Integrated transcriptomics and metabolomics analysis to characterize cold stress responses in *Nicotiana tabacum*

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**Abstract**

**Background:** CB-1 and K326 are closely related tobacco cultivars; however, their cold tolerance capacities are different. K326 is much more cold tolerant than CB-1.

**Results:** We studied the transcriptomes and metabolomes of CB-1 and K326 leaf samples treated with cold stress. Totally, we have identified 14,590 differentially expressed genes (DEGs) in CB-1 and 14,605 DEGs in K326; there was also 200 differentially expressed metabolites in CB-1 and 194 in K326. Moreover, there were many overlapping genes (around 50%) that were cold-responsive in both plant cultivars, although there were also many differences in the cold responsive genes between the two cultivars. Importantly, for most of the overlapping cold responsive genes, the extent of the changes in expression were typically much more pronounced in K326 than in CB-1, which may help explain the superior cold tolerance of K326. Similar results were found in the metabolome analysis, particularly with the analysis of primary metabolites, including amino acids, organic acids, and sugars. The large number of specific responsive genes and metabolites highlight the complex regulatory mechanisms associated with cold stress in tobacco. In addition, our work implies that the energy metabolism and hormones may function distinctly between CB-1 and K326.

**Conclusions:** Differences in gene expression and metabolite levels following cold stress treatment seem likely to have contributed to the observed difference in the cold tolerance phenotype of these two tobacco cultivars.

**Keywords:** Transcriptome, Metabolome, Tobacco, Cold stress

**Background**

Low temperature is one of the key environmental stresses that many plants have to cope with during their life cycle [1–3]. It can influence growth and development, as well as the yield, quality, and geographic distribution of crop plants. Cold stress can be classified as either chilling stress (0–15 °C) or freezing stress (<0 °C) [1, 4]. Generally, plants from temperate regions have the ability to cold acclimate, that is, to develop increased freezing tolerance after being exposed to low, nonfreezing temperatures [4]. However, many important crops, such as rice, maize, soybean, cotton, tomato, and tobacco, which originated in the tropics or subtropics, lack a cold acclimation mechanism and are thus sensitive to cold stress [5]. Conventional cross-breeding has used the cold-tolerant varieties as the main source to increase the cold tolerance of less tolerant cultivars [6]. However, the current lack of information about the detailed molecular mechanisms of cold stress biology limits the potential improvement of major crops [5].

Following the rapid progress in DNA microarray and sequencing technologies, whole transcriptome analysis is now being used frequently for transcriptome profiling [7–11]. Many studies investigating plant responses to cold stress have been reported [12–17]. Numerous cold-responsive genes have been identified from different species, including *Arabidopsis thaliana* [2, 18], *Oryza sativa* [19–22], *Zea mays* [23, 24], *Prunus persica* [25, 26], and many other species [13, 27–30]. At present, the best-understood genetics network associated with a role in cold acclimation is the *Arabidopsis* CBF cold response pathway [31]. Exposing *Arabidopsis* plants to low temperature results in rapid induction of a small family...
of genes that encode transcriptional activators known as C-repeat/dehydration responsive element (DREB)-binding factors (e.g. CBF1, −2, −3, −4, −6 or DREB1b, −c) [31]. These genes encode transcription factors that belong to the AP2/ERF domain family of DNA-binding proteins [32]. Furthermore, cold transcriptome analysis has shown that other cold regulatory gene networks also contribute to freezing stress tolerance and cold acclimation [33, 34]. Cold stress signaling integrates with various other signaling pathways to coordinate stress responses with plant growth and development. Cold can affect the expression of genes involved in gibberellin (GA) signaling [35]. Regulation of auxin homeostasis is known to be involved in cold stress responses. Several key components of the cytokinin signaling are also known to play roles in cold signaling. In addition to phytohormones, both the circadian clock and the physiological changes associated with the developmental transition to flowering are known to interact with cold signaling [2]. Despite these insights, many cold acclimation traits still cannot be explained based only on the differential expression change of single or multiple genes. Hence, much remains to be elucidated in the complicated gene network and in its interactions with various biomolecules including metabolites.

In addition to their complex gene regulatory networks, plants have multiple highly-regulated metabolic networks that can play important roles in their growth under cold treatment [34]. It is well established that cold treatment elicits the expression and/or activity of many biochemical pathways. For example, sugars and amino acids are widely accepted as related to cold response, as they are not only vital for the synthesis of functional proteins but also serve as precursors for a large array of metabolites with multiple functions in plant responses to cold treatment [31]. There have been only a limited number of metabolite studies conducted that seek to uncover the potential links between gene and metabolite networks during cold treatment.

Recent technical advances in genome sequencing, gene expression profiling, and metabolomics have made it possible to dissect the complex processes involved in plant responses to cold treatment [7, 8, 19, 33, 36, 37]. Using an integrated gene/metabolite approach, Hirai et al. and Monika et al. were able to identify new genes involved in glucosinolate metabolism in response to sulfur depletion [7, 38], and Tohge et al. (2005) identified novel flavonoid biosynthesis pathways in transgenic Arabidopsis plants overexpressing a MYB transcription factor [8]. Moreover, Kaplan et al. (2007) revealed sequential changes in both gene expression and metabolite profiles throughout the course of the cold acclimation process [37]. Thus, research using multiple-omics tools can potentially lead to the identification of useful genes that could help to improve yields in plants that must cope with cold environments.

