PuMYB21/PuMYB54 coordinate to activate PuPLDβ1 transcription during peel browning of cold-stored “Nanguo” pears

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Abstract
Refrigeration is commonly used to extend the storage life of “Nanguo” pears, but fruit in long-term refrigeration is prone to peel browning, which is related to membrane lipid degradation. To determine the mechanism of membrane lipid degradation, we identified two R2R3-MYB transcription factors (TFs), PuMYB21 and PuMYB54, from “Nanguo” pears, which were notably expressed in response to cold stress and during the peel-browning process. The results from yeast one-hybrid, electrophoretic mobility shift, and transient expression assays indicated that both PuMYB21 and PuMYB54 directly bind to the promoter of PuPLDβ1 (a key enzyme catalyzing the hydrolysis of membrane phospholipids) and activate its expression, which probably enhances the degradation of membrane phospholipids and eventually results in peel browning. Moreover, the overexpression of PuMYB21 and PuMYB54 can greatly activate the transcription of endogenous PuPLDβ1 in both “Nanguo” pear fruits and calli, and their silencing can inhibit its transcription. Furthermore, yeast two-hybrid, bimolecular fluorescence complementation, and pull-down assays verified that PuMYB21 interacts with PuMYB54 to enhance the expression of PuPLDβ1. In summary, we demonstrate that PuMYB21 and PuMYB54 may have roles in membrane lipid metabolism by directly binding to the downstream structural gene PuPLDβ1 during the low temperature-induced peel browning of “Nanguo” pears.

Introduction
Low temperature storage is routinely used to extend the postharvest life of fruit; however, various chilling injury (CI) symptoms commonly occur in fruits subjected to inappropriately low temperatures or long-term cold storage. “Nanguo” pears (Pyrus ussuriensis Maxim.) are specialty fruits in the Liaoning Province of China and are mainly produced in the Anshan region. “Nanguo” pears are typically harvested in the middle of September. Although the fruits can be stored for only ~20 d at room temperature, cold storage technology can extend their shelf life to several months. However, peel browning is one of the most typical CI symptoms in long-term low temperature-stored “Nanguo” pears, which seriously affects the quality and commercial value of the fruit. Thus, the exploration into the mechanism of peel browning in refrigerated “Nanguo” pears is of great significance.

The cytomembrane, a critical structure in cells, is important in providing a stable internal environment for the life activities of cells, nutrient exchange, and cell recognition. Furthermore, the cytomembrane is critical for cellular and subcellular compartmentalization, preventing contact between substrates and enzymes in the process of enzymatic browning. In addition, membrane lipids have significant roles in the adaptation of plants to cold stress. Phospholipids are fundamental molecules for establishing biological membranes and are highly conserved in bacteria, mammals, and seed plants. The degradation of membrane lipids can destroy the structural integrity of the cytomembrane and lead to the disruption of cellular compartmentalization, resulting in enzymatic browning caused by the contact of phenolics with polyphenol oxidase. It has been reported that phospholipase D (PLD), lipoxygenase (LOX), and lipase...
contribute to the degradation of membrane lipids. Among these enzymes, PLD catalyzes the hydrolysis of phospholipids, such as phosphatidylcholine (PC) and phosphatidylethanolamine (PE), producing a free head group and phosphatic acid (PA), which can cause membrane fusion and cell death due to the generation of hydroperoxides and free radicals. In addition, a number of studies have revealed that PLD has a crucial role in regulating the composition of membrane lipids during cold stress. Unsaturated fatty acids (USFAs) maintain membrane integrity upon chilling by reducing the melting temperature and improving membrane fluidity. An increase in polyunsaturated lipids was detected in cold-acclimated Arabidopsis thaliana. Our previous research showed that the browning of "Nanguo" pears is related to cellular membrane system damage, such as the degradation of cellular membrane phospholipids and the peroxidation of USFAs. Under cold stress, the lipid composition changes to become conducive to cytomembrane stabilization. In low temperature-stored bell peppers, increases in the percentage of PA and PLD activity were detected, which indicated that the PLD pathway had been activated and the membrane lipids had undergone remodeling during cold stress. There are many PLD family members in plants, including PLDα1, PLDβ1, PLDζ1, and PLDε. AtPLDα1 has been reported to hydrolyze structural phospholipids into PA, and the suppression of AtPLDα1-induced freezing tolerance in A. thaliana under cold stress. However, OsPLDα1, which is 79% identical to AtPLDα1 at the amino acid level, has opposite roles as those of AtPLDα1 in regulating cold stress responses. Genetic studies have shown that reducing the transcription of OsPLDα1 increased the chilling stress sensitivity of rice plants. In addition, the transcription level of bell pepper CaPLDε increased in response to cold stress. Recent research has shown that CaPLDε4 is regulated by the transcription factor (TF) CaNAC1, which binds to the promoter of CaPLDε4 and activates its transcription. The genetic knockout of AtPLDδ rendered A. thaliana plants more sensitive to freezing, whereas the overexpression of AtPLDδ increased its freezing tolerance, and lipid profiling revealed that the overexpression of AtPLDδ increased the production of PA species. It was reported that the knockdown of AtPLDβ enhanced the cold tolerance of A. thaliana. However, the transcriptional regulation of phospholipid degradation-related genes by upstream regulators was given little acknowledgment and is a scientific aspect that needs further exploration.

