A Genome-Wide Survey of MATE Transporters in Brassicaceae and Unveiling Their Expression Profiles under Abiotic Stress in Rapeseed

Cailin Qiao 1,2,†, Jing Yang 1,2,†, Yuanyuan Wan 1,2, Sirou Xiang 1,2, Mingwei Guan 1,2, Hai Du 1,2,‡, Zhihong Tang 1,2, Kun Lu 1,2,*, Jiana Li 1,2,* and Cunmin Qu 1,2,* 

1 Chongqing Engineering Research Center for Rapeseed, College of Agronomy and Biotechnology, Southwest University, No. 2 Tiansheng Road, Beibei, Chongqing 400715, China; qcl123@email.swu.edu.cn (C.Q.); yj18203807@email.swu.edu.cn (J.Y.); wyywinyuan@email.swu.edu.cn (Y.W.); x231915@email.swu.edu.cn (S.X.); ghhoo1770@email.swu.edu.cn (M.G.); dh20130904@swu.edu.cn (H.D.); tangzhlin@swu.edu.cn (Z.T.); drlukun@swu.edu.cn (K.L.)
2 Academy of Agricultural Sciences, Southwest University, Chongqing 400715, China
* Correspondence: ljn1950@swu.edu.cn (J.L.); drqucunmin@swu.edu.cn (C.Q.); Tel.: +86-23-6825-0701 (C.Q.)
† These authors contributed equally to this work.

Received: 16 July 2020; Accepted: 17 August 2020; Published: 20 August 2020

Abstract: The multidrug and toxic compound extrusion (MATE) protein family is important in the export of toxins and other substrates, but detailed information on this family in the Brassicaceae has not yet been reported compared to Arabidopsis thaliana. In this study, we identified 57, 124, 81, 85, 130, and 79 MATE genes in A. thaliana, Brassica napus, Brassica oleracea, Brassica rapa, Brassica juncea, and Brassica nigra, respectively, which were unevenly distributed on chromosomes owing to both tandem and segmental duplication events. Phylogenetic analysis showed that these genes could be classified into four subgroups, shared high similarity and conservation within each group, and have evolved mainly through purifying selection. Furthermore, numerous B. napus MATE genes showed differential expression between tissues and developmental stages and between plants treated with heavy metals or hormones and untreated control plants. This differential expression was especially pronounced for the Group 2 and 3 BnaMATE genes, indicating that they may play important roles in stress tolerance and hormone induction. Our results provide a valuable foundation for the functional dissection of the different BnaMATE homologs in B. napus and its parental lines, as well as for the breeding of more stress-tolerant B. napus genotypes.

Keywords: MATE transporters; Brassicaceae; Brassica napus; phylogenetic analysis; abiotic stress; expression profiles

1. Introduction

In nature, plants are exposed to various exogenous and endogenous toxins during their lifespans, and they have developed a series of adaptive response mechanisms to dispose of and detoxify toxic compounds [1]. A group of secondary transporters first reported in Vibrio parahaemolyticus and Escherichia coli [2,3], the MATE (multidrug and toxic compound extrusion or multi-antimicrobial extrusion)/DTX (detoxification efflux carrier) family, is a universal family of proteins involved in the metabolism of toxic compounds and organic acids. Members of the MATE family are composed of 400–700 amino acids and contain 12 transmembrane helices [1,4]. However, the MATE gene family lacks a conserved consensus sequence, and MATE proteins share only about 40% sequence similarity [1]. Numerous studies indicate that MATE family proteins play widespread roles in various biological processes in plants, such as secondary metabolite transport [5–10], disease resistance [11,12],
the detoxification of heavy metals [13–15], and the transport of plant hormones [16–18]. Numerous studies have characterized MATE genes from a wide range of living organisms, showing that they have even greater diversity in plant genomes than in mammalian genomes [19]. For example, 56 MATE genes have been isolated from Arabidopsis thaliana [13]. The MATE gene family has also been widely characterized in other plant species, including rice (Oryza sativa) (52) [20,21], maize (Zea mays) (49) [22], tomato (Solanum lycopersicum) (67) [4], upland cotton (Gossypium hirsutum) (72) [23], and soybean (Glycine max) (117) [24]. Recently, 67 MATE genes were identified in the Brassica rapa genome [25]. However, genome-wide analysis of the MATE family in Brassicaceae species has been limited, and no such studies have been reported in rapeseed (Brassica napus).

In the Brassicaceae, the relationships of three diploid species (B. rapa, AA, 2n = 20; B. nigra, BB, 2n = 16; and B. oleracea, CC, 2n = 18) and three amphidiploid plants (B. juncea; AABB, 2n = 36; B. napus, AACC, 2n = 38; and Brassica carinata, BBCC, 2n = 34) have been thoroughly described according to the "triangle of U" theory [26,27], which also provides an excellent evolutionary model to investigate the expansion of gene families [28,29]. During the course of evolution, the Brassicaceae have experienced α and β duplication events, as well as a trilicitation event specific to the Brassica clade compared with the model plant A. thaliana [30]. These whole-genome duplication (WGD) events, along with the merger of the two progenitor genomes, have resulted in copious gene duplication in the B. napus genome, followed by substantial gene loss [31]. Thus, Brassica is regarded as an ideal model for investigating the evolution of polyploidy [29]. However, no systematic and comprehensive study of this family is reported among Brassicaceae species. To date, with the completion of five Brassicaceae species (B. napus, B. oleracea, B. rapa, B. juncea, and B. nigra) genome sequencing [29,32–36], an investigation of the MATE gene family in Brassicaceae species has been feasible and can not only provide insights into the evolutionary mechanisms of hybridization (allopolyploidy) among these species, but also lay a theoretical basis for a clearer understanding of the mechanisms involved in plant stress defense.

In present study, we identified MATE family genes in the Brassicaceae genomes available in the Brassica database (BRAD) [37] using the 57 MATE sequences from A. thaliana as queries. Then, we performed a detailed phylogenetic analysis of the structure, motifs, chromosome distribution, and duplication of these genes, as well as an evolutionary analysis. We also investigated the expression profiles of MATE genes in different tissues of B. napus and during responses to hormone treatment and heavy metal stress. Our results provide important information about the origin and evolution of the MATE family in Brassicaceae and lay a foundation for the further functional characterization of MATE genes in B. napus.

2. Results

2.1. Identification of MATE Transporters in Brassicaceae Species

Using 57 AtMATE protein sequences as queries, we identified 499 putative MATE genes from various Brassicaceae species, including 124 BnaMATE, 85 BraMATE, 81 BolMATE, 79 BniMATE, and 130 BjuMATE genes. Based on their homology relationship with the respective A. thaliana genes, these genes were named BnaMATE1 to BnaMATE57 for B. napus, BraMATE1 to BraMATE57 for B. rapa, BolMATE1 to BolMATE57 for B. oleracea, BniMATE1 to BniMATE57 for B. nigra and BjuMATE1 to BjuMATE57 for B. juncea, respectively (Figure 1, Table S2). Eight AtMATE genes (AtMATE2, AtMATE10, AtMATE11, AtMATE20, AtMATE22, AtMATE26, AtMATE32, and AtMATE57) had no orthologs among the other five species, which was indicative of gene loss during evolution between A. thaliana and Brassica species.

Basic information about these MATE genes in Brassicaceae species, including name, locus, molecular weight, subcellular localization, and other related information, is listed in Table S2. In B. napus, we found that BnaMATE genes encode proteins of 124 (BnaMATE56d) to 971 (BnaMATE15c) amino acids, with an average length of 489 amino acids. The relative molecular weights (MWs) ranged from 13.07 kDa (BnaMATE56d) to 107.53 kDa (BnaMATE15c), and theoretical pI (isoelectric point)
values ranged from 4.89 ($BnaMATE40d$) to 10.55 ($BnaMATE17d$). Similarly, individual MATE proteins varied in length and physicochemical properties among the other five species (Table S2). These results suggest that MATE transporters might have changed between species to serve different functions.

**Figure 1.** Phylogenetic analysis of the multidrug and toxic compound extrusion (MATE) transporters. The rooted neighbor-joining phylogenetic tree was constructed using MEGA7.0 and visualized using Figure Tree v1.4.2. There are 556 MATEs in *A. thaliana* and five Brassicaceae species, including 57 AtMATEs, 124 BnaMATEs, 85 BraMATEs, 81 BolMATEs, 79 BniMATEs, and 130 BjuMATEs. These MATE transporters are divided into four subgroups, Groups 1–4, which are marked in yellow, green, red, and purple, respectively. The outer circle represents the four groups.

Previous studies have localized MATE transporters to the plasma membrane, chloroplast, cytoplasm, vacuole, and endoplasmic reticulum, as well as extracellularly [24,38,39]. Here, we found the highest proportion localized in the plasma membrane (109 genes), accounting for >80% of all *BnaMATE* genes in *B. napus*; four genes were expressed in chloroplasts and 11 genes were expressed in the vacuoles. The trend in the other four species was similar, with 91.8% (*B. rapa*), 81.5% (*B. oleracea*), 91.5% (*B. juncea*), and 83.5% (*B. nigra*) of MATE genes, respectively (Table S2), expressed in the plasma membrane, which was consistent with previous results [24,39]. This strong representation of MATEs in
the plasma membrane is consistent with their primary role of maintaining membrane integrity through the exclusion of toxins from plants.

2.2. Phylogenetic Analysis of the MATE Transporters

To investigate the evolutionary relationships among MATE transporters between *A. thaliana* and various Brassicaceae species, we divided the 556 MATE transporters into four categories, denoted Groups 1–4 (Table 1 and Figure 1). Group 2 was the largest clade, with 230 members (24 in *A. thaliana*, 51 in *B. napus*, 37 in *B. rapa*, 34 in *B. oleracea*, 44 in *B. juncea*, and 40 in *B. nigra*), and Group 3 was the smallest, with 59 members (6 in *A. thaliana*, 15 in *B. napus*, 10 in *B. rapa*, 7 in *B. oleracea*, 15 in *B. juncea*, and 6 in *B. nigra*); Groups 1 and 4 contained 137 and 130 members, respectively (Figure 1). All of the MATE transporters we identified in Brassicaceae were classified into previously defined subgroups along with *A. thaliana* MATE proteins, suggesting that they might share similar functions within the subfamilies. In addition, the classifications were consistent with those of soybean MATE transporters [1,23,24], indicating that the classification adopted in this study was accurate and conforms to previous findings about plant MATE proteins.