Tobacco (Nicotiana tabacum) is native to tropical and subtropical America, but it is now commercially cultivated worldwide. Moreover, it is sensitive to cold [18, 29]. Hence, the molecular mechanism of cold response has been of great concern in the production of this crop. Jin et al. tried to use proteomics to identify the important proteins which may be related to cold response in tobacco [39]. However, owing to the proteomic technique, only 101 proteins were found in their studies. Several works tried to increase the cold tolerance of tobacco using genes from other plants [18, 40]. However, the candidates were still very limited.

Here, our objectives were to explore, on a large scale, the changes that occur in the transcriptome and the metabolome of tobacco in response to low temperature. Moreover, we sought to reveal potential links between cold response genes and metabolites. The experiments that comprised this study included a comparison of the transcriptome and metabolite profiles of two tobacco ecotypes, CB-1 and K326; CB-1 was found to be more sensitive to low temperature than K326. Taken together, our results indicate that cold acclimation is associated with extensive changes in both the transcriptome and metabolome. In general, both the common cold-response genes and the common cold-responsive metabolites between CB-1 and K326 had a greater degree of change in K326 than in CB-1. The specific differentially accumulated transcripts/metabolites and different trends of secondary metabolites in K326 and CB-1 suggest that several cold responsive pathways exist in tobacco. In addition, energy metabolism may be different, and hormones appear to have distinct roles in CB-1 vs. K326.

Methods

Plant materials and cold treatments

The Nicotiana tabacum ecotypes K326 and CB-1 were used in this study. For the RNA-seq experiments, seeds were first germinated and cultivated in plastic pots under typical conditions until the plants grew to the six-leaf stage. Control plants were then grown in a growth chamber under normal conditions (28 °C with 16 h light and 24 °C with 8 h dark). For cold treatments, plants were transferred to 4 °C for one day with afore mentioned photoperiod; the one day treatment duration was chosen because the cold marker gene ICE1 was found in a pilot study to reach a maximum of expression after one day of such treatment (Additional file 1: Figure S1). Samples were harvested one day after the start of cold treatment (light period). The top two leaves of each plant were sampled. All leaf samples were immediately frozen in liquid nitrogen and stored at −80 °C. Each
sample consisted of leaves from six plants grown in the same condition.

RNA isolation and library preparation for transcriptomics analysis
Total RNA was isolated using TRIzol reagent (Invitrogen) according to the manufacturer’s instructions. Poly (A) + mRNA was purified with oligo (dT) beads, and then the mRNA was randomly segmented into small fragments by divalent cations in fragmentation buffer (Illumina) at 94 °C for 5 min. These short fragments were used as templates to synthesize first-strand cDNA using random hexamer primers. Second-strand cDNA was synthesized using RNaseH and DNA polymerase I. Short cDNA fragments were purified with a QiaQuick PCR extraction kit (Qiagen). The cDNA fragments were then connected with sequencing adapters according to an Illumina protocol. After agarose gel electrophoresis, the target fragments of 300–500 bp were selected for PCR amplification to create the final cDNA library.

Sequencing, assembly, and functional annotation
The RNA libraries were prepared using the TruSeq RNA Sample Preparation Kit v2, set A (RS-122-2001, Illumina) according to the product instructions [9–11]. All experiments were repeated twice. The quality and size of the cDNA libraries for sequencing were checked using the Agilent 2200 TapeStation system (Agilent). The libraries were run on single lanes for 100 cycles (paired-end) on HiseqTM 2000 (illumine Inc.), individually. Raw reads were analyzed by FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) for quality, and high-quality reads with Q > 20 were obtained using NGSToolkits (version: 2.3.3) [41]. The Trinity method was used for de novo assembly of the high-quality reads to generate unigenes [42]. Functions of the unigenes were annotated based on sequence similarities to sequences in the public UniProt database [43]. T-test was used for the statistics analysis.

DNA preparation and qRT-PCR
For cDNA preparation, approximately 1 μg of total RNA from each sample was used to synthesize first-strand cDNA in a 20 μl reaction solution using a Transcriptor First-Strand cDNA Synthesis Kit (Roche). The transcript levels for transcripts related to cold response were detected using quantitative real-time PCR (qRT-PCR) [9, 10]. The qRT-PCR amplification reactions were performed using a LightCycler ® 96 Real-Time PCR System (Roche). The reactions were conducted in a final volume of 20 μl containing 10 μl of SYBR Green premix (2X) (Roche). The PCR thermocycling program consisted of an initial incubation at 95 °C for 10 min, followed by 40 cycles of 95 °C for 10s, 58 °C for 20s, and 72 °C for 20s, and an additional final cycle for analysis of dissociation curves to ensure specific amplification. Relative quantification of the transcription level for each target gene was normalized to the signal for transcripts of the constitutively expressed tobacco 26S gene. T-test was used for statistic significant analysis. The gene specific primers used for qRT-PCR are listed in Additional file 1: Table S1.

