Multi-omics analysis reveals a link between Brassica-specific miR1885 and rapeseed tolerance to low temperature

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Abstract

Brassica crops include various edible vegetable and plant oil crops, and their production is limited by low temperature beyond their tolerant capability. The key regulators of low-temperature resistance in Brassica remain largely unexplored. To identify post-transcriptional regulators of plant response to low temperature, we performed small RNA profiling, and found that 16 known miRNAs were responsive to cold treatment in Brassica rapa. The cold response of seven of those miRNAs were further confirmed by qRT-PCR and/or northern blotting analyses. In parallel, a genome-wide association study of 220 accessions of Brassica napus identified four candidate MIRNA genes, all of which were cold-responsive, at the loci associated with low temperature resistance. Specifically, these large-scale data analyses revealed a link between miR1885 and the plant response to low temperature in both B. rapa and B. napus. Using 5' rapid amplification of cDNA ends approach, we validated that miR1885 can cleave its putative target transcripts, Bn.TIR.A09 and Bn.TNL.A03, in B. napus. Furthermore, overexpression of miR1885 in Semi-winter type B. napus decreased the mRNA abundance of Bn.TIR.A09 and Bn.TNL.A03, resulting in increased sensitivity to low temperature. Knocking down of miR1885 in Spring type B. napus led to increased mRNA abundance of its targets and improved rapeseed tolerance to low temperature. Together, our results suggested that the loci of miR1885 and its targets could be potential candidates for the molecular breeding of low temperature-tolerant Spring type Brassica crops.

INTRODUCTION

Brassica crop species include Brassica napus (rapeseed), Brassica rapa, Brassica oleracea, Brassica juncea, and Brassica nigra. Among them, B. napus arose approximately 7500 years ago by allopolyploidization between its ancestors, B. rapa and B. oleracea (Chalhoub et al., 2014). B. rapa and Arabidopsis thaliana share a common ancestor, but B. rapa has a larger genome than that of A. thaliana due to genome triplication (Wang et al., 2011). Both B. napus and B. rapa are important food sources. B. napus mainly provides edible plant oil and B. rapa is a valuable oil and vegetable crop. However, Brassica crops are negatively affected by low temperature, especially those that encounter frigid environment beyond their cold-tolerant capability. Worldwide, extensive economic losses of Brassica crops are attributed to low temperature, often in combination with other abiotic or biotic stresses (Sanghera et al., 2011). Brassica crops mainly have three ecotypes: winter, spring, and semi-winter types. The winter and semi-winter type Brassica require vernalization, which are more tolerant to low temperature as compared to spring type Brassica (Gomez-Campo and Prakash, 1999). Low temperature includes coldness ($>0$ and $<10$) and freeze ($<0$) (Ding et al., 2019). At the early seeding stage, long-term low temperature stress ($4{^\circ}C$) would restrict the growth of spring Brassica crops and even cause plant damage and yield loss (O’Neill et al., 2019). It is important
to identify key regulators contributing to plant response to cold stress for improving the cold resistance of spring type Brassica crops.

The vernalization of semi-winter type Brassica crops require moderate low temperature. However, long time exposure to low temperature in winter leads to plant damage at vegetative stage and even lead to yield loss (Liao and Guan, 2001; Zhang et al., 2015). The Brassica crops are usually sown in autumn after the harvest of rice in Yangtze River basin, China (Cong et al., 2019). In recent years, the delay of rice harvest usually led to the postpone of rapeseed sowing resulting in poor Brassica crop seedlings establishment due to low temperature. Lacking enough biomass of Brassica crops in winter cause the plants more susceptible to cold or freezing stress (Luo et al., 2019; Zhang et al., 2012). Moreover, some Spring type accessions would die due to long-term low temperature and decrease of seed yield (Ozer, 2003).

Previous studies have identified a number of genes involved in the plant response to low temperature (Ding et., 2019). Cold stress induced the expression of the transcription factor gene CBF (encoding C-repeat binding factor), and CBF triggered the expression of COR(COLD RESPONSIVE) genes (Shi et al., 2018). CBF/COR-dependent factors allowed plants to withstand subsequent freezing stress (McClung and Davis, 2010). In addition, many important genes, which contributed to low-temperature response in CBF-independent manner, were also found (Bolt et al., 2017; Liu et al., 2018). Although breeding for low-temperature resistance is possible, the genetic improvement of crops is still hindered by the lack of elite genetic resources related to low-temperature resistance.

MicroRNAs (miRNAs) are regulators of plant adaptation to abiotic or biotic stress (Biggar and Storey, 2015). They are approximately 20–24 nucleotides (nt) in length and originate from pre-miRNAs with stem-loop structures (Song et al., 2019). The mature miRNAs are loaded into the RNA-induced silencing complex that regulate target transcripts involved in various aspects of plant growth, development, and stress responses, including response to cold or freezing stress (Chen, 2009; Anjali and Sabu, 2020). For instance, overexpression of miR156 increased cold tolerance in rice, A. thaliana, and pine (Zhou and Tang, 2019). In rice, OsmiR319b down-regulated OsPCF6 and OsTCP21, resulting in enhanced cold tolerance (Wang et al., 2014), and overexpressing OsmiR1320 improved cold tolerance, although OsmiR1320 was repressed under cold stress (Sun et al., 2022). In A. thaliana, plants knocking-down miR165/166 displayed a drought- and cold-resistant phenotype (Yan et al., 2016). Other known cold-responsive miRNAs included miR172, miR396, miR845, and miR168 (Gupta et al., 2014; Zeng et al., 2018; Chen et al., 2019). However, miRNAs in response to low temperature stress have not been well studied in Brassica crops.

Plant R (RESISTANCE) gene encodes immune protein involved in the response to pathogen infection (Jones and Dangl, 2006). Most of the identified R genes contain a nucleotide-binding site-leucine-rich repeat (NBS-LRR) domain with a coiled-coil (CC) or a Toll/Interleukin-1 receptor (TIR) domain at the amino terminus (Dangl and Jones, 2001). Emerging evidence has shown that cold stress and pathogen attack induced the expression of some common genes, such as PR(PATHOGENESIS-RELATED) genes, which played roles in the response to pathogen infection and low temperature stress (Snider et al., 2000; Seo et al., 2008). Some R genes negatively regulated plants tolerance to low temperature. In A. thaliana, the rpp4-1d mutant plant with a gain-of-function mutation in RPP4 (TIR-NB-LRR protein PERONOSPORA PARASITICA 4) was chilling-sensitive (Huang et al., 2010). Mutations in HSP90 suppressed the chilling-sensitive phenotype of rpp4-1d mutant plant, and HSP90 interacted with the RPP4 protein in planta (Bao et al., 2014). Interestingly, a significant increase in HSP90 mRNA was detected in B. napus exposed to low temperature (Krishna et al., 1995). In addition, a gain-of-function mutant of CHS3, encoding a TIR-NB-LRR-LIM protein, resulted in enhanced defense responses and chilling-sensitivity phenotype (Yang et al., 2010). Other studies found that some R genes played positive roles in plants tolerance to low temperature. A point missense mutation of CHS1, encoding a TIR-NB protein, resulted in a chilling-sensitive phenotype (Yang et al., 2010; Zbierzak et al., 2013). R genes were also regulated by genes that contributed to cold tolerance. In tobacco, overexpression of SyRVE6 (REVEILLE) improved plant tolerance to low temperature and up-regulated the expression of genes encoding NB-LRR proteins (Chen et al., 2020).

