Genome-wide identification, phylogeny, evolution, and expression patterns of MtN3/saliva/SWEET genes and functional analysis of BcNS in Brassica rapa

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Abstract

Background: Members of the MtN3/saliva/SWEET gene family are present in various organisms and are highly conserved. Their precise biochemical functions remain unclear, especially in Chinese cabbage. Based on the whole genome sequence, this study aims to identify the MtN3/saliva/SWEETs family members in Chinese cabbage and to analyze their classification, gene structure, chromosome distribution, phylogenetic relationship, expression pattern, and biological functions.

Results: We identified 34 SWEET genes in Chinese cabbage and analyzed their localization on chromosomes and transmembrane regions of their corresponding proteins. Analysis of a phylogenetic tree indicated that there were at least 17 supposed ancestor genes before the separation in Brassica rapa and Arabidopsis. The expression patterns of these genes in different tissues and flower developmental stages of Chinese cabbage showed that they are mainly involved in reproductive development. The Ka/Ks ratio between paralogous SWEET gene pairs of B. rapa were far less than 1. In our previous study, At2g39060 homologous gene Bra000116 (BraSWEET9, also named BcNS, Brassica Nectary and Stamen) played an important role during flower development in Chinese cabbage. Instantaneous expression experiments in onion epidermal cells showed that the gene encoding this protein is localized to the plasma membrane. A basal nectary split is the phenotype of transgenic plants transformed with the antisense expression vector.

Conclusion: This study is the first to perform a sequence analysis, structures analysis, physiological and biochemical characteristics analysis of the MtN3/saliva/SWEETs gene in Chinese cabbage and to verify the function of BcNS. A total of 34 SWEET genes were identified and they are distributed among ten chromosomes and one scaffold. The Ka/Ks ratio implies that the duplication genes suffered strong purifying selection for retention. These genes were differentially expressed in different floral organs. The phenotypes of the transgenic plants indicated that BcNS participates in the development of the floral nectary. This study provides a basis for further functional analysis of the MtN3/saliva/SWEETs gene family.

Keywords: Chinese cabbage (Brassica rapa syn. Brassica campestris), MtN3/saliva/SWEETs, Gene family, BcNS, Nectary, Functional analysis

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Background

The MtN3/saliva/SWEET gene family widely exists in high-level eukaryotes and can be found in protozoa, metazoa, fungi, bacteria, and archaea [1, 2]. The first gene of this family was identified to be MtN3 (Medicago truncatula Nodul3), which is a root-nodule-related gene in M. truncatula and is associated with Rhizobium-induced nodule development [3]. Thereafter, a homologous gene of MtN3 was found in the saliva of Drosophila, and this gene is expressed in the salivary gland during embryonic development [4]. Since then, proteins with the conserved domains have been found in mice, humans, sea squirts, petunia, rice, Arabidopsis, and other plants. The conserved transmembrane domain is named as the MtN3/saliva domain; proteins containing this domain are assigned to the MtN3/saliva protein family [2]. A new family of sugar carriers that use sugar optical sensors have been found and named the SWEET family [5]. SWEET and the previously reported MtN3/saliva gene family belong to the same family [6]. MtN3/saliva/SWEET proteins can be classified into two types based on the number of MtN3/saliva domains: the first type consists of seven transmembrane helices that harbor two MtN3/saliva domains, and the second type comprises three transmembrane helices that harbor one MtN3/saliva domain [7]. These domains are highly conserved in the evolutionary process of the MtN3/saliva/SWEET gene family. The number of the members in this family differs among different species and is high in some plant species.

The evolution of the MtN3/saliva/SWEET gene family is highly conserved and these genes are involved in many different physiological functions in plants, including reproductive development, senescence, abiotic stress response, and host-pathogen interactions. The MtN3/saliva/SWEET family genes are associated with fertility in plants. Os11N3/SWEET14-knockout mutants exhibit reduced seed size and delayed reproductive development [8]. NEC1, an MtN3/saliva/SWEET gene, is specifically expressed in the nectary tube and stamen of Petunia [9]. When the gene was cosuppressed, the transgenic plants showed male sterility due to failure of anther dehiscence [10]. RPG1 (RUPTURED POLLEN GRAIN 1), also named AtSWEET8, is a necessary gene for anther development in Arabidopsis [11]; the gene was highly expressed in the tapetum and pollen mother cells from stage 5 to stage 7. The male fertility of rpg1 decreased significantly, and most of the pollen grains were degraded during development. The microspore membrane of rpg1 became dysfunctional in the tetrad stage, leading to sporopollenin deposition onto the microspore surface, which influenced pollen wall development and pollen fertility [11].

The MtN3/saliva/SWEET family genes are involved in senescence, abiotic stress response, and host-pathogen interactions in plants. The SAG29 (AtSWEET15) protein is located on the plasma membrane and regulates cell activity in a high-salt environment; the protein expression is induced by the ABA pathway, depending on osmotic pressure in plant senescent tissues; SAG29 mutants are less sensitive to high salt environments, whereas SAG29 35S-overexpressing transgenic plants are highly sensitive to high salt environments and exhibit accelerated aging phenotypes [12]. Suppressing the expression of either the dominant or recessive allele of Xa13/Os8N3/OsSWEET11 enhances the resistance to PXO99, which is an incompatible strain of xa13 [13].

Cruciferous plants occupies a very important niche in agricultural production, and nectaries play vital roles in the pollination and fertilization of the plants. However, the molecular mechanisms involved in nectary development are very limited. CRABS CLAW, BOP1 (BLADE-ON-PETIOLE1), BOP2 [14, 15], and CWINV4 (CELL WALL INVERTASE 4) were reported which involved in nectary development. In cwinv4 mutant plants, the nectary cannot secrete honeydew [16, 17]. There is a study that shows that many genes are involved in photosynthesis in turnips [18]. The another study assumed that nectaries could make carbohydrates themselves through photosynthesis [19]. The nectaries of cruciferous plants are small, and their structure is complicated, so they are difficulty to be studied. However, exploring nectary development will benefit our understanding of the plants’ reproductive process.

So far, the MtN3/saliva/SWEET gene family has been rarely reported in plants, and the molecular mechanism of the gene family remains poorly understood. This study aims to clarify the biological function of MtN3/saliva/SWEET family genes in Chinese cabbage. Thirty four genes belonging to the MtN3/saliva/SWEET family were identified in cabbage. Gene and protein sequences were evaluated and compared by analyzing the protein sequences, evolution of the gene family, protein transmembrane domain, gene chromosomal localization, and gene structure. The expression characteristics of MtN3/saliva/SWEET family genes were studied in different tissues and flower developmental stages to provide some clues of their function.

Methods

Identification and isolation of MtN3/saliva/SWEET family genes in Brassica rapa

To identify MtN3/saliva/SWEET family genes, we found candidate gene sequences with the reported MtN3/saliva/SWEET family genes in the Arabidopsis Genome Database TAIR (http://www.arabidopsis.org/), and then verified the obtained sequences in the conserved domain database of NCBI (National Center for Biotechnology Information) (http://www.ncbi.nlm.nih.gov/Structure/
Ultimately, 17 Arabidopsis MtN3/saliva/SWEET family members were found. Then, we obtained the candidate genes of the MtN3/saliva/SWEET family in the Brassica database (http://brassicadb.org/brad/searchGene.php) by analyzing Arabidopsis MtN3/saliva/SWEET family members’ amino-acid sequences and domain sequences (Additional file 1: Table S1) with BLASTP. The Pfam database (E-value is 1.0) (http://pfam.janelia.org/) and the Conserved Domain Database (CDD) of NCBI were used to determine whether each of the sequences each harbored one or more MtN3/saliva/SWEET conserved domains (MtN3_slv; PF03083.15). The genes that did not contain the known conserved domains of the gene families were excluded from further analysis. Eventually, we identified 34 genes of the MtN3/saliva/SWEET family genes in Chinese cabbage.

