Anthocyanins and anthocyanin-degrading enzymes in Kwai May and Wai Chee cultivars of litchis grown in Reunion Island and Spain

Marie-Noëlle DUCAMP-COLLIN*, Marc LEBRUN, Hassina RAMARSON, Guy SELF

* Correspondence and reprints

Abstract — Introduction. After harvest, litchi fruits (*Litchi chinensis* Sonn.) rapidly lose their bright red skin color. Peel browning of harvested litchi fruit has largely been attributed to rapid degradation of red anthocyanin pigments associated with the oxidation of phenolic compounds by polyphenol oxidase (PPO) and/or peroxidase (POD). An anthocyanase has been also identified in litchi peel. Our work aimed at characterizing two specific litchi varieties that differ greatly in their color and browning behavior. Materials and methods. The anthocyanins, polyphenol oxidase (PPO), peroxidase (POD) and anthocyanase in the pericarp of two litchi cultivars, Kwai May and Wai Chee, grown in Reunion Island and Spain, respectively, were studied. Results and discussion. The qualitative composition of major anthocyanins (cyanidin-3-rutinoside and cyanidin-3-glucoside) was identical for the two cultivars studied, but, quantitatively, the variety Kwai May had a lower concentration than Wai Chee (64% less) of cyanidin-3-rutinoside. This component represented more than 90% of total anthocyanins in both cultivars. The activity of PPO was 6 times greater in the variety Kwai May than in Wai Chee and the activity of POD 30 times greater. The activity of POD was greater than that of PPO in both varieties. These differences help to explain the different behaviors of the cultivars during subsequent postharvest studies.

Réunion / Spain / *Litchi chinensis* / fruits / pericarp / color / anthocyanins / catechol oxidase / peroxidases / phenolic content

Anthocyanines et enzymes de dégradation des anthocyanines dans le péricarpe des cultivars de litchi Kwai May et Wai Chee cultivés à l’île de la Réunion et en Espagne.

Résumé — Introduction. Après récolte, les fruits du litchi (*Litchi chinensis* Sonn.) perdent rapidement la couleur rouge lumineuse de leur peau. Le brunissement du litchi récolté a été en grande partie attribué à la dégradation rapide, dans la peau, des colorants rouges d’anthocyanine liés à l’oxydation des composés phénoliques par l’oxydase de polyphénol (PPO) et/ou la peroxydase (POD). Une anthocyanase a été également identifiée dans le péricarpe du litchi. Nos travaux ont cherché à caractériser deux variétés spécifiques de litchi différant considérablement quant à leur couleur et à leur brunissement. Matériel et méthodes. Les teneurs en anthocyanes, oxydase (PPO) de polyphénol, peroxydase (POD) et anthocyanase du péricarpe de deux cultivars de litchi, Kwai May et Wai Chee, développés respectivement à l’île de la Réunion et en Espagne ont été étudiées. Résultats et discussion. La composition qualitative des anthocyanines prépondérantes (3-rutinoside de cyanidol et 3-glucoside de cyanidol) a été identique pour les deux cultivars étudiés, mais, quantitativement, la variété Kwai May a eu une concentration en 3-rutinoside de cyanidol inférieure à celle Wai Chee (64 % en moins). Pour les deux cultivars, ce composant a représenté plus de 90 % des anthocyanines totales. L’activité de la PPO a été 6 fois plus grande pour la variété Kwai May que pour Way Chee et l’activité de la POD a été 30 fois plus grande. L’activité de la POD a été plus grande que celle de la PPO pour les deux variétés. Ces différences aident à expliquer les différents comportements des cultivars pendant les études après récoltes.

Réunion / Espagne / *Litchi chinensis* / fruits / péricarpe / couleur / anthocyanes / catéchol oxydase / peroxydase / teneur en phénols

* Correspondence and reprints
1. Introduction

Of Asian origin, the litchi (Litchi chinensis Sonn.) is prone to a number of postharvest problems, among which the most important is browning of the pericarp. This phenomenon occurs less than 72 h after harvest [1–3]. Although the organoleptic quality of the aril is not changed, the fruit loses commercial value due to the loss of the vibrant red color.

The red coloration of the litchi is due to the presence of anthocyanin pigments [4–5]. Cyanidin-3-rutinoside and cyanidin-3-glucoside have been identified as the major anthocyanins present in the litchi pericarp [5–8]. The intensity of the coloration of the anthocyanins depends on pH, which influences their structure. The flavylium form is stable at acid pH and is colored red, but, at basic pH, it is transformed into its carbinol base (colorless), into chalcone (yellow), or into its quinonic base (blue) [9].

