions

In this study, we used SMRT technology to determine the full-length transcriptome of N. tangutorum. A total of 21.83 Gb of data were obtained, of which 198,300 transcripts, 51,875 SSRs, 55,574 CDS, and 74,913 IncRNAs were identified. In addition, using this full-length transcriptome, we identified the key NHX genes that maintain ionic balance in plants, and we induced their expression under salt stress. Based on the full-length transcriptome data, we also cloned the NtNHX7 gene and found a high similarity to the predicted result through sequencing, which indicates that the gene prediction based on the full-length transcript is more accurate.

In summary, this study obtained high-quality, full-length transcript information on N. tangutorum by full-length transcription sequencing and provided abundant transcript information, which promotes omics and molecular research on N. tangutorum.

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| NHX          | Na+/H+ antiporter gene |
| SRMT         | Single-molecule real-time |
| CDS          | Coding sequence |
| SSR          | Simple sequence repeat |
| LncRNA       | Long non-coding RNA |
| CPC          | Coding potential calculator |
| CNCI         | Coding non-coding index |
| mRNA         | Messenger RNA |
| qRT-PCR      | Quantitative real-time PCR |
| TF           | Transcription factors |
| cDNA         | Complementary DNA |
| Pfam         | Protein family |
| NR           | Non-redundant protein sequence database |
| Swiss-Prot   | Swiss-Prot protein sequence database |
| GO           | Gene Ontology Consortium |
| COG          | Cluster of Orthologous Groups of proteins |
| KOG          | EuKaryotic Orthologous Groups |
| KEGG         | Kyoto Encyclopedia of Genes and Genomes |

Supplementary Materials: The following are available online at www.mdpi.com/DOI/10.3390/genes12060836/s1, Figure S1: The amino acid sequence comparison between predicted sequence and sequencing result of NtNHX7. The top row is the prediction result, and the second row is the sequencing result. Figure S2: The amino acid multiple fragment alignments of NtNHXs. Table S1: CDS length distribution and its frequency. Table S2: CNCI analysis for IncRNAs. Table S3: CPC analysis for IncRNAs. Table S4: Pfam analysis for IncRNAs. Table S5: KEGG annotation databases. Table S6: Prediction of NtNHX7 subcellular localization by ProtComp9.0 (http://linux1.softberry.com/berry.phtml). Table S7: Primers for gene cloning. Table S8: Primers for Quantitative RT-PCR. Table S9: Primers for subcellular localization. Table S10: amino acid sequence of NtNHXs.

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