**Chromosomal localization of the MtN3/saliva/SWEET genes**

We searched the start location and stop location on the B. rapa chromosomes of all MtN3/saliva/SWEET family genes in the Brassica database. Chromosomal localization of these genes was performed by MapChart software based on the relative location of each gene on each chromosome [20]. Tandem duplications were defined as genes located within 20 loci from each other [21, 22].

**Multiple sequence alignment and phylogenetic analysis of MtN3/saliva/SWEET family genes**

We downloaded the full-length DNA sequences and the cDNA sequences of MtN3/saliva/SWEETs genes from the Brassica database. GSDS (Gene Structure Display Server) (http://gsds.cbi.pku.edu.cn/index.php) was utilized to analyze the locations of introns and exons. Homologous sequence alignment of the 34 MtN3/saliva/SWEETs family genes was performed with ClustalW [23], and then the sequence alignment results were considered the basis for generating the unrooted phylogenetic tree of the MtN3/saliva/SWEET family members with MEGA (version 5.0) [24]. All parameters used were default parameters. Phylogenetic trees were generated with the value of the 1000 bootstrap samples by the neighbor-joining (NJ) method [24].

**Nonsynonymous substitution rate and synonymous substitution rate between orthologous SWEET genes between A. thaliana and B. rapa**

Synteny analysis of MtN3/saliva/SWEETs genes between were A. thaliana and B. rapa performed with PGDD (http://chibba.agtec.uga.edu/duplication/). The occurrence of duplication events and homologous genes divergence, as well as the selective pressure on duplicated genes, were estimated by calculating synonymous (Ks) and non-synonymous substitutions (Ka) per site between the orthologous genes using the LOCUS SEARCH utility of PGDD. The divergence time was calculated using the neutral substitution rate of 1.5 × 10\(^{-8}\) substitutions per site per year for the chalcone synthase gene (Chs) [25].

**Conserved short amino-acid sequence analysis of MtN3/saliva/SWEET in B. rapa**

We used MEME to analyze the common conservative motifs of 34 MtN3/saliva/SWEETs amino-acid sequences from Chinese cabbage [26]. Phylogenetic tree was generated with the value of 1000 bootstrap samples by the NJ method.

**Transmembrane helices and hydrophobicity analysis of the MtN3/saliva/SWEET family proteins in B. rapa**

The transmembrane helices of the MtN3/saliva/SWEET family proteins were predicted by the TMHMM (http://www.cbs.dtu.dk/services/TMHMM/) with their amino acid sequences [27]. Hydrophobicity analysis of MtN3/saliva/SWEET family proteins were analyzed with Protscal (http://web.expasy.org/protscal/) in accordance with the scoring criteria of Hphob/Kyte and Doolittle [28]. The GRAVY value for a peptide or protein was calculated as the sum of hydropathy values of all the amino acids divided by the number of residues in the sequence. A larger positive hydrophilic value indicated stronger hydrophilicity. By contrast, a smaller negative hydrophilic value indicated stronger hydrophillicity. Hydrophilic values taht ranged from +0.5 to −0.5 mainly corresponded to amphoterich amino acids. Furthermore, the isoelectric points (pIs) and molecular weights of the MtN3/saliva/SWEET proteins were predicted using ExPaSy (http://web.expasy.org/compute_pi/) [29].

**Cis-element analysis of the promoter sequences of MtN3/saliva/SWEET family genes in B. rapa**

To identify the cis-elements in the promoter sequences of the 34 MtN3/saliva/SWEET family genes, the sequences of these genes were used as query sequences for the BLASTN search against the Brassica database. A 1500 bp of genomic sequences upstream of the start codon of each MtN3/saliva/SWEET family gene was chosen. Afterward, PlantCare (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/) and a manual search were employed to predict the cis-elements [30].

**Quantitative reverse-transcription PCR (qRT-PCR) analysis of the MtN3/saliva/SWEET family genes in different tissues or different floral organs**

‘Chiifu-401’ is the species that was used in genome sequencing, so ‘Chiifu-401’ was used in our study for qRT-PCR analysis and grown in the experimental base of our laboratory. We then obtained roots, stems, leaves, flowers, pods, sepals, petals, stamens, pistils, nectar, big
buds (> 2.0 mm), and small buds (< 2.0 mm) in the flowering period. All the samples were collected and immediately frozen in liquid nitrogen and stored at –80 °C.

Total RNA was extracted from previous materials using the TRIzol reagent (Invitrogen, USA) in accordance with the manufacturer’s instructions. The first cDNA strand was generated following the manufacturer’s protocol using the Takara Reverse Transcription System (Japan). qRT-PCR was performed using the primer pairs listed in Additional file 2: Table S2. The primers were designed using the Primer program (version 5.0). The specificity of each primer to its corresponding gene was checked using the BLASTN program of the NCBI. A sample of cDNA (0.2 μg) was subjected to real-time PCR in a final volume of 20 μL containing 10 μL of SYBR Green Master Mix Reagent (Takara, Japan) and 0.8 μL of specific primers. Three biological and three technical replicates for each sample were performed in a real-time PCR machine (Bio-Rad CFX Manager). The heating program was as follows: 95 °C for 30 s, followed by 95 °C at 10 s for 40 cycles, and 52 °C at 30 s (varying with specific primers). To normalize the total amount of cDNA present in each reaction, the Chinese cabbage Cyp gene was amplified as an endogenous control for calibrating relative expression. The 2−ΔΔCT method of relative gene quantification recommended by Applied Biosystems (PE Applied Biosystems, USA) was used to calculate the expression level of different treatments [31].

In our previous study, we found that BcNS plays an important role in Chinese cabbage, so we studied the expression pattern of BcNS here. The expression in open flowers, I ~ V buds (I: < 1 mm, II: 1.2 ~ 1.6 mm, III: 1.8 ~ 2.2 mm, IV: 2.4 ~ 2.8 mm, V: 3.0 ~ 3.4 mm), tender pods, stems, and pistils in ‘Bajh97-01A/B’ sterile and fertile plants were examined 1 h, 8 h, and 24 h after pollinating with semi quantitative PCR, and the gene expression in sepal, petals, stamen, pistils and nectaries in ‘Bajh97-01A/B’ sterile and fertile plants were detected with semi-quantitative PCR and quantitative reverse-transcription PCR.

In situ hybridization

Flower buds at different developmental stages were fixed in 4% formaldehyde phosphate-buffered saline (PBS) solution with 0.1% Triton X-100 and 0.1% Tween-20, serially dehydrated, cleared with dimethyl benzene, and embedded in paraffin. Sections (8 μm) of flower buds were hybridized to specific digoxigenin-labelled RNA probes (Roche). Templates for the BcNS-specific probes were obtained from amplification with specific primers (upstream primer: 5′-TACGGATCCAGTCTGGCGTC TTTGCT-3′; downstream primer: 5′-CGCTCTAGA CCGTGGTGCTGTAGTTG-3′; the underlined section is the enzyme cut site). The sense and antisense probes were synthesized using an SP6/ T7 transcription kit (Roche) [32].

Subcellular localization of BcNS in the onion epidermal cell

We studied the subcellular localization of BcNS in the onion epidermal cell. First, the BcNS gene fragment was amplified from the nectaries of Chinese cabbage ‘Chifu-401’ with a high-fidelity enzyme (TOYBO, Japan). To overexpress the YFP (Yellow Fluorescent Protein) marker protein of the pB7YWG2.0-YFP, the stop codon of BcNS had to be replaced. Otherwise, the 5′-end of the upstream primer should add “CACC” when using the TOPO isomerase to connect to the entry vector. Thus, we designed the primer as follows: upstream (BcNS-GF), 5′-CAC CATGGTGTCATCAAGTTCATCAA-3′; downstream (BcNS-GR), 5′-GTAACCGGTGCTGGCTTGAGT-3′. The PCR reaction was programmed to heat the samples at 94 °C for 30 min, followed by 94 °C for 30 s in 29 cycles, 52 °C for 30 s (varying with specific primers), 68 °C for 90 s, and 68 °C extended to 7 min. The verified sequence was connect to the entry vector using the pENTR™ Directional TOPO® Cloning Kit (Invitrogen, USA). The target sequence was replaced into the expression vector through homologous replacement with the Gateway®LR Clonase™ Enzyme Mix (Invitrogen, USA). The constructed vector was transferred into an onion epidermal cell for transient expression by a gene gun.