2.3. Chromosomal Location of MATE Genes in Brassicaceae Genomes

Based on the physical positions of these MATE genes in the Brassicaceae genomes, 443 of the 499 full-length MATE genes were accurately mapped onto 64 chromosomes among the five Brassicaceae species (202 in the A∗ genome, 122 in the B∗ genome, and 119 in the C∗ genome) (Figure 2), whereas 56 MATE genes were assigned to random chromosomes (12), scaffold chromosomes (34), and contig chromosomes (10). Among them, 56 and 56 BnaMATE genes were distributed on the A∗ and C∗ subgenomes of *B. napus*; 61 and 57 BjuMATE genes were mapped to the Aju and Bju subgenomes of *B. juncea*; and 85 BraMATE genes were located on the Ara subgenomes of *B. rapa*, 65 BniMATE genes were located on the Bni subgenomes of *B. nigra*, and 63 BolMATE genes were located on the Col subgenomes of *B. oleracea* (Figure 2). However, the sum of the MATE genes from *B. rapa* (AA) and *B. oleracea* (CC) or from *B. rapa* (AA) and *B. nigra* (BB) in each subgenome was larger than that from *B. napus* (AACC) and *B. juncea* (AABB), suggesting that the MATE genes underwent a notable gene loss event during the evolution of *B. napus*, *B. juncea*, and their relatives. In addition, these genes were unevenly distributed on the chromosomes. For instance, regions dense in MATE genes were discovered on chromosomes A03, C01, and C07 in *B. napus* (Figure 2). In *B. oleracea*, chromosome C03 contained the greatest number of MATE genes, 12, whereas C04 and C05 contained three MATE genes. In *B. rapa*, chromosome A03 contained the most MATE genes, 15, whereas A04 contained two MATE genes. In *B. nigra*, chromosome B7 contained the most MATE genes, 17, and B6 contained one. In *B. juncea*, chromosome B03 contained the most MATE genes, 15, whereas A04 and A10 had only one (Figure 2), suggesting that they were unevenly distributed on each chromosome of Brassicaceae species.
| Type | Family     | A. thaliana | B. napus | B. oleracea | B. rapa | B. juncea | B. nigra |
|------|------------|-------------|----------|-------------|---------|-----------|---------|
| MATE1| AimMATE1   | 1           | /        | 1           | 2       | /         | /       |
| MATE1| AimMATE2   | /           | /        | /           | /       | /         | /       |
| MATE1| AimMATE3   | /           | 1        | /           | 1       | /         | /       |
| MATE1| AimMATE4   | /           | 1        | 1           | /       | /         | 1       |
| MATE1| AimMATE5   | /           | /        | /           | 1       | /         | 1       |
| MATE1| AimMATE6   | 2           | /        | 1           | 1       | /         | /       |
| MATE1| AimMATE7   | 1           | /        | 1           | 2       | 2         | /       |
| MATE1| AimMATE8   | 2           | 3        | 1           | 2       | 1         | /       |
| MATE1| AimMATE9   | 2           | 2        | 2           | 2       | 1         | /       |
| MATE1| AimMATE10  | /           | /        | /           | /       | /         | /       |
| MATE1| AimMATE11  | /           | /        | /           | /       | /         | /       |
| MATE1| AimMATE12  | 2           | 1        | 1           | /       | /         | /       |
| MATE1| AimMATE13  | /           | 2        | 2           | 5       | 3         | /       |
| MATE1| AimMATE14  | 4           | 5        | 4           | 7       | 5         | /       |
| MATE1| AimMATE15  | 4           | 2        | 1           | 2       | 2         | /       |
| MATE1| AimMATE16  | 4           | 2        | 3           | 3       | 2         | /       |
| MATE1| AimMATE17  | 4           | 2        | 2           | 3       | 3         | /       |
| MATE1| AimMATE57  | /           | /        | /           | /       | /         | /       |
| Group 1 | MATE2 | AimMATE18  | /        | /           | /       | 3         | /       |
| Group 2 | MATE2 | AimMATE19  | 3        | 2           | 2       | 2         | 5       |
| Group 2 | MATE2 | AimMATE20  | /        | /           | /       | /         | /       |
| Group 2 | MATE2 | AimMATE21  | 1        | 1           | 2       | 2         | 1       |
| Group 2 | MATE2 | AimMATE22  | /        | /           | /       | /         | /       |
| Group 2 | MATE2 | AimMATE23  | /        | /           | 1       | 1         | 1       |
| Group 2 | MATE2 | AimMATE24  | 4        | 2           | 2       | 3         | 1       |
| Group 2 | MATE2 | AimMATE25  | 2        | 2           | 2       | 3         | 2       |
| Group 2 | MATE2 | AimMATE26  | /        | /           | /       | /         | /       |
| Group 2 | MATE2 | AimMATE27  | 5        | 5           | 4       | 2         | 4       |
| Group 2 | MATE2 | AimMATE28  | 3        | 1           | 1       | /         | /       |
| Group 2 | MATE2 | AimMATE29  | 3        | 1           | 1       | 2         | 1       |
| Group 2 | MATE2 | AimMATE30  | /        | /           | 1       | /         | /       |
| Group 2 | MATE2 | AimMATE31  | 2        | /           | 1       | 2         | 2       |
| Group 2 | MATE2 | AimMATE32  | /        | /           | /       | /         | /       |
| Group 2 | MATE2 | AimMATE33  | 2        | 1           | 1       | 1         | 1       |
| Group 2 | MATE2 | AimMATE34  | 2        | 1           | /       | 2         | 1       |
| Group 2 | MATE2 | AimMATE35  | 4        | 3           | 3       | 5         | 3       |
| Group 2 | MATE2 | AimMATE36  | 3        | 3           | 2       | 2         | 2       |
| Group 2 | MATE2 | AimMATE37  | 4        | 2           | 2       | 2         | 3       |
| Group 2 | MATE2 | AimMATE38  | /        | 2           | 4       | 2         | 2       |
| Group 2 | MATE2 | AimMATE39  | 5        | 3           | 2       | 4         | 4       |
| Group 2 | MATE2 | AimMATE40  | 6        | 4           | 5       | 3         | 4       |
| Group 2 | MATE2 | AimMATE41  | 2        | 1           | 1       | 3         | 1       |
| Group 3 | MATE3 | AimMATE42  | 2        | 2           | 2       | 3         | 2       |
| Group 3 | MATE3 | AimMATE43  | 6        | 1           | 3       | 8         | 3       |
| Group 3 | MATE3 | AimMATE44  | 2        | /           | 1       | 2         | 1       |
| Group 3 | MATE3 | AimMATE45  | /        | 1           | 1       | /         | /       |
| Group 3 | MATE3 | AimMATE46  | 2        | 2           | 2       | 1         | /       |
| Group 3 | MATE3 | AimMATE47  | 3        | 1           | 1       | 1         | /       |
| Group 4 | MATE4 | AimMATE48  | 2        | 2           | 1       | 3         | 1       |
| Group 4 | MATE4 | AimMATE49  | 3        | 2           | 2       | 4         | 2       |
| Group 4 | MATE4 | AimMATE50  | 3        | 2           | 2       | 6         | 1       |
| Group 4 | MATE4 | AimMATE51  | 4        | 3           | 3       | 6         | /       |
| Group 4 | MATE4 | AimMATE52  | 4        | /           | 2       | 5         | /       |
| Group 4 | MATE4 | AimMATE53  | 4        | 2           | 2       | 4         | 2       |
| Group 4 | MATE4 | AimMATE54  | 4        | 2           | 2       | 3         | 2       |
| Group 4 | MATE4 | AimMATE55  | 3        | 2           | 2       | 5         | 2       |
| Group 4 | MATE4 | AimMATE56  | 5        | 4           | 2       | 4         | 2       |
2.4. Synteny and Gene Duplication Analysis of MATE Transporters

To explore the entire evolutionary history of each MATE genes, we performed comparative analysis of MATE genes between A. thaliana and five Brassicaceae genomes (Figure 3a) and determined their patterns of orthologous gene retention or loss based on their collinearity relationships. In total, 40 A. thaliana MATE genes showed collinear relationships with the five Brassicaceae species. Among them, we identified 30 collinear gene pairs between A. thaliana and B. napus, 17 collinear gene pairs between A. thaliana and B. oleracea, 29 collinear gene pairs between A. thaliana and B. rapa, 33 collinear gene pairs between A. thaliana and B. juncea, and 15 collinear gene pairs between A. thaliana and B. nigra (Table S3).

In addition, 52 gene pairs maintained their relative positions between the A\textsubscript{na} subgenome of B. napus and the A\textsubscript{ra} genome of B. rapa, and 44 gene pairs maintained their relative positions between the C\textsubscript{na} subgenome of B. napus and the C\textsubscript{ol} genome of B. oleracea. Furthermore, 29 gene pairs maintained their relative positions between the A\textsubscript{ju} subgenome of B. juncea and the A\textsubscript{na} genome of B. napus, and 42 gene pairs maintained their relative positions between the B\textsubscript{ju} subgenome of B. juncea and the B\textsubscript{ni} genome of B. nigra.

Figure 2. Cont.
Figure 2. Chromosomal distribution and analysis of duplication events in MATE family genes among Brassicaceae species. (a) Chromosomal distribution and analysis of duplication events in MATE family genes among A subgenome. (b) Chromosomal distribution and analysis of duplication events in MATE family genes among B subgenome. (c) Chromosomal distribution and analysis of duplication events in MATE family genes among C subgenome. Genes from the same subgroups are indicated by the same color, which is consistent with that for the corresponding family in the phylogenetic tree (Figure 1); genes located on the scaffold are not shown. The labels on the corresponding chromosomes indicate the source organism and the subgenome. The scales indicate the sizes of the various Brassica genomes (Mb). Bra, B. rapa; Bol, B. oleracea; Bni, B. nigra; Bra, B. napus; Bju, B. juncea.
Figure 3. Duplicated and orthologous genes in the Brassicaceae. (a) Circle plot of orthologous genes in A. thaliana (At), B. oleracea (Bol), B. rapa (Bra), B. napus (Bna), B. juncea (Bju), and B. nigra (Bni). (b) Density of Ks values of MATE gene pairs between B. napus and A. thaliana. (c) Duplicate gene pairs in B. napus. (d) Density of Ks values of duplicated MATE genes in B. napus. (e) Duplicate gene pairs in B. juncea.

