Effects of Storage Temperature on Fruit Quality and Expression of Sucrose Phosphate Synthase and Acid Invertase Genes in Japanese Pear

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Japanese pear (Pyrus pyrifolia Nakai) ‘Gold Nijisseiki’ fruit were stored for 28 days at 0, 4, 10, 15, or 22°C. The effects of storage temperature on sucrose metabolism and on the expression of genes for sucrose-metabolizing enzymes were investigated. Fruit firmness, skin color, and content of various sugars were affected by storage temperature. Fruit stored at 22°C underwent rapid fruit softening and skin color change. Fruit stored at 15 or 10°C had lower sucrose content and higher glucose and fructose content than those stored at 0, 4, or 22°C. To investigate whether sucrose loss is related to changes in the expression of sucrose-metabolizing enzymes (vacuolar acid invertase: AIV; and sucrose phosphate synthase: SPS), we examined the expression of 3 genes during storage, namely, those encoding AIV (PpAIV1 and PpAIV2) and SPS (PpSPS1). Storage at 10 and 15°C increased the expression of PpAIV2 within a week and that of PpAIV1 in 14 days, while storage at 0 and 4°C delayed increased expression of PpAIV1 and PpAIV2 until after 28 days. Changes in the gene expression of sucrose-metabolizing enzymes were followed by delayed responses in sugar content.

Key Words: acid invertase, cold storage, Pyrus pyrifolia Nakai, sucrose metabolism.

Introduction

The composition and concentration of sugars are important components of fruit quality. In pear, sorbitol is the major translocation carbohydrate form of photosynthates (Bieleski, 1977; Loescher, 1987; Webb and Burley, 1962), and is converted into fructose and glucose by NAD+-dependent sorbitol dehydrogenase (SDH) and sorbitol oxidase (SOX), respectively. Sucrose is then synthesized or degraded by the enzymes sucrose phosphate synthase (SPS), sucrose synthase (SS), and acid invertase (AIV) (Tanase and Yamaki, 2000; Yamaki and Moriguchi, 1989). For these reasons, sucrose, glucose, fructose, and sorbitol contents play a key role in determining the sweetness of Japanese pear (Pyrus pyrifolia Nakai) fruit (Kajiura et al., 1979), although there are large differences in sugar composition among cultivars (Moriguchi et al., 1992). Sucrose begins to accumulate in fruit tissues in the maturing stage. Some cultivars such as ‘Nijisseiki’ and ‘Chojuro’ accumulate sucrose as over 50% of the neutral sugar in ripening fruit (Kajiura et al., 1979). The cultivar differences in the accumulation of sucrose are regulated by the activity of enzymes catabolizing sucrose (Moriguchi et al., 1992).

Because most of the reactions in fruit are biochemical and thus temperature-dependent, storage temperature is the primary factor affecting fruit freshness, shelf life, and quality. At higher temperature following harvest, European pear as well as other temperate fruits rapidly become yellow and overripe (Knee et al., 1983; Porrit, 1964). Although storage at low temperature is a common practice for retarding the softening of fruit, long periods of cold storage are known to alter the taste rating of pear (Beutel, 1990; Ke et al., 1991). In our previous study, storage at 4°C for longer than 1 month caused the accumulation of hexoses and a decrease in sucrose in Japanese pear (Itai and Tanahashi, 2008) due to up-regulation of invertase genes (PpAIV1) and down-regulation of a SPS gene (PpSPS1). Moreover, the addition of treatment with 1-methylcyclopropene inhibited the accumulation of PpAIV1 transcripts and the decline in PpSPS1 transcripts, resulting in delayed sucrose loss.
However, it is unclear which storage temperatures specifically affect sucrose metabolism. Against this background, the present study aimed to determine the effect of 5 different storage temperatures on sucrose metabolism in Japanese pear fruit.

Materials and Methods

Plant materials and treatment

Over 300 fruit from 3 Japanese pear (Pyrus pyrifolia) ‘Gold Nijisseki’ trees grown at a commercial orchard in Tottori, Tottori Prefecture, Japan, were used for the present experiments. Fruit were harvested on 3 September, 2009, 141 days after full bloom, which is the optimal commercial harvest date for ‘Gold Nijisseki’. Fruit were packed in ventilated commercial boxes (W × D × H: 500 mm × 350 mm × 275 mm) and packed boxes of fruit were kept at 1 of 5 temperatures (0, 4, 10, 15, or 22°C) for up to 28 days.