Construction of the antisense expression vector of BcNS and molecular detection in transgenic plants

Flowering Chinese cabbage is a spring variety and can normally flower without vernalization. Using this species as the experimental material can improve the experimental efficiency. Our previous study found a 99.4% similarity of BcNS between B. campestris and B. rapa. Therefore, flowering Chinese cabbage is a suitable material for the study of BcNS function. We constructed the antisense expression vector pCAMIA1301-BcNS of the BcNS gene by enzyme digestion and other routine molecular biological methods. We then introduced the vector into Agrobacterium LBA4404 and EHA105 for genetic transformation.

One or two leaves were obtained from the transgenic plants and the controls. Dust was wiped away with 75% alcohol, and then DNA was extracted from the samples using the CTAB (Hexadecyltrimethylammonium bromide) method for PCR testing. The leaves and calluses of the transgenic plants and negative control were placed at X-Gluc solution at 37 °C for 24–36 h. Subsequently, the samples were placed at 95% ethanol for the dehydration treatment. Finally, sample staining was observed under a microscope. The morphological characteristics
of the transgenic plants were observed under an MZ16 FA Leica microscope.

Scanning electron microscopy observations
For the scanning electron microscopy of the nectaries in transgenic plants, we followed the protocol described by the Center of Electron Microscopy, Zhejiang University. Firstly, the nectaries of the transgenic plants and their control were fixed with glutaraldehyde in 4 °C overnight. Secondly, the sample was washed with phosphate buffer three times, each time for 15 min. Then, the sample was serially dehydrated in ethanol at different concentrations for 15 min at each concentration. The last two steps used 100% ethanol, and each step lasted 20 min. The samples were dried through the critical point drying method, and then the samples were pasted onto the metal sample table with a special double-sided adhesive. After coating the samples with a metallic membrane, the samples were placed on a desktop scanning electron microscope (TM-1000) to observe the nectary and stoma.

Results
Identification of MtN3/saliva/SWEET family genes in B. rapa
Thirty-four MtN3/saliva/SWEETs genes were obtained by BLASTP against the Arabidopsis Genome Database TAIR, NCBI databases and Brassica database. These genes were then named after their A. thaliana homologs (Table 1). Duplication clusters were identified and found to be BrSWEET1a/1b, BrSWEET2a/2b, BrSWEET3a/3b, BrSWEET4a/4b, BrSWEET5a/5b, BrSWEET7a/7b, BrSWEET11a/11bc, BrSWEET12a/12b, BrSWEET14a/14b, BrSWEET15a/15bc and BrSWEET16a/16b (Table 2). A total of 17 AtSWEETs were obtained, but homologs of Bra012412, which we named as BrSWEET 18, were not found in the annotated AtSWEETs (Table 1). The open reading frame (ORF) lengths of the MtN3/saliva/SWEETs genes ranged from 387 bp (BrSWEET2c) to 951 bp (BrSWEET15b). The encoded polypeptides ranged from 128 amino acids (BrSWEET2c) to 316 amino acids (BrSWEET15b), with molecular weights from 14.299 kDa (BrSWEET2c) to 35.814 kDa (BrSWEET15b). The theoretical pI ranged from 7.59 (BrSWEET2) to 9.62 (BrSWEET7), and most of the pIs were more than 8.50 except BrSWEET2a, BrSWEET3b, BrSWEET5b, BrSWEET5c and BrSWEET15c (Table 1). The MtN3/saliva/SWEET genes were distributed on 10 chromosomes. BrSWEET13 was distributed on a scaffold. A total of 31 MtN3/saliva/SWEETs members contained two conserved domains, whereas the others contained only one conserved domain (Table 1).

Chromosomal localization of the MtN3/saliva/SWEET genes
By chromosomal localization analysis, the 34 MtN3/saliva/SWEETs genes were determined to be distributed among ten chromosomes. By contrast, BrSWEET13 was distributed on a scaffold. In particular, 7 genes were distributed on the 3rd chromosome; 5 genes on chromosomes 2 and 6; 4 genes on chromosomes 1; and 3 genes on chromosomes 8 and 9; 2 genes were distributed on the fifth and tenth chromosome, while the fourth and seventh chromosome only had one MtN3/saliva/SWEET gene. The MtN3/saliva/SWEET family genes were scattered across genome, not concentrated on a few chromosomes (Fig. 1).

Phylogenetic tree analysis of the MtN3/saliva/SWEET in B. rapa, A. thaliana and O. sativa
The SWEET genes of A. thaliana, O. sativa, and Lycopersicum esculentum have been classified into four classes, namely, Type-A1, Type-A2, Type-A3, and Type-B1 [33]. As we known, the genome of B. rapa has undergone a whole-genome triplication (WGT), therefore, to investigate the retention and loss of the MtN3/saliva/SWEET family genes through evolution, we constructed a phylogenetic tree of all B. rapa, A. thaliana and O. sativa MtN3/saliva/SWEET genes (Fig. 2). Given the existing classification method of Arabidopsis MtN3/saliva/SWEET genes, the phylogenetic tree were also classified into four classes. In this study, the MtN3/saliva/SWEETs genes in the green, blue, yellow, and purple clades were termed as Type-A1, Type-A2, Type-A3, and Type-B1, respectively, and the clades contained 8, 4, 7, and 15 Brassica MtN3/saliva/SWEETs family members, respectively. Twelve pairs of protein sequences (one pair means they have the same homologous gene in Arabidopsis), for example, BrSWEET1a/BrSWEET1b, exhibited high similarity to each other and were located in closely related branches. These properties indicated the functional redundancy of such sequences. Furthermore, BrSWEET7, BrSWEET9, BrSWEET8, and BrSWEET10 showed a relatively low homology with the other MtN3/saliva/SWEET family genes. One branch only contained the B. rapa MtN3/saliva/SWEET genes (Fig. 2, black arrows) and one branch only contained the Arabidopsis MtN3/saliva/SWEET genes (Fig. 2, red arrows). To analyze the number of nodes in the phylogenetic tree, we found 17 nodes with high confidence values (>70%). This result indicated the existence of at least 16 supposed ancestor genes of A. thaliana and B. rapa (red dots); six nodes with the same standard mean the possible ancestor genes of A. thaliana, B. rapa and O. sativa (Fig. 2, black dots).