TFs are proteins that have the capacity to regulate gene transcription by binding the specific nucleotide upstream sequence of the 5'-end of a gene upon sensing abiotic stressors, including low temperature. Commonly, the nuclear localization signal, transcription-activation domain, DNA-binding domain, and oligomerization sites are the four domains that constitute TFs. In addition, TFs affect many multigene family members, greatly increasing the complexity of transcriptional regulation. Higher plants contain a wide range of TFs, among which MYB TFs constitute one of the largest TF families, accounting for ~9% of all plant TFs and participating in the regulation of many aspects of plant growth, development, metabolism, and stress responses. Most MYB family members share a highly conserved MYB domain in the N terminus. The DNA-binding domain in MYB TFs is typically composed of 50–53 amino acid residues, which each form a domain consisting of three α-helices. According to an analysis of the three-dimensional structure of the MYB domain, the second and third helices of each repeat have three regularly spaced tryptophan (W) residues forming a helix-turn-helix (HTH) structure with a hydrophobic core, whereas the third helix directly interacts with the major groove of the target DNA. Based on the number of repeats in MYB domains, MYB superfamily members are classified into four primary subfamilies: MYB-related proteins (1R-MYB, one repeat), R2R3-MYB proteins (2R-MYB, two repeats), R1R2R3-MYB proteins (3R-MYB, three repeats), and 4R-like MYB protein (4R-MYB, four repeats). Among these types of MYB TFs, R2R3-MYB is the major subfamily. R2R3-MYB TFs are characterized by various functions in regulating several aspects of biological processes, and many MYB TFs are involved in regulating responses to abiotic stresses, including cold stress. In apple, the transcription of MdMYB23 is induced by cold stress, and the overexpression of MdMYB23 increases the cold tolerance of transgenic apple calli and A. thaliana by activating the expression of C-repeat binding factors (CBFs), including MdCBF1 and MdCBF2. In addition, MdMYB88 and MdMYB124 positively regulate cold hardiness and cold-responsive gene expression under cold stress through CBF-dependent and CBF-independent pathways. Peel browning is caused by cold stress; however, the involvement of MYB TFs in peel browning has been largely unexplored.

In this work, we report the functions of PuMYB21 and PuMYB54 in “Nanguo” pears, which are both induced by cold stress. Biochemical experiments showed that both PuMYB21 and PuMYB54 bind directly to the PuPLDβ1 promoter in vivo and in vitro and then activate the expression of PuPLDβ1. Therefore, we propose a model of increasing abiotic tolerance based on a membrane lipid metabolism feedback loop in response to cold stress. This study provides insight into the regulatory mechanisms of MYB TFs under cold stress in plants.

Results

Peel browning

To estimate the severity of CI, we calculated the BI and peel-browning incidence of cold-stored “Nanguo” pears during the refrigeration and subsequent shelf life periods.
As shown in Fig. 1a, b, no peel browning was observed in the fruits stored for 60 d during the entire shelf life period, whereas the fruit stored for 120 d showed browning on the 3rd day of the shelf life period. The browning incidence and BI index increased greatly with the extension of shelf life time.

Changes in membrane phospholipids
The fruit refrigerated for 120 d showed obvious browning on the 6th day of the shelf life period, whereas no significant differences in ripeness were observed between the fruits refrigerated for 60 and those refrigerated for 120 d. We detected five key phospholipids, namely, phosphatidic acid (PA), phosphatidylcholine (PC), phosphatidylethanolamine (PE), phosphatidylserine (PS), and phosphatidylglycerol (PG), in the fruit stored for different periods (Table 1). The total phospholipid content level in the pears refrigerated for 120 d was much lower than it was in the short-term-refrigerated pears, which indicated that low temperature stress promotes the degradation of phospholipids in the peel tissues and may be closely associated with the peel browning of the refrigerated “Nanguo” pears. Furthermore, the distribution of phospholipids in the samples refrigerated for 60 and 120 d was different. The content levels of PA and PS in the samples refrigerated for 120 d were lower than they were in the samples refrigerated for 60 d; however, their percentages were significantly higher in pears refrigerated for 120 d. In addition, the content levels and percentages of PG, PC, and PE were reduced after long-term refrigeration.

The relative proportion of phospholipids is the percentage of the dry weight of each polar lipid with respect to the total dry weight of all the lipids. LT (60 d) and LT (120 d) indicate pears kept at low temperature (0 ± 0.5 °C) for 60 and 120 d, respectively. The values represent the mean and corresponding SD (n = 3). Different letters a and b indicate a 5% level of significance between different cold storage times as determined by pairwise Student’s t-test.

Changes in the expression of PLD family members in “Nanguo” pears stored for different times
To analyze the effects of low temperature stress on different PLD family members, this study tested the changes in PLD activity and expression of PLD family members, including PuPLDα1, PuPLDα4, PuPLDβ1, PuPLDδ, and PuPLDζ, in “Nanguo” pears stored at room temperature after different periods of refrigeration. As shown in Fig. 2a, the PLD activity level in the samples stored for 120 d was significantly higher than that in the samples stored short-term, except on the first day of shelf life. In terms of the expression of PLD, the performance of  

Table 1 Phospholipid composition and proportion of “Nanguo” pears on shelf life 6 d after being in cold storage for 60 and 120 d

| Lipid class | Lipid/dry weight (nmol mg⁻¹) | Relative proportion to total phospholipids (%) |
|-------------|-----------------------------|---------------------------------------------|
|             | LT (60 d)                   | LT (120 d)                                  | LT (60 d) | LT (120 d) |
| PA          | 1.55 ± 0.02a                 | 1.41 ± 0.02b                               | 26.40 ± 3.27a | 36.51 ± 2.58a |
| PG          | 0.91 ± 0.01a                 | 0.58 ± 0.02b                               | 15.47 ± 2.15a | 15.09 ± 2.17a |
| PS          | 0.11 ± 0.01a                 | 0.10 ± 0.01a                               | 1.95 ± 0.56b | 2.54 ± 0.89a  |
| PC          | 1.97 ± 0.03a                 | 1.02 ± 0.01b                               | 33.56 ± 4.33a | 26.35 ± 1.96b |
| PE          | 1.33 ± 0.02a                 | 0.75 ± 0.02b                               | 22.65 ± 4.12a | 19.48 ± 2.25b |
| Total lipids | 5.88 ± 0.33a                | 3.86 ± 0.42b                               | 100a       | 100a        |