Although many studies have investigated the molecular mechanisms underlying low-temperature resistance,
the roles of miRNAs in low-temperature resistance in Brassica crops are largely unknown. In this study, we aimed to identify cold-responsive miRNAs in winter or semi-winter Brassica and applied them to improving cold resistance of Spring type Brassica crops. To do this, we constructed and sequenced small RNA libraries from B. rapa under cold stress and performed a genome-wide association study (GWAS) related to the phenotype of cold resistance using a B. napus population. Combining the results from both small RNA sequencing and GWAS, we found that Brassica-specific miR1885 was responsive to cold stress in both B. rapa and B. napus. Previous studies have revealed that heat, as well as Turnip mosaic virus (TuMV) and Plasmodiophora brassicae infection, induced miR1885 accumulation (He et al., 2008; Yu et al., 2011, Paul et al., 2021). Here, we further demonstrated that miR1885 was also involved in the response to low-temperature stress through its target genes and functioned as a negative regulator of B. napus in low temperature tolerance, providing potential genetic candidates for cold-resistant rapeseed breeding.

RESULTS

Genome-wide analysis of small RNAs responsive to cold stress in B. rapa

To study the cold response of B. rapa, the degree of cold tolerance was estimated by evaluating morphological traits under cold stress. After 7 days of cold treatment (4 degC), the plants stopped growing and their leaves drooped (Figure 1A). Therefore, to identify genome-wide cold-responsive miRNAs, we performed cold treatment for 7 days on 3-week-old seedlings of B. rapa in a growth chamber, and plants grown at 22 degC was used as control for small RNA sequencing.

Totally, 17.96 million and 17.17 million raw reads were obtained from these two small RNA sequencing samples treated with 22 degC and 4 degC, respectively (Figure 1B). Among them, most of small RNA sequences were shared between these two libraries, while 2.82 million small RNA sequences were specific in the seedling under normal temperature, and 2.23 million small RNA sequences were specific in the seedling under 4 degC (Figure 1B). Small RNA length distribution analysis showed that 24-nt small RNAs were most abundant, and 21-nt small RNAs were second most abundant (Figure 1C).

To learn the genomic loci that produced these small RNAs, 11.46 million (22 degC) and 11.36 million (4 degC) clean reads were mapped to the B. rapa reference genome (http://brassicadb.cn/#/). We found that the small RNAs were generated from exons, introns, miRNAs, rRNAs, snRNAs, snoRNAs, tRNAs, and repeat regions (Table S1). Notably, focusing on small RNAs from non-coding gene regions, most abundant small RNAs were produced from ribosomal RNAs (rRNAs) (22 degC, 44.67% and 4 degC, 48.10%), unknown RNAs (22 degC, 27.65% and 4 degC, 25.22%), and miRNA precursors (22 degC, 8.50% and 4 degC, 8.71%) (Figure S1).

Identification of B. rapa miRNAs responsive to cold stress

To identify known miRNAs in our sequence libraries, we aligned the small RNA sequences with mature miRNAs in miRBase and the B. rapa reference genome, allowing no more than two mismatches. In total, 77 mature miRNAs related to 184 conserved miRNA precursors were found in our libraries. Among them, 52 miRNAs were conserved with those in Arabidopsis, while 16 miRNAs were Brassica-specific miRNAs (Table S2). To further identify novel miRNA candidates, the flanking sequences of unknown small RNAs in genome were extracted for RNA secondary structure prediction using RNAfold. Based on the plant miRNA criteria (Axtell et al., 2018, see the detail in the method), we identified 11 highly credible novel miRNAs (Table S3). Among them, novel-miR1 and novel-miR11 were identified with more than 100 reads.

Among the 77 known miRNAs, 16 miRNAs were differentially expressed between the sample under cold treatment and the control (>1.5-fold) (Table 1). Among these mature miRNAs, bra-miR157a~c, bra-miR161, bra-miR393f, bra-miR396b, bra-miR399c, bra-miR156k, bra-miR1140, bra-miR158a and bra-miR156a~f were down-regulated; and bra-miR168d, bra-miR403, bra-miR390a~b, bra-miR398b~c, bra-miR408, and bra-miR1885 were up-regulated under cold treatment. Among them, the Brassica-specific bra-miR1885 was induced more than four-fold by cold stress. Among the novel miRNAs, only novel-miR11 was cold-responsive. Taken together, the small RNA profiling provided potential candidates of cold-responsive miRNAs in B. rapa.
Validation of cold-responsive mature miRNAs in *B. rapa*

To validate the cold-responsive miRNAs detected by small RNA sequencing, we further analyzed the cold responsiveness of mature bra-miR1885, bra-miR398, bra-miR408, bra-miR161, bra-miR157, bra-miR393, and bra-miR168 using qRT-PCR. We found that the abundance of bra-miR1885, bra-miR398 and bra-miR168 were significantly up-regulated by cold stress, whereas bra-miR161, bra-miR157 and bra-miR393 were significantly down-regulated under cold stress, as compared to normal temperature at the same growth stage. These results were consistent with the small RNA sequencing data (Figure 2A). Among these miRNAs, bra-miR398 and bra-miR168 were conserved miRNAs that were induced more than three-fold by cold treatment (Figure 2A). We then performed northern blotting analyses to further confirm the cold responsiveness of bra-miR398, bra-miR168 and bra-miR1140. Consistently, the accumulation of bra-miR398 and bra-miR168 was much higher in 4 degC than in 22 degC, and *Brassica*-specific miR1140 was repressed after cold treatment (Figure 2B). In addition, the results of qRT-PCR analyses further confirmed the abundance of mature bra-miR1885 was increased in leaf under cold stress as compared to normal condition (Figure 2C).