The gains and losses, as well as the copy number changes, during the evolution of the Arabidopsis and B. rapa MtN3/saliva/SWEET genes in different clades were analyzed in this study (Additional file 3: Figure S1). The numbers of Arabidopsis and B. rapa MtN3/
| Group | Arabidopsis thaliana | Brassica rapa | Number of domain |
|-------|----------------------|---------------|----------------|
|       | Gene name | Accession number | ORF(bp) | Deduced polypeptide | Gene name | Accession number | Chromosome | Subgenome | ORF(bp) | Deduced polypeptide | Protein length (aa) | MW(kD) | PI |
|       |           |                |         |                    |           |                |            |           |         |                    |                      |        |    |
| I     | AtSWEET1  | AT1G21460      | 744     | 247                | BrSWEET1a| Bra016421      | 8           | MF1       | 756     | 251                 | 27.818               | 9.27   | 2  |
|       |           |                |         | 27.268             | BrSWEET1b| Bra017916      | 6           | LF        | 741     | 246                 | 27.185               | 9.3    | 2  |
|       | AtSWEET2  | AT3G14770      | 711     | 236                | BrSWEET2a| Bra021577      | 1           | MF1       | 711     | 236                 | 26.567               | 7.59   | 2  |
|       |           |                |         | 26.467             | BrSWEET2b| Bra027314      | 5           | LF        | 648     | 215                 | 24.078               | 9.16   | 2  |
|       |           |                |         |                    | BrSWEET2c| Bra027312      | 5           | LF        | 387     | 128                 | 14.299               | 9.15   | 1  |
|       | AtSWEET3  | AT5G53190      | 792     | 263                | BrSWEET3a| Bra022636      | 2           | MF2       | 780     | 259                 | 29.328               | 8.71   | 2  |
|       |           |                |         | 29.537             | BrSWEET3b| Bra003075      | 10          | LF        | 579     | 192                 | 21.593               | 7.64   | 1  |
| II    | AtSWEET4  | AT3G28007      | 756     | 251                | BrSWEET4a| Bra025317      | 6           | LF        | 747     | 248                 | 27.488               | 8.97   | 2  |
|       |           |                |         | 27.815             | BrSWEET4b| Bra033034      | 2           | MF1       | 738     | 245                 | 24.078               | 8.71   | 2  |
|       | AtSWEET5  | AT5G62850      | 723     | 240                | BrSWEET5a| Bra035879      | 9           | MF2       | 723     | 240                 | 27.3     | 8.94   | 2  |
|       |           |                |         | 27.21              | BrSWEET5b| Bra029245      | 2           | MF1       | 723     | 240                 | 26.971               | 8.15   | 2  |
|       |           |                |         |                    | BrSWEET5c| Bra029244      | 2           | MF1       | 723     | 240                 | 26.826               | 8.14   | 2  |
|       | AtSWEET6  | AT1G66770      | 786     | 261                | BrSWEET6a| Bra035270      | 9           | MF1       | 750     | 249                 | 27.362               | 8.63   | 2  |
|       |           |                |         | 28.867             | BrSWEET6b| Bra000725      | 3           | LF        | 537     | 178                 | 19.581               | 8.97   | 2  |
|       | AtSWEET7  | AT4G10850      | 777     | 258                | BrSWEET7a| Bra035270      | 9           | MF1       | 750     | 249                 | 27.362               | 8.63   | 2  |
|       |           |                |         | 28.531             | BrSWEET7b| Bra000725      | 3           | LF        | 537     | 178                 | 19.581               | 8.97   | 2  |
|       | AtSWEET8  | AT5G40260      | 720     | 239                | BrSWEET8a| Bra025595      | 4           | MF2       | 813     | 270                 | 30.125               | 8.18   | 2  |
|       |           |                |         | 26.886             | BrSWEET8b| Bra000116      | 3           | MF2       | 813     | 270                 | 30.125               | 8.18   | 2  |
| III   | AtSWEET9  | AT3G39060      | 777     | 258                | BrSWEET9a| Bra022761      | 3           | MF1       | 870     | 290                 | 33.041               | 8.92   | 2  |
|       |           |                |         | 28.716             | BrSWEET9b| Bra022761      | 3           | MF1       | 870     | 290                 | 33.041               | 8.92   | 2  |
|       | AtSWEET10 | AT5G50790      | 870     | 289                | BrSWEET10a| Bra018039      | 6           | LF        | 870     | 289                 | 32.068               | 8.92   | 2  |
|       |           |                |         | 32.763             | BrSWEET10b| Bra019560      | 6           | MF2       | 858     | 285                 | 31.584               | 9.32   | 2  |
|       |           |                |         |                    | BrSWEET10c| Bra029914      | 1           | MF1       | 873     | 290                 | 32.343               | 9.04   | 2  |
|       | AtSWEET11 | AT3G48740      | 870     | 289                | BrSWEET11a| Bra009700      | 6           | LF        | 867     | 288                 | 31.781               | 9.07   | 2  |
|       |           |                |         | 31.921             | BrSWEET11b| Bra026487      | 9           | MF2       | 834     | 277                 | 30.494               | 9.13   | 2  |
|       |           |                |         |                    | BrSWEET11c| Bra026487      | 9           | MF2       | 834     | 277                 | 30.494               | 9.13   | 2  |
|       | AtSWEET12 | AT5G23660      | 858     | 285                | BrSWEET12a| Bra040489      | 1           | LF        | 885     | 294                 | 32.747               | 9.05   | 2  |
|       |           |                |         | 32.504             | BrSWEET12b| Bra009700      | 6           | LF        | 867     | 288                 | 31.781               | 9.07   | 2  |
|       | AtSWEET13 | AT5G50800      | 885     | 294                | BrSWEET13| Bra000116      | 3           | MF2       | 819     | 272                 | 29.916               | 9.18   | 2  |
|       |           |                |         | 30.069             | BrSWEET14a| Bra013854      | 1           | LF        | 822     | 273                 | 30.105               | 9.29   | 2  |
|       | AtSWEET14 | AT4G25010      | 846     | 281                | BrSWEET14b| Bra010477      | 8           | MF2       | 819     | 272                 | 29.916               | 9.18   | 2  |
|       |           |                |         | 30.969             | BrSWEET14c| Bra019197      | 3           | MF1       | 717     | 238                 | 26.408               | 9.08   | 2  |
|       | AtSWEET15 | AT5G13170      | 879     | 292                | BrSWEET15a| Bra023394      | 2           | MF2       | 894     | 297                 | 33.156               | 8.26   | 2  |
|       |           |                |         | 32.936             | BrSWEET15b| Bra006185      | 3           | MF1       | 951     | 316                 | 35.814               | 8.86   | 2  |
|       |           |                |         |                    | BrSWEET15c| Bra008850      | 10          | LF        | 630     | 209                 | 23.314               | 7.74   | 2  |
| Group | Arabidopsis thaliana | Brassica rapa | Number of domain |
|-------|----------------------|--------------|-----------------|
|       | Gene name | Accession number | ORF(bp) | Deduced polypeptide | Protein length (aa) | MW(kD) | PI | Protein length (aa) | MW(kD) | PI | |
|       |           |               |         |                 |                      |        |    |                      |        |    | |
| N     | AtSWEET16 | AT3G16690 | 693 | 230 | 25.744 | 5.1 | BrSWEET18 | Bra012412 | 7 | MF2 | 528 | 175 | 20.093 | 9.38 | 2 |
|       |           |               |         |                 |                      |        |    |                      |        |    | |
|       | AtSWEET17 | AT4G15920 | 726 | 241 | 26813 | 6.1 | BrSWEET17a | Bra012752 | 3 | MF1 | 723 | 240 | 26.444 | 8.76 | 2 |
|       |           |               |         |                 |                      |        |    |                      |        |    | |
Table 2  The Non-synonymous substitution rate and synonymous substitution rate between orthologous SWEET gene pairs between *A. thaliana* and *B. rapa*