Fig. 1 Changes in peel browning. a Peel-browning incidence. b Peel-browning index (BI) of “Nanguo” pears during the refrigeration and subsequent shelf life periods. LT (60 d) and LT (120 d) indicate pears maintained at low temperature (0 ± 0.5 °C) for 60 and 120 d, respectively. The values represent the mean and corresponding SD (n = 3). Different letters indicate a 5% level of significance as determined by pairwise Student’s t-test.
different family members was significantly different. PuPLDδ and PuPLDζ2 demonstrated similar trends (Fig. 2e, f). On the day the samples were removed from cold storage, the expression level in the short-term stored pears was significantly higher than that in the long-term-stored samples. However, when the pears were transferred to shelf life, the expression level decreased rapidly and vacillated in the long-term-refrigerated samples. The differences in the expression in PuPLDα1 in the pears during different storage periods were mainly reflected in the middle and late stages of shelf life, with the expression of PuPLDα1 in the middle stage of the shelf life after long-term refrigeration being higher than that after short-term storage, and the trend in the later shelf life period was exactly the opposite (Fig. 2b). The expression of PuPLDα4 in the early and middle shelf life period was higher in the long-term-refrigerated fruit than it was in the fruit stored for 60 d, and in the later shelf life period, the expression
of PuPLDα4 was significantly higher in the long-term-stored fruit than it was in the short-term-stored fruit (Fig. 2c). In contrast to other PLD family members, the expression of PuPLDβ1 in the short-term-stored pears remained low throughout the shelf life period, whereas the expression of PuPLDβ1 in the pears after long-term storage gradually increased throughout the shelf life period, especially in the middle and late stages (Fig. 2d).

**Isolation and motif analysis of the PuPLDβ1 promoter**

To identify the cis-elements in the promoter sequences, a 700 bp region upstream of the translation start site in the PuPLDβ1 gene was isolated based on the homologous gene of the Chinese white pear and identified using PlantCARE. In addition to the core cis-acting elements, such as a CAAT and TATA box motif, an MYB-binding site (MBS) with a core sequence (CAACTG), WRKY-binding site W-Box, and bHLH-binding site G-Box were identified in the PuPLDβ1 promoter, suggesting the possible involvement of MYB, WRKY, and bHLH TFs in regulating PuPLDβ1. Binding sites for AREB/ABF TFs were also identified (Table 2).

**Interaction of PuMYB21 and PuMYB54 with the PuPLDβ1 promoter**

To verify the binding of the candidate transcription factor proteins to the PuPLDβ1 promoter, based on the transcription factor-binding site on the PuPLDβ1 promoter and our transcriptome results, 17 transcription factors from 3 families (PuMYB4, PuMYB6, PuMYB21, PuMYB24, PuMYB44, PuMYB54, PuMYB66, PuMYB84, PuMYB86, PuMYB91, PuMYB306, PuMYB1R1, PubHLH13, PubHLH15, PubHLH21, PubHLH66, and PuWRKY20) were selected for use in the yeast one-hybrid (Y1H) assay. Only the yeast strains with PuMYB21 and PuMYB54 proteins were able to grow on the cultivation medium containing AbA (Fig. 3a); the other TFs did not interact with PuPLDβ1 (Supplementary Fig. S1).

To investigate the specific binding sites, the full-length PuMYB21 and PuMYB54 proteins were purified, and EMSA was conducted with a biotin-labeled PuPLDβ1 promoter fragment containing the MYB-binding motif used as the labeled probe. PuMYB21 and PuMYB54 bound to the PuPLDβ1 promoter, and when an unlabeled probe containing mutated nucleotides was added as a competitor, the binding of PuMYB21 and PuMYB54 to the PuPLDβ1 promoter was not affected (Fig. 3b). This result indicates that PuMYB21 and PuMYB54 can physically bind to the MBS motif in the PuPLDβ1 promoter.

PuPLDβ is transcriptionally activated by PuMYB21 and PuMYB54

These results demonstrate that PuMYB21 and PuMYB54 can bind to the promoter of PuPLDβ1. To further examine whether the transcriptional activity of PuPLDβ1 was activated or suppressed by PuMYB21 and PuMYB54, a GUS transactivation experiment was performed in wild tobacco (Nicotiana benthamiana) leaves. The CDSs of PuMYB21 and PuMYB54 were inserted into a pRI101 vector, and the promoter fragments of PuPLDβ1 were inserted into a pBI101 vector. A significant increase in GUS activity level was observed, compared with the control, when 35S::PuMYB21 + ProPuPLDβ1::GUS and 35S::PuMYB54 + ProPuPLDβ1::GUS were coexpressed (Fig. 4a). Furthermore, we found that there was significantly higher relative GUS activity in the coexpressed 35S::PuMYB21 + 35S::PuMYB54 + ProPuPLDβ1::GUS tobacco leaves than in the others. These results suggest

| Factor or site name | Signal sequence | Probable function |
|---------------------|----------------|-------------------|
| ABRE                | CACGTG, ACGTG, TACGTG, CACGTA | Abscisic acid-responsive element |
| as-1                | TGACG           | Transcription factor-binding site |
| CAAT-box            | CCAAT           | Common cis-acting element in the promoter and enhancer regions |
| ERE                 | ATTTTTAAA       | ERF (ethylene response factor) binding site |
| G-box               | CACGTT          | Involved in light responsiveness |
| MBS                 | CCACTG          | MYB-binding site |
| TATA-box            | TATA; TATAA; TATAA | Core promoter element approximately –30 from the transcription start site |
| TCA-element         | CCATCCITT       | Salicylic acid-responsive element |
| TGAGC-motif         | TGAGC           | MeJA-responsive element |
| W-box               | TTGACCC         | WRKY-binding site |
| WRE3                | CCACTC          | WRE transcription factor-binding site |
| WUN-motif           | AAATTTCTT       | WUN transcription factor-binding site |
that PuMYB21 and PuMYB54 enhance the transcription of \textit{PuPLD\textbeta 1} by directly interacting with its promoter.