Of the 101,040 genes in the B. napus genome, 997 genes (0.99%) appeared to have undergone tandem duplication and 74,183 genes (73.4%) appeared to have undergone WGD or segmental duplication (Table S4), which showed less tandem duplication and more WGD or segmental duplication compared with the results of a previous study [40]. Furthermore, 113 out of 124 BnaMATE genes (91.1%) were derived from WGD or segmental duplication, which was a number slightly larger than the average percentage at the whole-genome level (73.4%). Therefore, it appears that WGD or segmental duplication...
played an important role in the expansion of the MATE family in *B. napus*. Likewise, we also examined the MATE gene expansion patterns in *B. rapa, B. oleracea, B. nigra,* and *B. juncea,* and we showed that all of the *BnaMATE* genes (100%; 85/85) and most of the *BolMATE* (95.1%; 77/81), *BniMATE* (93.7%; 74/79), and *BjuMATE* genes (90.8%; 118/130) were derived from segmental duplication, followed by dispersed duplication (0% in *B. rapa, 3.7% in B. oleracea, 3.8% in B. nigra,* and 7.7% in *B. juncea*) (Table S4).

In addition, we used the synonymous base substitution rates (*Ks* values) as a proxy for time to estimate the approximate ages of the segmental duplication events [1]. Then, we predicted the *Ks* values for all collinear gene pairs. These ranged from 0.3322 to 0.9765 between *B. napus* and *A. thaliana,* with a mean of 0.4903 (Figure 3b). The divergence time ranged from 11.07 to 32.55 million years ago (MYA), with a mean of 16.34 MYA, indicating that the MATE genes of *B. napus* and *A. thaliana* diverged approximately 16 MYA. This is consistent with the likely timing of the recent whole-genome triplication event in *Brassica,* which has been estimated at approximately 9–15 MYA or even 28 MYA [41]. In addition, we identified 36 and 26 duplicate gene pairs in *B. napus* and *B. juncea,* respectively (Figure 3c,e). Then, we estimated the timing of the WGD event based on the distribution of *Ks* values, which ranged from 0.0192 to 0.4286 (mean, 0.135) in *B. napus* duplicate gene pairs and from 0.0310 to 0.2591 (mean, 0.1684) in *B. juncea* duplicate gene pairs. In *B. napus,* the corresponding duplication time range was 0.64–14.29 MYA (mean, 3.45 MYA), whereas in *B. juncea,* it was 1.03–8.64 MYA (mean, 5.61 MYA) (Table S5). One peak (0.06–0.09) in *Ks* values was observed in *B. napus,* representing the duplication time of these genes, which occurred during the formation of *B. napus* 7500–12,500 years ago (Figure 3d). Therefore, the processes of Brassicaceae speciation and whole-genome triplication likely played an important role in the divergence of the duplicated MATE genes among these species.

We also calculated the rates of nonsynonymous substitutions (*Ka*) and synonymous substitutions (*Ks*) and the *Ka/Ks* ratios of the MATE gene pairs to identify the evolutionary constraints acting on the MATE gene pairs, and we found that the *Ka/Ks* ratios of the duplicates were commonly less than 1 (Table S5), indicating that these genes were subject to purifying selection.

### 2.5. Analysis of MATE Gene Structures and Conserved Motifs between *A. thaliana* and *B. napus*

The allotetraploid Brassicaceae species rapeseed (*B. napus*) is one of the main oilseed crops worldwide, serving as a source of edible oil, biodiesel, and protein-rich animal feed. To provide a clear functional characterization of the *BnaMATE* genes, as well as to enable the manipulation of traits related to rapeseed metabolism, we analyzed the intron–exon structures of these genes using GSDS v2.0 (http://gsds.cbi.pku.edu.cn/index.php); the results are displayed along with the subgroups of phylogenetic tree (Figure 4a). Notably, the lengths and numbers of exons and introns were more similar within the same subgroup (Figure 4b). For example, the number of exons ranged from 5 to 11 in the largest group, Group 2, and the maximum number of exons (11–14 exons) was found in Group 3, except for *BnaMATE43d* and *BnaMATE47c,* which have only 5 exons. The genes in Group 4 generally had the smallest number of exons, 1–3, and the MATE genes in Group 1 had 4–10 exons. Our findings were in agreement with previous studies reporting that MATE genes from different groups were generally distinct, with each group sharing a common gene structural layout [22]. In addition, similar genes structures were observed among *B. rapa, B. oleracea, B. nigra,* and *B. juncea,* which were divided into the same subfamilies (Figure 1, Table S2).

Next, we identified the conserved motifs of MATE sequences using the MEME online software (Figure 4c, Figure S1). In most cases, the protein architecture was remarkably conserved within specific subgroups, providing further support to the phylogenetic analysis based on MATE domains (Figure S2) and suggesting that there are functional similarities within the groups. In addition, the types and sequences of the motifs are similar among Group 1, 2, and 4, which contain all 12 conserved motifs, whereas the most distinctive pattern of motifs was found in Group 3 proteins, which contain a smaller number of motifs. These findings suggest that these conserved motifs may have crucial roles in subfamily-specific functions.
Figure 4. Cont.
2.6. Expression Profiles of BnaMATE Genes in Various Rapeseed Tissues

Based on the transcriptome sequencing datasets from *B. napus* ZS11 (BioProject ID PRJNA358784), we characterized the relative transcript abundances of *BnaMATE* gene transcripts in 15 different tissues of rapeseed (Figure 5, Table S6). Within Group 1, most *BnaMATE* genes showed low or no expression in the tested tissues and organs, such as *BnaMATE8a, BnaMATE8b, BnaMATE9a, BnaMATE9b, BnaMATE14a, BnaMATE14b, BnaMATE14c*, and *BnaMATE17a*, which were expressed at low levels; in contrast, *BnaMATE1, BnaMATE6a, BnaMATE6b, BnaMATE7, BnaMATE15c*, and *BnaMATE17d* were barely expressed in the tested tissues and organs. However, *BnaMATEs* also displayed the specifically expression patterns; for example, *BnaMATE12a* and *BnaMATE12b* were more highly expressed in seeds and embryos than in other tissues; *BnaMATE15a* and *BnaMATE15b* were more highly expressed in seeds (40 DAF) and seed coats (40 DAF) than in other tissues; *BnaMATE16a* and *BnaMATE16b* were more highly expressed in the tissues of the whole flower, leaves, and silique pods compared with other tissues; and *BnaMATE17b* and *BnaMATE17c* were more highly expressed in flowers and seed pods than in other tissues (Figure 5).
Figure 5. Expression patterns of all 124 BnaMATE genes in 15 different tissues at different developmental stages in B. napus. The color bar represents log2 expression levels (fragments per kilobase per million mapped fragments, or FPKM). Red, blue, and white indicate high, low, and medium expression levels, respectively, and black indicates no expression (FPKM = 0). The abbreviations above the heatmap indicate the different tissues and organs/developmental stages of B. napus ZS11 (listed in Table S1).

In Group 2, BnaMATE19a and BnaMATE19c were more highly expressed in the tissues of flowers and late developmental tissues (at 40 and 49 DAF), including seeds, embryos, and silique pods, than in other tissues; BnaMATE24a, BnaMATE24b, BnaMATE24c, and BnaMATE24d were specifically expressed in seeds, embryos, and seed coat; BnaMATE35a, BnaMATE35b, BnaMATE35c, and BnaMATE35d were specifically expressed in flower, seed coat, and silique pod tissues; and BnaMATE40a, BnaMATE40c, and BnaMATE40f were highly expressed in seeds and embryos and (for the latter two) in leaves, anthocaulus, calyx, and silique pods. BnaMATE33a, BnaMATE33b, BnaMATE37a, BnaMATE37b, BnaMATE37c, and BnaMATE37d were expressed in all tissues at different developmental stages, and they were more highly expressed in flower tissues (anthocaulus, calyx, petal, pistils, stamens, anthers, filaments, and the tops of main inflorescence flowers). Finally, BnaMATE19b, BnaMATE31a, BnaMATE31b, BnaMATE28b, BnaMATE36c, and BnaMATE39d were barely expressed in any tissue (Figure 5).

Among Group 3 genes, most were expressed in at least one tissue, with the exception of BnaMATE43b, BnaMATE43d, and BnaMATE43f, which were barely detected in any tissue or organ. BnaMATE44a, BnaMATE44b, and BnaMATE46b were highly expressed throughout plant development, whereas BnaMATE43e was highly expressed in the seed coat as compared with other tissues (Figure 5).
In Group 4, BnaMATE55a, BnaMATE55b, and BnaMATE55c were specifically expressed in flower tissues (anthocaulus, calyx, petal, pistils, stamens, anthers, filaments, and the tops of main inflorescence flowers), and BnaMATE56a, BnaMATE56b, BnaMATE56c, BnaMATE56d, and BnaMATE56e were specifically expressed in seeds and seed coat (at 21, 30, and 40 DAF). In addition, the four BnaMATE52 genes and four BnaMATE53 genes were barely expressed in any tissues (Figure 5).

Overall, 11 BnaMATE genes were expressed in all tissues at different developmental stages, whereas 29 exhibited almost no expression, and 84 were specifically expressed in particular tissues, suggesting that they may play multiple or tissue-specific roles in the tested tissues and organs of B. napus.

2.7. Expression Patterns of BnaMATE Genes under Heavy Metal Stress

Previous studies showed that the MATE genes have important roles in resistance to various abiotic stresses [13–18]. Thus, we investigated the expression patterns of the BnaMATE genes under As\(^{3+}\) and Cd\(^{2+}\) stress using RNA-seq. Under normal conditions, the expression patterns of BnaMATEs were similar to those identified in the absence of metal ion stress, with different expression profiles detected among different rapeseed varieties (Figure 6, Table S7). For instance, BnaMATE33a, BnaMATE33b, BnaMATE37a, BnaMATE37b, BnaMATE37c, and BnaMATE37d were highly expressed in all tissues, whereas BnaMATE31a, BnaMATE31b, and the members of BnaMATE52 and BnaMATE53 were expressed minimally or repressed. In general, the expression of BnaMATE genes in Groups 1, 2, and 4 was clearly induced by As\(^{3+}\) or Cd\(^{2+}\) treatment, whereas that of Group 3 genes was suppressed, compared with their expression under normal conditions (Figure 6).