A number of authors have linked browning of the pericarp to a degradation of anthocyanins by the action of polyphenol oxidase [10–14]. Enzymes such as polyphenol oxidase (PPO) and peroxidase (POD) are primarily membrane-bound, whereas anthocyanins are vacuolar ([6]). Contact between the enzyme and substrate cannot therefore take place unless there is disruption at the cellular level. Thus, browning can follow different types of stress, such as climatic conditions preceding maturation of the fruit, disease, desiccation and thermal shock [15].

The principal postharvest treatment currently used for litchi is fumigation with sulfur dioxide (SO$_2$). SO$_2$ is a strong antioxidant that blocks oxidation, causing skin browning. Furthermore, it acidifies the cellular contents of the pericarp, thus stabilizing the anthocyanins. SO$_2$ is also an antifungal agent [16]. Nevertheless, the treatment has limitations. Fruits do not recover their initial red coloration after treatment [17] and, more importantly, sulfur residues can reach the aril, where they represent a health risk for people allergic to sulfur. In the USA, the application of SO$_2$ is banned on all food products, except grapes [3, 18].

Several alternative treatments to SO$_2$ fumigation have been proposed: for example, soaking in a solution of N$_6$-benzyladeneine [19], the combined use of glutathione and citric acid [20] and hot-water brushing [21], but no alternative method has been established commercially as yet.

The first objective of our work was to optimize the extraction methods and measurement of the pericarp enzymes implicated in browning (PPO, POD and anthocyanase) and of the anthocyanins. These different components were then measured and compared in two distinct varieties to obtain baseline data for storage trials of different modified-atmosphere and acid-dip treatments as alternatives to SO$_2$ fumigation, that will be reported elsewhere.

2. Materials and methods

2.1. Plant material

The litchi fruits used were of two varieties: ‘Kwai May’ came from Reunion Island and ‘Wai Chee’ from Spain. In these two varieties, the red coloration of the fresh fruit and the structure of the pericarp are very different (Figure 1). The fruits were harvested at maturity from orchards of the same age,
Compounds involved in the browning of litchis

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pre-cooled immediately and shipped in cold conditions (12 °C). The sample lots (20 kg of each variety) were selected in the orchard for uniformity of size and color, and were free of signs of fungal infection and insect attack. The sample lots were split into four batches of 5 kg. The fruits were immediately peeled and the pericarp flash-frozen upon arrival at the laboratory.

2.2. Measurement of anthocyanins by HPLC

2.2.1. Anthocyanin extraction

Frozen pericarp (2 g) (from a mixture of 40 homogenized fruits) was ground with 20 mL of extraction solution [(2.5% hydrochloric acid in methanol (v/v))][6]. After stirring for 20 min, the mixture was filtered through sintered glass (porosity n°1). The residue was re-extracted with 20 mL of the same extraction solution.

The total filtered extract was dried under vacuum at 30 °C in a rotary vacuum evaporator (RE 100, Bibby Sterilin Ltd., UK), and then solubilized in 2 mL of methanol. This was then filtered through a hydrophilized PTFE membrane (Filtre Millex 0.45 µm, Millipore, France) and stored at –20 °C. The extracts were diluted 20 times in extraction solution prior to HPLC analysis.

2.2.2. High-pressure liquid chromatography

The extracts were analyzed with a Spectra System P1000XR chromatograph (Thermo Separation Products, US) equipped with a 20-µL injection loop, an automatic gradient controller and a column oven, all controlled by PC1000 software.

Anthocyanins were separated on a Lichrospher 100RP18E column (5 µm, 250 mm × 4.6 mm i.d., Merk Eurolab S.A., France). The column oven temperature was 30 °C. The mobile phase was a mixture of solution A [2% formic acid in water (v/v)] and solution B [2% formic acid in water and 80% acetonitrile (v/v)], used with the following gradients [%A:%B]: at 0 min [97:3], at 4 min [97:3], at 52 min [65:35], at 57 min [20:80], at 61 min [20:80] at 62 min [97:3] and at 70 min [97:3]. The flow rate was 1 mL·min⁻¹. The eluted compounds were detected by measuring absorbance at wavelengths between 450 nm and 600 nm. Quantification was achieved using appropriate standards.