| Orthologous gene pairs | Non-synonymous substitution rate (Ka) | Synonymous substitution rate (Ks) | Ka/Ks |
|------------------------|--------------------------------------|----------------------------------|-------|
| AT1G21460 Bra016421    | BrSWEET1a 0.0400                     | 0.4300                           | 0.0930|
| Bra017916 BrSWEET1b    | 0.0400                                 | 0.4000                           | 0.1000|
| AT3G14770 Bra021577    | BrSWEET2a 0.0700                      | 0.3700                           | 0.1892|
| Bra027314 BrSWEET2b    | 0.0500                                 | 0.2700                           | 0.1852|
| AT5G53190 Bra022636    | BrSWEET3a 0.0900                      | 0.3600                           | 0.2500|
| Bra003075 BrSWEET3b    | 0.1200                                 | 0.3700                           | 0.3243|
| AT3G28007 Bra025317    | BrSWEET4a 0.0500                      | 0.3300                           | 0.1515|
| Bra033034 BrSWEET4b    | 0.0600                                 | 0.3900                           | 0.1538|
| AT5G62850 Bra035879    | BrSWEET5a 0.0600                      | 0.3800                           | 0.1579|
| Bra029245 BrSWEET5b    | 0.0800                                 | 0.3700                           | 0.2162|
| AT4G10850 Bra035270    | BrSWEET7a 0.1200                      | 0.5800                           | 0.2069|
| Bra000725 BrSWEET7b    | 0.1500                                 | 0.5300                           | 0.2830|
| AT5G40260 Bra025595    | BrSWEET8 0.2100                       | 1.2400                           | 0.1694|
| AT2G39060 Bra000116    | BrSWEET9 0.0600                       | 0.2800                           | 0.2143|
| AT5G50790 Bra022761    | BrSWEET10 0.1000                      | 0.5100                           | 0.1961|
| AT3G48740 Bra018039    | BrSWEET11a 0.0500                     | 0.2800                           | 0.1786|
| Bra019560 BrSWEET11b   | 0.0500                                 | 0.2900                           | 0.1724|
| AT5G23660 Bra009700    | BrSWEET12a 0.0600                     | 0.3400                           | 0.1765|
| Bra026487 BrSWEET12b   | 0.0400                                 | 0.2200                           | 0.1818|
| AT4G25010 Bra013854    | BrSWEET14a 0.0700                     | 0.3300                           | 0.2121|
| Bra010477 BrSWEET14b   | 0.0700                                 | 0.2900                           | 0.2414|
| AT5G13170 Bra023394    | BrSWEET15a 0.1100                     | 0.3700                           | 0.2973|
| Bra006185 BrSWEET15b   | 0.1100                                 | 0.3200                           | 0.3438|
| Bra008850 BrSWEET15c   | 0.0900                                 | 0.3100                           | 0.2903|
| AT3G16690 Bra01638     | BrSWEET16a 0.1600                     | 0.4600                           | 0.3478|
| Bra021190 BrSWEET16b   | 0.0800                                 | 0.3500                           | 0.2286|
| AT4G15920 Bra038060    | BrSWEET17b 0.0600                     | 0.3000                           | 0.2000|

saliva/SWEET genes in clade type A1 were 5 and 8, respectively, moreover, the number of putative common ancestor genes was 4; the number of genes gained were 4 and 1, respectively, and the number of genes lost were 0 and 0, respectively. Clade type A2 was the branch with the lowest numbers of MtN3/saliva/SWEET genes at 2 and 4 for *Arabidopsis* and *B. rapa*. These genes had 2 ancestor genes. The *Arabidopsis* MtN3/saliva/SWEET genes did not show gains and losses, whereas *B. rapa* gained 2 MtN3/saliva/SWEET genes.

Fig. 1 Locations of the MtN3/Saliva/SWEETs genes on the chromosomes of *B. rapa*. The black triangle denotes the transcriptional direction.
saliva/SWEET genes. *Arabidopsis* and *B. rapa* attained 3 and 7 MtN3/saliva/SWEET genes in clades type A3, and these genes had 3 putative common ancestor genes and gained 4 and 0 genes, respectively, and lost 0 and 0 genes, respectively. Clade type B1 was the largest group with 15 *B. rapa* MtN3/saliva/SWEET genes and 7 *Arabidopsis* MtN3/saliva/SWEET genes. They have 8 supposed ancestor genes, with gaining 7 genes gained in *B. rapa* and 1 gene lost in *A. thaliana*.

**Ka and Ks calculation of orthologous SWEET genes between *B. rapa* and *A. thaliana***

The ratio of the nonsynonymous substitution rate to the synonymous substitution rate is a significant indicator of the selection pressure acting on genes being assessed. Ka/Ks > 1, =1, and < 1 indicate positive Darwinian selection, neutral selection, and purification selection, respectively [34]. We calculated the Ka/Ks ratios of the orthologous SWEET genes between *B. rapa* and *A.*
**Structural analysis of MtN3/saliva/SWEET genes**

The results of the intron–exon location analysis showed that the number and distribution of the introns and exons of MtN3/saliva/SWEET genes were highly conserved (Fig. 4). A total of 26 members of the MtN3/saliva/SWEET family harbored six exons. BrSWEET2c has three exons. BrSWEET3b, BrSWEET7b, BrSWEET15c and BrSWEET18 contained four exons, BrSWEET7a BrSWEET14c and BrSWEET 15b contained five exons. BrSWEET17b comprised a special structure with six exons, three short introns and two long introns. The amino acid sequences of the 29 MtN3/saliva/ SWEET family genes were aligned with ClustalW. The results reveal that the intramembranous areas of the MtN3/saliva/SWEET domain were highly conserved. Furthermore, the transmembrane regions were relatively conserved, and the two highly conserved regions may have each contained a serine phosphorylation site (Additional file 4: Figure S2). Hence, the intramembranous areas of the MtN3/saliva/SWEET family protein may correspond to important functional or regulatory regions of the protein structure. The regulatory methods may be reversible phosphorylation/dephosphorylation pathways controlled by a number of protein kinases/phosphatases. This information can be an important entry point for the functional analysis of Arabidopsis MtN3/saliva/ SWEET family genes.

**Promoter sequences analysis of MtN3/saliva/SWEET genes**

To further understand the transcriptional regulation and potential function of the MtN3/saliva/SWEET genes, we analyzed the cis-elements in the promoter sequences (Additional file 5: Table S3). There are more than half cis-acting elements involved in light response elements in the development associated elements, these underlined elements are all light response elements. Most of MtN3/saliva/SWEET genes contained one or more cis-elements involved in photoresponse, these are ACE, Box4, BoxI, G-box, GAG-motif, GT1-motif, MRE, Sp1, TCCC-motif, ATCT-motif, TCT-motif, GAG-motif, and CATT-motif. Therefore, we hypothesized that the MtN3/saliva/SWEET family genes are involved in plant photosynthesis. Some MtN3/saliva/SWEET genes contained the cis-elements involved in hormonal response. A total of 14 MtN3/saliva/SWEET genes contained the ABRE element involved in ABA response. Then, 15 MtN3/saliva/SWEET genes contained the GARE-motif involved in gibberellin response. Meanwhile, 25 genes harbored the TGACG-motif and CGTCA-motif involved in the MeJA response. A total of 13 genes contained the TGA-element involved in auxin response. In contrast, 22 genes contained the TCA-element involved in the salicylic acid response. Otherwise, the scope of the stress-response-related cis-acting elements was relatively broad and included high-temperature stress, low-temperature stress, drought-induced stress, anerobic induction, and other related components. A total of 26 genes contained HSE involved in the high-temperature
stress response, whereas 17 genes contained LTR involved in the low-temperature stress response. Statistical analyses suggested that more than half of the cis-elements were light-responsive elements. About one-third were related to stress, and the rest were associated with growth and development. Hence, the MtN3/saliva/SWEET genes may be regulated by light signals. The cis-elements present in the promoter regions of the MtN3/saliva/SWEET genes revealed their essential role in growth and development, as well as in mediating responses to abiotic stresses.

**Analysis of the secondary structure of the MtN3/saliva/SWEET proteins in B. rapa**

NPSA-MLRC ([https://npsa-prabi.ibcp.fr/cgi-bin/npsa_automat.pl?page=/NPSA/npsa_mlrc.html)] [35] was employed to predict the secondary structure of the MtN3/saliva/SWEET proteins. The main secondary structures of the MtN3/saliva/SWEET proteins were the α-helix, irregular curl, and extended strand. Most of MtN3/saliva/SWEET proteins exhibited some similarities in the proportion of constituent secondary structural elements. That is, approximately 15%–40% were α-helices, approximately 40%–50% were irregular curls, and approximately 20%–40% were extended strands (Additional file 6: Table S4).