Transient expression assays with tobacco leaves using a dual-luciferase reporter were also performed. The CDSs of PuMYB21/54 were cloned into pGreen II 62-SK, and the promoter fragment of \textit{PuPLD\textbeta 1} was inserted into pGreen II 0800-LUC. The data showed that the interaction of PuMYB21 with \textit{PuPLD\textbeta 1} led to a nearly 2.2-fold increase in the relative LUC/REN ratio, and the interaction of PuMYB54 with \textit{PuPLD\textbeta 1} led to a nearly fourfold increase in the relative LUC/REN ratio (Fig. 4b). Interestingly, there was a 7.4-fold relative LUC/REN ratio in the tobacco leaves injected with PuMYB54 + PuMYB21 + \textit{PuPLD\textbeta 1}. These results indicate that PuMYB21 and PuMYB54 are transcriptional activators that regulate the expression of \textit{PuPLD\textbeta 1}. In addition, we speculated that PuMYB21 and PuMYB54 most likely interacted with each other.

To elucidate the positive regulation of PuMYB21 and PuMYB54 in activating \textit{PuPLD\textbeta 1}, we monitored the expression of \textit{PuPLD\textbeta 1} by real-time PCR analysis in “Nanguo” pear fruit and calli transiently transformed with MYB overexpression or silencing constructs. As shown in Fig. 4c, the expression of \textit{PuPLD\textbeta 1} was significantly higher in both the PuMYB21-OE and PuMYB54-OE “Nanguo” pears and calli. In contrast, \textit{PuPLDB1} expression was significantly lower in both PuMYB21-Anti and PuMYB54- Anti “Nanguo” pears and calli. These results suggest that both PuMYB21 and PuMYB54 can positively regulate the transcription of endogenous \textit{PuPLD\textbeta 1} in “Nanguo” pears.

**Isolation and sequence analysis of PuMYB21 and PuMYB54**

The increased expression of \textit{PuPLD\textbeta 1} in long-term cold-stored fruit suggested that the action of low temperature in accelerating peel browning involves transcriptional regulation. Since the \textit{PuPLD\textbeta 1} promoter has MYB-binding motifs, the PuMYB21 and PuMYB54 transcription factors were isolated from “Nanguo” pears. After performing a NCBI BLAST search, we found that PuMYB21 and PuMYB54 were homologous genes of PbrMYB21 (XM_009359681.2) and MdMYB54 (XM_017337313.2), respectively. The PuMYB21 gene is 720 bp and encodes a 239 amino acid polypeptide, and the PuMYB54 gene is 813 bp and encodes a 269 amino acid polypeptide. Multiple sequence alignment indicated that both genes belong to the R2R3-MYB subfamily, as they have two conserved MYB domains: an R2 MYB domain (red box) and an R3 MYB domain (green box) (Fig. 5a, b). A phylogenetic analysis indicated that PuMYB21 and PuMYB54
PuMYB54 have high similarity to PbrMYB21 and MdMYB54, respectively (Fig. 5c, d).

Expression of PuMYB21 and PuMYB54 in “Nanguo” pears during the shelf life period after refrigeration

The relative expression levels of the candidate genes PuMYB21 and PuMYB54 were tested during the refrigeration and subsequent shelf life periods. The expression of PuMYB21 in pears during different refrigeration periods increased throughout the shelf life time, whereas the expression in the samples under long-term refrigerated storage were much higher than those under short-term storage during the middle and late periods (Fig. 6a). The expression of PuMYB54 in the pears...
refrigerated for 60 d gradually increased throughout the shelf life time. In contrast, the increase range and expression level of PuMYB54 in pears refrigerated for 120 d were significantly higher than in the short-term-stored pears (Fig. 6b). The reverse-transcription quantitative PCR (RT-qPCR) results showed that the transcription levels of PuMYB21 and PuMYB54 were upregulated in response to cold stress and had an expression pattern similar to that of PuPLDβ1. From our previous RNA-seq study, the expression of other detected MYBs following long-term cold treatment is shown in Supplementary Table S1.

The subcellular localization of PuMYB21 and PuMYB54

To confirm the subcellular localization of PuMYB21 and PuMYB54 in vivo, the CDS regions (without the stop codon) were fused to the eGFP reporter gene in the pRI101 vector. The constructs 35S::PuMYB21-eGFP and 35S::PuMYB54-eGFP were used to infect onion epidermal cells using the Agrobacterium-mediated transformation method with 35S::eGFP used as the negative control. The results showed that fluorescence signals from the 35S::PuMYB21-eGFP and 35S::PuMYB54-eGFP fusion proteins were exclusively localized to the nucleus, whereas 35S::eGFP was ubiquitously distributed throughout the cells (Fig. 7).

PuMYB21 interacts with PuMYB54

To verify that PuMYB21 can interact with PuMYB54, a Y2H assay was performed. We constructed the pGBKTT7-PuMYB54 bait vector and determined its self-activating activity. The results showed that PuMYB54 had self-activating activity; therefore, PuMYB54 was separated into
two domains and inserted into a pGBK7 vector. As shown in Fig. 8a, PuMYB54C also showed self-activating activity, whereas PuMYB54N did not exhibit self-activating activity. Therefore, PuMYB54N was used to identify the interaction between PuMYB54 and PuMYB21. Yeast cells transformed with pGBK7-PuMYB54N + pGADT7-PuMYB21 grew well and turned blue in SD/-Trp/-Leu/-His/-Ade/+X-alpha-gal, indicating that PuMYB21 interacted with PuMYB54 in yeast cells. Moreover, the N-terminal regions of PuMYB54 interacted with PuMYB21 (Fig. 8b). Then, the recombined polyhistidine-tagged PuMYB21 (PuMYB21-His) and recombinant glutathione S-transferase-tagged PuMYB54 (PuMYB54-GST) fusion proteins were purified, and a pull-down assay was conducted to confirm the interaction between PuMYB21 and PuMYB54. The results showed that PuMYB21 interacted with PuMYB54 (Fig. 8c). Finally, a BiFC assay was conducted using tobacco leaves to verify the interaction between PuMYB21 and PuMYB54 in vivo (Fig. 8d). PuMYB54 was fused to the N-terminal fragment of yellow fluorescent protein (YFP) to form PuMYB54-YFPN, and PuMYB21 was fused to the C-terminal fragment of YFP as PuMYB21-YFPc. The results showed that a yellow fluorescence signal was observed in the onion
Fig. 8 Interaction between PuMYB21 and PuMYB54. a The self-activating activity of PuMYB54. I pGBKT7 empty vector. II pGBKT7-PuMYB54 in Y2H. III pGBKT7-PuMYB54C in Y2H. IV pGBKT7-PuMYB54N in Y2H. b The N terminus of PuMYB54 interacted with PuMYB21. Y2H Yeast strain containing the bait plasmid PuMYB54N was transformed into PuMYB21. QDO is medium without Leu, Trp, His, or Ade. X-α-gal is QDO with X-α-gal. SV40 and P53 served as the positive control, and the others were negative controls. c A pull-down assay verified the interaction between PuMYB21 and PuMYB54. Escherichia coli-expressing GST or PuMYB54-GST fusion proteins were incubated with cobalt chelate affinity resin containing the immobilized histidine-tagged MdBT2 protein. The protein mixtures were purified using a HIS purification kit. d Bioluminescence complementation assays showing the interaction between PuMYB21 and PuMYB54 in the nuclei of onion epidermal cells.
cells coexpressing PuMYB54-YFP\textsuperscript{N} and PuMYB21-YFP\textsuperscript{C}, suggesting that PuMYB54 physically interacted with PuMYB21 in the nucleus.