![Figure 6](image_url). Expression analysis of BnaMATE family genes in different rapeseed tissues under As\(^{3+}\) and Cd\(^{2+}\) treatment. The color bar represents log\(_2\) expression levels (FPKM). Red, blue, and white indicate high, low, and medium expression levels, respectively, and black indicates no expression (FPKM = 0). The abbreviations above the heatmap indicate the different cultivars of B. napus tested (listed in Table S7).
2.8. Expression Patterns of BnaMATE under Hormone Treatments

In *A. thaliana*, MATE family proteins have significant roles in the transport of hormones, including salicylic acid (SA) and abscisic acid (ABA) [11,16–18]. To explore the temporally and spatially specific expression patterns of the BnaMATE genes under hormone treatments, we used the transcriptome sequencing datasets to analysis their expression patterns over the course of treatment with the hormones indoleacetic acid (IAA), aminocyclopropane carboxylic acid (ACC), ABA, gibberellic acid (GA3), and 6-benzyladenine (6BA) (Figure 7). The Group 2 MATE genes showed relatively higher expression after the various hormone treatments than genes in the other three groups, whereas the Group 4 MATE genes were the most weakly expressed and in many cases showed no expression (Figure 7). Some BnaMATEs were up-regulated under multiple hormone treatments, including BnaMATE15b, BnaMATE19b, BnaMATE28c, BnaMATE39a, and BnaMATE43a, whereas others were up-regulated by specific hormone treatment: for example, BnaMATE16c and BnaMATE16d were highly expressed under ABA treatment, whereas BnaMATE39a, BnaMATE48a, and BnaMATE48b seemed more sensitive to IAA and ABA. Other BnaMATE genes were down-regulated by hormones treatment, such as BnaMATE12a, BnaMATE37a, BnaMATE37b, BnaMATE37c, BnaMATE37d, BnaMATE50a, BnaMATE50b, and BnaMATE51c. Overall, these results suggest that many BnaMATE genes may play important roles in processes related to hormones.

![Figure 7](image_url)
3. Discussion

Multidrug and toxic compound extrusion (MATE) is a present categorized multidrug efflux transporter family in almost all prokaryotes and eukaryotes, which is a large family of secondary active transporters in all kingdoms of life [4,22]. Recently, numerous studies of the MATE family have been carried out in plants, in species including rice [20,21], maize [22], tomato [4], upland cotton [23], soybean [24] and potato [38], but they have rarely been studied in the Brassicaceae. However, the Brassicaceae, given their complex history of genome duplication, are a particularly good model for investigating polyploidy in evolution, which is a major evolutionary process in eukaryotes [29,33]. Thus, the completion of Brassicaceae genome sequencing provides an excellent opportunity for the genome-wide characterization of the MATE family in the major Brassica plants, which include three diploid (B. rapa, B. nigra and B. oleracea) and two allotetraploid species (B. juncea and B. napus). Based on our results, we identified 124, 85, 81, 79, and 130 MATE genes in B. napus, B. rapa, B. oleracea, B. nigra, and B. juncea, which in all cases exceeds the 57 found in A. thaliana [13] and is consistent with the past genome duplications. In addition, the phylogenetic tree was supported by the intron–exon structures and motif compositions, as these showed considerable conservation within each group and greater differences between groups, particularly in the numbers and sizes of introns and exons (Figure 2, Table S2). The exonization of intronic sequences and pseudoexonization of exonic sequences play major roles in the divergence of gene structure, which is another likely cause for the functional diversification of MATE genes [43]. Introns are thought to alter gene activities, and the presence of introns in the genome is believed to impose a substantial burden on the host. Interestingly, the only intronless MATE genes were found in Group 4, whose members contains a maximum of only two introns, which is consistent with the patterns in cotton and maize [22,39]; this strongly supports the likelihood that the expansion of the MATE genes in Brassicaceae was also governed by intron loss and gain.

Gene duplication is a major driving force in gene family evolution and expansion [44]. In this study, 113 out of 124 BnaMATE genes (91.1%) were derived from WGD or segmental duplication, suggesting that such duplication events were the main contributors to the expansion of the BnaMATE genes. To establish strong evidence for homology, we assessed microsynteny within the Brassicaceae genomes, as well as syntenic genes between the Brassicaceae and A. thaliana genomes. There were 57 MATE genes in A. thaliana, which would be expected to have 171 MATE genes in B. rapa and B. oleracea given the whole-genome triplication (WGT) that occurred in Brassica species [33,41], and 342 MATE genes in B. napus, which was generated through hybridization of the progenitor species B. rapa and B. oleracea 7500–12,500 years ago [29]. In actuality, we found 85 BraMATE, 81 BolMATE, and 124 BnaMATE genes, indicating that duplicated genes were lost after WGT. Since the divergence of A. thaliana and Brassica, about 35% of genes in Brassica have been lost via deletion [45]. In total, we identified 29 collinear gene pairs between A. thaliana and B. rapa, but only 17 collinear gene pairs between A. thaliana and B. oleracea, although this may be due to assembly errors in the currently available B. oleracea genome information. B. napus, as an allopolyploid, was created through the hybridization of its diploid progenitors B. oleracea and B. rapa. The number of MATE genes (124) in B. napus was far less than the sum of the MATE genes in B. oleracea (81) and B. rapa (85), indicating that MATE gene loss that occurred during the evolution of B. napus. B. juncea is descended from a hybridization between B. rapa and B. nigra. Gene loss occurred among the MATEs of B. juncea as well, such that the number of MATE genes (130) in B. juncea is likewise much lower than the sum of those in B. rapa (85) and B. nigra (79). When we determined the \( K_s \) and \( K_a \) values, an important tool in investigating the type of selection pressure acting on protein-coding genes [46], we found that, interestingly, the \( K_d/K_a \) ratio was <1 for all gene pairs of A. thaliana and B. napus or B. napus and B. napus, indicating that the BnaMATE genes have undergone large-scale purifying selection.

Our synten analysis between B. napus and its diploid progenitors B. oleracea and B. rapa indicated that most Brassica MATE genes were located in syntenic regions, with 52 gene pairs shared between the A\(_{na}\) subgenome of B. napus and the A\(_{ra}\) genome of B. rapa, 44 gene pairs shared between the C\(_{na}\) subgenome of B. napus and the C\(_{ol}\) genome of B. oleracea, 29 gene pairs shared between the
AtMATE41, which may also be involved in the transport of phenolic compounds, along with AtMATE50, where different species had different numbers of chromosomes, and the A, B, and C subgenomes have undergone rearrangements, during which genes were presumably lost in the process of polyploidization.

According to our phylogenetic analysis, these MATE genes were classified into four groups, which is congruent with previous findings [1,23,24], suggesting similar evolutionary trajectories among these plants. In contrast, the previous studies reported that five MATE subfamilies were identified in potato [36], and seven were identified in maize [22], indicating a functional diversification among plant species in the complex plant processes. In addition, phylogenetic analysis revealed that all of the MATE family genes from Brassicaceae are closely associated with AtMATE genes within the same groups (Figure 1) and exhibited similar exon–intron structures (Figure 4), indicating that members within the subfamily have highly conserved and might have similar functions in the same groups [47]. Group 1 contains 26 BnaMATE genes along with AtMATE1 from A. thaliana, whose protein product mediates the efflux of plant-derived or exogenous toxic compounds from the cytoplasm, conveying resistance to Cd$$^{2+}$$ [13], while other members of Group 1 were rarely reported. Group 2 contains 51 BnaMATE genes, along with A. thaliana genes whose products are reported to mediate the transport of alkaloids and phenolic compounds and influence multidrug resistance, such as AtMATE19, AtMATE33, AtMATE35, and AtMATE41. AtMATE19, also known as ALF5 (ABERRANT LATERAL ROOT FORMATION 5), encodes a protein involved in multidrug resistance [48]. The protein product of AtMATE35, also known as FFT (FLOWER FLAVONOID TRANSPORTER), was considered to be a flavonoid transporter [49]; however, other evidence suggests that AtMATE35, as well as AtMATE33, functions as a vacuolar chloride channel involved in cell turgescence during stomatal movements, root hair elongation, and pollen germination [50]. AtMATE41 is also known as TT12 (TRANSPARENT TESTA 12), and previous studies raised the possibility that the functional of TT12 is a flavonoid transporter in Arabidopsis, which is highly specific expressed in developing seeds [7,51,52]. Among the B. napus MATE genes in Group 2, BnaMATE19 genes, because of their close relation with characterized transporters, seem to be the best candidates for conveying the protection of roots against inhibitory compounds; the BnaMATE33 and BnaMATE35 genes most likely encode proteins that function as chloride channels essential for turgor regulation, which may also be involved in the transport phenolic compounds, along with BnaMATE41. Group 3 contains 15 BnaMATE genes along with AtMATE42, AtMATE43, and AtMATE47 from A. thaliana and BolMATE42a from B. oleracea. AtMATE42 encodes the root citrate transporter, which, together with the root malate transporter ALMT1, is involved in the primary mechanism of aluminum (Al) tolerance [53]. AtMATE43 (FRD3, FERRIC REDUCTASE DEFECTIVE 3) encodes a protein responsible for loading citrate, a chelator of Fe, into the root xylem for efficient Fe translocation [54,55]. The AtMATE47 (EDS5, ENHANCED DISEASE SUSCEPTIBILITY) protein product participates in SA-dependent signaling for disease resistance [12,56]. BolMATE42a, also called BolMATE (KF031944), is the homolog of an A. thaliana gene encoding a protein that activates citrate efflux upon exposure to Al$$^{3+}$$, leading to enhanced Al tolerance [15].

Overall, these findings strongly suggest that BnaMATE42, BnaMATE43, and BnaMATE47 may be related to Fe and Al homeostasis. Additionally, Group 4 contains 32 BnaMATE genes along with AtMATE48, AtMATE50, and AtMATE51 from A. thaliana. AtMATE48 (BCD1, BUSH, AND CHLORORIC DWARF 1) is involved in maintaining cellular Fe homeostasis by secreting excess Fe [57], whereas AtMATE50 acts as an ABA efflux transporter [18]; AtMATE51 (ADS1, ACTIVATED DISEASE SUSCEPTIBILITY 1) negatively regulates the accumulation of the plant immune activator SA for disease resistance [11]. Thus, members of this group, especially BnaMATE48, BnaMATE50, and BnaMATE51, encode proteins involved in ABA and SA efflux and Fe tolerance mechanisms.

