Grape berry development may be divided into two major stages of growth, separated by a lag phase (Coombe, 1976). During stage I berry pericarp growth is rapid, at first due to cell division and expansion, and later due to cell expansion alone (Harris et al., 1968). Berries accumulate organic acids but little sugar during stage I, and remain green and hard. Stage II is referred to as the lag phase of development, as berry growth slows. Rapid berry growth, as a result of cell enlargement, resumes with the initiation of stage III. During stage III sugar and color accumulate rapidly, and the concentration of organic acids declines.

Light influences the growth and composition of a wide variety of fruit, including grapes. Grape berries exposed to sunlight are generally higher in sugars, anthocyanins, and phenolics, and lower in titratable acidity, malate, and pH, compared to berries ripened in canopy shade (Kliewer, 1977; Kliewer and Antcliff, 1970; Kliewer and Lider, 1968; Morrison, 1988; Reynolds et al., 1986). Previous studies have shown that the effects of temperature (Hale and Buttrose, 1974) and vine water status (Hardie and Considine, 1976) on berry size and composition vary during fruit ontogeny. The purpose of this investigation was to determine the influence of cluster light exposure during specific stages of berry development on fruit growth and composition.

Materials and Methods

Plant materials and growth conditions. Three-year-old own rooted grapevines of Vitis vinifera L., cultivars ‘Cabernet Sauvignon’ and ‘Pinot noir’ were used in the study. Plants were grown in 12-L plastic pots containing 1 soil : 1 sand : 1 peat medium (by volume). Vines were grown outdoors under full sunlight for two seasons before their use. At the initiation of the experiment, 22 dormant vines of each cultivar were pruned to four, two-bud spurs and placed in the stationary phytotron at the Univ. of California, Davis. The phytotron is illuminated with natural sunlight, and provided a diurnal temperature range between 28 ± 2.4 °C (day) and 16 ± 1.8 °C (night). Relative humidity was maintained at 40%. Shoot growth was initiated 10 d after the vines were placed in the phytotron. When shoot length was about 0.3 m, each vine was thinned to four cluster bearing shoots. Shoots were vertically trained on bamboo stakes anchored in the soil of the pot.

Treatments and experimental design. Anthesis occurred about one month after budbreak. At fruit set, 16 vines of each cultivar were selected on the basis of uniformity of shoot growth and cluster development. Extra vines were removed from the phytotron. Shoots were thinned to one cluster, and four treatments (one per shoot) randomly assigned to each vine as follows:

1) Clusters exposed to light during stages I, II, and III (experimental control).
2) Clusters grown without light during stages I and II, and exposed to light during stage III.
3) Clusters exposed to light during stages I and II, and exposed without light during stage III.
4) Clusters grown without light during stages I, II, and III.
In this study fruit development stages I and II refer to the period between berry set (2 to 3 mm in diameter) and the initiation of berry softening, while stage III refers to the period between fruit softening and maturity. Each vine received all four treatments; thus, each plant served as an experimental unit. The vines of each cultivar were arranged in a randomized complete-block design consisting of four, four vine blocks.

Clusters grown in the absence of sunlight were placed in 17×20-cm aluminum-coated, white paper bags. Basal leaves on all shoots were removed to facilitate the fastening of bags to the shoot; contact between the bags and clusters was prevented. To encourage ventilation, one end of an 18 cm long black plastic tube (0.8 cm inner diameter) was inserted into each bag next to the cluster. Tubing outside the bag was wrapped in a ring to prevent light from entering the tube. The tops and bottoms of each bag were sealed tightly with black adhesive tape. Bags were opened briefly at night to apply fungicides.

A LI-190S quantum sensor attached to a LI-185 quantum meter (LI-COR, Lincoln, Neb.) was used to ensure that the interior of each bag was free of detectable light at the initiation of the experiment. A CR21 data-logger (Campbell Scientific, Logan, Utah) with four quantum sensors was used to monitor photosynthetically active radiation (PAR) incident to one control cluster in each experimental block. Quantum sensors were positioned horizontally, immediately above the clusters. PAR in this region was about 20% of ambient. Spectroradiometer measurements (LI-1800) revealed that the spectral composition of sunlight and light in the phytotron was similar.

The temperature of light-exposed and nonexposed fruit was determined by inserting thermocouples into randomly selected berries. The berries were discarded after the 24-h measurement period was completed. Measurements were taken on four sets of 'Cabernet Sauvignon' berries and two sets of 'Pinot noir' berries at various times throughout the experiment. The temperature of light-exposed and nonexposed fruit was similar, and exceeded air temperature by 1 to 3 °C.