Fruit quality assessments

Ten fruit per treatment were sampled at 0, 7, 14, and 28 days after harvest. Fruit color and flesh firmness of the 10 individual fruit were determined on opposite sides of each fruit. Flesh firmness was determined by using a rheometer (RT-3010D; Sun Scientific Co., Tokyo, Japan) to measure the force required to penetrate each peeled fruit to a depth of 1 cm. Puncture tests were performed using an 8-mm probe on a drill base with crosshead speed set at 50 mm·min⁻¹. Skin color at harvest was greenish, but fruit stored at 0, 4, 10, or 15°C demonstrated loss of firmness at 7 days after harvest, but was maintained at around 13 N even at 28 days after harvest. No significant differences were observed in firmness among fruit stored at 0, 4, 10, or 15°C within the 28 days of storage.

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RNA extraction and real-time quantitative RT-PCR

Total RNA was extracted by the hot borate method (Wan and Wilkins, 1994). First-strand cDNA was synthesized from 1 μg of total RNA from fruit using M-MLV reverse transcriptase (ReverTra Ace; Toyobo, Tokyo, Japan). Expression levels were then analyzed using the Mini Opticon real-time polymerase chain reaction (PCR) system from Bio-Rad. Each sample was tested in duplicate in a 48-well plate (Bio-Rad, Hercules, CA, USA). The reaction mix (20 μL final volume) consisted of 10 μL of FastStart SYBR Green mix (Roche Applied Science, Mannheim, Germany), 1 μL of each primer (200 nM final concentration) described below, 2 μL of H₂O, and 3 μL of a 1/10 dilution of the cDNA preparation. The absence of genomic DNA in RNA samples was checked by real-time PCR before cDNA synthesis (minus RT control). A blank (no template) control was incorporated in each assay. The thermocycling program consisted of initial holding at 95°C for 3 min, followed by 40 cycles of 10 s at 95°C, 20 s at 60°C, and 20 s at 72°C. After completion of these cycles, melting-curve data were collected to assess PCR specificity, contamination, and the absence of primer dimers. Furthermore, PCR products were checked by electrophoresis on a 2% agarose gel and staining with ethidium bromide. The threshold cycle (Ct) values were measured using the gene expression module of the BIORAD CFX Manager software by the ΔΔCq method.

The PpSPSI gene was amplified from cDNA by PCR using the following oligonucleotide primers: sense 5'-TGACTTAGATTACCGTTCGCA-3' and antisense 5'-GGGAATTCACTAGTGGATCT-3', based on the PpSPSI sequence (Itai and Tanahashi, 2008). The primers for PpAIV1 and PpAIV2 were designed as follows (Itai and Tanahashi, 2008). The sense primer was 5'-GTGGTTGATCATCGTCATTCT-3', while the antisense primer was 5'-GGTAGCCTTGGACATCCA-3', and antisense primer was 5'-CTAAAGCTTGCGTTGCTTGTC-3' for PpAIV1. The sense primer was 5'-CATCAACAATGGGTTCCTCCA-3', while the antisense primer was 5'-ATGGTAGGAGGATATCAACCC-3' and antisense 5'-CATGTCATCCACGATTTGCTCATT-3', based on the PpACT1 sequence.

Results

Fruit assessment during storage

The firmness of ‘Gold Nijisseki’ flesh was 16.4 N at harvest (Fig. 1). The flesh firmness of fruit stored at 22°C decreased rapidly to 7.6 N at 28 days after harvest. Fruit stored at 0, 4, 10, or 15°C demonstrated loss of firmness at 7 days after harvest, but was maintained at around 13 N even at 28 days after harvest. No significant differences were observed in firmness among fruit stored at 0, 4, 10, or 15°C within the 28 days of storage considered in the present study.

Skin color at harvest was greenish, but fruit stored at 10, 15, or 22°C rapidly turned yellow within 14 days. However, cold storage, particularly at 0 or 4°C, delayed these color changes, as evidenced by the changes in value of the hue angle (Fig. 2).

