Biochemical Changes in *Vitis vinifera* Buds between Dormancy and Forced Bursting: A Case Study of Three Portuguese White Varieties

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Abstract: The transition between the dormancy and budburst is accompanied by physiological and biochemical changes, which include variations in the carbohydrates, enzymatic activity, as well as hormones present inside the buds. The present study aimed to evaluate the biochemical changes that occur in dormant buds and forced bursting of three *Vitis vinifera* L. white varieties (Alvarinho, Fernão-Pires, and Loureiro) from two vineyards located in the Vinhos Verdes Demarcated Region, NW Portugal. For this purpose, the contents of malondialdehyde (MDA), proline, carbohydrates, and gibberellic acid (GA₃), as well as antioxidant enzymes activity were analysed. The results showed that the MDA, proline, total soluble sugars, and GA₃ contents tended to be higher at budburst. In contrast, the starch content was higher at dormancy. The enzymatic activities of ascorbate peroxidase (APX), peroxidase (POX), and superoxide dismutase (SOD) were lower at budburst, compared with dormancy, while the reverse was observed in catalase (CAT). Significant differences were also observed between varieties for most biochemical parameters analysed, with the exception of CAT activity. Alvarinho stood out from other varieties for presenting higher MDA, proline, soluble sugars, and GA₃ content, while Fernão-Pires displayed higher values of APX and POX activities and starch content. To our knowledge, the present work is the first attempt to examine the biochemical changes in dormant buds and bursting, which determine the vegetative and reproductive continuity of grapevines. This information contributes to better vineyard management, particularly important in light of the climate change scenarios, which can affect the biochemical mechanisms of the buds and compromise the yield and quality of grapevines.

Keywords: carbohydrates; enzymatic activity; grapevine; lipid peroxidation; gibberellic acid

1. Introduction

In temperate climates, such as NW Portugal, the decrease in temperature and photoperiod induces grapevines to enter a period of dormancy, to survive due to unfavourable environmental conditions [1,2]. To ensure their survival during this cold period, grapevines set some special mechanisms, such as suspension of visible growth, reduced metabolic and enzymatic activity, accumulating carbohydrates (soluble sugars and starch), changes in endogenous plants growth regulators, and decreased in tissue water content [1,3–6]. However, at this stage, buds do not remain completely inactive, and the essential metabolic and respiratory activities continue to occur but at a lower intensity [4–6]. After a period of chilling accumulation, dormancy is broken, and buds start to burst. Budburst marks the visible, physiological ‘awakening’ (active period) of grapevines after a period of dormancy. This process is environmentally mediated by variety-specific responses to temperature and the photoperiod and by intrinsic factors [7–9]. This stage involves metabolic and physiological changes—namely, in respiration rate, growth regulators, carbohydrate metabolism, and tissue water content that can intensify the formation of reactive oxygen species (ROS) [10].
In recent years, biochemical mechanisms related to dormancy and budburst in different regions and plant species such as walnut [11], apricot [12], apple [13], grapevine [14], and figs [15] have been reported. All these studies describe changes in the activity of antioxidant enzymes (CAT and APX), sugar and starch content, and endogenous hormones such as GA_3.

To eliminate ROS and minimize oxidative damage, grapevines have an effective antioxidant control system, involving nonenzymatic antioxidants and enzymic antioxidants that protect cellular structures, neutralizing and/or removing free radicals and converting them into less reactive species [16,17]. Proline plays an important role in plant response to abiotic stress since it contributes to cellular osmotic adjustment, removal of ROS, protection of membrane integrity, and stabilization of enzymes and proteins [18–20]. Carbohydrates stored during the previous growing season constitute sources of energy for metabolic activity during dormancy and budburst. Soluble sugars are also important for the osmotic properties of buds and for conferring freezing tolerance, as they are essential for their winter survival [21]. In winter, starch is the main reserve of carbohydrates, but during spring, part of the starch reserves is hydrolysed into soluble sugars, providing energy for buds’ growth and development. At this stage, consumption of soluble sugars increases due to increased respiration rates and metabolic activity [22,23].