Clusters from one randomly selected vine in each block were harvested 14, 32, 46, and 63 ('Cabernet Sauvignon') or 14, 25, 42, and 56 ('Pinot noir') d after berry set. These dates coincided with the following stages of berry development for each cultivar: middle of stage I (harvest 1); 20% berry softening (harvest 2); middle of stage III (harvest 3); and the end of stage III (harvest 4). To maintain a constant light environment, harvested vines remained in the phytotron until the completion of the study.

**Fruit analysis.** Harvested clusters were placed in plastic bags and transported on ice to the laboratory. Berries were removed from the rachis, weighed, and their diameters recorded with a caliper. On the second sample date the number of softened berries (i.e., berries with compressible flesh), and the number of berries with 25% or more of their surface colored, were recorded. Berries were then randomly separated into two equal subsamples. One sample was used for pH, sugar, and acid determinations, and the other for anthocyanin and total phenolic determinations. The samples were stored in sealed plastic bags at –20 °C.
until analyzed. Sample size was about 20 to 25 berries for ‘Cabernet Sauvignon’, and 25 to 30 berries for ‘Pinot noir’.

Frozen berries were thawed at room temperature in 250-mL breakers. Berries were crushed and skins macerated with a pestle. Juice pH was determined after 30 min of settling. Following pH determinations, juice samples were quantitatively transferred to their original beaker. Beakers were covered with a watch glass and boiled for 30 min with frequent stirring. Samples were cooled and filtered through Whatman no. 2 filter paper into a 250-mL volumetric flask. Following two 25-mL rinses with boiling distilled water, the filter paper and solid material were discarded, the filtrate was brought to volume with distilled water. An aliquot of this solution was retained for sugar and acid analyses. The hexose sugar and organic acid fractions of the samples were determined by high-performance liquid chromatography [pump model 1330, differential refractometer model 1770, and organic acid analysis column HPX-87H (300 × 7.8 mm), Bio-Rad Laboratories, Richmond, Calif.] according to the procedure of McCord et al. (1984).

Skin discs (4 mm in diameter) were removed from the equator of 20 frozen berries in each sample. The discs were placed in a polystyrene tube containing 50 mL of acidified methanol (1% HCl v/v) and extracted in darkness. After 48 h, the samples were mixed and allowed to settle. Absorbance was determined at 520 nm. Anthocyanin concentration (anthocyanin/cm² of berry skin) was determined using a molar absorbance of 28,000 and a molecular weight of 529 (Amerine and Ough, 1980). Total phenolics were determined on an aliquot of the above extract using the automated method of Slinkard and Singleton (1977), and were expressed as mg gallic acid equivalents (GAE)/cm² berry skin.

Results

Berry growth. Differences in berry size among the treatments in both cultivars were established before berry softening, and generally maintained throughout ripening (Fig. 1). The weight and diameter of berries grown without light during stages I and II, or stages I, II, and III, were similar and significantly lower compared to the control. In contrast, the growth of control berries and berries grown without light during stage III were similar. An exception was noted for ‘Pinot noir’ on the final sample date, as the weight of berries grown without light during stage III was lower compared to the control.

Berry composition. Berry hexose sugar concentrations were similar among the treatments in both cultivars following fruit set, with total sugar concentration (glucose + fructose) ranging between 10 to 15 mg·g⁻¹ fresh weight (Fig. 2). Berry softening was delayed when clusters were grown without light during stages I and II (Table 1). Following fruit softening, however, the sugar concentrations of all treatments increased sharply. Immediately after fruit softening, control fruit of ‘Cabernet Sauvignon’ had greater hexose sugar concentrations than the remaining treatments. On the final sample date, ‘Cabernet Sauvignon’ control berries and berries grown without light during stages I and II had similar hexose sugar concentrations. Berries of this cultivar grown without light during stages I, II, and III, as well as those grown without light during stage III, were lower in sugar than the control on the final sample date. Following fruit softening in ‘Pinot noir’, berries grown without light during stages I, II, and III were lower than the control, while the sugar concentrations of berries grown without light during stages I and II, or stage III, were similar to the control.

Following berry set, the malate concentration of the control and berries grown without light during stages I and II were similar for ‘Cabernet Sauvignon’ (Fig. 3). After berry softening in this cultivar, malate declined more rapidly in the control than in the other treatments. On the final sample date, however, only berries grown without light during stages I and II, and III were greater in malate than the control. Berries of ‘Pinot noir’ grown without light during stages I and II were lower in malate following berry set, and at berry softening, compared to the control. Immediately following fruit softening in ‘Pinot noir’, berries grown without light during stages I and II, or stage III, were greater in malate than the control. On the final sample date, control berries of this cultivar were lower in malate compared to the remaining treatments. Cluster light exposure had no effect on the berry tartaric concentration or juice pH of either cultivar (Fig. 3).