2.3. Measurement of PPO and POD

The present protocol was adapted from several published methods. The principal parameters that were optimized were the quantity of polyclar AT, the substrate and the temperature optimum.

Frozen pericarp (10 g) from 50 fruits was homogenized (Ultra Turrax T 25 Basic, Ika Labortechnik, Germany) with 50 mL of 0.05-M phosphate buffer (pH 6.8) and 3 g Polyclar AT (insoluble polyvinylpyrrolidone). After centrifugation at 4800 rpm and 4 °C for 20 min (CR 4.12 Jouan, USA), the supernatant was filtered through a 0.45-µm Millipore filter and constituted the enzyme extract.

To measure the activity of PPO, 0.2 mL of enzyme extract was added to 0.85 mL of phosphate buffer (pH 6.8) and mixed with 0.6 mL of 100 mM catechol freshly dissolved in phosphate buffer in a 1.5-mL cuvette. PPO activity was measured at 400 nm for 3 min (25 °C) with a UVIKON 933 spectrophotometer (Kontron Instruments, Italy) and expressed as ΔOD₄₀₀nm·min⁻¹·g⁻¹ dry matter.

To measure POD activity, 0.1 mL of enzyme extract was mixed with 0.5 mL of 10 mM galactol solution. The reaction was started immediately by the addition of 1 mL of 10 mM H₂O₂. Activity was measured, at ambient temperature, at 470 nm for 3 min as before and expressed as ΔOD₄₇₀nm·min⁻¹·g⁻¹ dry matter.

2.4. Measurement of anthocyanase activity

The method used was that of Zhang et al. [22], except for the buffer and the centrifugation speed and duration. Frozen pericarp (4 g) was homogenized with 10 mL of sodium acetate buffer (pH 4.0) and Polyclar AT (insoluble polyvinylpyrrolidone at 10% of pericarp weight). The homogenate was
centrifuged at 4800 rpm and 4 °C for 20 min, the supernatant representing the crude enzyme extract.

One mL of enzyme extract was added to 4 mL of 0.05 mM cyanidin-3-glycoside (Extra-synthèse, France) dissolved in sodium acetate buffer (pH 4.0). After incubation for 10 min at 40 °C, 5 mL of 0.1 M HCl was added to the mixture and the absorbance recorded at 510 nm for 3 min. Anthocyanase activity was expressed as $\Delta \text{OD}_{510\text{nm}} \cdot \text{min}^{-1} \cdot \text{g}^{-1}$ dry matter.

2.5. Determination of dry matter

Twenty litchi fruits were peeled and between 2 g and 5 g of pericarp were pre-dried at 70 °C for 24 h in an ULM 500 oven (Memmert, Germany), then dried under vacuum at the same temperature in a 45003 oven (Bioblock Scientific, US) for between 24 h and 48 h until the mass did not vary by more than 1 mg. The dry matter content was expressed as a percentage of the fresh weight.

3. Results and discussion

It is important to note that the two varieties, though harvested from orchards of the same age, were grown in very different conditions: Kwai May in a tropical climate (Reunion Island) and Wai Chee (Spain) in a Mediterranean climate. The coloration and structure of the peel were totally different in the two varieties.

3.1. Anthocyanin content

The two litchi cultivars studied had the same HPLC profile and identical major constituents. Nevertheless, Wai Chee was richer in the amount of anthocyanins than Kwai May, in which the concentration of cyanidin-3-rutinoside was only 36% of that in Wai Chee (figure 2).

It was found that cyanidin-3-rutinoside was 12.3 times more abundant than cyanidin-3-glucoside in Wai Chee, but in Kwai May, it was only 3.6 times more abundant. In the cultivar Kwai May, Le Roux et al. [23] found cyanidin-3-rutinoside and cyanidin-3-glucoside present in a ratio of 4 to 1. For the cultivar Wai Chee, Zhang et al. [8] reported that cyanidin-3-rutinoside was the major anthocyanin present in the pericarp, representing 97% of the total (32.3 times more abundant than all other anthocyanins). Cyanidin-3-rutinoside therefore appears to be particularly abundant in the cultivar Wai Chee, with the maximum proportion of anthocyanins as cyanidin-3-rutinoside 80% in Kwai May [6] and 75% in the cultivar Brewster [7].