Gene expression analysis is widely considered a valuable tool for obtaining fundamental information on the possible functions of genes under study [22,39]. To further analyze the temporally
and spatially specific expression patterns of BnaMATE genes, we defined their transcription levels in 15 tissues at different developmental stages. We found that most showed differing expression in different tissues (Figure 5), implying that individual MATE genes may play specific roles in the development of individual tissues under normal conditions. The transcription levels of Group 2 BnaMATE genes were higher than those for the other three groups, indicating that the Group 2 genes may be important throughout B. napus development. It is well established that tissue-specific gene expression plays a crucial role in developmental and other physiological processes in specific tissue [58,59]. Among the BnaMATE genes, BnaMATE19a and BnaMATE19c seemed to have high expression levels late in development (Figure 5), and both were orthologous to AtMATE19, which is involved in multidrug resistance [48]. BnaMATE33a, BnaMATE33b, and BnaMATE35c were specifically expressed at high levels in floral organs (Figure 5), and they were orthologous to AtDTX33 and AtDTX35, respectively, which are thought to function as vacuolar chloride channels involved in cell turgescence [50]; AtMATE35 is also considered to be a flavonoid transporter [48]. BnaMATE35b and BnaMATE35d were highly expressed in floral organs, seed coat, and siliques (Figure 5), suggesting that those proteins may have the same function as AtMATE35. BnaMATE41a and BnaMATE41b had high expression levels during the prometaphase of seed development (Figure 5), which is in accordance with previous studies [7], and they were orthologous to AtMATE41 (TT12), implying that the proteins could play a similar role in flavonoid transport [7,50,51]. BnaMATE37c and BnaMATE37d were highly expressed in all of tissues tested, and BnaMATE24 genes were strongly specifically expressed in seeds, embryos, and seed coat, but no information is available on the functions of these genes or their homologs. Interestingly, we found that BnaMATE47a and BnaMATE47b had a high expression in siliques (at 30, 40, and 46 DAF) and were probably necessary for SA-dependent signaling, given their close phylogenetic relationship with A. thaliana AtMATE47 [12,56]. We also found tissue-specific gene expression in Group 4 genes: BnaMATE56a, BnaMATE56b, BnaMATE56c, BnaMATE56d, and BnaMATE56e were preferentially expressed in seeds and seed coat. These BnaMATE56 genes were orthologous to AtMATE56, which encodes a protein that participates in increasing CO2 and stomatal closure [60]. It seems that the function of the BnaMATE56 genes has been diverted over evolutionary time. Overall, the results revealed that most duplicated BnaMATE gene pairs had diverse expression patterns, suggesting that the functions of duplicated genes have tended to diversify, conforming to a major theme of long-term evolution.

Having established these basic patterns of MATE gene expression, we next sought to assess how these are influenced by plant exposure to abiotic stress. Heavy metal (Cd2+ and As3+) pollution in agricultural soils is a critical problem affecting crop production and quality, and the expression of some MATE genes is reported to increase in response to various abiotic stresses in plants [13–15,61–63]. Therefore, we carried out transcriptome expression profiling in B. napus seeds under As3+ and Cd2+ stress, which showed expression changes in BnaMATE genes in response to As3+ and/or Cd2+ stress. For instance, BnaMATE1 showed higher expression under Cd2+ treatment (Figure 6); this was consistent with its status as the homolog of AtMATE1, which is involved in Cd2+ detoxification [13], and it strongly suggests that BnaMATE1 might play an important role in Cd2+ tolerance. BnaMATE6a and BnaMATE6b showed expression patterns similar to that of BnaMATE1, implying that they may have similar functions. Furthermore, 18 and 26 genes were observably up-regulated (to >2-fold) under As3+ or Cd2+ treatment, respectively, compared to control conditions (CK), and 25 and 19 genes were down-regulated (to <0.5-fold); thus, all of these genes may have specific roles in As3+ and Cd2+ stress response. We also noted that some pairs of duplicated genes showed a distinct divergence of expression levels: for example, BnaMATE43b was down-regulated under As3+ or Cd2+ treatment, whereas BnaMATE43c was up-regulated. These results implied that these duplicated genes have developed diverse functions in response to As3+ and Cd2+ stress in the evolutionary history.

Previous studies have confirmed that MATE genes act in hormone transport [16–18]. We analyzed the expression profile of the BnaMATE genes in plants treated with five separate hormones for different periods of time through transcriptome expression profiling (Figure 7). We found a variety of expression...
patterns among BnaMATE family genes. For instance, BnaMATE27c, BnaMATE35c, BnaMATE42a, BnaMATE42b, BnaMATE44a, BnaMATE44b, BnaMATE46a, BnaMATE46b were not induced in response to hormone treatments, whereas BnaMATE12a, BnaMATE37a, BnaMATE37b, BnaMATE37c, BnaMATE37d, BnaMATE50a, BnaMATE50b, and BnaMATE51c were down-regulated, and BnaMATE15b, BnaMATE19b, BnaMATE39a, BnaMATE28c, and BnaMATE43a were up-regulated. Interestingly, BnaMATE16c and BnaMATE16d were specifically up-regulated under ABA treatment, implying that they may function in IAA and ABA than to other hormones (Figure 7). Since BnaMATE50a and BnaMATE50b were orthologous to AtMATE50, which acts as an ABA efflux transporter [18], and BnaMATE51c was orthologous to AtMATE51, which negatively regulates the accumulation of the plant immune activator SA for disease resistance [11], we believe that these BnaMATEs also have specific roles in responses to different hormones. Our results showed the first comprehensive analysis of MATE family genes and investigated expression profiles of BnaMATEs under heavy metals or hormones treatment, laying the foundation for further elucidating the roles of MATE genes in the export of toxins and other substrates.

4. Materials and Methods

4.1. Identification of MATE Transporters in Brassicaceae Species

The MATE transporter gene family has at least 56 members in the A. thaliana genome [13], and a 57th gene, AT2G04066 (Gene ID: 814942), is described by the term “MATE efflux family protein” in the NCBI (National Center for Biotechnology Information) database [66] and was therefore included in this study. The protein sequences of the 57 MATE transporters were downloaded from The Arabidopsis Information Resource (TAIR) database (Table S2) [67]. Using 57 AtMATE sequences as queries, a BLASTp analysis [68] was performed against whole-genome sequences in the BRAD database [37]. The candidate sequences were chosen based on an E-value of ≤1 × 10−50 and further analyzed using PfamScan (http://www.ebi.ac.uk/Tools/pfa/pfamscan/) to confirm the presence of a MATE domain (PF01554.18, MATE). BLAST analysis of the MATE protein sequences in the B. napus genome database was performed using Geneious 4.8.5 software (http://www.geneious.com/); Biomatters, Auckland, New Zealand). The candidate genes were named using two- or three-letter abbreviations (italicized) denoting the source organism, the family name, and the positions in the subtribe, e.g., BnaMATE1a. The A. thaliana sequences were named following the generic system proposed for A. thaliana as AtMATE1 to AtMATE57, respectively [13].

Genomic, protein, and coding sequences from A. thaliana were downloaded from TAIR and those from B. napus (v5), B. rapa (v3.0), B. oleracea (v1.1), B. juncea (v1.5), and B. nigra (v1.1) were downloaded from the BRAD database. The physicochemical properties of the deduced proteins, including the molecular weight (MW) and isoelectric point (pI), were determined using the ExPASy-ProtParam tool (http://web.expasy.org/protparam/) [69].

4.2. Subcellular Location Analysis of Brassicaceae MATE Transporters

Subcellular location prediction of the candidate MATE proteins was performed with the online TargetP1.1 (http://www.cbs.dtu.dk/services/TargetP/) server [70] and Protein Prowler Subcellular Localization Predictor version 1.2 (http://bioinf.scmb.uq.edu.au/Pprowler_webapp_1-2/) [71]. Validation and determination of the possible cell compartmentalization, as suggested by the two software programs, was done with WoLF PSORT (https://wolfpsort.hgc.jp) [72].

4.3. Phylogenetic Analysis of Brassicaceae MATE Transporters

The protein sequences of the MATE proteins from A. thaliana and B. napus were used for multiple protein sequence alignments selected with MUSCLE [73] by Molecular Evolutionary Genetics Analysis (MEGA) 7.0 software (Arizona State University, Tempe, Arizona, USA) [74]. Phylogenetic trees were built using the neighbor-joining (NJ) method with 1000 replicates for the bootstrap test [75], and the
evolutionary distances were computed using the Poisson correction method [76]. The phylogenetic trees were visualized using FigTree v1.4.2 (http://tree.bio.ed.ac.uk/software/figtree/).

4.4. Chromosomal Distribution, Gene Duplication, and Evolutionary Analysis of Brassicaceae MATE Genes

All MATE genes were mapped to the chromosomes of Brassicaceae species according to their physical distances in the GFF genome files, which were downloaded from BRAD [37]. The physical chromosome maps were visualized with MapChart 2.2 (Plant Research International, Wageningen, Netherlands) [77]. Multiple collinear scanning toolkits (MCScanX) (University of Georgia, Athens, GA, USA) with the default parameters were used to view collinearity and analyze the gene replication events between A. thaliana and the five Brassicaceae species [78].

TBtools was used to construct a schematic diagram of the putative duplications of the MATE genes [79] and calculate the ratio of the nonsynonymous substitution rate (K_a) to the synonymous substitution rate (K_s) with the NG (Nei and Gojobori) method [80]. Divergence time was inferred using the formula T = K_a/2R, where R is 1.5 × 10^{-8} synonymous substitutions per site per year [81].

4.5. Gene Structure and Conserved Motif Analysis of BnaMATE Genes

The Gene Structure Display Server (GSDS 2.0: http://gsds.cbi.pku.edu.cn/) [42] was used to analyze the coding regions and genomic DNA sequences of the candidate MATE genes from A. thaliana and B. napus to determine their exon–intron structures. The conserved domains of the MATE protein family were predicted by the NCBI conserved domain search (www.ncbi.nlm.nih.gov) [82] and further confirmed by conducting a Pfam database search. Conserved motifs were determined using MEME 5.0.1 (http://meme-suite.org/) with the following parameters—Motif width, between 6 and 300 residues; Number of repetitions, any; maximum number of motifs, 12 [19] and visualized using TBtools (https://github.com/CJ-Chen/TBtools) [79].