Next, to identify the potential cold-regulated *cis*-elements in the bra-MIR1885 promoter, we extracted the promoter sequence of bra-MIR1885 using Promoter 2.0 Prediction Server (https://services.healthtech.dtu.dk/service.php?Promoter-2.0), and analyzed the *cis*-elements in the putative bra-MIR1885 promoter sequence using PlantCare (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/). These analyses revealed that the promoter contained light-responsive, defense-responsive, stress-responsive, and low temperature-responsive elements (LTR: CCGAAA) (Figure S2). A previous study has revealed that LTR motif (CCGAAA) within the promoter region of CsINV5 was the core *cis* element in response to low temperature (Qian et al., 2018). This result suggested that the bra-MIR1885 promoter contained a low temperature-responsive element that potentially regulated the expression of bra-MIR1885 under cold stress.

Identification of low temperature resistance-associated loci containing MIRNA genes in *B. napus* using Genome-wide association study

*B. napus* is closely related to *B. rapa* and both are *Brassica* crops. To identify MIRNA genes associated with low temperature stress in *Brassica* crops, we constructed a *B. napus* population with 220 accessions. The phenotype of low-temperature resistance was divided into 5 levels (Grade 1 indicates most cold-resistant, while Grade 5 indicates most cold-sensitive) and was measured in the field in January of 2017 and 2018, and the phenotypic data of rapeseed population showed a nearly skewed distribution (Xu et al., 2021). The genome re-sequencing of these 220 *B. napus* accessions generated 2.13 Tb clean reads, and single nucleotide polymorphism (SNP) genotyping of *B. napus* population identified 3.80 million highly quality SNPs (MAF > 0.05, geno < 0.2) (Xu et al., 2021). Overall, the SNPs were not evenly distributed across the whole genome. The SNP density in the A sub-genome (5.25 SNP/Kb) was higher than that in the C sub-genome (4.74 SNP/Kb) (Figure S3A). A principal component analysis (PCA) was used to assess the genetic relationship in the association panel. Based on the PCA, the *B. napus* population was roughly divided into three groups corresponding to winter (W), semi-winter (SW), and spring (S) ecotypes (Figure S3B). We further analyzed the linkage disequilibrium (LD) throughout the genome. The LD was estimated as *r*² (the squared Pearson correlation coefficient) between all pairs of SNP markers. The average distance over which LD decayed to *r*² =0.2 of its maximum value was 100 kb, indicating that SNPs located in the 200-kb genomic region around each peak SNP represented a GWAS-quantitative trait locus (QTL) (Figure S3C).

The GWAS was performed using a mixed linear model, and the SNPs significantly associated with low-temperature resistance were identified at a threshold of *p* < 10⁻⁶. As a result, we identified 32 highly credible GWAS-QTLs (Figure 3A). To discover candidate MIRNA genes within these QTLs, we collected 163 pre-miRNA sequences specific to the Cruciferaceae family from four species (*B. napus*, *B. rapa*, *B. oleracea*, and *Arabidopsis*) according to previous reports (Jones-Rhoades and Bartel, 2004; Alves-Junior et al., 2009; Yu et al., 2011; He et al., 2018; Zhang et al., 2018; Li et al., 2019). All the pre-miRNA
sequences were mapped to the *B. napus* reference to identify their location (Table S4). We found that four candidate MI RNA genes, *bna-MIR166C*, *bna-MIR1885*, *bna-MIR168A* and *bna-MIR845A*, were in the QTLs associated with low-temperature resistance.

A previous study proved that knock-down of miR165/166 expression conferred a cold-resistant phenotype in *A. thaliana* (Yan et al., 2016). Conserved miR845a was known to be differentially expressed in leaves of winter turnip rape under cold stress (Zeng et al., 2018). To further determine whether these four miRNAs were responsive to low temperature, we measured the abundance of mature miRNAs in *B. napus* under cold stress by qRT-PCR. From this analysis, we found that all of them were cold-responsive. bna-miR1885 and bna-miR168a were up-regulated under cold stress compared to under normal condition at the same growth time points, while bna-miR845a and bna-miR166c were down-regulated (Figure 3B and Figure 4A).

To identify if any genetic variation in cis-regulatory element contribute to the accumulation of mature miRNAs in the population of rapeseed, we studied the LD of SNPs in *bna-MIR1885*, *bna-MIR166C*, *bna-MIR168A* and *bna-MIR845A* gene locus, respectively. We found that the SNPs in their corresponding genes exhibited significant linkage (Figure S3D, F, G, H). Importantly, we identified 22 SNPs and 7 INDELS in *bna-MIR1885* promoter region, which were found to be significant associated with cold resistance. We further investigated the five variants located in the 200-bp genomic region around the low temperature-responsive (LTR: CCGAAA) element identified above. The *B. napus* accessions were classified into three haplotype groups based on the genotypes of these variants (Figure 3C). We found that 179 accessions contained the H1 haplotype, which variants were identical with the genome reference, and this haplotype was associated with low temperature-sensitive phenotype. 24 accessions carried H2 haplotype, which contained mutated LTR cis-element and was associated with low temperature-resistant phenotype. Consistently, H3 haplotype had an average resistance degree, which indicated more resistant than H1 haplotype, and less resistant than H2. Next, we randomly selected 4 accessions with low temperature-resistant variant alleles and low temperature-sensitive variant alleles, respectively, to measure their abundance of miR1885 after cold treatment by qRT-PCR. The qRT-PCR data indicated that the abundance of miR1885 was significantly higher in low temperature-sensitive accesses with normal LTR in the promoter (H1 haplotype), as compared to low temperature-resistant accesses with mutated LTR in their promoter (H2 haplotype) (Figure S3E). In total, these results suggested that genetic variation in the LTR region at *MIR1885* promoter in the rapeseed population potentially affected the miR1885 expression and were associated with cold resistance.

**Identification of miR1885 target genes and their cold responsivity in *B. napus***

Because bra-miR1885 and bna-miR1885 shared the same mature sequence (Figure 4B), we used miR1885 to represent the two miRNAs in *Brassica* crops hereafter. We first predicted the candidate target genes of miR1885 in *B. napus* using a plant small RNA target analysis tool (psRNATarget). Two *R* genes, *Bn.TIR.A09* (*BnaA09g14980D*) and *Bn.TNL.A03* (*BnaA03g56180D*), were identified as highly credible targets, and the miR1885 target sites were within the TIR domain. To verify the miR1885 cleavages of the predicted target genes, we performed a 5' rapid amplification of cDNA ends (5'-RACE) assay in *B. napus* and detected miR1885 cleavage sites frequently within *Bn.TIR.A09* and *Bn.TNL.A03* (Figure 4C and Figure S4), indicating that they were targeted by miR1885 in *B. napus*.