Total sugar content was hardly changed at all temper-
tures throughout the experiment (data not shown). Of the soluble sugars in the fruit, fructose and sucrose were predominant at harvest (Fig. 3). Sucrose content was maintained at all temperatures until 14 days after harvest, but decreased at 10, 15, or 22°C storage by 28 days after harvest. In particular, the sucrose content of fruit kept at 10 or 15°C decreased more rapidly than that of fruit stored at 22°C. Fructose content increased in fruit stored at 15°C by 1 week after harvest, and the fructose content of fruit stored at 10, 15, or 22°C increased rapidly 28 days after harvest. Fructose content was highest in fruit stored at 15°C. Glucose and sorbitol contents in fruit were lower than those of sucrose and fructose. Glucose content followed a similar trend to fructose content. Higher storage temperatures (10, 15, or 22°C) resulted in higher glucose content 28 days after harvest. Colder storage treatments (0 and 4°C) resulted in little change in glucose content. However, colder storage (0 and 4°C) resulted in higher sorbitol content at 28 days after harvest.

Expression of SPS and AIV during storage

The expression profiles of \(PpSPS1\), \(PpAIV1\), and \(PpAIV2\) were investigated in fruit exposed to different storage temperatures using real-time RT-PCR (Fig. 4). The transcript levels of the genes were standardized relative to the constitutive expression level of the actin gene. \(PpSPS1\) was expressed at nearly constant levels during the entire experimental period, except for 14 days after harvest and storage at 15°C (Fig. 4A). The level of \(PpAIV1\) transcripts began to increase sharply after 14 days of storage at 0, 10, 15, or 22°C, reaching its maximum value at 10°C and then remaining constant until 28 days of storage at 10, 15, or 22°C, whereas its expression of 0°C increased after 28 days after harvest (Fig. 4B). A delayed increase in the level of \(PpAIV1\) transcripts with 4°C storage was observed after 28 days after harvest. On the other hand, \(PpAIV2\) transcripts responded quickly to exposure to 10 or 15°C (Fig. 4C). At 10°C, the \(PpAIV2\) mRNA increased rapidly and reached its maximum level within 7 days after harvest and then remained constant. Under the 15°C treatment, \(PpAIV2\) showed higher expression at 7 days after harvest, and then decreased again until 28 days. \(PpAIV2\) expression under the 22°C treatment remained at basal levels during the entire experimental period. In contrast, under cold storage (0 or 4°C), \(PpAIV2\) transcript accumulated and reached its maximum value at 28 days of 4°C storage.

Discussion

Pear fruit were stored at different temperatures (0, 4, 10, 15, or 22°C) for 28 days and then analyzed for the expression of genes for sucrose-metabolizing enzymes and for sugar levels. Softening proceeded more quickly in fruit stored at 22°C. Low and moderate temperatures helped fruit retain firmness during a month in storage. However, skin colors changed more rapidly in fruit stored at temperatures of 10 to 22°C. Color change is a major aspect of fruit ripening, along with softening. A delayed change in skin color was observed upon cold storage. In pear, as in other fruit species, cold storage slows the aging processes related to flavor, color, and texture.

Various studies have examined the effects of different storage temperatures on sugar content in other plant products (Ding et al., 1998; Huang et al., 1999). During storage at cold temperatures, potato tubers accumulate free reducing sugars derived from the breakdown of starch to sucrose, which is then converted into glucose and fructose (Blenkinsop et al., 2004; McKenzie et al., 2013; Samotus et al., 1974). This metabolic process is known as cold-induced sweetening (Sowokinos, 2001). Zrenner et al. (1996) reported that the soluble acid in-
vertase gene (INV19) is involved in the accumulation of reducing sugars during cold storage in potato tubers. We previously found the same cold-induced accumulation of reducing sugars in pear fruit (Itai and Tanahashi, 2008). Sucrose content was almost zero 1 month after 4°C storage in ‘Gold Nijisseiki’ and 2 months after 4°C storage in ‘Hosui’. Fruit accumulate sugars in the vacuole, and sucrose in the vacuole is converted into hexose by vacuolar acid invertase. Additionally both neutral invertase and apoplastic acid invertase activities are significantly lower than that of vacuolar acid invertase and negligible in Japanese pear fruit (Yamada et al., 2006). Therefore, we cloned AIV (PpAIV1 and PpAIV2) and SPS (PpSPS1) from ripe pear fruit and examined their expression during storage (Itai and Tanahashi, 2008). The deduced amino acid sequences of PpAIV1 and PpAIV2 are almost identical to PsS-AIV1 and PsS-AIV2, respectively, which are considered to be vacuolar acid invertases with a WECVD motif (Itai and Tanahashi, 2008; Yamada et al., 2007). The SPS and AIV constitute a multigene family (Ruan, 2014), so other isoforms should be clonable from other tissues. The expression of 2 vacuolar acid invertase genes was found to play an important role in sucrose degradation during cold storage. However, which specific temperatures had an influence on sucrose metabolism in pear fruit was unknown. Therefore, we studied the changes in the sucrose metabolism at 5 different storage temperatures.