GC-MS sample preparation
The freeze-dried tissue were ground to a uniform powder, filter using a 40-mesh sieve, and stored at -80 °C until the metabolic study. The tissue powder (20 mg) was added to a 2 ml Eppendorf tube and soaked in 1.5 ml of an extraction solvent containing isopropanol/acetonitrile/water(3/3/2,v/v/v) with 15 μlB. (2 mg/mL) of trideca-nonic acid as an internal standard. All extracts were sonicated for 1 h and centrifuged for 10 min (1,4000 rpm, 4 °C). Then the supernatant (1.3 mL) was removed to a new 1.5 ml centrifuge tube. After centrifugation at 1,5000 rpm for 10 min, 4 °C, 350 μL of supernatant was collected to a conical insert of a 2 mL glass vial and dried under nitrogen flow on an N-EVAP Nitrogen Evaporator. The silylation reaction for increasing the volatility of the metabolites was performed by adding 100 μL of methyl-trimethyl-silyl-trifluoroacetamide(MSTFA) to the sample and incubating it for 60 min at 60 °C.

GC/MS data analysis
Metabolomic analysis was performed on an Agilent 7683B series injector (Agilent, Santa Clara, CA) coupled to an Agilent 6890 N series gas chromatograph system and a 5975 mass selective detector (MSD) (Agilent, Santa Clara, CA). Agilent DB-5MS column (0.25 μm, 0.25 mm × 30 m, Agilent Technologies, Inc., Santa Clara, CA) was used. The column temperature was 70 °C for the first 4 min and then increased at 5 °C/min to 310 °C for 15 min. The injection temperature was set as 300 °C, and the injection volume was 1 μl with a 10:1 split ratio. Helium (99.9995%, China) was applied as a carrier gas. The column flow was 1.2 mL/min, and the column was equipped with a linear velocity control model. Prior to the instrumental analysis, the mass spectrometer was tuned using perfluorotributylamine (PFTBA) to obtain optimum performance. A simultaneous full scan-selected ion monitoring mode (Scan-SIM) was used to acquire the data. The mass spectra scanning scope was set to 33–600 m/z in the full scan mode and 90 chromatographic peaks of 25 groups were set in the selected ion monitoring. The scan speed is 2.59 scan s-1 and the solvent cut time is 5.0 min. The temperatures of the interface and the ion source were adjusted to 280 and 230 °C, respectively. The detector voltage was maintained at 1.2 kV, and the electron impact (EI) model was selected to achieve ionization of the metabolites at 70 eV.
samples were nearly dead, whereas around 2–3 survived the cold treatment. Totally, 10 out of 12 CB-1 more cold-tolerant than CB-1, for which more plants phenotype shown in Fig. 1a, it is clear that K326 is much checked after one day of cold treatment. From the cold responses. The phenotype of CB-1 and K326 was the leaf tissues of CB-1 and K326 were chosen to study Phenotypic and physiological responses to cold stress

Results
Phenotypic and physiological responses to cold stress
The leaf tissues of CB-1 and K326 were chosen to study cold responses. The phenotype of CB-1 and K326 was checked after one day of cold treatment. From the phenotype shown in Fig. 1a, it is clear that K326 is much more cold-tolerant than CB-1, for which more plants survived the cold treatment. Totally, 10 out of 12 CB-1 samples were nearly dead, whereas around 2–3 out of 12 K326 samples were nearly dead (Fig. 1).

Gene network construction and visualization
The interaction between genes was retrieved from several public protein-protein interaction (PPI) databases, including BioGrid [44], IntAct [45], BOND [46], DIP [47], MINT [48], HPRD [49], MIPS [50], and TAIR [51]. The interaction between genes and metabolites was calculated as Pearson-correlation coefficients, with a threshold value of 0.9 for inclusion in the network. Cytoscape [52] was used to visualize the final interaction network.

LC-MS sample preparation
The freeze-dried leaf tissues were ground to a uniform powder, filtered with a 40-mesh sieve, and stored at -80 °C. 2 mg tissue powder was added to a 2 mL Eppendorf tube to which was added 1.5 mL of an extraction solvent containing a 75% methanol with 10 μg/ ml Umbelliferon as an internal standard. All extracts were sonicated for an hour and were centrifuged for 10 min (14,000 rpm, 4 °C). 1 ml of supernatant was collected for LC – MS analysis.

LC-MS data analysis
LC – MS analysis was performed on an Agilent 1290 lipid chromatograph system and a 6540 6540 UHD Q-TOF equipped with an electrospray ionization source (Agilent, Santa Clara, CA). Aglient SB-C18 column (RRHD,1.8 μm, 2.1 × 100 mm, Agilent Technologies, Inc., Santa Clara, CA) was used. The samples were analyzed in positive ion mode. The gradient was as follows: 2 min, 30% B; 10 min, 85% B; 15 min, 90% B; 17 min, 100% B; 20 min, 100% B. Mobile phases A and B were 95% water: 5% acetonitrile with 0.1% formic acid and 5% water: 95% acetonitrile with 0.1% formic acid, respectively. The flow rate was 0.3 mL/min. 5 μL aliquots of sample were loaded for every individual analysis. The capillary voltage and spray shield were set to 3500 V. The sheath gas was set to 10 L/min at a temperature of 350 °C. The nebul gas was set to 12 L/min at 350 °C. Spectra were acquired over the m/z 50–1200 range. The collision energy in the MS/MS mode was set to 20 V.