Next, we examined the transcript levels of these two target genes under cold stress, using three low temperature-responsive marker genes (*Bn.CBF1.C03*, *Bn.HSP70.A01* and *Bn.ERF105.C09*) as positive controls (Wang and Hua, 2009; Bolt et al., 2017; Liu et al., 2018). The target genes were down-regulated under cold stress in *B. napus* (Figure 4D), while the expression of *Bn.CBF1.C03* and *Bn.HSP70.A01* were induced and *Bn.ERF105.C09* were repressed in samples under cold treatment as compared to those under normal condition, consistent with those reported in previous studies (Figure 4E).

**Overexpression of miR1885 in *B. napus* led to down-regulation of its target genes *Bn.TIR.A09* and *Bn.TNL.A03***

Genomic synteny analysis using TBtools (https://github.com/CJ-Chen/TBtools) suggested that bna-miR1885 in *B. napus* is inherited from the genome of *B. rapa* through genome merging (Figure S5), and
bna-miR1885 and bra-miR1885 shared the same mature sequence. To further verify the function of miR1885, we constructed a vector with the bra-MIR1885 gene under the control of the AA6 promoter (Patent WO 2007/069894) and transformed it into B. napus cv. K407 (Figure 5A). The strategy using B. napus system for functional analysis of B. rapa genes have been used in several studies (Liu et al., 2014; Tian et al., 2018; Cui et al., 2020). In total, seven positive transgenic lines (T0) were screened by PCR (Figure 5B and C). The expression level of pri-miR1885 was confirmed by qRT-PCR. We found that pri-miR1885 was overexpressed in the leaves of X2, X8, X11, X12, and X15 lines (Figure 5D). Compared with WT, there were no obvious morphological abnormalities among miR1885-OE lines at seeding stage (Figure 5B). We further checked the abundance of mature miR1885 in the transformed lines by qRT-PCR and northern blotting analyses. As expected, mature miR1885 was found to be over-accumulated in the X2, X8, X11, X12, and X15 lines (Figure 5E and 5F). And then, we conducted qRT-PCR analyses to validate the influence of the miR1885 on the expression level of the two target genes in the transgenic lines. As expected, transcript levels of Bn.TIR.A09 and Bn.TNL.A03 were lower in the transgenic lines than in wild type (Figure 5G and 5H).

Next, we checked the RNA levels of miR1885 and its targets in miR1885-OE lines and WT with or without cold treatment by qRT-PCR. As expected, the abundance of miR1885 was increased in miR1885-OE lines and WT after cold treatment (Figure 6A). In addition, the RNA levels of Bo.TIR.A09 were further decreased in miR1885-OE under cold stress (Figure 6B), but those of Bn.TNL.A03 were unchanged (Figure 6C), suggesting that there was additional cold-triggered regulation of Bn.TNL.A03 at the transcriptional level, while the cold response of Bo.TIR.A09 was majorly contributed by miR1885-mediated post-transcriptional regulation. Given that CBF/CORs played critical roles in plant response to low temperature, we investigated whether miR1885 can affect the cold induction of CBF/COR factors. To do this, we analyzed the expression levels of Bn.CBF1.CO3, Bn.CBF2.Ann and Bn.COR15.A03 in the miR1885-OE under cold treatment. However, there were no significant differences at the expression levels of Bn.CBF1.CO3, Bn.CBF2.Ann and Bn.COR15.A03 after cold treatment between WT and miR1885-OE lines (Figure 6D-F). This result suggested that miR1885 regulated low temperature tolerance in Brassica through CBF/CORs-independent pathway.

miR1885 negatively regulated low temperature tolerance in Brassica

A previous study reported that knockdown of CHS1, encoding a TIR-NB protein, led to a chilling-sensitive phenotype in A. thaliana (Zbierzak et al., 2013). We found that two TIR-NB transcripts were the targets of miR1885 and both miR1885 and these two TIR-NB genes were cold-responsive. Therefore, we further tested the effects of miR1885 and its targets on low-temperature resistance in B. napus. Semi-winter type WT K407 and miR1885-OE plants were grown for 6 weeks at 22 degC, and then were transferred to 4 degC for 30 days or 2 degC for 7 days. We found that the miR1885-OE plants were more sensitive as compared to WT under 2 degC for 7 days (Figure 7A) but showed no apparent phenotypic difference under 4 degC (Figure S6A). This result indicated miR1885 functioned as negative regulator in rapeseed response to low temperature. We further performed phenotypic studies using WT K407 and miR1885-OE seeds during post-germination under normal and cold environment. The seeds were harvested at the same season and the cold environment was 6 degC. The hypocotyl length (NHL) of WT K407 and miR1885-OE showed no difference after growing for 2 days on the germination bed without cold treatment. However, after 7 days cold treatment, the miR1885-OE exhibited shorter NHL compared with WT K407 (Figure S6B-D), indicating that miR1885 also played critical role in cold response in the early developmental stage, when no significant phenotypic difference between WT and miR1885-OE was observed under normal condition.

To test that whether the cold-responsive miR1885 can be applied to improve low temperature tolerance of Spring type Brassica, the miR1885 knockdown plants (STTM1885) generated in Sping type Brassica napus cv. ‘Westar’ were used for further study (Cui et al., 2020). The qRT-PCR results showed that the accumulation of miR1885 was reduced in STTM1885 plants. Compared with those in wildtype Westar, the expression levels of Bn.TNL.A03 and Bn.TIR.A09 were increased in STTM1885 plants (Figure 7B).
study morphological responses to cold stress, we then moved the Westar and STTM1885 plants into 2 degC environment. After 14 days cold treatment, a cold-tolerant phenotype was observed in STTM1885 plants (Figure 7C). This finding further suggested that attenuating miR1885 resulted in enhanced cold tolerance of rapeseed potentially via their targets.