4.6. Expression Patterns of BnaMATE Genes among Tissues and Developmental stages

To obtain a better understanding of the functions of the BnaMATE genes, we investigated their transcriptional levels in the different tissues and organs (roots, stems, leaves, anthocaulus, calyx, petal, pistils, stamens, anthers, filaments, the top of main inflorescence flowers, seeds, embryos, seed coat, and silique pods) of B. napus Zhongshuang No. 11 (ZS11) at different developmental stages. The resulting data were deposited in BioProject with ID PRJNA358784. The fragments per kilobase per million mapped fragments (FPKM) values were normalized using log2 transformation, and then the similarities and differences of the BnaMATE genes were visualized with heatmaps drafted by using Heatmap Illustrator 1.0 (Heml 1.0) (Huazhong University of Science and Technology, Wuhan, Hubei, China) [82].

4.7. Transcriptome Analysis of BnaMATE Genes in Response to Abiotic Stresses

The expression profiles of BnaMATE genes under heavy metal (As^{3+} and Cd^{2+}) [83] and hormone (BioProject ID PRJNA608211) treatments were investigated by RNA sequencing (RNA-seq). The plant materials and heavy metal and hormone treatments were as described in previous work [83–85]. The expression profiles of candidate BnaMATE genes were calculated based on FPKM values and visualized as heatmaps by using Heml 1.0 [82].

Supplementary Materials: Supplementary Materials can be found at http://www.mdpi.com/2223-7747/9/9/1072/s1. Table S1. The tissues and organs from B. napus ZS11 used in this study; Table S2. List of the identified MATE genes in B. napus genome; Table S3. MATE collarine genes in A. thaliana, B. napus, B. oleracea, B. rapa, B. juncea and B. nigra; Table S4. Gene duplication pattern in B. napus, B. oleracea, B. rapa, B. nigra and B. juncea; Table S5. The K_a/K_s values duplication time for duplicated MATE gene pairs in B. napus; Table S6. Heat map of expression levels of B. napus MATE family genes at the indicated growth periods and tissues and organs; Table S7 Comparison expression analysis of BnaMATE family genes in rapeseed between the CK and As^{3+}/Cd^{2+} treatments; Table S8. Comparison expression analysis of BnaMATE family genes in rapeseed between the CK and hormones treatment; Figure S1.
The detailed information of motif logos are obtained from the MEME Suite website, Figure S2: The detailed information of conserved domains of MATE proteins between B. napus and A. thaliana.

**Author Contributions:** C.Q. (Cunmin Qu) and J.L. conceived and designed the experiments; C.Q. (Cailin Qiao) and J.Y. conducted the experiments; Y.W. and S.X. collected and analyzed the data; C.Q. (Cailin Qiao) and M.G. carried out the experiments and performed software; H.D. and Z.T. analyzed the transcriptome data and collected the plant materials; C.Q. (Cailin Qiao), J.Y. and C.Q. (Cunmin Qu) wrote the manuscript; K.L. and J.L. reviewed the manuscript. All authors have read and agreed to the published version of the manuscript.

**Funding:** This work was supported by National Key R&D Program of China (2018YFD0100500), the National Natural Science Foundation of China (31830067), Fundamental Research Funds for the Central Universities (XDJK2020B030, XDJK2019C099), and the “111” Project (B12006), Modern Agro-industry Technology Research System (CARS-13), Chongqing Special Postdoctoral Science Foundation (cstc2019jcj-bshl0102).

**Acknowledgments:** We would like to thank Meichen Zhu and Xintong Zhou for data analysis in this study.

**Conflicts of Interest:** The authors declare no conflict of interest.

**References**

1. Wang, L.; Bei, X.; Gao, J.; Li, Y.; Yan, Y.; Hu, Y. The similar and different evolutionary trends of MATE family occurred between rice and Arabidopsis thaliana. *BMC Plant Biol.* 2016, 16, 207. [CrossRef] [PubMed]
2. Morita, Y.; Kodama, K.; Shiota, S.; Mine, T.; Kataoka, A.; Mizushima, T.; Tsuchiya, T. NorM, a Putative Multidrug Efflux Protein, of *Vibrio parahaemolyticus* and Its Homolog in *Escherichia coli*. *Antimicrob. Agents Chemother.* 1998, 42, 1778–1782. [CrossRef] [PubMed]
3. Brown, M.H.; Paulsen, I.T.; Skurray, R.A. The multidrug efflux pump NorM is a prototype of a new family of transporters. *Mol. Microbiol.* 1999, 31, 394–395. [CrossRef] [PubMed]
4. Santos, A.L.d.; Chaves-Silva, S.; Yang, L.; Maia, L.G.S.; Chalfun-Júnior, A.; Sinharoy, S.; Zhao, J.; Benedito, V.A. Global analysis of the MATE gene family of metabolite transporters in tomato. *BMC Plant Biol.* 2017, 17, 185. [CrossRef] [PubMed]
5. Takanashi, K.; Shitan, N.; Yazaki, K. The multidrug and toxic compound extrusion (MATE) family in plants. *Plant Biotechnol.* 2014, 31, 417–430. [CrossRef]
6. Shoji, T.; Inai, K.; Yazaki, Y.; Sato, Y.; Takase, H.; Shitan, N.; Yazaki, K.; Goto, Y.; Toyooka, K.; Matsuoka, K.; et al. Multidrug and Toxic Compound Extrusion-Type Transporters Implicated in Vacular Sequestration of Nicotine in Tobacco Roots. *Plant Physiol.* 2009, 149, 708–718. [CrossRef]
7. Debeaujon, I.; Peeters, A.; Léon-Kloosterziel, K.; Koornneef, M. The TRANSPARENT TESTA12 gene of Arabidopsis encodes a multidrug secondary transporter-like protein required for flavonoid sequestration in vacuoles of the seed coat endothelium. *Plant Cell* 2001, 13, 853–871. [CrossRef]
8. Pérez-Díaz, R.; Rnyagajo, M.; Pérez-Díaz, J.; Peña-Cortés, H.; Casaretto, J.A.; González-Villanueva, E.; Ruiz-Lara, S. *VvMATE1* and *VvMATE2* encode putative proanthocyanidin transporters expressed during berry development in *Vitis vinifera*. *Plant Cell Rep.* 2014, 33, 1147–1159. [CrossRef]
9. Gomez, C.; Terrier, N.; Torregrosa, L.; Vialat, S.; Fournier-Level, A.; Vialet, S.; Fournier-Level, A.; et al. Grapevine MATE-Type Proteins Act as Vacular H+-Dependent Acylated Anthocyanin Transporters. *Plant Physiol.* 2009, 150, 402–415. [CrossRef]
10. Gomez, C.; Conejero, G.; Torregrosa, L.; Cheynier, V.; Terrier, N.; Ageorges, A. In vivo grapevine anthocyanin transport involves vesicle-mediated trafficking and the contribution of anthoMATE transporters and GST. *Planta* 2011, 67, 960–970. [CrossRef]
11. Sun, X.; Gilroy, E.M.; Chini, A.; Nurmberg, P.L.; Hein, I.; Lacomme, C.; Birch, P.R.J.; Hussain, A.; Yun, B.W.; Leoke, G.J. *ADSL* encodes a MATE-transporter that negatively regulates plant disease resistance. *New Phytol.* 2011, 192, 471–482. [CrossRef] [PubMed]
12. Ishihara, T.; Sekine, K.T.; Hase, S.; Kanayama, Y.; Seo, S.; Ohashi, Y.; Kusano, T.; Shibata, D.; Shah, J.; Takahashi, H. Overexpression of the Arabidopsis thaliana EDS5 gene enhances resistance to viruses. *Plant Biol.* 2008, 10, 451–461. [CrossRef] [PubMed]
13. Li, L.; He, Z.; Pandey, G.K.; Tsuchiya, T.; Luan, S. Functional cloning and characterization of a plant efflux carrier for multidrug and heavy metal detoxification. *J. Biol. Chem.* 2002, 277, 5360–5368. [CrossRef] [PubMed]
14. Furukawa, J.; Yamaji, N.; Wang, H.; Mitani, N.; Murata, Y.; Sato, K.; Katsuhara, M.; Takeda, K.; Ma, J.F. An Aluminum-Activated Citrate Transporter in Barley. *Plant Cell Physiol.* 2007, 48, 1081–1091. [CrossRef]
15. Wu, X.; Li, R.; Shi, J.; Wang, J.; Sun, Q.; Zhang, H.; Xing, Y.; Qi, Y.; Zhang, N.; Guo, Y.D. *Brassica oleracea* MATE Encodes a Citrate Transporter and Enhances Aluminum Tolerance in *Arabidopsis thaliana*. *Plant Cell Physiol.* 2014, 55, 1426–1436. [CrossRef]

16. Serrano, M.; Wang, B.; Aryal, B.; Garcia, C.; Abou-Mansour, E.; Heck, S.; Geisler, M.; Mauch, F.; Nawrath, C.; Métraux, J.P. Export of Salicylic Acid from the Chloroplast Requires the Multidrug and Toxin Extrusion-Like Transporter EDS5. *Plant Physiol.* 2013, 162, 1815–1821. [CrossRef]

17. Yamasaki, K.; Motomura, Y.; Yagi, Y.; Nomura, H.; Kikuchi, S.; Nakai, M.; Shiina, T. Chloroplast envelope localization of EDS5, an essential factor for salicylic acid biosynthesis in *Arabidopsis thaliana*. *Plant Signal Behav.* 2013, 8, e23603. [CrossRef]

18. Zhang, H.; Zhu, H.; Pan, Y.; Yu, Y.; Luan, S.; Li, L. A DTX/MATE-Type Transporter Facilitates Abscisic Acid Efflux and Modulates ABA Sensitivity and Drought Tolerance in *Arabidopsis*. *Mol. Plant* 2014, 7, 1522–1532. [CrossRef] [PubMed]