Transcriptome analysis for CB-1 and K326
We used RNA-seq to enable comparison of the transcriptomes of CB-1 and K326 in response to cold treatment. Around 100 million high quality reads of 90 base pairs (bp) were generated for each library. Table 1 shows an overview of the RNA-seq reads for the eight libraries; a total of 664,432,026 high-quality reads were obtained from all libraries. Using the Trinity method with default parameters [42], these high-quality reads were finally assembled into 135,148 unigenes with 230,735 isoforms. All unigenes were longer than 200 bp. The N50 of the final assembled transcripts was 1479 bp. For an overview of the assembly results refer to Additional file 2: Data Set 1.1. The unigenes were annotated by performing BLASTX searching against the UniProt database [43]. Among the 135,148 non-redundant unigenes, 23,246 had at least one hit via BLASTX [53] searching with an E-value <= 1e-3 and at least 30% sequence identity, highlighting that there are still many uncharacterized genes or tobacco specific genes. In order to calculate the expression level for the assembled transcripts, we first mapped reads onto assembled unigenes using bowtie [54]. RSEM (RNA-seq by Expectation-Maximization) [55] was used to estimate the abundance of assembled transcripts and to calculate the expression level.

To verify the RNA-seq data, 28 genes were chosen for qPCR validation, including DREBs, CORs, and SIZ1, which have already been reported to be related to cold responses. The results of the RNA-seq and qPCR analyses were similar (Fig. 2), showing the same general expression trends (more than 90% correlation), such as DREB (c79563_g1).

Identification of differentially expressed genes under cold stress
Differentially expressed genes (DEGs) among samples were defined using fold change values by the expression of assembled transcripts. We defined DEGs with |log2(−fold change) > 1| and corrected p-value < 0.05. In total, we identified 7447 up-regulated DEGs between the control and cold-treated CB-1 samples, with 7143 down-regulated DEGs. For K326, there were 6398 up-regulated DEGs, with 8207 down-regulated DEGs (Fig. 1b, c). Gene expression profiles in CB-1 and K326 changed under cold stress to differing degrees; however, there were some overlapping DEGs that were regulated by cold stress in both tobacco cultivars, with 41.3% for up-regulated DEGs and 50.7% for down-regulated DEGs. To compare the transcriptomes in CB-1 and K326 under cold treatment, a heat map was generated to present the transcript abundance for all DEGs under cold stress (Fig. 1d). The results also showed that a series of changes in gene expression in CB-1 and K326 when plants subjected to cold treatment.
Cold response genes common between CB-1 and K326

Following cold treatments, 4046 cold-induced transcripts and 5167 cold-repressed transcripts were identified in both CB-1 and K326 (Fig. 1b, c). We analyzed these groups, trying to investigate the mechanisms that function in responses to cold stress. Functional classification of DEGs was achieved using a gene ontology (GO) analysis via InterproScan [56]. These DEGs were assigned to three classes of GO: biological processes, molecular functions, and cellular compartments. Additional file 1: Figure S2 illustrates the distinct distribution of the main GO categories. The GO terms “response to stimulus”, “signaling” and “metabolic process” are the most highly enriched among both the up-regulated and down-regulated gene sets, which is in agreement with previous findings [17, 57, 58]. GO terms also represented diverse functional activities corresponding with the mentioned biological processes. The highest numbers of DEGs were categorized in “catalytic and binding activities”, which have also been reported as over-

Table 1 Summary of sequencing data quality

| Sample | reads | HQ reads | Total bps | Total bps in HQ reads | Q20 (%) | N (%) | GC (%) |
|--------|-------|----------|-----------|-----------------------|---------|-------|--------|
| CT1    | 55,643,444 | 52,340,970 | 5,619,987,844 | 5,286,437,970 | 98.52   | 0.89  | 42.54  |
| CT2    | 117,283,874 | 110,687,372 | 11,845,671,274 | 11,179,424,572 | 98.48   | 1.02  | 42.86  |
| CC1    | 90,303,872 | 85,210,840 | 9,120,691,072 | 8,606,294,840 | 98.53   | 0.89  | 43.81  |
| CC2    | 102,439,604 | 96,792,568 | 10,346,400,004 | 9,776,049,368 | 98.53   | 0.89  | 43.11  |
| KT1    | 81,708,490 | 77,877,970 | 8,252,557,490 | 7,865,674,970 | 98.87   | 0.89  | 42.46  |
| KT2    | 85,728,846 | 81,281,390 | 8,658,613,446 | 8,209,420,390 | 98.78   | 0.85  | 42.59  |
| KC1    | 61,961,226 | 58,117,664 | 6,258,083,826 | 5,869,884,064 | 98.73   | 0.85  | 42.53  |
| KC2    | 107,703,362 | 102,123,252 | 10,780,039,562 | 10,314,448,452 | 98.72   | 0.87  | 43.10  |
| total  | 702,772,718 | 664,432,026 | 70,980,044,518 | 67,107,634,626 | 98.65   | 0.89  | 42.88  |

CC refers to the control group of the CB-1 sample; CT refers to the cold treatment group of the CB-1 samples; KC refers to the control group of the K326 samples; KT refers to the cold treatment group of the K326 samples; 1 and 2 refers to the two replicates
represented terms under treatment with a variety of stresses [16, 33]. In contrast with previous studies, which highlighted the role of ABA-related events under cold stress [1], no such events were detected in our study. Interestingly, the GO terms “immune system process”, “negative regulation of biological process”, and “growth” were enriched in the GO list of cold down-regulated genes.