The endogenous gibberellins (GAs) regulate several aspects of plant growth and development processes, including seed germination, cell division and elongation, and floral induction [24]. Among the classes of gibberellins biologically active in breaking dormancy is gibberellic acid (GA_3). Physiological and transcriptomic studies showed the involvement of endogenous GA in regulating and breaking dormancy in different perennial species—namely, sweet cherry [25], grapevine [26], and the Japanese apricot [27].

An irregular and variable budburst is a significant obstacle to normal bud development, with a potentially severe impact on bud fruitfulness that can result in substantial yield losses due to decreased bunch number. Alvarinho, Fernão-Pires, and Loureiro are three of the most relevant white grapevine varieties for wine production in the Vinhos Verdes Demarcated Region, VVDR (Portugal’s prominent wine appellation). Despite their importance for the region, to date, no research has been developed on biochemical and physiological processes that occur in buds. Consequently, there is a need for better understanding and knowledge about the underlying biochemical changes in the buds, to allow improved vineyard management (e.g., bud load adjustment), particularly in the scenario of climate change [28]. To this end, this study aimed to investigate changes in antioxidant enzymatic activity, MDA, proline, GA_3, and carbohydrate content in buds during dormancy and budburst of these varieties from two vineyards of the VVDR.

2. Materials and Methods

2.1. Plant Material and Growth Conditions

The plant material was collected in two commercial vineyards, located in Celorico de Basto (41°25′15″ N, 7°58′10″ W, 243 m, Sub-Region of Basto) and Penafiel (41°12′24″ N 8°17′56″ W, 199 m, Sub-Region of Sousa) in the VVDR, NW Portugal (Figure 1). The region has a warm temperature climate with warm and dry summer (Cs), according to the Köppen–Geiger climate classification. The soil at Celorico de Basto is classified as umbric lepto soil, while at Penafiel is cumulic anthrosol [29].

After leaf fall (late October 2017), canes, with two retained buds, were randomly removed from 1-year-old wood of three Vitis vinifera L. white varieties—Alvarinho, Loureiro, and Fernão-Pires. These varieties are recognized for their adaptability to local edaphoclimatic conditions and high oenological potential for the production of ‘Vinho Verde’, distinguished by its aromatic, acidity, and flavour characteristics. The former two varieties are also extensively grown in NW Spain.
were placed in a walk-in growth chamber (FitoClima 10,000 EHHF, Aralab, Rio de Mouro, Portugal) for 15 days to force budburst. During this period, the chamber environmental conditions were: temperature = 25 °C/18 °C (day/night), photoperiod = 16 h, relative humidity = 60%, and a photosynthetic photon flux density = 300 µmol m\(^{-2}\)s\(^{-1}\). The canes were irrigated twice a week, up to field capacity, using tap water.

When the buds reached the phenological stage C (Baggiolini scale, 1952), the budburst percentage was calculated, according to the following equation: Budburst (%) = ((number of burst buds)/(total number of buds)) \times 100. The buds were then collected, immediately frozen in liquid nitrogen, and then stored at −80 °C for further biochemical analysis.

2.2. Biochemical Determinations

2.2.1. Preparation of Extracts

Buds (500 mg) were previously ground into a fine powder in liquid nitrogen using a chilled mortar and pestle and then homogenised in 8 mL extraction buffer (50 mM potassium phosphate buffer, pH = 7.8, containing 2 mM EDTA). The extracts were heated at 48 °C for 15 min and then centrifuged (4000 rpm) at 4 °C for 15 min. The supernatant was transferred into new tubes and stored at −20 °C for quantification of protein, malondialdehyde, and proline contents. The antioxidant enzymatic activity of bud extracts was also measured. Each analysis was performed in triplicate and determined against blank in 96-well microplates (Costar 3590, Corning, New York, USA).

Malondialdehyde Content

The level of lipid peroxidation was determined through the quantification of malondialdehyde (MDA). MDA content was measured by its reaction with thiobarbituric acid (TBA), following the procedure proposed by Ali et al. [30]. Then, 200 µL of the extract was added to 500 µL of 0.25% TBA in 10% of Trichloroacetic acid. The reaction mixture was heated at 100 °C for 30 min and then cooled in an ice bath. After centrifugation at 4000 rpm for 15 min, the absorbance of the supernatants was measured at 532 and 600 nm. MDA was calculated according to the following equation: MDA (%) = ((absorbance of sample - absorbance of blank)/(absorbance of standard - absorbance of blank)) \times 100.
calculated by subtracting the absorbance (A532–A600) and using the extinction coefficient of 155 mM$^{-1}$cm$^{-1}$. The results were expressed as µg g$^{-1}$ of fresh weight (FW).