We further tested whether overexpression of miR1885 affected plant growth at freezing temperature compared to its background winter ecotype K407, while the spring ecotype Westar was used as negative control (Figure S7A). The abundance of mature miR1885 in wild-type B. napus under freezing stress was examined by real-time PCR. Based on previous studies, we set the freezing temperature at -10 degC (Xu and Cai, 2019; Ljubej et al., 2021). We found that miR1885 was up-regulated under freezing stress, and its expression level was approximately 2.4-fold than in wild type at 2.5 h (Figure S6B). The wild type and miR1885-OE lines were subjected to a -10 degC treatment for 2.5 h, then allowed to recover at 22 degC for 7 days. The survival rate of wild type K407 was 78.00%, and those of miR1885-OE were much lower, ranging from 22.33% to 55.33%, whereas 100% of spring ecotype Westar died (Figure S7A and Figure S7C). Accumulation of free proline (Pro) during cold or freezing stress is thought to protect plants (Xin and Browse, 1998). To test whether miR1885-OE had altered Pro content, we determined the Pro content in wild type and miR1885-OE with or without freezing treatment (Figure STD). As we expected, the Pro content was higher in wild type and miR1885-OE grown at -10 degC than in those grown at 22 degC. In plants grown at -10 degC, more Pro accumulated in the wild type plants than in the miR1885-OE. This finding was consistent with the freezing-sensitive phenotype of transgenic lines subjected to a -10 degC treatment. These results demonstrated that overexpression of miR1885 increases plant sensitivity to freezing at the seedling stage.

**DISCUSSION**

At present, many MIRNA genes functioning in stress response have been identified by small RNA profiling and RNA-seq (Ahmed et al., 2020; Ma et al., 2022; Zhu et al., 2022). However, the MIRNA genes that function in the response to cold stress in crops remain largely unexplored. In this study, we used small RNA profiling and GWAS to identify cold-responsive candidate miRNAs or MIRNA genes associated with low-temperature resistance. The results from these two methods provide mutual support for the roles of miR1885 in the cold response of *Brassica* crops.

We identified up-regulated or down-regulated miRNAs in small RNA sequencing libraries of *B. rapa* and candidate MIRNA genes from GWAS loci associated with low-temperature resistance in *B. napus*. The small RNA profiling suggested that miR1885 was drastically triggered by cold stress. Using GWAS, we found *MIR1885* gene was in the loci associated with low-temperature resistance. We further confirmed that miR1885 was induced by cold stress in both *B. rapa* and *B. napus*. We used the newest version of *B. rapa* and *B. napus* genome reference to analyze their genomic synteny. As expected, the *B. napus* miR1885 (ChrA06) is inherited from the genome of *B. rapa* (ChrA06 genome) through genome merging (Fig. S5). Additionally, For the cold responsive miRNAs in this study, we found some miRNAs have been reported to regulate flowering, such as miR156/7, miR168 and miR158. This suggested that some the cold-responsive miRNAs also play roles in vernalization.

Mature miR1885 is a *Brassica*-specific miRNA that targets the *R* gene family. In *A. thaliana*, the TIR-NB-LRR protein RPP4 and CHS1 were involved in plant resistance to low temperature stress (Huang et al., 2010; Zbierzak et al., 2013). Here, two *R* genes, *Bn.TIR.A09* and *Bn.TNL.A03*, were predicted as targets for cleavage by miR1885 in *B. napus*. These *R* genes belonged to two different subgroups: *Bn.TIR.A09* encoded a protein with the TIR motif lacking NBS-LRR domain; while *Bn.TNL.A03* encoded a protein with the entire TIR-NBS-LRR structure. We found that these two targeted genes were down-regulated in wild-type *B. napus* under cold treatment, opposite to the expression pattern of miR1885 (Figure 6). These results demonstrated that miR1885 negatively regulated these *R* genes under cold temperatures.

Overexpression of miR1885 in semi-winter type *B. napus* attenuated rapeseed sensitivity to low temperature at the seedling stage. While knocking down of miR1885 in Spring type *B. napus* (STTM1885 lines) improved plant tolerance to low temperature (Figure 7). Additionally, we did not find obvious differences between
knock-down (STTM1885) lines and control lines at the early seeding stage. But STTM1885 did delay the flowering time and increased the plant height at a later stage. The RNA levels of miR1885-cleaved R genes Bn.TIR.A09 and Bn.TNL.A03, which increased in STTM1885 rapeseed and decreased in miR1885-OE lines, may contribute to rapeseed response to cold stress. Although a link between miR1885 and cold response has been revealed, the hiding mechanism is still uncovered. In addition, we have checked some CBF-dependent genes in miR1885-OE lines, including CBFs and CORs, and found that the transcript levels of these genes in miR1885-OE in response to cold stress were no different with that of wild type, suggesting that miR1885-targets regulated plant resistance to low temperature independent on CBFs pathway.

In this study, we revealed a link between miR1885 and cold response in Brassica through its targeted R genes. However, how miR1885 and these R genes involve in CBF-independent cold response in rapeseed requires further examination. In Arabidopsis, 22-nt miRNAs can trigger the production of 21-nt phased secondary siRNAs (Cuperus et al., 2010; Li et al., 2012). 22-nt miR482/2118, which widely present in seed plants, triggered phasiRNA biogenesis from its targeted R genes (NBS-LRR), forming another layer of regulation to reinforce its silencing effect (Fei et al., 2013; Liu et al., 2020). Therefore, 22-nt miR1885 in rapeseed possibly triggers phasiRNAs biogenesis from its targeted R genes, and these phasiRNAs may target unknown downstream genes to regulate plant response to low temperature stress. Additionally, co-immunoprecipitation (co-IP) followed by mass spectrometry can be performed to identify proteins interacting with miR1885-regulated R proteins under cold stress, which may play role in cold signal transfer and resistance to cold stress.

In rice, OsmiR1320 was repressed by cold stress. However, overexpressing OsmiR1320 enhanced plant tolerance to cold stress, indicating that OsmiR1320 played positive role in cold response (Sun et al., 2022). On the contrary, we found that, as a negative regulator of cold response, miR1885 was induced under cold stress potentially via its low-temperature response (LTR) cis-element in MIR1885 promoter. Therefore, MIRNA transcription could be regulated incoherently by cold stress. The genetic mutation in and close to this LTR in the rapeseed population may contribute to their diversity of cold response (Figure 3C and Figure S3E). Genetic editing in this LTR could be used for detaching the cold response of MIR1885, and therefore enhance rapeseed resistance to cold stress. Previous works showed that over-expression of miR1885 affected development traits and biotic response, including flowering time and disease resistance. The abundance of miR1885 may regulated by different signals through potentially different promoter cis elements and played a very important role in regulating both plant growth and development.

In summary, using multi-omics data, we identified miR1885 involved in plant response to low temperature stress by small RNA sequencing and GWAS. Overexpression of miR1885 in B. napus not only increased plant sensitivity to low temperature, while knockdown of miR1885 improved plant tolerance to low temperature. Our findings suggested that MIR1885 and its target genes can be used as genetic resource for improving resistance to low temperature in the breeding of Brassica crops.