KEGG (Kyoto Encyclopedia of Genes and Genomes) pathway analysis provides classifications that are valuable for studying the complex biological functions of genes [17]. KEGG analysis showed that “Ribosome”, “Plant hormone signal transduction”, and “Estrogen signaling pathway” were significantly enriched by DEGs under cold treatment (Fig. 3, Additional file 2: Dataset 1.2). “MAPK signaling pathway” and “cAMP signaling pathway” were also enriched. Previously, it was reported that different protein kinase families (such as MAPKs) are activated by osmotic stresses [1, 2, 33]. These protein kinases may be important components in the signal transduction pathway of various environment signals in plants under cold stress.

Furthermore, by checking the detailed expression levels for these DEGs, many genes were induced to much higher in K326 than CB-1, especially for known cold related genes such as the DREB genes (Additional file 1: Table S2, Fig. 1d). This may be one of the major reasons that K326 is much more cold-tolerant than CB-1 as indicated by our gross phenotypic analysis.

** Cultivar specific differentially expressed genes (DEGs) **

In total, 5377 DEGs were only identified in CB-1 in the cold treatment, with 3401 up-regulated and 1976 down-regulated (Fig. 1b, c). And 5392 DEGs were identified in K326, with 2352 up-regulated and 3040 down-regulated (Fig. 1b, c). Similar with common DEGs between CB-1 and K326, KEGG enrichment analysis for the K326-specific DEGs revealed that “Ribosome”, “Plant hormone signal transduction”, and “MAPK/Estrogen/FoxO signaling pathway” were highly enriched, which suggested the ability for cold tolerance of K326 was strengthened (Fig. 4b). In addition, K326-specific DEGs were enriched in “2-Oxocarboxylic metabolism”, “Caffeine metabolism”, “Carbon metabolism” in K326, which CB-1-specific DEGs were enriched in “Alanine, aspartate and glutamate metabolism”, “Butanoate metabolism”, and “Methane metabolism” in CB-1. (Fig. 4a, b, Additional file 1: Figure S3).

** Metabolome reprogramming under cold treatment in CB-1 and K326 **

In an attempt to generate a comprehensive picture of metabolite reprogramming that occurs in response to cold treatment, we performed metabolite profiling of CB-1 and K326 using GC-MS and LC-MS. We reproducibly detected a total of 198 primary metabolites and 4269 secondary metabolites, with 4045 un-target metabolites and 224 target metabolites (Additional file 2: Dataset S1.3). Among these identified primary metabolites and secondary
Fig. 3 Top 30 enriched pathways for overlapping cold responsive genes between K326 and CB-1.

Fig. 4 Top 30 enriched pathways for specific cold responsive genes of CB-1 and K326. a Top 30 enriched pathways for CB-1 specific cold responsive genes. b Top 30 enriched pathways for K326 specific cold responsive genes.
metabolites, 200 metabolites showed differential accumulation (Fold change >1.5) in CB-1, with 161 increased and 39 decreased under cold treatment compared with the control (Fig. 5a, b). In contrast, 194 differentially accumulated metabolites were identified in K326, with 75 increased and 119 decreased (Fig. 5a, b).

Of the metabolites with differential accumulation during cold treatment, several of them had previously been shown to increase in Arabidopsis plants upon exposure to low temperature [31], including several amino acids, which was found in our data (Fig. 5c, d). However, these amino acids increased to a much greater extent in K326 than in CB-1; several of them, including glycine, phenylalanine, and alanine, even decreased in CB-1 (Additional file 2: Dataset S1.3). Similar results were also found for sugars (glucose, fructose, inositol, galactinol, raffinose, and sucrose); and trehalose, putrescine, and ascorbate (Additional file 2: Dataset S1.3) [31, 37]. As in the transcriptome data, these changes may help to explain how it is that K326 is much more cold-tolerant than CB-1.

Plant hormone signals appear to have different influences on CB-1 and K326. Unfortunately, ABA was not detected in our study. However, ethylene and salicylic acid (SA) were among the differentially-accumulated metabolites. Ethylene is negatively correlated with the cold tolerance of both CB-1 and K326 (Fig. 6a), which has already been reported in previous studies [35]. In contrast, different from previous reports, SA was in our results positively correlated with CB-1, whereas it was negatively correlated with K326 (Fig. 6b).

Another interesting finding is that we observed a lot of differential accumulation of secondary metabolites. Besides previously-reported metabolites such as flavonoids and lipid related metabolites, a fragrance (benzene) and nicotine (nicotine, Nicotinic acid) were also found to be regulated by cold treatment in our study. However, most of the identified secondary metabolites were up-regulated by cold stress in CB-1, while K326 showed the opposite trend (Fig. 6c, Additional file 1: Figure S4). More work will need to be done to investigate the reasons for these differences.