Intermediate temperatures (10 and 15°C) induced higher expression of PpAIV2 at 1 week after harvest and of PpAIV1 at 14 days after harvest. Increased expression of these genes was followed by a decrease in sucrose and an increase in reducing sugar content at 28 days after harvest. There is a time lag between the changes in the levels of expression of the genes analyzed and the changes in the content of each sugar analyzed. The expression of PpAIV1 was induced by 22°C treatment, whereas that of PpAIV2 was not. These data indicate that these 2 vacuolar acid invertase genes respond differently to temperature. Meanwhile, the expression of both PpAIV1 and PpAIV2 was increased at

Fig. 3. Changes in total and individual sugar contents of ‘Gold Nijisseiki’ pears stored at different temperatures (0, 4, 10, 15, or 22°C). (A) Glucose content, (B) fructose content, (C) sucrose content, (D) sorbitol content. Vertical bars indicate the standard error from each mean value (n = 5).
changed upon cold storage. Our previous study revealed that sucrose content had reached almost zero 1 month after 4°C storage in ‘Gold Nijisseiki’ (Itai and Tanahashi, 2008). It is unclear how the difference between the previous and present studies was generated. It may have resulted from annual variation in fruit physiological status. High levels of $PpAIV1$ and $PpAIV2$ expression on 28 days at 4°C storage would cause lower sucrose and higher reducing sugar content later. Yamada et al. (2007) investigated the relationship between the expression of 2 acid invertase genes and the enzyme activity throughout fruit development and found that the acid invertase activity seemed to correlate with the $PsS-AIV1$ transcript level during the developmental stages. It is not known whether the acid invertase activity was correlated with increased expression of 2 vacuolar invertase genes during storage. However, differences in the expression level of 2 vacuolar acid invertase genes between different temperatures appeared to be associated with reducing sugar accumulation. Villalobos-Acuna and Mitcham (2008) reported that exposure to intermediate temperature (10°C) stimulated the capacity to produce adequate levels of ethylene during ripening more quickly than did exposure to low (0°C) or high (20°C) temperatures in ‘Bartlett’ pear. The results of the present and other studies indicate that ripened pear fruit may have differential sensitivity to low, intermediate, and high temperatures.

In our previous study (Itai and Tanahashi, 2008), we investigated the effect of two different temperatures (4°C and 25°C) and 1-MCP treatment on the sucrose metabolism of stored fruit and suggested that $PpAIV1$ could be responsible for sucrose degradation during storage in Japanese pear. In the present study, in addition to two different temperatures, we examined the effect of intermediate temperatures on sucrose metabolism during storage and suggested an important role for both $PpAIV1$ and $PpAIV2$ in sucrose degradation during storage. The effect of different temperatures on sucrose metabolism was variable in other species and tissues. The invertase transcript did not accumulate in potato tubers stored at 10°C, but in tubers stored at 1°C, the invertase transcript level increased markedly within 7 days (Zhou et al., 1999). Invertase activities and reducing sugar concentration also significantly increased in roots of sweet potato kept at a low temperature (Huang et al., 1999). Sucrose loss was shown not to be increased by cold storage in loquat fruit (Ding et al., 1998), while sucrose content was largely unaffected during cold storage in apple (Ackermann et al., 1992). These reports provide evidence that sugar content during storage may be regulated by variability in the temperature-dependent expression of sucrose-metabolizing genes in various species.

In summary, storage temperature significantly influenced the expression of vacuolar acid invertase genes in fruit of Japanese pear. The expression of 2 vacuolar
acid invertase genes had different responses to storage temperatures, especially at low (0 and 4°C) and intermediate temperatures (10 and 15°C).

**Literature Cited**