It should be noted that results could vary according to the methods of extraction and measurement used, and the geographical origin of the material, as well as the age of the tree, the position of the fruit in the tree, the soil type and the climate. For HPLC analysis, Lee and Wicker [5] found a ratio of cyanidin-3-rutinoside to cyanidin-3-glucoside of 6.7 for the cultivar Brewster grown in Florida, while Riviera-Lopez [7] found a ratio of 4.4 for the same cultivar grown in Mexico.

3.2. PPO, POD and anthocyanase activity

The activities of PPO, POD and anthocyanase were greater in the cultivar Kwai May than in the cultivar Wai Chee (figures 3, 4). PPO activity was 6 times greater in Kwai
May than in Wai Chee, and the activity of POD 30 times greater. In both cultivars, it was found that POD activity was greater than that of PPO. These observations tend to confirm those of Zauberman et al. [16] and Zhang et al. [24].

The [POD / PPO] ratios obtained here are lower than those reported in the literature: [14.1 $\Delta$OD$_{470}$ nm·min$^{-1}$·g$^{-1}$ dry matter / 30 $\Delta$OD$_{400}$ nm·min$^{-1}$·g$^{-1}$ dm] for Kwai May [16] and [2.2 $\Delta$OD$_{470}$ nm·min$^{-1}$·g$^{-1}$ dm / 7.2 $\Delta$OD$_{400}$ nm·min$^{-1}$·g$^{-1}$ dm] for Wai Chee [24]. This is probably because these different authors used 4-methyl catechol as a substrate for the measurement of PPO, whereas we used catechol, which has a greater affinity for the PPO present in litchi pericarp [12].

The activity of anthocyanase was almost twice as great in the cultivar Kwai May as in Wai Chee (figure 4). The degradation of anthocyanins and its relation to browning in the litchi pericarp is complex. PPO and POD have, in fact, low affinities for anthocyanins, probably due to stearic hindrance caused by sugar moiety [25]. Zhang et al. [22] demonstrated an anthocyanase activity in litchi pericarp and it is thought that this enzyme, also known as anthocyanin-β-glucosidase, removes the sugar moiety from anthocyanins to form anthocyanidins, which are then oxidized by PPO and POD in the presence of phenolic compounds to form colored products.

Differences in enzyme activities may therefore only partly explain differences in browning, because other factors such as membrane stability and the metabolism of other polyphenols are important factors in anthocyanin degradation. The results presented here will be used in a follow-up to this study, which will test modified atmosphere storage and acid dips to prevent pericarp browning in the two cultivars.

References


Resumen — Introducción. Tras la cosecha, los frutos del lichi (*Litchi chinensis* Sonn.) pierden rápidamente el color rojo luminoso de su cáscara. Se atribuyó el oscurecimiento del lichi cosechado, en gran parte, a la rápida degradación, en la cáscara, de los colorantes rojos de antocianidina relacionados con la oxidación de los componentes fenólicos mediante la oxidasa de polifenol (PPO) y/o la peroxidasa (POD). Asimismo se identificó una antocianosis sin el pericarpio del lichi. Nuestros estudios pretendieron caracterizar dos variedades específicas de lichi diferenciándose considerablemente en cuanto a su color y a su oscurecimiento. **Material y métodos.** Se estudiaron los contenidos de antocianidinas, oxidasa (PPO) de polifenol, peroxidasa (POD) y antocianosis del pericarpio de dos cultivares de lichi, Kwai May y Wai Chee, desarrollados respectivamente en la isla de la Reunión y en España. **Resultados y discusión.** La composición cualitativa de las antocianinas preponderantes (3-rutinosido de cianidol y 3-glucósido de cianidol) fue idéntica en ambos cultivares estudiados. Sin embargo cuantitativamente la variedad Kwai May tuvo una concentración de 3-rutinosido de cianidol inferior que la variedad Wai Chee (el 64 % menos). Para ambos cultivares este componente representó más del 90 % de las antocianidinas totales. La actividad de la PPO fue 6 veces mayor para la variedad Kwai May que para Wai Chee y la actividad de la POD fue 30 veces mayor. La actividad de la POD fue mayor que la actividad de la PPO para ambas variedades. Estas diferencias ayudan a explicar los diferentes comportamientos de los cultivares durante los estudios tras las cosechas.

**Reunión / España / Litchi chinensis / frutas / pericarpio / color / antocianinas / catecol oxidasa / peroxídasas / contenido fenólico**