In a PCA model based on differentially accumulated genes and metabolites, the first two principal components clearly separated all samples, both control and cold treatment for both CB-1 and K326, and explained around 80% of the total variation in the entire data set (Fig. 7). Specifically, PC1 of transcriptome and metabolism analysis may reflect cold-responsive genes and metabolites, most of which are shared by CB-1 and K326, such as amino acids and sugars. However, PC2 may largely reflect the reprogrammed parts between CB-1 and K326, especially on secondary metabolites (Fig. 7).

**Interaction network analysis between cold regulated genes and metabolites**

Gene-metabolite interaction networks can be used to help understand functional relationship and to aid in identifying new regulatory elements [33]. Here, we constructed an interaction network based on the differentially expressed genes (Fig. 1b, c) and metabolites (Fig. 5a, b). Gene-gene
interactions were retrieved from known PPI databases. All genes harboring potential interaction relationships are shown in Additional file 2: Dataset S1.4 (Additional file 1: Figure S5).

From the clusters shown in Additional file 1: Figure S5, it is apparent that most of the secondary metabolites closely connect together, including flavonoid families. In order to check the interaction relationships, we also showed the detailed networks for sugar metabolites (Additional file 1: Figure S6A) and target secondary metabolites (Additional file 1: Figure S6B). Furthermore, the sugar-related network can be divided into two clusters, one with sucrose, trehalose, tagatose, and another with cellobiose, fructose, and raffinose. The interaction relationships in these two sugar clusters suggest not only that the CBF/DREB and ZAT families play some role with the biosynthesis of sugar metabolites, but also suggest that TFs, including those of the ARR, GT, UP-BEAT, VIP, and MYB families, may also contribute to sugar biosynthesis during plant responses to cold treatment. These potential new markers for cold stress could be useful for cold resistance studies in the future.

Analysis of cold-responsive transcription factors
Transcription factors (TF) play a key role in the regulation of gene expression under abiotic and biotic stresses in plants. RNA-seq results showed that many TFs were regulated under cold stress in CB-1 and K326 (Additional file 2: Dataset S1.5, Table 2). In total, more than 20 families of TFs were differentially expressed, implying their important roles in regulating genes involved in cold responses.

In particular, families showing especially strong responses to cold treatment included WRKY, AP2-EREBP, NAC, MYB, HSF, bHLH, UPB, and bZIP (Additional file 2: Dataset S1.5, Table 2). For instance, some genes from these families were up-regulated by more than 30 times during cold treatment. For example, most of our predicted CBF
TFs presented dramatic activation (especially c64765_g1 with nearly 60 times up-regulation); this gene is known to play a direct role in activating the expression of downstream cold-responsive genes, except for c71296_g1. Another is a famous known cold regulator ICE1, which encodes a MYC-type basic helix-loop-helix (bHLH) TF [2, 3] that directly binds to MYC cis-elements in the promoter of CBF/DREBs. The RNA-seq results showed that the transcript level of only one MYC-type bHLH (c71107_g1) increased under cold stresses, suggesting that DREBs may act as its target.

Discussion
In plants subjected to cold treatment, to enhance adaptability to low temperature, more sugars and other energy carriers (e.g., ATP) are consumed to produce lipids, amino acid, membrane components, and other molecules to further promote cell membrane fluidity and structural rearrangement [19, 33]. In our study, a lot of sugar components showed up-regulation under cold treatment in both CB-1 and K326 (Figs. 8 and 9a), including glucose, fructose, inositol, galactinol, and raffinose, but excepting trehalose and sucrose in CB-1. Meanwhile, some sugars that have not been widely-reported to be related to cold responses were also detected in our study, such as tagatose, mannose, and cellobiose. It seems evident that the sugar metabolism is much more active in K326 than in CB-1 in response to cold treatment (Figs. 8 and 9a). However, for lipids, another alternative of energy metabolism, especially linolenic acid (18:3), was down-regulated in K326 but up-regulated in CB-1 (Fig. 9b). CB-1 and K326 have no significant differences in the accumulation of some amino acids including valine, leucine, and tyrosine, whereas serine and L-aspartate were accumulated to much higher extents in K326 (Fig. 8, Additional file 2: Dataset 1.6). Together, these results suggest that CB-1 and K326 may employ different aspects of energy metabolism as they respond to cold stress.

Hormones are signaling molecules that play key roles in regulating gene expression under cold stress [27, 35, 59]. RNA-seq analysis revealed that many genes related to abscisic acid, ethylene, auxin, salicylic acid, and gibberellin were regulated by cold stress in CB-1 and K326 (Additional file 2: Dataset 1.7). The KEGG pathway, "plant hormone signal transduction", was significantly enriched among the differentially expressed genes in both CB-1 and K326.