19. Omote, H.; Hiasa, M.; Matsumoto, T.; Otsuka, M.; Moriyama, Y. The MATE proteins as fundamental transporters of metabolic and xenobiotic organic cations. *Trends Pharmacol. Sci.* 2006, 27, 587–593. [CrossRef] [PubMed]

20. Hvorup, R.N.; Winnen, B.; Chang, A.B.; Jiang, Y.; Zhou, X.F.; Saier, M.H. The multidrug/oligosaccharidyl-lipid/poly saccharide (MOP) exporter superfamily. *Eur. J. Biochem.* 2003, 270, 799–813. [CrossRef] [PubMed]

21. Tiwari, M.; Sharma, D.; Singh, M.; Tripathi, R.D.; Trivedi, P.K. Expression of OsMATE1 and OsMATE2 alters development, stress responses and pathogen susceptibility in *Arabidopsis*. *Sci. Rep.* 2014, 4, 3964. [CrossRef] [PubMed]

22. Zhu, H.; Wu, J.; Jiang, Y.; Jin, J.; Zhou, W.E.I.; Wang, Y.U.; Han, G.; Zhao, Y.; Cheng, B. Genomewide analysis of MATE-type gene family in maize reveals microsynteny and their expression patterns under aluminum treatment. *J. Genet.* 2016, 95, 691–704. [CrossRef] [PubMed]

23. Xu, L.; Shen, Z.; Chen, W.; Si, G.; Meng, Y.; Guo, N.; Sun, X.; Cai, Y.; Lin, Y.; Gao, J. Phylogenetic analysis of upland cotton MATE gene family reveals a conserved subfamily involved in transport of proanthocyanidins. *Mol. Biol. Rep.* 2019, 46, 161–175. [CrossRef] [PubMed]

24. Liu, J.; Li, Y.; Wang, W.; Gai, J.; Li, Y. Genome-wide analysis of MATE transporters and expression patterns of a subgroup of MATE genes in response to aluminum toxicity in soybean. *BMC Genom.* 2016, 17, 223. [CrossRef] [PubMed]

25. Julião, M.H.M.; Silva, S.R.; Ferro, J.A.; Varani, A.M. A Genomic and Transcriptomic Overview of MATE, ABC, and MFS Transporters in *Citrus sinensis* Interaction with *Xanthomonas citri* subsp. *citri*. *Plants* 2020, 9, 794.

26. Nagaharu, U.; Nagaharu, N. Genome analysis in *Brassica* with special reference to the experimental formation of *B. napus* and peculiar mode of fertilization. *J. Japan Bot.* 1935, 7, 389–452.

27. Kim, J.; Lee, J.; Choi, J.P.; Park, I.; Yang, K.; Kim, M.K.; Lee, Y.H.; Nou, I.S.; Kim, D.S.; Min, S.R.; et al. Functional innovations of three chronological mesohexaploid *Brassica rapa* genomes. *BMC Genom.* 2014, 15, 606. [CrossRef] [PubMed]

28. Yu, J.; Hu, F.; Dossa, K.; Wang, Z.; Ke, T. Genome-wide analysis of UDP-glycosyltransferase super family in *Brassica rapa* and *Brassica oleracea* reveals its evolutionary history and functional characterization. *BMC Genom.* 2017, 18, 474. [CrossRef] [PubMed]

29. Challhoub, B.; Denoeud, F.; Liu, S.; Parkin, I.A.; Tang, H.; Wang, X. Early allopolyploid evolution in the post-Neolithic *Brassica napus* oilseed genome. *Science* 2014, 345, 950–953. [CrossRef] [PubMed]

30. Jenčovské, E.; Chèvre, A.M.; Alik, K. Chromosomal and Gene Expression Changes in *Brassica* Allopolyploids. In *Polyploid and Hybrid Genomics*; Chen, Z.J., Birchler, J.A., Eds.; John Wiley & Sons, Inc.: Oxford, IA, USA, 2013; pp. 171–186.

31. Lee, T.-H.; Tang, H.; Wang, X.; Paterson, A.H. PGDD: A database of gene and genome duplication in plants. *Nucleic Acids Res.* 2013, 41, D1152–D1158. [CrossRef] [PubMed]

32. Sun, F.; Fan, G.; Hu, Q.; Zhou, Y.; Guan, M.; Tong, C.; Li, J.; Du, D.; Qi, C.; Jiang, L.; et al. The high-quality genome of *Brassica napus* cultivar ‘ZS11’ reveals the introgression history in semi-winter morphotype. *Plant J.* 2017, 92, 462–468. [CrossRef] [PubMed] [PubMed]

33. Liu, S.; Liu, Y.; Yang, X.; Tong, C.; Edwards, D.; Parkin, I.A.P.; Zhao, M.; Ma, J.; Yu, J.; Huang, S.; et al. The *Brassica oleracea* genome reveals the asymmetrical evolution of polyploid genomes. *Nat. Commun.* 2014, 5, 3930. [CrossRef] [PubMed]
34. Wang, X.; Wang, H.; Wang, J.; Sun, R.; Wu, J.; Liu, S.; Bai, Y.; Mun, J.H.; Bancroft, I.; Cheng, F.; et al. The genome of the mesopolyploid crop species Brassica rapa. Nat. Genet. 2011, 43, 1035–1039. [CrossRef] [PubMed]

35. Yang, J.; Liu, D.; Wang, X.; Ji, C.; Cheng, F.; Liu, B.; Hu, Z.; Chen, S.; Pental, D.; Ju, Y.; et al. The genome sequence of alloployploid Brassica juncea and analysis of differential homoeolog gene expression influencing selection. Nat. Genet. 2016, 48, 1225–1232. [CrossRef]

36. Perumal, S.; Koh, C.S.; Jin, L.; Buchwaldt, M.; Higgins, E.; Zheng, C.; Sankoff, D.; Robinson, S.J.; Kagale, S.; Navabi, Z.; et al. High contiguity long read assembly of Brassica nigra allows localization of active centromeres and provides insights into the ancestral Brassica genome. BioRxiv 2020, 932665. [CrossRef]

37. Cheng, F.; Liu, S.; Wu, J.; Fang, L.; Sun, S.; Liu, B.; Li, P.; Hua, W.; Wang, X. BRAD, the genetics and genomics database for Brassica plants. BMC Plant Biol. 2011, 11, 136. [CrossRef]

38. Li, Y.; He, H.; He, L.F. Genome-wide analysis of the MATE gene family in potato. Mol. Biol. Rep. 2019, 46, 403–414. [CrossRef] [PubMed]

39. Lu, P.; Magwanga, R.O.; Guo, X.; Kirungu, J.N.; Lu, H.; Cai, X.; Zhou, Z.; Wei, Y.; Wang, X.; Zhang, Z.; et al. Comparative Genomics of Brassica oleracea and parental species: Cloning, evolution, and differential homoeolog gene expression influencing selection. Mol. Genet. Genom. 2018, 281, 138–159. [CrossRef] [PubMed]

40. Wei, L.; Zhu, Y.; Liu, R.; Zhang, A.; Zhu, M.; Xu, W.; Lin, A.; Lu, K.; Li, J. Genome-wide identification and characterization of the MATE transporter gene family member, AtMATE, in Arabidopsis. Plant Physiol. 2015, 167, 109–123. [CrossRef] [PubMed]

41. Cheng, F.; Wu, J.; Wang, X. Genome triplication drove the diversification of Brassica plants. Hortic. Res. 2014, 1, 1024. [CrossRef] [PubMed]

42. Hu, B.; Jin, J.; Guo, A.Y.; Zhang, H.; Luo, J.; Gao, G. GSDS 2.0: An upgraded gene feature visualization server. Bioinformatics 2015, 31, 1296–1297. [CrossRef]

43. Fan, K.; Wang, M.; Miao, Y.; Ni, M.; Bibi, N.; Yuan, S.; Li, F.; Wang, X. Molecular Evolution and Exploration Analysis of the NAC Transcription Factor in Zea mays. PLoS ONE 2014, 9, e111837. [CrossRef] [PubMed]

44. Taylor, J.S.; Raes, J. Duplication and Divergence: The Evolution of New Genes and Old Ideas. Annu. Rev. Genet. 2004, 38, 615–643. [CrossRef] [PubMed]

45. Town, C.D.; Cheung, F.; Maiti, R.; Crabtree, J.; Haas, B.J.; Wortman, J.R.; Hine, E.E.; Althoff, R.; Arbogast, T.S.; Tallon, L.J.; et al. Comparative Genomics of Brassica oleracea and Arabidopsis thaliana Reveal Gene Loss, Fragmentation, and Dispersal after Polyploidy. Plant Cell 2006, 18, 1348–1359. [CrossRef] [PubMed]

46. Zhang, Z.; Li, J.; Zhao, X.Q.; Wang, J.; Wong, G.K.; Yu, J. KaKs_Calculator: Calculating Ka and Ks through model selection and model averaging. Genom. Proteom. Bioinf. 2006, 4, 259–263. [CrossRef]

47. Sun, H.; Fan, H.J.; Ling, H.Q. Genom-wide identification and characterization of the bHLH gene family in tomato. BMC Genom. 2015, 16, 9. [CrossRef]

48. Diener, A.C.; Gaxiola, R.A.; Fink, G.R. Arabidopsis ALF5, a multidrug efflux transporter gene family member, confers resistance to toxins. Plant Cell 2001, 13, 1625–1637. [CrossRef] [PubMed]

49. Thompson, E.P.; Wilkins, C.; Demidchik, V.; Davies, J.M.; Glover, B.J. An Arabidopsis flavonoid transporter is required for anthocyanin and pollen development. J. Exp. Bot. 2010, 61, 439–451. [CrossRef] [PubMed]

50. Zhang, H.; Zhao, F.; Tang, R.; Yu, Y.; Song, J.; Wang, Y.; Li, L.; Suan, S. Two tonoplast MATE proteins function as turgor-regulating chloride channels in Arabidopsis. Proc. Natl. Acad. Sci. USA 2017, 114, E2036–E2045. [CrossRef] [PubMed]

51. Chai, Y.; Lei, B.; Huang, H.; Li, J.; Yin, J.; Tang, Z.; Wang, R.; Chen, L. TRANSPARENT TESTA 12 genes from Brassica napus and parental species: Cloning, evolution, and differential involvement in yellow seed trait. Mol. Genet. Genom. 2009, 281, 109–123. [CrossRef]