**Discussion**

**The expression of PuPLDβ1 is related to peel browning during the refrigeration and subsequent shelf life periods**

Cold storage is a valid method to extend the shelf life of postharvest fruits and vegetables, but it can cause some extent of CI\textsuperscript{3}\textsuperscript{5}. Peel browning is a main CI manifestation in refrigerated “Nanguo” pears during its shelf life, affecting the quality and reducing the commercial value of the pears\textsuperscript{3}\textsuperscript{5}. The biomembrane is not only a barrier that protects cells from injury but also a primary location for sensing cold signals\textsuperscript{10}. An increasing number of studies have reported that one of the main causes of CI is changes in membrane lipid composition, structure, and metabolic processes, which ultimately result in the degradation of cell membrane lipids\textsuperscript{10}. In bell pepper samples, the level of total membrane lipids declined notably because of CI\textsuperscript{6}. The authors of this previous found that the percentage of PA was higher in the CI samples than in the fresh samples and that the content level and percentage of PC exhibited an extreme decreasing trend, whereas the content of other phospholipids, such as PE, PG, PS, and PI, decreased slightly. Here, we found that the content level of all the membrane phospholipids in “Nanguo” pears refrigerated for 120 d was significantly lower than that it was in the fruit stored for 60 d. Furthermore, the percentage of PA in the peel of the browning pears stored for 120 d was significantly higher than that in the short-term refrigerated pears. In addition, the content levels and percentages of PC and PE in the long-term-stored fruit were significantly lower than those in the short-term refrigerated samples. PLD can catalyze the hydrolysis of phosphodiester bonds and produce secondary messengers, such as inositol triphosphate, diester glycerol, acetylcholine, and PA, which can cause a series of secondary reactions by changing intracellular protein kinase K and Ca\textsuperscript{2+} levels, thus completing the cell response to cold signals\textsuperscript{15}. Under cold stress, PLD activity increases significantly, which facilitates the process of membrane lipid metabolism. In the current study, the PLD activity level in the fruit refrigerated for 120 d increased significantly during the shelf life period compared with the pears stored for 60 d. These results suggested that low temperature stress stimulated an increase in PLD enzyme activity and increased the possibility of membrane lipid degradation, which might be one of the main reasons for the formation of peel browning. Plant PLD is composed of a family of heterogeneous enzymes with distinguishable catalytic and regulatory properties, endowing them with diverse physiological functions\textsuperscript{34}. In cucumber, higher PLD activity at 2 °C was due to increased PLD mRNA levels\textsuperscript{35}. The transcription level of *Arabidopsis* PLDβ1 was significantly increased by bacterial and fungal pathogen infection\textsuperscript{36}. A similar study was reported 10 years later, in which PLDβ1 was found to be a negative regulator of salicylic acid-dependent resistance to bacterial Pst DC3000 but a positive regulator of the jasmonic acid-dependent pathway and resistance to the fungal pathogen *Botrytis cinerea*. PLDβ1 could also be induced by wounding stress\textsuperscript{38}. The suppressed expression of rice PLDβ1 inhibited the germination of rice seed\textsuperscript{39}. In addition, the suppression of OsPLDβ1 could also activate defense responses and increase disease resistance\textsuperscript{40}. These results indicate that PLDβ1 has a crucial role in plant growth, development, and the response to stress. In this study, the expression patterns of *PuPLD* were investigated. The expression of *PuPLDα1* and *PuPLDα4* in the two groups of pears fluctuated during the shelf life period. The expression of *PuPLDδ* and *PuPLDC2* in the long-term-refrigerated pears exhibited a downregulated trend after refrigeration. However, the expression of *PuPLDβ1* was significantly increased after transfer from long-term cold storage, and its transcription patterns were consistent with PLD activity. Thus, we speculate that *PuPLDβ1* has an important role in cold-induced membrane lipid metabolism and peel browning in “Nanguo” pears.