Alternations of ethylene levels by cold stress treatment have been observed in many plants [35]. However, the role of ethylene in cold tolerance varies in different plant species, and can even vary in Arabidopsis plants that are grown in different environments [35]. For example, the cold tolerance of soil-grown Arabidopsis seedlings treated with 1-aminocyclopropane-1-carboxylated (ACC) is enhanced [60] but is reduced in vitro; cold tolerance
is increased when aminoethoxyvinylcyclohexane (AVG) is applied [61]. Ethylene positively affects the cold tolerance of tomato (Lycopersicon esculentum) [62]. In contrast, ethylene levels are negatively correlated with the cold tolerance of Medicago truncatula [27]. Previous studies have shown that ethylene also negatively influences the cold tolerance of tobacco (Nicotiana tabacum) [40], which was confirmed by our finding that ethylene levels are negatively correlated with the cold tolerance in both CB-1 and K326, with a greater magnitude of down-regulation in K326. Therefore, the role of ethylene on cold tolerance is possibly species dependent. ERF TFs, which are located downstream of the ethylene signal pathway, play important roles in cold responses via recognizing dehydration-responsive elements (DRE) [40, 61, 62]. This family can be further divided into ERF (Ethylene-responsive transcription factor ERF), CRF (Ethylene-responsive transcription factor CRF), ABR (Ethylene-responsive transcription factor ABR), RAP (Ethylene-responsive transcription factor RAP) and EIN (ETHYLENE INSENSITIVE protein) (Additional file 1: Table S3, Additional file 2: Dataset S1.7). From the expression pattern for ERFs in our study, we found that more than half of them were up-regulated under cold treatment, with 44 in K326 and 27 in CB-1 (Additional file 1: Table S3). The most highly responsive ERFs are members of ABR (c74525_g2: around 40 times up-regulation) and CBF (c64865_g1: around 30 times up-regulation). These results suggest that ERFs may be involved in plant response to cold stress as regulated by ethylene levels.

Different from ethylene, salicylic acid (SA) shows a different pattern in CB-1 vs. K326, with up-regulation in CB-1 and down-regulation in K326. Previous studies showed that cold temperatures promote the accumulation of endogenous free SA and glucosyl SA in Arabidopsis shoots, wheat, and grape berry [28, 59, 63, 64]. Additionally, it has been reported that high concentrations and the continual application of SA cause severe growth damage and a decrease in cold tolerance capacity [59, 65–67]. Plants from seeds imbibed in a high concentration of SA (1 mM) did not show any alternation of cold tolerance, whereas low concentrations of SA (0.1–0.5 mM) promoted tolerance to cold stress in bean and tomato [68]. In summary, these data suggest that the sporadic application of SA may enhance the cold tolerance but that continual application may decrease this tolerance. Hence, the increasing of SA in CB-1 may be another reason for its relatively weaker cold tolerance. SA is synthesized via two distinct pathways, the isochorismate (IC) pathway and the phenylalanine ammonia-lyase (PAL) pathway [67]. Although we do not yet have direct measurements of the level of chorismate, many secondary metabolites were up-regulated in CB-1, which may be an upstream indication relating to the high level of SA in CB-1. These results suggest that the effect of SA on cold tolerance may be dependent on the organism, concentration, and period of application.

### Conclusions

CB-1 and K326 are two closely related tobacco cultivars, but their tolerance to cold is quite different. In summary, the large-scale-"omics" technologies utilized here to investigate the overall transcriptome and metabolome changes associated with cold treatment are surprisingly informative in uncovering novel regulatory elements and

### Table 2

| gene_id | CC  | CT  | KC  | KT  | FC(K326) | name   |
|---------|-----|-----|-----|-----|----------|--------|
| c78380_g1 | 0.733 | 6.628 | 1.314 | 7.238 | 5.895 | WRKY33 |
| c64765_g1 | 0.000 | 2.709 | 0.000 | 5.693 | 2.709 | DREB1F |
| c74525_g2 | 0.000 | 5.085 | 0.344 | 5.873 | 5.085 | ABR1   |
| c40491_g1 | 0.293 | 2.973 | 0.035 | 5.456 | 2.973 | HBP-1b |
| c77368_g4 | 0.000 | 4.180 | 0.000 | 5.395 | 4.180 | HSFA4B |
| c73830_g2 | 1.097 | 5.904 | 1.211 | 6.379 | 4.807 | WRKY33 |
| c64865_g1 | 0.000 | 4.614 | 0.000 | 5.127 | 4.614 | CRF6   |
| c79891_g2 | 0.743 | 2.194 | 0.496 | 5.479 | 1.452 | ERF109 |
| c81375_g1 | 0.567 | 4.541 | 0.358 | 5.193 | 3.975 | ERF053 |
| c74235_g2 | 0.195 | 3.704 | 0.345 | 4.962 | 3.509 | WRKY40 |
| c83950_g2 | 0.180 | 3.968 | 0.397 | 4.863 | 3.788 | WRKY6  |
| c73886_g1 | 0.000 | 2.621 | 0.000 | 4.253 | 2.621 | DREB2A |
| c72670_g1 | 1.984 | 5.685 | 2.190 | 6.424 | 3.701 | WRKY11 |
| c71056_g2 | 0.000 | 2.704 | 0.000 | 4.217 | 2.704 | DREB2H |
| c52393_g1 | 0.588 | 4.436 | 0.000 | 4.165 | 3.847 | DREB2H |
| c69854_g3 | 2.768 | 6.470 | 2.503 | 6.573 | 3.702 | ERF053 |
| c4821_g1   | 0.288 | 3.422 | 0.000 | 3.913 | 3.422 | MYB306 |
| c81178_g1  | 0.000 | 2.706 | 0.000 | 4.390 | 2.706 | ABR1   |
| c83320_g1  | 0.470 | 3.990 | 0.473 | 4.347 | 3.519 | CRF6   |
| c112787_g1 | 0.288 | 4.104 | 0.301 | 5.354 | 4.104 | ERF053  |
| c58192_g1  | 0.000 | 3.259 | 0.000 | 3.814 | 3.259 | HSFA6b |
| c73618_g1  | 0.000 | 2.962 | 0.000 | 3.714 | 2.962 | GT-3A  |
| c72172_g1  | 1.011 | 4.721 | 1.536 | 4.261 | 3.710 | HSF1C  |
| c69854_g2  | 1.541 | 5.161 | 1.664 | 5.352 | 3.621 | NAC002 |
| c81178_g3  | 0.000 | 2.725 | 0.000 | 3.649 | 2.725 | NAC002 |
| c68026_g1  | 3.930 | 4.090 | 0.792 | 3.613 | 4.090 | TCP19  |
| c69854_g1  | 1.792 | 4.978 | 2.347 | 5.835 | 3.187 | CRF6   |
| c67824_g2  | 1.961 | 4.702 | 1.968 | 5.439 | 2.741 | WRKY40 |
| c76976_g2  | 0.000 | 1.908 | 0.139 | 3.517 | 1.908 | ERF109 |
| c82962_g2  | 4.150 | 4.153 | 4.123 | 0.783 | 4.153 | MYB39  |