52. Marinova, K.; Pourcel, L.; Weder, B.; Schwarz, M.; Barron, D.; Routaboul, J.M.; Debeaujon, I.; Klein, M. The Arabidopsis MATE Transporter TT12 Acts as a Vacular Flavonoid/H+-Antiporter Active in Proanthocyanidin-Accumulating Cells of the Seed Coat. Plant Cell 2007, 19, 2023–2038. [CrossRef]

53. Liu, J.; Luo, X.; Shafi, J.; Liang, C.; Jia, X.; Li, Z.; Magalhaes, J.; Kochian, L.V. A promoter-swap strategy between the AtAtLMT and AtMATE genes increased Arabidopsis aluminum resistance and improved carbon-use efficiency for aluminum resistance. Plant J. 2012, 71, 327–337. [CrossRef] [PubMed]

54. Green, L.S.; Rogers, E.E. FRD3 Controls Iron Localization in Arabidopsis. Plant Physiol. 2004, 136, 2523–2531. [CrossRef] [PubMed]
55. Durrett, T.P.; Gassmann, W.; Rogers, E.E. The FRD3-Mediated Efflux of Citrate into the Root Vasculature Is Necessary for Efficient Iron Translocation. *Plant Physiol.* **2007**, *144*, 197–205. [CrossRef] [PubMed]

56. Nawrath, C.; Heck, S.; Parinthawong, N.; Métarraux, J.P. EDS5, an Essential Component of Salicylic Acid–Dependent Signaling for Disease Resistance in *Arabidopsis*, Is a Member of the MATE Transporter Family. *Plant Cell* **2002**, *14*, 275–286. [CrossRef] [PubMed]

57. Seo, P.J.; Park, J.; Park, M.J.; Kim, Y.S.; Kim, S.G.; Jung, J.H.; Park, C.M. A Golgi-localized MATE transporter mediates iron homeostasis under osmotic stress in *Arabidopsis*. *Biochem. J.* **2012**, *442*, 551–561. [CrossRef]

58. Ingram, G.C.; Boisnard-Lorig, C.; Dumas, C.; Rogowsky, P.M. Expression patterns of genes encoding HD-ZipIV homeo domain proteins define specific domains in maize embryos and meristems. *Plant J.* **2000**, *22*, 401–414. [CrossRef]

59. Gallie, D.R.; Geisler-Lee, J.; Chen, J.; Jolley, B. Tissue-specific expression of the ethylene biosynthetic machinery regulates root growth in maize. *Plant Mol. Biol.* **2009**, *69*, 195–211. [CrossRef]

60. Tian, W.; Hou, C.; Ren, Z.; Pan, Y.; Jia, J.; Zhang, H.; Bai, F.; Zhang, P.; Zhu, H.; He, Y.; et al. A molecular pathway for CO₂ response in *Arabidopsis* guard cells. *Nat. Commun.* **2015**, *6*, 6057. [CrossRef]

61. Yokosho, K.; Yamaji, N.; Fuji-Kashino, M.; Ma, J.F. Functional Analysis of a MATE Gene OsFRDL2 Revealed its Involvement in Al-Induced Secretion of Citrate, but a Lower Contribution to Al Tolerance in Rice. *Plant Cell Physiol.* **2016**, *57*, 976–985. [CrossRef]

62. Liu, J.; Magalhaes, J.V.; Shaff, J.; Kochian, L.V. Aluminum-activated citrate and malate transporters from the MATE and ALMT families function independently to confer *Arabidopsis* aluminum tolerance. *Plant J.* **2009**, *57*, 389–399. [CrossRef] [PubMed]

63. Wang, J.; Hou, Q.; Li, P.; Yang, L.; Sun, X.; Benedito, V.A.; Wen, J.; Chen, B.; Mysore, K.S.; Zhao, J. Diverse functions of multidrug and toxin extrusion (MATE) transporters in citric acid efflux and metal homeostasis in *Medicago truncatula*. *Plant J.* **2017**, *90*, 79–95. [CrossRef] [PubMed]

64. Cai, S.; Chen, G.; Wang, Y.; Huang, Y.; Marchant, D.B.; Wang, Y.; Yang, Q.; Dai, F.; Hills, A.; Franks, P.J.; et al. Evolutionary Conservation of ABA Signaling for Stomatal Closure. *Plant Physiol.* **2017**, *174*, 732–747. [CrossRef] [PubMed]

65. Xie, Y.; Mao, Y.; Zhang, W.; Lai, D.; Wang, Q.; Shen, W. Reactive Oxygen Species-Dependent Nitric Oxide Production Contributes to Hydrogen-Promoted Stomatal Closure in *Arabidopsis*. *Plant Physiol.* **2014**, *165*, 759–773. [CrossRef] [PubMed]

66. Lu, S.; Wang, J.; Chitsaz, F.; Derbyshire, M.K.; Geer, R.C.; Gonzalez, N.R.; Gwadz, M.; Hurwitz, D.I.; Marchler, G.H.; Song, J.S.; et al. CDD/SPARCLE: The conserved domain database in 2020. *Nucleic Acids Res.* **2020**, *48*, D265–D268. [CrossRef]

67. Reiser, L.; Subramaniam, S.; Li, D.; Huala, E. Using the *Arabidopsis* information resource (TAIR) to find information about *Arabidopsis* genes. *Curr. Protoc. Bioinform.* **2017**, *60*, 1.11.1–1.11.45. [CrossRef]

68. Altschul, S.F.; Madden, T.L.; Schäffer, A.A.; Zhang, J.; Zhang, Z.; Miller, W.; Lipman, D.J. Gapped BLAST and PSI-BLAST: A new generation of protein database search programs. *Nucleic Acids Res.* **1997**, *25*, 3398–3402. [CrossRef]

69. Gasteiger, E.; Gattiker, A.; Hoogland, C.; Ivanyi, I.; Appel, R.D.; Bairoch, A. ExPASy: The proteomics server for in-depth protein knowledge and analysis. *Nucleic Acids Res.* **2003**, *31*, 3784–3788. [CrossRef] [PubMed]

70. Emanuelsson, O.; Brunak, S.; von Heijne, G.; Nielsen, H. Locating proteins in the cell using TargetP, SignalP and related tools. *Nat. Protoc.* **2007**, *2*, 953–971. [CrossRef]

71. Boden, M.; Hawkins, J. Prediction of subcellular localization using sequence-biased recurrent networks. *Bioinformatics* **2005**, *21*, 2279–2286. [CrossRef] [PubMed]

72. Horton, P.; Park, K.J.; Obayashi, T.; Fujita, N.; Harada, H.; Adams-Collier, C.J.; Nakai, K. WoLF PSORT: Protein localization predictor. *Nucleic Acids Res.* **2007**, *35*, W585–W587. [CrossRef] [PubMed]

73. Edgar, R.C. MUSCLE: Multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Res.* **2004**, *32*, 1792–1797. [CrossRef] [PubMed]

74. Kumar, S.; Stecher, G.; Tamura, K. MEGA7: Molecular Evolutionary Genetics Analysis Version 7.0 for Bigger Datasets. *Mol. Biol. Evol.* **2016**, *33*, 1870–1874. [CrossRef] [PubMed]

75. Saitou, N.; Nei, M. The neighbor-joining method A new method for reconstructing phylogenetic trees. *Mol. Biol. Evol.* **1987**, *4*, 406–425. [PubMed]

76. Zuckerkandl, E.; Pauling, L. Evolutionary divergence and convergence in proteins. In *Evolving Genes and Proteins*; Bryson, V., Vogel, H.J., Eds.; Academic Press: New York, NY, USA, 1965; pp. 97–166.
77. Voorrips, R.E. MapChart: Software for the graphical presentation of linkage maps and QTLs. *J. Hered.* 2002, 93, 77–78. [CrossRef]

78. Wang, Y.; Tang, H.; Debarry, J.D.; Tan, X.; Li, J.; Wang, X.; Lee, T.H.; Jin, H.; Marler, B.; Guo, H.; et al. MCScanX: A toolkit for detection and evolutionary analysis of gene syteny and collinearity. *Nucleic Acids Res.* 2012, 40, e49. [CrossRef]

79. Chen, C.; Chen, H.; Zhang, Y.; Thomas, H.R.; Frank, M.H.; He, Y.; Xia, R. TBtools: An Integrative Toolkit Developed for Interactive Analyses of Big Biological Data. *Mol. Plant.* 2020, 13, 1192–1204. [CrossRef]

80. Nei, M.; Gojobori, T. Simple methods for estimating the numbers of synonymous and nonsynonymous nucleotide substitutions. *Mol. Biol. Evol.* 1986, 3, 418–426.

81. Koch, M.A.; Haubold, B.; Mitchell-Olds, T. Comparative evolutionary analysis of chalcone synthase and alcohol dehydrogenase loci in *Arabidopsis, Arabis*, and related genera (Brassicaceae). *Mol. Biol. Evol.* 2000, 17, 1483–1498. [CrossRef]

82. Zhang, Z.; Deng, W.; Wang, Y.; Liu, Z.; Cheng, H.; Xue, Y. Heml: A Toolkit for Illustrating Heatmaps. *PLoS ONE* 2014, 9, e111988.

83. Wan, Y.; Wang, Z.; Xia, J.; Shen, S.; Guan, M.; Zhu, M.; Qiao, C.; Sun, F.; Liang, Y.; Li, J.; et al. Genome-Wide Analysis of Phosphorus Transporter Genes in *Brassica* and Their Roles in Heavy Metal Stress Tolerance. *Int. J. Mol. Sci.* 2020, 21, 2209. [CrossRef] [PubMed]

84. Guo, P.; Wen, J.; Yang, J.; Ke, Y.; Wang, M.; Liu, M.; Ran, F.; Wu, Y.; Li, P.; Li, J.; et al. Genome-wide survey and expression analyses of the GRAS gene family in *Brassica napus* reveals their roles in root development and stress response. *Planta* 2019, 250, 1051–1072. [CrossRef] [PubMed]

85. Ke, Y.Z.; Wu, Y.W.; Zhou, H.J.; Chen, P.; Wang, M.M.; Liu, M.M.; Li, P.F.; Yang, J.; Li, J.N.; Du, H. Genome-wide survey of the bHLH super gene family in *Brassica napus*. *BMC Plant Biol.* 2020, 20, 115. [CrossRef] [PubMed]