Chlorophyll Fluorescence as a Nondestructive Indicator of Freshness in Harvested Broccoli

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Abstract. Chlorophyll “a” fluorescence (Fvar) was compared with respiration and vitamin C content of broccoli [Brassica oleracea L. (Botrytis group)] during storage at 1°C. The amplitude of the Fvar maxima declined in a similar manner as respiration and vitamin C content. Fvar was highly correlated with respiration (r = 0.83, P > |r| = 0.0001). The correlation of Fvar with vitamin C content was weaker (r = 0.42, P > |r| = 0.0002). The results demonstrate that Fvar is an indicator of postharvest changes in broccoli and that Fvar can be used as a nondestructive indicator of early changes in tissue condition (i.e., degree of freshness) of broccoli in storage.

Rapid, nondestructive methods are needed to assess changes in quality of vegetables (Watada, 1989). An important component of quality in many cases is the degree of freshness. Loss of freshness may not itself reflect large measurable changes in quality, but it precedes visible and readily measurable tissue deterioration. Any deterioration or decline of tissue from a freshly harvested state is considered a decline in freshness (Schwerdtfeger, 1979). Early changes of various tissue characteristics in vegetables have been identified, including respiration, free amino acids, vitamin C, and chlorophyll content (Perrin and Gaye, 1986; Schwerdtfeger, 1979; Solomos, 1983; Thimann, 1987). These changes are closely correlated, and they occur either in parallel or sequentially as the tissue begins to subsequently senesce (Thimann, 1987). Measurement of these characteristics involves a considerable amount of time to either process samples and/or perform analyses.

Two methods involving measurements of chloroplast function have been used for assessment of health of plant tissue. Delayed light emission (DLE) and chlorophyll “a” fluorescence (Fvar) are sensitive to changes at the cellular level; however, chlorophyll fluorescence generally has been found to be more useful (Havaux and Lannoye, 1985). DLE transients consist of many components and are affected by temperature, whereas Fvar analysis is relatively insensitive to temperature and the interpretation of transients is simpler (Jursinic, 1986; Krause and Weis, 1984). Fvar has been used to monitor ripening and chilling injury in tropical fruits (Smillie et al., 1987) and determine post-storage viability of conifer seedlings (Vidaver et al., 1989). Compared with DLE, the fluorescence assessment is simple to use, portable, and relatively inexpensive. Measurement of a single sample can take as little as 10 sec, and data interpretation can be done almost immediately.

This work was done to determine if chlorophyll fluorescence changes could be used to assess changes in tissue condition of broccoli, as has been done with established methods such as respiration measurement and vitamin C analysis. Changes in respiration rate and vitamin C content are considered sensitive indicators of changes in tissue condition after harvest (Perrin and Gaye, 1986; Solomos, 1983).

Broccoli used in this study was grown at the Agriculture Canada Research Station, Agassiz, B.C.; it was harvested on 28 Sept. 1990 and placed into storage at 1°C, relative humidity 97% ± 2%. During the course of storage, samples were taken for fluorescence, respiration, and vitamin C analysis.

The experiment was a randomized complete-block design. In all experiments, 24 replicates were selected at each of three times during storage. Three heads per replicate were subsampled for fluorescence and respiration measurement. For the latter, these three subsamples were bulked into one respiration chamber, and for Fvar measurement, three samples were averaged into one value. Only one head was sampled for each replicate in the vitamin C analysis.

Samples were placed into a dark room for 15 min, before the Fvar measurement, which was performed with a model SF30 Plant Productivity Fluorometer (Richard Brancker Res., Ottawa). The excitation light level was set at 650 W·m⁻² and measurement was for 10 sec. Stray light was corrected for by use of a color standard that approximated the reflectance of the broccoli. Fvar was expressed as maximum amplitude of the variable component relative to the nonvariable component of chlorophyll “a” fluorescence (Toivonen and Vidaver, 1988).

Respiration measurements were performed at 4°C. Samples were placed in 12-liter polyethylene pails with tight-fitting lids. The pails were sealed and flushed with gas free of CO₂, and at time zero a measurement was taken. A subsequent sample was taken = 1 h later by flushing a gas sample bulb with sample atmosphere. The CO₂ was determined on a Shimadzu GC-9A gas chromatograph (TekScience, Oakville, Ont.), fitted with a 8 × 0.0032 m Porapak Q (Supelco, Oakville, Ont., 80/100 mesh) column. The oven temperature was 75°C and the helium carrier flow set at 50 ml·min⁻¹. A methanizer (at 350°C) converted the CO₂ to methane, which was then measured by an FID detector (at 250°C).

Vitamin C analyses were performed on tissue samples, including floral portions and stem tissue, that are normally considered to

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Correlation of respiration and relative fluorescence (Fvar) of broccoli kept for 24 days at 1°C and 97% ± 2% relative humidity. The equation of the fitted line is $y = 16.80x + 7.27$.

Table 1. Changes in respiration, vitamin C content, and maximum Fvar amplitude of broccoli kept at 1°C and 97% ± 2% relative humidity.

<table>
<thead>
<tr>
<th>Time in storage (days)</th>
<th>Respiration rate (mg CO/kg h)</th>
<th>Vitamin C (mg/l00 g fresh wt)</th>
<th>Fvar (relative units)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>36.6</td>
<td>81.8</td>
<td>1.67</td>
</tr>
<tr>
<td>12</td>
<td>26.5</td>
<td>73.8</td>
<td>1.20</td>
</tr>
<tr>
<td>20-24</td>
<td>23.6</td>
<td>73.0</td>
<td>0.98</td>
</tr>
</tbody>
</table>

Significance

- Linear: ***
- Quadratic: NS

Average of 24 replicates.

**Significant at $P = 0.01$ or 0.001 or not significant, respectively.

Fig. 1. Correlation of respiration and relative fluorescence (Fvar) of broccoli kept for 24 days at 1°C and 97% ± 2% relative humidity.

Linear

Quadratic

Respiration rate and vitamin C content were significantly correlated ($P > |r| = 0.0002$), the relationship was rather weak ($r = 0.42$). After an initial decline from day 4 to 12, the vitamin C content did not change appreciably, while respiration and Fvar continued to decline (Table 1). The plot of respiration against Fvar and their correlation is shown in Fig. 1.

Measurements of respiration and Fvar provide direct information on the functioning of the mitochondria and chloroplast, respectively. These organelles are very sensitive to early stages of deterioration in plant tissue (Dalling and Nettleton, 1986; Solomos, 1983). While vitamin C is very labile (Perrin and Gaye, 1986), its tissue content is determined by the complements of other components of an oxidant protective system and the rate of production of active oxygen species by the mitochondria and chloroplast (Salin, 1987). Thus, the vitamin C content is determined by a complex biochemical matrix and the state of mitochondrial and chloroplast function; therefore, it may not completely reflect changes in either organelle’s function. The relatively weak correlations of vitamin C with respiration ($r = 0.37$) and Fvar ($r = 0.42$) bears out this reasoning.

The results show that Fvar analysis can be used to obtain freshnes assessments similar to those provided by measurement of respiration. Vitamin C provided parallel information during the early period of storage. Fvar is the least expensive and time-consuming of the three methods. It was capable of monitoring early stages of deterioration in a green vegetable because it produced information that was consistent with another established method. The method is also practical for many samples, since each measurement takes $= 10$ sec to complete. Hence, Fvar has great potential as a nondestructive method for assessing decline in freshness of green vegetables, such as broccoli, during cold storage.

Literature Cited