**Transmembrane regions and hydrophobicity analysis of the MtN3/saliva/SWEET proteins**

Using TMHMM software to predict the transmembrane regions of the 34 MtN3/saliva/SWEET proteins, we found that 24 members of the encoded protein harbored seven transmembrane regions. Five members contained six transmembrane regions, one harbored five transmembrane regions, and two members contained four transmembrane regions. These results imply that the MtN3/saliva/SWEET family proteins likely localize in the membrane or membrane receptors (Fig. 5).
The hydrophobicity analysis revealed that all the MtN3/saliva/SWEET proteins were hydrophilic proteins (Additional file 7: Table S5).

**Signal peptide analysis and subcellular localization analysis of the MtN3/saliva/SWEET family proteins**

Signal peptide analyses of the 34 MtN3/saliva/SWEET proteins were performed on the basis of the neural network model of the SignalP 4.1 Server. A total of 34 MtN3/saliva/SWEET proteins did not contain a signal peptide sequence.

To understand the location of the 34 MtN3/saliva/SWEET proteins in the cell, we performed subcellular localization. BrSWEET2a, BrSWEET16a, BrSWEET16b, BrSWEET17a, and BrSWEET17b are likely positioned at the vacuolar membrane, whereas BrSWEET1a, BrSWEET1b,
BrSWEET3a and BrSWEET3b may be located in the chloroplast. The rest are more likely located in the plasma membrane except BrSWEET2c, which may be located in extracellular domain (Additional file 8: Table S6).

**Conserved short amino-acid sequence analysis of the MtN3/saliva/SWEET proteins in B. rapa**

MEME analysis showed seven conserved motifs of MtN3/saliva/SWEET existing in B. rapa (Fig. 6 and Additional file 9: Figure S3). Most members in the Clade type A1, typeA2 and typeA3 have 6 motifs except motif 6, while most genes in Clade type B1 have 7 motifs. BrSWEET2c only has motif 1, 4 and motif 7; BrSWEET3b only has motif 3 and motif 4; BrSWEET7b only has motif 2, 4 and motif 5. These genes only have two or three motif, because they just have one domain. The result show that TypeA and TypeB have its own conservative motif and our classification is reasonable.

**Expression characterization of the MtN3/saliva/SWEET genes**

Tissue-specific and developmental stage-related expression data provides us with clues about the function of the MtN3/saliva/SWEET genes in different tissues and floral organs. Thus, we obtained an initial picture of the functions of the MtN3/saliva/SWEET genes during vegetative and reproductive development. To achieve this goal, we analyzed the transcript levels of the genes in the roots, stems, leaves, flowers, immature silique, sepals, petals, stamens, pistils, nectaries, and small and large buds by qRT-PCR.

From the qRT-PCR results of samples from different tissues (Fig. 7), we found that the expression patterns of most of the MtN3/saliva/SWEET genes were tissue specific, except for the amplified products from primers named 021577, 016421, 026487,025595, 009700, and 035879. Most of the expression differences were more than 100 times different. The transcripts of BrSWEET11c and BrSWEET11a were specifically expressed in the floral axis. Thus, the genes may be related to floral axis development. Meanwhile, the expression patterns of BrSWEET13, BrSWEET14a, BrSWEET14c, BrSWEET10, BrSWEET7, BrSWEET9, BrSWEET14a, BrSWEET5c, and BrSWEET15b were relatively specific in the flower. Thus, these genes may serve roles in the reproductive developmental process of the Chinese cabbage. BrSWEET15a, BrSWEET7, BrSWEET18, and BrSWEET1b were specifically expressed in silique and may be involved in reproductive development. In summary, the MtN3/saliva/SWEETs gene family may be involved in the reproductive developmental process of Chinese cabbage.
qRT-PCR of the MtN3/saliva/SWEET genes from the different floral organs showed that the gene expression levels of the different MtN3/saliva/SWEET genes evidently differed among the different flower organs (Fig. 7). The transcripts of *BrSWEET11c* and *BrSWEET11a* were expressed at a significantly higher level in sepals, petals, stamens, and nectaries than in the other parts of the flower. By contrast, *BrSWEET13a* was relatively highly expressed in the sepals, petals, stamens, and small flower buds. Some genes were specifically expressed in different floral organs. *BrSWEET13, BrSWEET7, BrSWEET1a, and BrSWEET9* were specifically expressed in the nectary. *BrSWEET14a, BrSWEET14c, and BrSWEET10* were specifically expressed in the stamen. *BrSWEET15b* was specifically expressed in the sepals. *BrSWEET15c* and *BrSWEET1b* were specifically expressed in the large and small flower buds. *BrSWEET4a* and *BrSWEET8* exhibited relatively high expression levels in various floral organs. By contrast, *BrSWEET12a, BrSWEET5a, BrSWEET12b, and BrSWEET18* revealed almost no expression in various flower organs.
Subcellular localization analysis and overexpression analysis of BcNS in B. rapa

To explore the subcellular localization of BcNS, we constructed a pB7YWG2.0-BcNS-YFP overexpression vector and transiently expressed the gene in onion epidermal cells via gene-gun bombardment. The transient expression of BcNS in onion epidermal cells showed that the protein was located on the cell membrane (Fig. 8). This result is consistent with the predicted results (Additional file 8: Table S6; Additional file 10: Figure S4).

Results of semi-quantitative PCR showed that BcNS was expressed in V buds and opened flowers and there was not significant difference between sterile plants and fertile plants (Fig. 9a). To further understand the expression pattern of the BcNs, we detected in expression in different flower organs. The results revealed that BcNS was specifically expressed in the stamen and nectary (Fig. 9b). These results are consistent with quantitative reverse-transcription PCR (Fig. 9c). Meanwhile, result of in situ hybridization indicated that BcNS was expressed in nectary tissue (Fig. 9d).

We also constructed the antisense expression vector pCAMIA1301-BcNS and transformed this vector into B. rapa by Agrobacterium-mediated transformation to study the biological function of BcNS (Additional file 11: Figure S5). In this study, the transgenic plants were detected by PCR, X-Gluc, and fluorescence quantitative PCR. These methods provided reliable materials for functional verification (Additional file 12: Figure S6). We obtained 45 transgenic germ lines and 22 positive lines after PCR detection. The positive rate was approximately 50%. We detected the calluses and leaves of the transgenic plants with the X-Gluc histochemical assay. The calluses were stained blue, and the leaves were stained with difficulty. Hence, the antisense BcNS gene fragment was integrated into the genomic DNA in B. rapa. The transgenic plants detected by PCR and GUS staining were analyzed by qRT-PCR. Expression in the transgenic plants decreased relative to that in the negative control.

We observed the flower organ development of the transgenic plants, controls, and wild types. Unlike in the wild type, the petals, stamens, and pistils in the transgenic plants and the controls were normal. Nevertheless, we found splitting phenomena at the base of the lateral nectarines (Fig. 10 and Additional file 13: Figure S7), and the proportion of affected nectaries exceeded 70% (Table 3).

Discussion

MtN is M. truncatula Nodulin and was first reported in 1996 [3]. A new sugar transporter family was found by sugar optical sensors and named the SWEET family in 2010 [7]. Afterward, SWEETs and the gene family MtN3/saliva were reported to be the same [6]. Currently, relatively large-scale studies are available on MtN3/saliva/SWEET involved in plant pathogen invasion and reproductive development. The MtN3/saliva/SWEET family proteins are involved in anther dehiscence, pollen wall formation, and other gametophyte developmental processes in Arabidopsis, petunia, and rice [9–11].