**PuMYB21 and PuMYB54 directly upregulate the expression of *PuPLDβ1***

Our research showed that low temperature can induce the expression of *PuPLDβ1* and promote the degradation of membrane lipids, thereby mediating the peel browning of cold-stored “Nanguo” pear. However, the mechanisms involved in how *PuPLDβ1* is regulated during cold stress remain to be studied. TFs are regulators that have important roles in many biological processes in plants by regulating spatiotemporal gene expression through recognizing specific DNA sequences in target gene promoters\textsuperscript{41}. In this study, two candidate MYB TFs were identified that directly bind to the *PuPLDβ1* promoter and activate the expression of the *PuPLDβ1* gene. A comparison of the PuMYB21 and PuMYB54 amino acids with MYB proteins from other plant species revealed that PuMYB21 and PuMYB54 may be clustered into the typical R2R3-type MYB. Furthermore, phylogenetic analysis suggested that PuMYB21 and PuMYB54 shared a specific sequence similarity to PbrMYB21 of *Pyrus betulaefolia* and MdMYB54 of *Malus domestica*, respectively, indicating that PuMYB21 and PuMYB54 are putative MYB homologs of *Pirus ussuriensis*. The overexpression of PbrMYB21 improved drought tolerance in *P. betulaefolia* by upregulating the expression of stress-related genes, including ADC, PS5CS, and LEA\textsuperscript{52}. OsMYB30 is a cold-response R2R3-type
MYB gene, and the overexpression of OsMYB30 increases cold sensitivity, whereas the osmyb30-knockout mutant showed increased cold tolerance. Moreover, OsMYB30 interacted with OsJAZ9, which had a significant role in suppressing the transcriptional activation of OsMYB30 and in repressing the BMY genes mediated by OsMYB30. This repression of BMY increased the content level of maltose, which might contribute to cold tolerance as a compatible solute. In the current study, the expression patterns showed that PuMYB21 was significantly increased by long-term refrigeration and markedly boosted on the sixth day of shelf life after refrigeration. Similar to PuMYB21, the expression of PuMYB54 exhibited a continuous increase regardless of the previous length of refrigeration or subsequent shelf life. These results indicated that PuMYB21 and PuMYB54 were also induced by cold stress, and their transcriptional patterns were similar to those of PuPLDβ1. Considering these results, we speculated that PuMYB21 and PuMYB54 might bind to the PuPLDβ1 promoter and regulate its expression, thereby mediating and participating in the peel browning in refrigerated “Nanguo” pears. The MYB cis-acting element contains a MYB recognition site with the TAACTG sequence, called MBS, in the target gene promoter. Previous studies showed that many MYB TFs can recognize and bind to the TAACTG consensus sequence to modulate the transcription of their downstream target genes, such as Glycine max GaMYB1, Oryza sativa OsMYB2, P. betulaefolia PbrMYB21, and Malus domestica MdMYB1. In the current study, sequence analysis showed that PuPLDβ1 contains MBS in its promoter region. Furthermore, Y1H and EMSA assays demonstrated that the PuMYB21 and PuMYB54 proteins can specifically bind to the PuPLDβ1 promoter by recognizing the MBS cis-element with the conserved TAACTG sequence; this result is consistent with that of previous studies. In addition, because the expression levels of PuMYB21, PuMYB54, and PuPLDβ1 were upregulated during the process of cold-induced peel browning, we speculated that the occurrence of peel browning in refrigerated “Nanguo” pears was induced by the positive regulation of PuMYB21 and PuMYB54 on PuPLDβ1. To verify our speculation, GUS and dual-luciferase reporter assays were conducted. The results indicated that PuMYB21 and PuMYB54 can indeed activate the transcription of a reporter gene downstream of the PuPLDβ1 promoter. These results were also verified by analyzing the expression of PuPLDβ1 in over-expressed and silenced “Nanguo” pears and calli that have transiently expressed the PuMYB21 and PuMYB54 genes. These findings are the first to demonstrate that PuMYB21 and PuMYB54 can activate PuPLDβ1 expression to mediate membrane lipid metabolism.

The interaction of PuMYB21 and PuMYB54 promotes the upregulation of PuPLDβ1 expression

The regulation of gene expression by TFs is a complex process in which one or more TFs or other proteins may participate in coordinated expression. In banana, the expression of MaMYB4 alone significantly repressed the transcription of MaFAD3-1, MaFAD3-3, MaFAD3-4, and MaFAD3-7; however, this repression was further enhanced when MaMYB4 was coexpressed with MaHDA2, which interacts with MaMYB4. Two MYB TFs, GmMYB76 and GmMYB177, were shown to enhance the freezing tolerance of A. thaliana by activating the expression of freeze-tolerance-responsive genes. In this study, we found that the transcription level of reporter genes downstream of the PuPLDβ1 promoter was enhanced by the simultaneous existence of PuMYB21 and PuMYB54. Therefore, we suspect that this phenomenon may be induced by the interaction between PuMYB21 and PuMYB54 or the overlapping individual effects of PuMYB21 and PuMYB54. To verify whether there is an interaction between PuMYB21 and PuMYB54, Y2H, BIFC, and pull-down assays were performed. These assays proved that there was indeed an interaction between PuMYB21 and PuMYB54. These results help us further understand the regulation of membrane lipid metabolism in refrigerated “Nanguo” pears. Furthermore, this is the first study to research the interaction between two MYBs during cold storage in fruit. However, the specific synergistic model of PuMYB21 and PuMYB54 needs further study.

In this study, we identified two MYB TFs, PuMYB21 and PuMYB54, expressed in the peel of browning “Nanguo” pears after refrigeration. The two MYB TFs belong to the R2R3-MYB family and are closely related to PbrMYB21 (XM_009359681.2) from Pyrus × bretschneideri and MdMYB54 (XM_017337313.2) from Malus domestica. They bind to the promoter regions of PuPLDβ1 and coregulate its expression. Correlating with the increased expression of PuMYB21 and PuMYB54, the expression level of PuPLDβ1 was elevated during the pear refrigeration and subsequent shelf life periods. Our discovery led to new regulatory targets and metabolic pathways of PuMYB21 and PuMYB54 proteins and complemented the current understanding of the coregulatory network of the membrane lipid degradation mechanism, providing a new perspective for revealing the molecular mechanism of membrane lipid degradation-induced peel browning in cold-stored “Nanguo” pears. It remains to be investigated whether other TFs bind to the PuPLDβ1 promoter and regulate membrane lipid degradation in plants. The members of these TF families can participate in more metabolic pathways, and these results suggest that different TF families may be recruited in the same way to commonly
regulate peel browning of “Nanguo” pears through different metabolic pathways.