Color initiation was delayed in both cultivars when berries were grown without light during stages I and II (Table 1). In both cultivars, treatments grown without light during stages I and II were lower in anthocyanins compared to the control (Fig. 4). On the final harvest date in both cultivars, anthocyanins were greatest for control berries and lowest for berries grown without light during stages I, II, and III. Berries grown without light during stage III were also lower in anthocyanins than the control, but had more pigment than berries grown without light during stages I, II, and III. Treatment differences in phenolics followed the same relationship as described for anthocyanins (Fig. 4). In both cultivars, phenolics were greatest for control berries and lowest for berries grown without light during stages I, II, and III.

Discussion

The temperature of grape berries is directly related to their incident radiation (Smart and Sinclair, 1976); thus, daytime temperatures of sunlight-exposed berries may be 11 °C or more greater than nonexposed berries, depending on time of day and solar conditions (Kliewer and Lider, 1968; Reynolds et al., 1986). It is therefore difficult to separate the effects of light and temperature on fruit development in the field. In the present study, berry temperatures among the treatments were similar. Reported differences in berry growth and composition are therefore attributed to

Table 1. Influence of cluster light exposure on the softening and coloration of ‘Cabernet Sauvignon’ and ‘Pinot noir’ grapes.²

| Cultivar            | Days after berry set | Treatment               | Softened berries (%) | Colored berries (%) |
|---------------------|----------------------|-------------------------|-----------------------|---------------------|
| Cabernet Sauvignon  | 31                   | Control                 | 32 a⁵                 | 18 a                |
|                     |                      | No light stages I and II| 4 b                   | 0 b                 |
| Pinot noir          | 26                   | Control                 | 25 a                  | 18 a                |
|                     |                      | No light stages I and II| 6 b                   | 0 b                 |

²Control clusters received about 20% of ambient photosynthetically active radiation. Clusters grown without light were placed in aluminum foil bags at berry set.
³Means for each cultivar followed by different letters within columns are significantly different at P = 0.05 (Duncan’s multiple-range test).
The direct effects of light, rather than differences in fruit temperature as a result of cluster light exposure.

Berries grown without light during the initial stages of fruit development had lower weights and diameters than berries exposed to light during this same period. These results are contrary to previous studies, which reported that sunlight-exposed berries were smaller than berries grown in paper bags or canopy shade (Crippen and Morrison, 1986; Kliewer and Lider, 1968; Reynolds et al., 1986). Reynolds et al. (1986) suggested that the higher berry temperatures associated with sunlight exposure may inhibit fruit growth. The optimum temperature for grape berry growth ranges between 25 to 30 °C (Hale and Buttrose, 1974). Kliewer and Lider (1968) and Reynolds et al. (1986) reported the daytime temperature of sunlight-exposed berries often exceeds 30 °C, while the temperature of nonexposed berries is maintained within the optimum range for growth. The transpiration rate of sunlight-exposed fruit is also greater compared to nonexposed fruit (Blanke and Leyhe, 1987), and several investigators have associated the decreased size and higher soluble solids of sunlight-exposed fruit to their lower turgor and higher transpiration rates (Crippen and Morrison, 1986; Reynolds et al., 1986). Plants used in this experiment were well watered at all times, thus diurnal or seasonal fluctuations in vine water status were negligible.

During the initial stages of development, grape pericarp growth is a result of cell division and expansion (Harris et al., 1968). Light has been reported to activate cell division and induce cell expansion in leaves of *Phaseolus vulgaris* L. (Humphries and Wheeler, 1960; Verbelen and De Greef, 1979). Blanpied and Wilde (1968) reported that cell division in nonexposed (bagged) apple fruit was lower than in sunlight-exposed fruit shortly after fruit set. The authors concluded that these differences were at least partially due to the lower temperature of nonexposed fruit. In the current experiment, differences in berry size between light-exposed and nonexposed fruit appear to be due to light-mediated effects on cell division and/or cell enlargement, particularly during the initial stages of growth. While berries grown in the absence of light during stages I and II were significantly smaller than the control, berries grown without light during stage III were similar in size (‘Pinot noir’), or only slightly smaller (‘Cabernet Sauvignon’), compared to the control. In addition, berries grown without light during stages I and II, and exposed to light during stage III, were similar in weight and diameter to berries grown without light during stages I, II, and III. Thus, reductions in berry size resulting from sunlight exclusion during stages I and II could not be reversed via subsequent light exposure in stage III.