The genus Brassica is known for its important economic crops, such as Chinese cabbage, cabbage, broccoli, cauliflower, rutabaga, turnip and some seeds used in the production of canola oil and condiment mustard. To date, there are a number of gene families that have been identified in Brassica [22, 36, 37]. In this study, we successfully identified 34 MtN3/saliva/SWEET genes based on the Arabidopsis Genome Database TAIR and Brassica database. There genes were termed as Type-A1, Type-A2, Type-A3, and Type-B1in the same way as rice and tomato [38, 39]; the clades respectively contained 8, 4, 7, and 15 members in B. rapa, respectively. Physiological and biochemical property analyses revealed that the total lengths of the coding regions of the genes were 350–1000 bp, the molecular weights were 14–35 kDa,
and the pIs were 7.59–9.62. The 34 genes were distributed on various chromosomes. After analyzing the cis-acting elements of the promoters of these genes, we found that most were related to optical response. Therefore, we hypothesized that the MtN3/saliva/SWEET are likely involved in plant photosynthesis. Gene Ontology analysis of the turnip nectary ESTs, floral organ ESTs, root ESTs, and 12,448 tomato nectary ESTs showed that the photosynthesis-related lateral nectary ESTs (MLN ESTs) were even richer than the median nectary ESTs (MMN ESTs) [18]. Although numerous floral organ ESTs and tomato nectary ESTs exist, other biological processes are more prominent than the photosynthetic process. Most turnip nectary-specific genes are involved in photosynthesis. Therefore, the results of the cis-acting element analysis are highly meaningful in the study of Brassica nectar carbohydrates.

After predicting the transmembrane regions of the MtN3/saliva/SWEET proteins, we found that all the members comprised transmembrane regions. That is, 24 members contained seven transmembrane regions, and the extracellular regions of these proteins were generally very short. The intramembrane areas were highly conserved regions, whereas the transmembrane regions were relatively conserved. The two highly conserved regions may harbor amino acid phosphorylation sites. These results indicate that the intramembrane areas are important functional areas of the protein. Furthermore, the regulatory method may be reversible phosphorylation/dephosphorylation pathways controlled by a number of protein kinases/phosphatases. These findings can be an important entry point for the functional analysis of the MtN3/saliva/SWEET family genes.

Evolutionary analysis showed that some genes with known physiological functions exist in a relatively distant branch on the evolutionary tree. These genes possibly play a separate role in the developmental process. Our results showed MtN3/saliva/SWEET gene family of B. rapa diverged from A. thaliana at around 10 MYA to 13 MYA, which is a little later than a previous study, which have confirmed the genome of B. rapa diverged from A. thaliana lineage at least 13 ~ 17 MYA [40–42]. In the previous study, the result was based on the whole genome, while our results was gained from the MtN3/saliva/SWEET gene family which only have 34 members. As we known, A. thaliana and Brassicaceae taxa have undergone three times (γ, β and α) of the whole genome duplication or triplication (WGD or WGT) [43]. After the α duplication, Arabidopsis lineage separated from its Brassica ancestor approximately 43.2 MYA [40]. In the long river of species evolution, the difference of the divergence time is acceptable. These results hint that the MtN3/saliva/SWEET gene family is a good molecular reference candidate to be utilized in the study of plant evolution analysis. Our results further confirmed the hypothesis that Brassica genomes have undergone another whole genome triplication (WGT) after speciation from A. thaliana. The existence of gene duplications in Chinese cabbage and Arabidopsis that occurred during 3R also demonstrated their conservation during the long-term evolution [40, 44]. Meanwhile, another obvious distribution peak of the relative Ks appeared 1.24, which was responsible for the 3R event before the speciation between Brassica and Arabidopsis genomes. In addition, MtN3/saliva/SWEET family genes involve different physiological functions in plants, including reproductive
development, senescence, abiotic stress response, and host–pathogen interactions. At least seven members (AtSWEET2, AtSWEET4, AtSWEET7, AtSWEET8, AtSWEET10, AtSWEET12, and AtSWEET15) can be induced by the genus Pseudomonas DC3000 in Arabidopsis [45]. Moreover, AtSWEET4, AtSWEET15, AtSWEET17, AtSWEET11, and AtSWEET12 can be induced by other elicitor types [45, 46]. RPG1 (AtSWEET8) is involved in the pollen wall development process [11]. SAG29 (AtSWEET15) is mainly expressed through the abscisic acid pathway depending on the osmotic pressure in plant senescent tissues [12]. Silencing Xa13/Os8N3/OsSWEET11 can significantly increase the resistance to PXO99 (Xa13 nonaffinity strains) in rice; thus, Xa13 is essential for PXO99 in rice [7]. These findings

Table 3 The result of abnormal nectary development between the transformants and their controls

| Materials         | No. of flower with normal nectary | No. of flower with abnormal nectary | Ratio of flower with abnormal nectary(%) |
|-------------------|-----------------------------------|-------------------------------------|------------------------------------------|
| WT                | 34.67 ± 4.16                      | 6.33 ± 0.58                         | 15.50% ± 1.19%                          |
| CK                | 35.00 ± 2.94                      | 7.00 ± 0.81                         | 16.70% ± 1.76%                          |
| Transformants of A-12 | 12.33 ± 2.52 | 31.67 ± 4.73 | 71.80% ± 6.23%                          |
| Transformants of A-13 | 13.33 ± 3.21 | 35.00 ± 2.65 | 72.49% ± 5.91%                          |

Three plantlets and about 15 flowers per plantlet have been employed to investigate the abnormal nectary development, respectively. Thereinto, the transformant of A-20 damped off before flowering, and did not get the observation data.

Fig. 10 Scanning electron microscopy observation of the nectaries in the transgenic plants and their control in B. rapa ssp. chinensis var. parachinesis. a–c are wild types, d–g are pCAMBIA transgenic plants, h–i are pCAMBIA-BcNS transgenic plants. The arrow in H refers to the forking of the nectary development. The arrows in b, e, and G refer to the stoma of the nectary. The results indicate that not all of the nectaries were constantly open (f); some were closed (i) or half-closed (c)
provide information for future biological function research on other MtN3/saliva/SWEETs family genes.

The gene expression levels of the different MtN3/saliva/SWEET genes evidently differed among different tissues and flower organs. Many genes were specifically expressed in different floral organs. The MtN3/saliva/SWEET gene family was found to be involved in reproductive development in Chinese cabbage. RPG1 (RIP-TURED POLLEN GRAIN1/Sweet8) encodes a membrane protein in Arabidopsis; this protein is essential for small exine formation [11]. One study showed that RPG1 affects cell integrity, whereas the fertility of mutations are partially repaired during late reproductive development. The authors speculated that the gene was partially supplemented by RPG2. These results indicated that RPG1 and RPG2 interact in pollen wall sediment formation and consequently affect pollen wall formation [47]. In A. thaliana and rice, expression pattern analysis revealed additional genes from the MtN3/saliva/SWEET family that were involved in reproductive development. At least six SWEET genes were highly expressed in flowers at different developmental stages in rice. A number of six SWEET genes were highly expressed in flowers at that were involved in reproductive development. At least additional genes from the MtN3/saliva/SWEET family are closely related to reproductive development in Chinese cabbage. This result may be due to the neofunctionalization or subfunctionalization of the genes.

Protein function is closely related to protein subcellular localization [54]. To determine the location of genes related to nectary development in B. rapa, we studied the subcellular localization characteristics of the BcNS gene by the YFP-labeling method based on the Chinese cabbage cultivar ‘Chiifu-401.’ The protein was located on the cytoplasmic membrane; hence, the gene may play a role in the stability of the cell membrane or the control of protein transport and other physiological and biochemical processes.