METHODS AND MATERIALS

Plants materials, cold treatment, and freezing tolerance assay

The wild-type B. rapa variety ‘Bre’ was grown under a 16-h light/8-h dark (22 degC) photoperiod for 3 weeks. To analyze the miRNAs abundance in ‘Bre’ under cold stress, some ‘Bre’ plants were moved to 4 degC condition for 7 days cold treatment and some were grown at 22 degC for additional 7 days as control. The aboveground parts of treated and untreated were then harvested, frozen in liquid nitrogen, and stored at -80 degC until extraction of total RNA. Small RNA sequencing (one biological replicate) was performed using the Illumina GAI1 sequencer at BIG (ShenZhen, China).

The wild-type B. napus variety ‘K407’, a Semi-winter ecotype, was grown under a 16-h light/8-h dark (22 degC) photoperiod with light intensity of 250 μmol m\(^{-2}\)s\(^{-1}\). For the cold resistance assay, the plants were grown under standard conditions (22 °C) for 6 weeks before transferring to a cold environment (4 °C or 2 °C, 16-h/8-h light/dark photoperiod). The similar methods for studying cold stress in Brassica crops have been used in previous reports (Raza et al., 2021; Hussain et al., 2022). After cold treatment, the cold resistance
of plants was determined. The freezing tolerance assay was performed in a freezing chamber, after cold acclimation for 24 h, in which wild-type and transgenic lines were subjected to a -10 degC treatment for 2.5 h. The plants were then allowed to recover at 22 degC for 7 days before determining the survival rate. The Pro content was measured immediately after the freezing treatment using a commercial assay kit (A107-1-1, Nanjing JianCheng Bioengineering Institute, Nanjing, China), according to the manufacturer’s instructions.

The wild-type *B. rapa* cultivar “Bre”, *B. napus* cultivar ‘K407’ are all inbred lines.

**Identification of known and novel miRNAs in *B. rapa***

We aligned the clean reads with known miRNAs in the miRBase (https://www.mirbase.org/), allowing two mismatches using TopHat (http://ccb.jhu.edu/software/tophat/index.shtml). The potential miRNAs were mapped to *B. rapa* reference genome (http://brassicadb.cn/#/) to extract the 600-bp flanking sequence. The flanking sequence of miRNAs were also aligned with known pre-miRNAs to confirm their identities of miRNA family. The flanking sequence of unknown small RNAs was also used to predict secondary structures using RNAfold software (http://rna.tbi.univie.ac.at/cgi-bin/RNAWebSuite/RNAfold.cgi). To identify novel miRNA, Small RNAs that showed less than 2 mismatches with known miRNAs in miRbase were filtered. The potential miRNAs that had fewer than 10 hits in the reference genome were retained. 600-bp flanking sequences of small RNAs were used to predict their secondary structures using RNAfold software with default parameters. Novel miRNAs were identified according to the plant miRNA criteria (Axtell et al., 2018, Yu et al., 2011). In brief, a key judgement was made by determining whether the potential miRNAs and their miRNA* were on the stem with fewer than four bulges, and the miRNA candidate were most abundant among small RNAs generated from given miRNA precursors. After Blast these novel miRNA precursors in NCBI, we found them were only detected in Brassica, therefore, they were defined as Brassica-specific miRNAs.

**Real time PCR**

Total RNA was isolated with TRIzol reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s instruction. High-quality total RNA (A260/280 > 2.00) was reverse-transcribed into cDNA. Small RNA qRT-PCR was conducted using a miRNA First Strand cDNA Synthesis kit (stem-loop method) (Sangon Biotech, Shanghai, China). A total of 3 μg total RNA was used for first-strand cDNA synthesis with the stem-loop primer and U6 reverse primer in a 20-μL reaction volume. The U6 snoRNA was used as an internal control. For qRT-PCR analysis of coding genes, 2 μg total RNA treated with DNase I was reserve-transcribed into cDNA using M-MuLV (Takara, Otsu, Japan) with oligos (dT). The internal control was ACT7. qRT-PCR was performed using SYBR mix (YEASEN, China) on a Bio-CFX96 instrument. Three biological replicates and three technological replicates were analyzed. Real-time PCR and reverse transcriptase (RT)-PCR were performed using gene-specific primer pairs (Table S5).

**Northern blot analyses**

A 30–45 μg aliquot of high-quality total RNA (A260/280 >2.00) was separated on a 19% polyacrylamide denaturing gel, and then transferred to a Hybond membrane (GE healthcare) for 2 h at 200 mA at 4 °C. In brief, after crosslinking for 5 min with ultraviolet irradiation, the membrane was hybridized overnight at 45 °C with DNA probes (Yu et al., 2011). Washing and autoradiography of the Hybond membrane were performed according to the instructions of the North2South Chemiluminescent Hybridization and Detection kit (No.17097, Pierce, Rockford, IL).

**Genome-wide association analysis (GWAS)**

A diversity panel, consisting of 220 *B. napus* inbred lines was used for the GWAS in this study. The accessions were grown using a randomized complete block design with two replicates. Each accession was grown in plots with four rows. For the phenotypic data about low temperature resistance of rapeseed population showed a nearly skewed distribution. For more details about field experiment design and the distribution of phenotype were described in our previous reports (Xu et al., 2021). More specifically, the population included winter, semi-winter and spring oilseed rape. The accessions originated from Asia, Europe, and
North America. The B. napus population was grown at the Farm Station in Yangling, Shaanxi province, China (107°59′–108°08′E; 34°14′–34°20′N), where the local temperature ranged from -15 to 10 degC in winter (https://lishi.tianqi.com/xianyang/202101.html). The reference genome of B. napus “Dar- mor” (version 4.1) was used to call variants, which was downloaded from BRAD (http://brassicadb.cn/#/) (Chalhoub et al., 2014). Low-temperature resistance was measured in the field in January of 2017 and 2018. The grade of low-temperature resistance was evaluated with criteria including grade 1 (the plants are normal, and the leaves are green), grade 2 (a small part of leaves are necrotic), grade 3 (a half of leaves are necrotic), grade 4 (most of leaves are necrotic), and grade 5 (all leaves are necrotic). The association analysis for low-temperature resistance was carried out using a MLM with the software package TASSEL 5 (https://www.softpedia.com/get/Science-CAD/TASSEL.shtml). The population structure was calculated using PCA. The software PopLDdecay (https://github.com/BGI-shenzhen/PopLDdecay) was used to calculate LD. The parameter $r^2 = 0.2$ was read as LD length. The threshold of the GWAS was set to $P$-value $< 10^{-6}$.