Materials and methods
Plant material and treatment
“Nanguo” pears were harvested on September 11, 2018, at an orchard located in Anshan, Liaoning Province, China. The picked fruits were transported in <3 h to the laboratory and then selected based on uniform color, size, and absence of pests and damage induced mechanically or by disease. The fruits were randomly allocated into two triplicate groups, each with 300 fruits, which were placed at room temperature (20 ± 1 °C) for a 4-d pre-ripening period and covered with newspaper to prevent water loss50,51. After pre-ripening, all of the fruits were pre-cooled at 0 ± 0.5 °C for 24 h and then placed in reusable plastic boxes lined with 0.04-mm-thick polyethylene bags. There were 50 pears in each plastic bag, and each bag was tied and stored at 0 ± 0.5 °C with 80–85% relative humidity (RH). After storing for 60 and 120 d, the fruits were moved to room temperature for the shelf life study (20 ± 1 °C, 80–85% RH, natural light). Fifty fruits were used to evaluate peel-browning indexes and browning incidence. The peel tissues of the remaining fruits were frozen in liquid nitrogen every 3 d and stored at −80 °C until the biochemical analysis was performed.

The calli of “Nanguo” pears were used for A. tumefaciens infiltration. The calli were cultured on solid Murashige and Skoog (MS) medium containing 2.0 mg of 2,4-dichlorophenoxyacetic acid and 1.5 mg of 6-benzylaminopurine at 28 °C in the dark (Supplementary Fig. S2).

Incidence and index of peel browning
The incidence of peel browning was characterized as the proportion of browning fruit to total fruit. The value is expressed as a percentage.

The browning index (Bl) was calculated as previously described5. The severity of browning was assessed based on the following four-grade scale: 0 = no peel browning, 1 = slight (browning area < 1/3 of the total area), 2 = moderate (1/3 < browning area < 2/3 of the total area), and 3 = severe (browning area > 2/3 of the total area). The result was calculated using the following formula: Bl (%) = Σ [(browning scale) × (number of fruit at that scale)]/(3×total number of fruit) × 100.

Membrane phospholipid content
The extraction of membrane phospholipids was performed as previously described16. The solution containing membrane phospholipids was evaporated under nitrogen. Membrane lipid extracts were dissolved in 1 mL of chromatographic grade methanol, and then detection was conducted using automated electrospray ionization-tandem mass spectrometry as previously described52.

Determination of phospholipase D activity
The activity of PLD in the peel tissue was determined as previously described53. The pellet containing PLD was dissolved in 1 mL of acetone, and the activity of PLD was determined using a spectrophotometer (TU-1810 DSPC, Beijing Puxi Instrument Co., Beijing, China) at 520 nm. One unit of PLD activity was defined as a change in absorbance of 0.01 per hour at 520 nm. The specific PLD activity was based on the protein mass; it is expressed as U mg⁻¹ fresh weight.

RNA extraction and RT-qPCR analysis
Total RNA was extracted from the “Nanguo” pear peel using the RNAPure plant kit (DNase I) (CW0559S, CWBIO, China) following the manufacturer’s instructions. Total RNA concentration and purity were quantified using a Microplate reader (Eon; BioTek, USA) at OD260/OD280 and OD260/OD230, and the integrity of the total RNA was detected using 1.0% agarose gel electrophoresis. The RNA obtained was used as the template to synthesize cDNA using a HiFiScript cDNA synthesis kit (CW2569, CWBIO, China) according to the kit instructions. RT-qPCR was performed using the UltraSYBR mixture kit (CW0957M, CWBIO, China) in 20 μL of volume, according to the kit instructions, on a QuantStudio™ 5 Real-Time PCR system (ThermoFisher, USA) with the following protocol: denaturation step of 94 °C for 5 min, followed by 40 cycles of 95 °C for 10 s and 60 °C for 1 min. Dissociation curves were generated for gene primers to detect the presence of nonspecific amplification. The expression of the target gene was normalized to that of the internal reference gene (PuActin) using the 2⁻ΔΔCt method (the comparative Ct method). The specific primer sequences that were designed by using Primer Premier 6.0 software and synthesized by GENEWIZ Biotechnology Synthesis Lab (Jiangsu, China). The primers used are listed in Supplementary Table S2. A total of three biological replicates were included.

Promoter isolation and analysis
A NuClean Plant Genomic DNA kit (CW0531S, CWBIO, China) was used to extract genomic DNA from “Nanguo” pear peel tissue following the manufacturer’s instructions. The PuPLDβ1 promoter fragment was isolated according to the sequence (ID:103966461) in Chinese white pear (primers are listed in Supplementary Table S3, and the promoter sequences are listed in Supplementary Table S4). The conserved cis-element motifs of the PuPLDβ1 promoter sequence were predicted using PlantCARE online tools (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/)54.

Yeast one-hybrid (Y1H) assays
To construct the prey and bait vectors, the full-length CDSs of the TFs were cloned (the primers are listed in
Supplementary Table S3, and the sequences are listed in Supplementary Table S4) and inserted into the pGADT7 vector, and the promoter of PuPLDβ1 was inserted into a pAbAi vector. The Y1H assay was conducted using a Matchmaker Gold yeast one-hybrid system (Clontech) as previously described.55.

Electrophoretic mobility shift assay (EMSA)
The full-length CDSs of PuMYB21 and PuMYB54 were inserted into a pET-SUMO vector to construct expression vectors, which were introduced into Escherichia coli strain BL21 (DE3) cells. To purify the proteins, the cells were incubated in 0.5 mM isopropyl-β-D-thiogalactopyranoside (IPTG) for 3 h at 30 °C.

The EMSA assay was performed using a LightShift chemiluminescent EMSA kit (20148, ThermoScientific, Illinois, United States) as previously described. PuPLDβ1 promoter fragments with MYB-binding sites was synthesized and labeled with biotin probes (Gene Create). The EMSA assay was performed using as previously described. A small part of the PuPLDβ1 promoter fragments that contains PuMYB21 and PuMYB54 were constructed and inserted into the pGBKT7 vector as bait. The CDSs of PuMYB54 were inserted into a pGBK7T vector as prey. PuMYB21 was inserted into the pGBKT7-53 vector as insert. The bait was transformed into Y2H yeast cells and grown on synthetic medium without tryptophan (Trp) and containing 200 μM aureobasidin A and 20 mg mL⁻¹ chromogenic substrate X-α-gal, which turns blue upon transactivation.