The expression pattern of BcNS in different stages of flower development showed that the gene was expressed in fifth buds. There was no significant difference between sterile and fertile lines, which indicates that the gene may not be involved in pollen fertility. To further prove our conjecture, we examined the expression of the gene in different flower organs of ‘Beajh97-01A/B’ and ‘Chiifu-401’. The results showed that the gene is specifically expressed in nectaries and stamens. One MtN3/saliva family gene, NECI, is also expressed in nectaries and stamens, and plays an important role during anther development. We had an assumption that BcNS may be related to nectaries and anther development in Chinese cabbage. The antisense genetic method has been used to studying the function of the MtN3/saliva family genes [10, 11, 55]. We cloned an MtN3/saliva gene and named the gene as BcNS. This gene is highly expressed in the nectaries and stamen. We then constructed an antisense expression vector pCAMIA1301-BcNS and obtained transgenic plants. Morphological observations revealed a splitting phenomenon at the base of the lateral nectaries of the transgenic plants. However, detailed mechanisms of the phenotype remains to be further studied. Nectaries secrete nectar to attract insects for pollination [56]. The nectary is necessary for seeds formation, and the removal of the nectary from the flower of Christmas
bells caused an increase in nectar production, but a decrease in the ability to produce seeds [57]. Many studies suggest that nectary development is dependent on hormonal signaling. For example, PIN6, an auxin efflux transporter factor, positively regulated nectar production [58]. Jasmonic acid (JA) can promote nectar secretion [59]. Additionally, overexpression of GA2ox6 (GA 2-oxidase6) increased nectar sugar in nectaries, suggesting that GA negatively regulates nectar production [60].

Previous studies similarly reported on the nectarine split. NEC1 had differential expression in petunia with high expression in anther dehiscence cells, nectaries, and upper stamen filaments. Nec1 possesses an early another dehiscence forward phenotype [9]. The YABBY transcription factor genes CRABS CLAW (CRC) plays a role in the early development of nectaries and carpels in Arabidopsis [61]. Some genes that regulate CRC expression are also involved; examples include the B-type gene (APETALA3 and PISTILLATA) and the C-type gene (AGAMOUS and SEPELLAATA). The study showed that CRC expression was very high in the early stage of development and throughout the whole secretory period [62]. Therefore, CRC plays an important role in the development and secretion of nectar in nectaries. In addition, bop1 and bop2 double mutants could not form a normal nectary, but the expression of CRC in these mutants is normal [15]. We speculated that the coordinated interaction of the three genes is the basis for the production of a normal nectary. In future research, we intend to focus on analyzing whether CRC, BOP1, BOP, and C-type genes (AGAMOUS and SEPELLAATA) are related to the changes in the nectary structures of antisense BcNS transgenic plants. The results of such study could provide important clues to clarify the molecular mechanism in Chinese cabbage.

Conclusion

In this study, 34 MtN3/saliva/SWEETs genes were identified in B. rapa. We found that the full lengths of the coding regions ranged from 387 bp to 951 bp and encoded 128 amino acids to 316 amino acids. The Ka/Ks ratio of orthologous SWEET genes between B. rapa and A. thaliana implies that the genes suffered strong purifying selection for retention. Ks values concentrated location between 0.30 and 0.40, after calculating, B. rapa diverged from A. thaliana at around 10 MYA to 16 MYA. qPCR results indicated that the MtN3/saliva/SWEETs genes family may be involved in the reproductive developmental process of Chinese cabbage. Our study shows that BcNS play an important role in the development of the floral nectary in plant.

Additional files

**Additional file 1:** Table S1. The MtN3/saliva domain sequences of A. thaliana. (XLSX 27 kb)

**Additional file 2:** Table S2. Primer sequences for qRT-PCR analysis. (XLSX 11 kb)

**Additional file 3:** Figure S1. Copy number changes in the B. rapa and Arabidopsis MtN3/saliva/SWEET genes in Clades A–D. The numbers in ellipses and rectangles represent the numbers of MtN3/saliva/SWEET genes in the extant and ancestral species, respectively. The numbers on the branches with plus and minus symbols represent the numbers of gene gains and losses, respectively. (TIF 953 kb)

**Additional file 4:** Figure S2. Multiple alignment analysis of the MtN3/saliva/SWEET gene family in B. rapa. The black box represent the MtN3/saliva/SWEETs domain. The thick lines represent the transmembrane domain. The thick gray line represent the conservative intracellular region. (TIF 577 kb)

**Additional file 5:** Table S3. Putative cis-elements in promoter region of the 34 MtN3/saliva/SWEETs genes in B. rapa. (XLSX 14 kb)

**Additional file 6:** Table S4. Secondary structural prediction of the MtN3/saliva/SWEETs proteins of B. rapa. (XLSX 52 kb)

**Additional file 7:** Figure S5. Hydrophobic character prediction based on the Kyte–Doolittle scale for the deduction of MtN3/saliva/SWEETs proteins in B. rapa. (XLSX 59 kb)

**Additional file 8:** Table S5. Protein subcellular localization prediction of MtN3/saliva/SWEETs proteins in B. rapa. (XLSX 11 kb)

**Additional file 9:** Figure S3. WebLogo of the most conserved consensus motifs of the amino acids of B. rapa. (TIF 1081 kb)

**Additional file 10:** Figure S4. Prediction of transmembrane helices in B. rapa. (GIF 11 kb)

**Additional file 11:** Figure S5. Obtained transgenic plantlets of BcNS. A: Sowm seeds; B: seedling at 4–5 days; C: cotyledon–hypocotyl explants during preculture; D: E: cotyledon–hypocotyl explants during differentiation; F: grown seedling; G: H: roots induced from the HygR seedling; and I: regenerated plants transferred to the plot. (TIF 13426 kb)

**Additional file 12:** Figure S6. Positive detection of transgenic Chinese cabbage (B. campestris ssp. chinensis var. parachinensis), A, PCR amplification detection. Lane 1, marker; Lane 2, amplification results of the negative control, water; Lane 3, amplification results of the positive control, 35S-pCAMBIA1301; Lanes 4–8, amplification results of the positive control, 35S-pCAMBIA1301. Lanes 9–19 amplification results of the 35 s-BcNS transformants. B and C, X-Gluc histochemical staining detection of calluses and leaves. D, Fluorogenic quantitative PCR detection; WT, wild type; CK, negative control; A-12, A-13, and A-20, antisense expression plant. (TIF 1610 kb)

**Additional file 13:** Figure S7. Morphology observation of the flowers of B. rapa ssp. chinensis var. parachinensis transgenic plants. A–D are wild types, E–H are pCAMBIA transgenic plants, and I–L are pCAMBIA-BcNS transgenic plants. The arrows refer to the nectaries. (TIF 1003 kb)

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**Abbreviations**

- BcNS: Brassica Nectary and Stamens
- F: Flower
- GSDS: Gene Structure Display Server
- L: Leave
- LB: large bud
- MtNS: Medicago truncatula Nodulin15
- NCBI: National Center for Biotechnology Information
- Nt: nectary
- Pe: petal
- Pi: pistil
- qRT-PCR: Quantitative reverse-transcription PCR
- R: root
- SB: small bud
- Se: sepal
- Si: Silique
- St: stamen
- St: stem
- WGT: whole-genome triplication

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Availability of data and materials
Plant materials are available under request to the respective owner institutions. The datasets supporting the results of this article are included within the article and its additional files.

Authors’ contributions
LMM, XYL, LH, and QZC carried out the experiment, analyzed the data, and drafted the manuscript. LJK, CQC, SYW, JBL, and JL performed the qRT-PCR analysis and in situ hybridization. LMM, FH2, and JSC conducted the sequence analysis and wrote the manuscript. XLY designed and supervised the research and gave final approval of the version to be published. All the authors read and approved the final manuscript.

Ethics approval and consent to participate
Not applicable. This study has not directly involved humans, animals. The Ethics approval and consent to participate approved the final manuscript and gave final approval of the version to be published. All the authors read and approved the final manuscript.

Consent for publication
Not applicable.

Competing interests
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