5'-RACE

The 5’modified RACE was performed using the FirstChoice RLM-RACE Kit (Invitrogen). In brief, total RNA was isolated from young leaves of B. napus. Then, 8–10 μg high-quality total RNA (A260/280 [≥ 2.00]) was directly ligated to the 5'-RACE adapter without calf intestinal phosphatase and tobacco acid pyrophosphatase treatment. The first-strand cDNA was synthesized using M-MLV reverse transcriptase according to the manufacturer’s instructions. The PCR amplification was performed using the 5’ outer primer and gene-specific outer primer, and 0.1 μL PCR product was used as the template for nested PCR. Nested PCR amplification was performed using the 5’ inner primer and gene-specific inner primer. The PCR products were gel-purified and then cloned and sequenced.

Plasmid construction, genetic transformation, and phenotypic analysis

The plant overexpression vector was generated using pCAMBIA1301 harboring the AA6 promoter and the tAA6 terminator. The bra-MIR1885 genomic DNA fragment was isolated from ‘Bre’ seedlings and cloned into the pCAMBIA1301 binary vector. The binary construct was introduced into Agrobacterium tumefaciens strain GV3101. Genetic transformation of the bra-MIR1885 gene construct into B. napus was performed as described elsewhere (Bhalla and Singh, 2008). Seedlings exhibiting resistance to hygromycin were transplanted and grown in a greenhouse at 22 °C under a 16-h/8-h light/dark photoperiod. The genomic DNA extracted from T0 plants was used for PCR amplification with specific primers. The seeds (T1) were harvested separately from positive plants for further analyses.

The wild-type and transgenic lines (T2~T3 generations homozygote lines) were grown in the field, and the genomic DNA extracted from plants was used for PCR amplification with specific primers. Each transgenic line was grown in a row of 10 plants, with 30-cm spacing between plants and 40 cm spacing between rows. The phenotypes of transgenic and wild-type plants were measured in the field.

AUTHOR CONTRIBUTIONS

P.X., W.Z., X.Y., and Y.H. conceived the project; P.X., W.Z. and Y.H. designed the experiments; P.X., X.W. and Y.Z. performed the experiments; P.X. and W.Z. analyzed and interpreted the data; P.X., W.L., X.Y., and Y.H. wrote the manuscript.

ACKNOWLEDGEMENTS

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CONFLICT OF INTEREST
The authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT
All the data and materials that support the findings of this study are available upon request from the corresponding author. The sequencing data generated in this study have been deposited in the National Center for Biotechnology Information Sequence Read Archive database under the accession PRJNA748869 and PRJNA745269.

TABLES
Table 1. The miRNAs that were down- or up-regulated by more than 1.5-fold change under cold treatment (4 °C) as compared to normal temperature (22 °C).

| RPM: Reads per million. |

FIGURE LEGENDS
Figure 1. Phenotypes of B. rapa after cold treatment and global view of small RNA reads in sequencing libraries.

(A) Phenotypes of wild-type B. rapa plants grown in soil at 22 °C (left) and 4 °C (right) for 7 days. Red scale bar represents 5 cm. (B) Number of total reads in small RNA library of B. rapa. Shared small RNAs were presented in both libraries. Specific small RNAs were those detected only in normal temperature (22 °C) or low temperature (4 °C) library. (C) Length distribution of unique and raw reads in the two small RNA libraries.

Figure 2. Abundance of differentially expressed miRNAs between cold-stress and normal conditions.

(A) qRT-PCR analysis of seven differentially expressed miRNAs in libraries with normal temperature and cold temperature. (B) Relative abundance of three differentially expressed miRNAs under normal condition and cold condition detected by northern blotting. (C) Relative abundance of miR1885 of B. rapa plants under 22 °C and 4 °C condition for 0, 7, 14, 21 and 28 days, respectively. SAM: shoot apical meristem. “D” represents days. Error bar represents mean ± SE; n = 3. U6 was used as the endogenous reference gene in expression analyses. * indicates P < 0.05; ** indicates P < 0.01; Student’s t-test.

Figure 3. Genome-wide association study of low-temperature resistance in a panel of B. napus accessions.

(A) Manhattan plots and Q-Q plots for genome-wide association study of low temperature resistance phenotype measured in Year 2017 (top) and Year 2018 (bottom), respectively. (B) qRT-PCR analysis of the abundance of three candidate miRNAs under 22 °C and 4 °C condition for 0, 10, 20 and 30 days, respectively. Error bar represents mean ± SE; n = 3. U6 was used as the endogenous reference gene. * indicates P < 0.05; ** indicates P < 0.01; Student’s t-test. (C) Haplotypes of candidate SNPs within MIR1885 promoter for low-temperature resistance in B. napus population. LTR represents low-temperature element (CCGAAA). Five variants, including four SNPs and the one-bp deletion, which were significantly associated with low-temperature resistance, were marked with dark bold vertical lines. The five variants located in the 200-bp genomic region around LTR were used for analysis. ‘Hap.’ denotes the haplotypes of MIR1885; ‘Number’ denotes the number of lines belonging to each haplotype group. ‘Degree’ denotes the average grade of low-temperature resistance.

Figure 4. qRT-PCR analysis of the abundance of miR1885 and its targets under cold stress in B. napus.

(A) qRT-PCR analysis of the abundance of miR1885 in B. napus under 22 °C and 4 °C condition. U6 was used as endogenous reference gene. (B) Alignment of miR1885 mature sequence between B. rapa and
B. napus. Short black line represents the perfect matches. (C) Validation of miR1885 target genes and cleavage sites using 5'-RACE PCR. Predicted TIR, NBS, and LRR domains encoded by target genes are labeled (Left). Reverse complementary matches of miR1885 and their target sites were showed and vertical arrows indicated 5 termini of miRNA-guided cleavage products, as identified by 5'-RACE, with frequency of clones shown (Right). (D) The transcript levels of Bn.TIR.A09 and Bn.TNL.A03 in B. napus under cold treatment. “D” represents days. (E) Relative transcript levels of cold-related genes in wild-type B. napus under 22 degC and 4 degC condition. ACT7 was used as endogenous reference gene. Error bar represents mean +- SE; n = 3. * indicates P < 0.05; ** indicates P < 0.01; Student’s t-test.

Figure 5. Transgenic B. napus plants overexpressing miR1885.