Several mechanisms may be responsible for the photocontrol of berry growth, including light-mediated effects on fruit photosynthesis and carbon metabolism. Based on published assimilation rates (Kriedemann, 1968), fruit photosynthesis contributed <1% of the estimated dry weight difference between light-exposed and nonexposed fruit in this study. Hole and Scott (1981) suggested that, despite its small contribution to total fruit dry weight at maturity, fruit photosynthesis during the initial stages of growth may be critical to subsequent fruit development. Light may also be essential for regulating the importation or metabolism of carbon and other assimilates in young fruit. Schapendonk and Brouwer (1984) reported a 50% reduction in the accumulated dry weight of cucumber fruit as a result of fruit shading. These authors estimated that fruit photosynthesis contributed only 2% to 7% of the total dry weight in sunlight-exposed fruit, and concluded that the elimination of fruit photosynthesis was not fully responsible for the dry weight differences between light-exposed and shaded fruit. They suggested that the sink strength of shaded fruit was reduced, and that sink function may be partially regulated by a growth factor synthesized in the light. Hole and Scott (1981) reported that the yield of pea was significantly reduced by fruit shading, but concluded that only a small portion of this reduction could be attributed to an absence of fruit photosynthesis. Their results also suggest that shading young fruit may permanently reduce sink strength. Light may also influence the concentration or activity of phytohormones which regulate fruit growth and development (Coombe, 1973).

The results of this study are in agreement with previous reports that sugar accumulation is greater for light-exposed fruit than for nonexposed fruit (Brown and Coombe, 1985; Crippen and Morrison, 1986; Kliewer and Lider, 1968, Reynolds et al., 1986). Compared to the control fruit softening was delayed, and sugar accumulation
reduced, when fruit were grown without light during stages I, II, and III. Based on the available data it is unclear whether the lower sugar content of these fruit resulted from a delay in berry softening, or if a reduction in the rate of glucose and fructose accumulation following fruit softening also occurred. The activity of at least one enzyme regulating sugar metabolism, invertase, is believed to be photoregulated in grape berries (Kliewer and Smart, 1989). A reduction in the concentration or activity of invertase, or other enzymes responsible for the regulation of sugar accumulation, may have been responsible for the lower sugar concentration of nonexposed fruit. Sugar accumulation is also inhibited when berry growth during stage III is restricted (Coombe, 1973). Perhaps due to their smaller size, or to alterations in their pericarp cell walls, the ability of berries grown without light during stages I, II, and III to expand and accumulate solutes during ripening may have been limited.

Under field conditions, light-exposed fruit are generally lower in malate than nonexposed fruit (Crippen and Morrison, 1986; Reynolds et al., 1986). These differences are normally attributed to the higher temperatures and respiration rates of light-exposed fruit (Kliewer and Lider, 1968; Lakso and Kliewer, 1978). In this study, light-exposed ‘Pinot noir’ berries had higher malate levels before fruit softening, and lower malate levels following fruit softening, compared to nonexposed berries. Since light is not known to influence malic acid accumulation in grape tissues (Ruffner, 1982), differences in malate accumulation and degradation in this study were likely due to treatment effects on fruit maturation rate.

Light had no effect on fruit tartaric acid concentration or juice pH in this study. That fruit tartaric acid concentration was not affected by light is in agreement with previous reports (Crippen and Morrison, 1986; Kliewer and Lider, 1968). However, field investigations indicated that sunlight-exposed fruit had significantly lower juice pH than nonexposed fruit (Smart, 1985). The results of the current study suggest that the effects of light on berry pH may be indirect, and perhaps related to the concomitant effects on berry temperature. The results also lend support to the suggestion that, with regard to juice pH, foliage light exposure is more important than cluster light exposure (Morrison, 1988).

Light-exposed fruit are generally higher in anthocyanins and phenolics compared to nonexposed fruit (Kliewer, 1977; Kataoka et al., 1984). Roubelakis-Angelakis and Kliewer (1986) found that the activity of phenylalanine ammonia-lyase, a key enzyme in secondary metabolism, increased when grape berries were exposed to light. Although grape berries do not accumulate pigments during this period, light exposure during stages I and II appears necessary for maximum pigment production during stage III. Exposing fruit to light before the onset of pigment production may increase the initial concentration or activity of one or several anthocyanin biosynthetic enzymes (Takeda et al., 1988). Once pigment accumulation is initiated, light is needed to maintain the maximum activity of these enzymes during ripening.

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