A toxicity test indicated that the full-length PuMYB54 protein does not cause toxic effects in the yeast host cell (data not shown). The Y2H yeast strains were cotransformed with PuMYB54 + PuMYB21, PuMYB54N + PuMYB21 or PuMYB54C + PuMYB21. PuMYB54 + pGADT7 or pGBK7T-53 + pGADT7-T was used as a negative or positive control.

Bioluminescent assays (BIFC)
BIFC assays were conducted in onion tissue to confirm the interaction between PuMYB54 and PuMYB21. The CDS of PuMYB54 was fused to the N-terminal fragment of yellow fluorescent protein (YFP) in a 35S-pSPYNE-YFP vector to generate PuMYB54-YFP, and the CDS of PuMYB21 was fused to the C-terminal fragment of YFP in a 35S-pSPYNE-YFP vector to generate vector PuMYB21-YFP. The transformed Agrobacterium was mixed (1:1, v/v) and used to infect onion epidermal cells. After 3 d, the YFP fluorescent signal was detected with a laser scanning confocal microscope (TCS SP8-SE; Leica, Wetzlar, Germany).

Pull-down assays
The CDS of PuMYB54 was recombined into a GST-tagged PGEX-6P-1 vector, and the CDS of PuMYB21 was inserted into the His-tagged pET-SUMO vector. The reconstruction vectors of PuMYB54-GST and PuMYB21-His were transformed into BL21 to induce proteins. The pull-down assays were performed using a His-tagged protein purification kit (Clontech, Palo Alto, CA, USA) according to the manufacturer’s instructions. The eluted samples were detected by western blotting using HIS and GST antibodies (Abcam, Cambridge, UK).

Vector construction and the generation of transgenic “Nanguo” pears and calli
The CDSs of PuMYB21 and PuMYB54 were constructed into the pRI101 vector to form PuMYB21-OE and PuMYB54-OE vectors. For RNAi constructs, the interference fragments of PuMYB21 (386 bp) and PuMYB54 (387 bp) were obtained using the primers (Supplementary Table S5) and inserted into a pCAMBIA2301-ky vector to form PuMYB21-Anti and PuMYB54-Anti vectors (interference fragments are listed in Supplementary Table S6). The vectors were transformed into A. tumefaciens strain.
was repeated three times for three biological replicates. To overexpress or silence PuMYB21 or PuMYB54 in "Nanguo" pears, 100 μL of infection suspension was injected into pears at a depth of 0.5 cm. Ten fruits were infiltrated with each construct. Three fruits were used as biological replicates. To overexpress or silence PuMYB21 or PuMYB54 in "Nanguo" pear calli, the “Nanguo” pear calli were dipped into the infection suspension for 25 min at room temperature for transfection. The infected calli were then grown in the dark at 28 °C on solid MS medium containing 2,4-D and 6-BA for 8 d and subsequently used for RNA extraction, as described above. Each transfection was repeated three times for three biological replicates.

Bioinformatics analysis of PuMYB21 and PuMYB54
The full CDS sequences of PuMYB21 and PuMYB54 were amplified according to the accession numbers XM_009359681.2 and XM_017337313.2, respectively. The comparisons of the amino acid sequences for homologous PuMYB21 and PuMYB54 were performed using ClustalX 1.83 (http://align.genome.jp/). A phylogenetic tree was constructed using the neighbor-joining method in MEGA5.0.

Subcellular localization of PuMYB21 and PuMYB54
To investigate the subcellular localization of PuMYB21 and PuMYB54, the full-length coding sequences (CDSs) of PuMYB21 and PuMYB54 without terminate codons were inserted into the pRI101-eGFP vector using NdeI and KpnI restriction enzyme sites to form the fusion expression vectors PuMYB21-eGFP and PuMYB54-eGFP under the regulation of the cauliflower mosaic virus (CaMV) 35S promoter. The fusion expression vectors were transferred into A. tumefaciens strain GV3101 and used to transform onion epidermal cells as previously described59. Laser scanning confocal microscopy (TCS SP8-SE; Leica, Wetzlar, Germany) was used to detect fluorescence signals.

Statistical analyses
All experiments were repeated at least three times. Statistical analyses were performed using SPSS 19.0 software (IBM Inc., Chicago, IL, USA). Two groups of data were compared using Student’s t-test, and a difference was considered to be significant when P<0.05.

Acknowledgements
We sincerely thank Dr. Yujin Hao (College of Horticulture Science and Engineering, Shandong Agricultural University, Tai-An, Shandong, China) for the generous gift of the expression vectors 35S-pSPYCE-YFPβ, 35S-pSPYNE-YFPβ, pGreen II 0800, and pGreen II 62-SK. We thank Dr. Aide Wang (College of Horticulture, Shenyang Agricultural University, Shenyang, China) for the gift of Y1H and Y2H Gold strains and pGADT7-pGBK7, pRI101, and pBl101 vectors. We also thank Zhihong Zhang (College of Horticulture, Shenyang Agricultural University, Shenyang, China) for providing the pRI101-eGFP vector. This work was supported by the National Natural Science Foundation of China (No. 31971698).

Author contributions
H-J.S. and S-J.J. participated in the experimental design and the improvements made of the manuscript, M-LL, Y-YS, W-YG, and M-MY. performed the experiments. X.Z. and Q.Z. analyzed the data. The authors of the study have reviewed the content of the manuscript to ensure accuracy.

Conflict of interest
The authors declare that they have no conflict of interest.

Supplementary Information accompanies this paper at (https://doi.org/10.1038/s41438-020-00356-3).

Received: 24 February 2020 Revised: 2 June 2020 Accepted: 6 June 2020
Published online: 01 September 2020

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