(A) Construct with bra-MiR1885 under control of AA6 promoter. pCaMV35S, CaMV35S promoter; Hyg R, hygromycin-resistance gene; pAA, AA6 promoter; LB, left border of T-DNA; t35S, CaMV35S terminator; tAA6, AA6 terminator. Eco RI, Nco I and Bst EII are restriction sites. (B) Plants of T1 transgenic lines and wild type in the field. Red scale bar = 10 cm. X2–X16, transgenic plants. (C) Detection of transgene in rapeseed transgenic lines by PCR using specific primer spanning AA6 promoter and MiR1885 fragment. M, DNA marker. (D) Real-time PCR showing expression of pri-miR1885 in transgenic lines. ACT7 was used as endogenous reference gene. (E) Relative abundance of mature miR1885 in transgenic lines. U6 was used as endogenous reference gene. (F) Northern blot analysis of accumulation of mature miR1885 in MIR1885 -OE lines. U6 was used as endogenous reference gene. (G-H) Relative transcript levels of Bn.TIR.A09 (G) and Bn.TNL.A03 (H) in miR1885-OE lines as compared to WT. ACT7 was used as endogenous reference gene. Error bar represents mean +- SE; n = 3. * indicates P < 0.05; ** indicates P < 0.01; Student’s t-test.

Figure 6. Relative transcript levels of miR1885 and its targets in transgenic and wild-type plants with or without cold treatment.

(A-C) qRT-PCR analysis of RNA levels of miR1885 (A), Bn.TNL.A03 (B) and Bn.TIR.A09 (C) in transgenic plants after 22degC and 4 degC cold treatment for 30 days. Error bar represents mean +- SD; n = 3. * indicates P < 0.05 and ** indicates P < 0.01 under normal condition; + indicates P < 0.05 and ++ indicates P< 0.01 under 4 degC. Student’s t-test. U6 and ACT7 were used as endogenous reference gene for checking miR1885 and targets expression levels, respectively.

Figure 7. Effects of overexpressing and knocking down of miR1885 in B. napus after cold treatment. (A) Overexpression of miR1885 increased the sensitivity of cold in B. napus. The cold treatment was 2 degC for 7 days. (B) Expression of miR1885 and its targets in miR1885 knock down plants (STTM1885). Error bar represents mean +- SD; n = 3. * indicates P < 0.05; ** indicates P < 0.01; Student’s t-test. U6 and ACT7 were used as endogenous reference gene for measuring miR1885 and targets expression levels, respectively. (C) Knocking down of miR1885 increased the resistance to cold in B. napus as compared to its background. The cold treatment was 2 degC for 14 days. Red scale bar was 5 cm.

Supplementary datasets

Table S1. Distribution of small RNAs among different categories of genomic loci. tRNA: ribosomal RNA; snRNA: small nuclear RNA; snoRNA: small nucleolar RNA; tRNA: transfer RNA; unann: small RNAs without any specific annotation.

Table S2. Differential expression analysis of identified known miRNAs in Brassica rapa by small RNA sequencing. RPM: reads per million.

Table S3. Putative target genes of novel miRNAs in B. rapa. The miRNA target genes were predicted by psRNATarget.

Table S4. Location information of pre-miRNAs in Brassica napus reference genome.

Table S5. List of primers used in this study.

Supplementary Figures
Figure S1. Proportion of small RNAs generated from different categories of genomic loci in the two small RNA libraries of *Brassica rapa*.

Figure S2. Cis-element analysis of *bra-MIR1885* promoter. G-box, GA-motif, P-box, TC, TCT, AAAC, ATCT, Box 4, I-box and LTR represent cis-elements. A06: chrA06. 24219516 and 24221516: Location information. Red or green bar represent position.

Figure S3. Analysis of the single nucleotide polymorphism (SNP) density distribution, principal component analysis (PCA), and linkage disequilibrium (LD) of *B. napus* population. (A) SNP density distribution (number of SNPs in 0.1 Mb sliding windows across each chromosome). (B) PCA analysis. Blue, green, and red points represent winter (W), semi-winter (SW) and spring (S) ecotypes, respectively. (C) Genome-wide average LD decay estimated from *B. napus*. (D) LD heatmap of SNPs in *bna-MIR1885* promoter. Numbers indicate physical position where *bna-MIR1885* promoter locates. Deeper red color indicates stronger linkage relationship. (E) Comparison of the miR1885 relative expression abundance between the two groups with extremely different phenotypes. ES: Low-temperature sensitive accessions; ER: Low-temperature resistant accessions. *U6* was used as an internal control. Data are presented as boxplot. Significant differences were determined by Student’s t-test: * indicates p < 0.05. (F-I) The local LD heatmap of SNPs in *bna-MIR166C*, *bna-MIR168A* and *bna-MIR845A* gene loci.

Figure S4. 5'-RACE assays of *Bn. TIR.A09* (*BnaA09g14980D*) and *Bn. TNL.A03* (*BnaA03g56180D*). PCR products of *BnaA09.TIR* and *BnaA03.TNL* were separated in 2% agarose gel. DNA ladders are labeled on the side. PCR products were cloned into T vectors for Sanger sequencing. Red line represents target fragment; black line represents inner primer.

Figure S5. Genomic synteny analysis between *B. rapa* and *B. napus* species.

Figure S6. Phenotypes of transgenic plants overexpressing miR1885 after cold treatment. (A) Phenotypes of transgenic lines grown at 22 degC or 4 degC. Plants were grown 22 degC for 6 weeks (upper panel), and then transferred to 4 degC for an additional 30 days (middle panel), and then allowed to recover at 22 degC for 7 days. (B) Post-germination phenotypes of miR1885-OE and WT K407 after cold treatment. Seeds were grown on the germination bed at 22 degC for 2 days and then transferred to 6 degC for 7 days. (C) Seeds were grown on the germination bed at 22 degC for 7 days. (D) Quantitative analysis of net hypocotyl lengths from WT K407 and MIR1885-OE lines. n = 10-15 plants. Similar results were obtained in three independent experiments.

Figure S7. Freezing sensitivity of miR1885 over-expressing transgenic plants.

(A) Phenotypes of transgenic lines grown at 22 degC and -10 degC. Plants were grown at 22 degC for 6 weeks (upper panel), and then transferred to -10 degC for 2.5 h (middle panel), and then allowed to recover at 22 degC for 7 days. (B) Real-time PCR showing the expression pattern of miR1885 under freezing treatment. *U6* was used as endogenous reference gene. Error bar represents mean ± SD; n = 3. (C) Survival rates of transgenic lines and wild-type plants after freezing treatment (-10 degC in a growth chamber for 2.5 h, n = 15–20 plants). * indicates P < 0.05; ** indicates P < 0.01 (Student’s t-test). (D) Proline content in transgenic plants and wild type after freezing treatment. Error bar represents mean ± SD. * indicates P < 0.05, significant difference from plants subjected to -10 degC before freezing treatment (0 h). + indicates P < 0.05, significant difference from plants subjected to -10 degC for 2.5 h (Student’s t-test).

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4021.


Table 1. The miRNAs that were down- or up-regulated by more than 1.5-fold change under cold treatment.

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