The value is Log2 RESM value for each assembled TF. FC is the Log2 fold change in the treatment sample as compared to wild type
Fig. 8 The energy metabolism through sugar and amino acids metabolic pathways under cold treatment. *Red color* represents the fold change between wild type and cold stress in CB-1. *Blue color* represents the fold change between wild type and cold stress in K326.

Fig. 9 The levels of sugar metabolites and fatty acid metabolites. Y-axis represents the relative quantitate level of metabolites. **a** The relative metabolite levels for sugars. **b** The relative metabolite levels for fatty acids.
genes. Consequently, we found that transcriptional regulators/TFs, which are persistently changed during cold treatment, can be responsible for maintaining the cold acclimated status via complex regulatory networks. Moreover, both the transcriptomes and metabolomes indicated the higher responsive ability on common genes/metabolites might be one of the major reasons to explain the cold tolerance for K326. In addition, there may exist other cold regulatory pathways in K326 that are inactive or not present in CB-1. The results of our study contribute to a deeper understanding of the highly complex regulatory mechanisms that function in plants during cold treatment.

Additional files

Additional file 1: Figure S1. The expression levels for cold responsive genes ICE1 under cold treatment. 1: 2 h control; 2: 2 h treatment; 3: 6 h control; 4: 6 h treatment; 5: 12 h control; 6: 12 h treatment; 7: 1 d control; 8: 1 d treatment; 9: 2 d control; 10: 2 d treatment; 11: 3 d control; 12: 3 d treatment; 13: 4 d control; 14: 4 d treatment. Figure S2. Gene classification based on gene ontology (GO) for differentially expressed genes by cold up-regulated (induced) gene and cold down-regulated (repressed) gene. Blue color represents enriched GO terms by cold induced genes; Red color represents enriched GO terms by cold repressed genes. Figure S3. Gene classification based on gene ontology (GO) for differentially expressed genes (DEGs) of CB-1 and K326. Blue color represents the enriched GO terms based on DEGs of CB-1. Red color represents the enriched GO terms based on DEGs of K326. Figure S4. Boxplot for levels of secondary metabolites in CC, CT, KC and KT. Figure S5. Visualization of the final gene-metabolite network. Blue color represents genes; Red color represents secondary metabolites; cyan color represents primary metabolites. Figure S6. Visualization of interaction networks of sugar metabolites and identified secondary metabolites. (A) The interaction networks of identified secondary metabolites and associated genes. (B) The interaction networks of identified secondary metabolites and associated genes. Table S1. Primer sequences used for qRT-PCR analysis. Table S2. The detailed expression levels for known cold responsive genes and DREB family genes. Table S3. The number of up-regulated, down-regulated and unchanged genes for ethylene biosynthesis family. (PP TX 788 kb) Additional file 2: Dataset S1. RNA-seq assembly results and expression analysis from various tissues. (XLSX 38411 kb)

Abbreviations

ABA: Abscisic acid; CBF: CCAAT-binding transcription factor; CRT: C-repeat; DEG: Differentially expressed gene; DREB: Dehydration responsive element; EkHa: Ethylene insensitive protein; ERF: Ethylene responsive transcription factor; ET: Ethylene; GA: Gibberellin; GOG: Gene ontology; IC: Isochorismate; ICE1: Inducer of CBF Expression 1; KEGG: Kyoto Encyclopedia of Genes and Genomes; PAL: Phenylalanine ammonia lyase; SA: Salicylic acid

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Availability of data and materials

The raw RNA-seq data supporting the result of this article is NCBI, with accession numbers GSE94105.

Authors’ contributions

JJ and PJC conceived and designed the experiments. HZ and JFZ prepared plant materials and carried out qPCR analysis. JJ, ZFL, YLX, and PL performed the RNA sequencing and bioinformatics data processing. PP and XC did the GC-MS and LC-MS experiments. JJ, HZ, JFZ, and PJC wrote the manuscript and all authors read and approved the final version.

Competing interests

The authors declare that they have no competing interests.

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