Proline Content

Proline content was determined according to Ábraháim et al. [31]. In summary, 200 µL of the extract was added to 100 µL of 3% sulfosalicylic acid, 200 µL glacial acetic acid, and 2.5% acid ninhydrin solution. The mixture was heated at 96 °C for 60 min and then cooled in an ice bath. The absorbance values were recorded at 520 nm. A standard calibration curve prepared with proline was performed and the results were expressed as mg g$^{-1}$ of FW.

Total Protein Content

The total protein content of the extract was quantified according to the method referred by Lowry et al. [32], using bovine serum albumin (BSA) as standard. Two solutions were prepared: solution A (50 mL of 2% sodium carbonate with 50 mL of 0.1 N NaOH) and solution B (10 mL of 1.56% copper sulphate with 10 mL of 2.37% sodium potassium tartrate). Finally, 2 mL of solution B was added to 100 mL solution A and designed as the working solution. Then, the 0.4 mL of sample was mixed with 4 mL of working solution and 0.4 mL of 1 N Folin–Ciocalteau. The reaction mixture was incubated in the dark, at room temperature, for 30 min. The absorbance was measured at 660 nm, and the protein content was calculated from a BSA standard calibration curve and expressed as mg g$^{-1}$ FW.

2.2.2. Antioxidant Enzyme Activities

Ascorbate Peroxidase Activity

The activity of ascorbate peroxidase (APX) was assayed according to Amako et al. [33]. The reaction mixture contained 50 mM phosphate buffer (pH = 7.0), 0.25 mM ascorbic acid, and 10 µL of enzyme extract. The reaction was initiated by adding 5 µL of 200 mM hydrogen peroxide (H$_2$O$_2$) to 185 µL of the reaction mixture. Absorbance was measured and monitored by oxidation of NADPH at 290 nm for 3 min. The APX activity was calculated using the molar extinction coefficient $\epsilon$ = 2.8 mM$^{-1}$.cm$^{-1}$, and the results were expressed as enzyme units (U) mg$^{-1}$ of protein.

Peroxidase Activity

The peroxidase (POX) activity was determined as oxidation of guaiacol according to the method of Maehly and Chance [34], with some modifications. The reaction mixture consisted of 200 µL of 8 mM guaiacol in 50 mM potassium phosphate buffer (pH = 7.0) and 50 µL of enzyme extract. The reaction was initiated by adding 50 µL of 1% H$_2$O$_2$ and enzymatic activity was detected by measuring the increase in absorbance at 430 nm for 5 min. POX activity was calculated using the extinction coefficient $\epsilon$ = 6.39 mM$^{-1}$.cm$^{-1}$ and expressed as enzyme units (U) mg$^{-1}$ of protein.

Catalase Activity

Catalase (CAT) activity was measured according to the method described by Aebi [35]. The reaction mixture consisted of 10 µL enzyme extract and 50 mM potassium phosphate buffer (pH = 7.0). The reaction was initiated by the addition of 100 µL of 3% H$_2$O$_2$. The decrease in H$_2$O$_2$ was monitored at 240 nm for 3 min. The activity of CAT was calculated using the extinction coefficient $\epsilon$ = 43.6 mM$^{-1}$.cm$^{-1}$ and expressed as enzyme units (U) mg$^{-1}$ of protein.

Glutathione-S-Transferase Activity

The activity of Glutathione-S-Transferase (GST) was assayed by the procedure of Habig et al. [36]. GST activity was measured by adding 10 µL enzyme extract, 50 mM potassium phosphate buffer (pH = 7.0), 1 mM of glutathione (GSH), and 1 mM of CDNB (1-chloro-2,4-dinitrobenzene). The absorbance was measured at 340 nm, immediately and
after 2 min. The enzyme activity of GST was calculated using the extinction coefficient \( \varepsilon = 9.6 \text{ mM}^{-1}\text{cm}^{-1} \) and expressed as enzyme units (U) mg\(^{-1}\) of protein.

**Superoxide Dismutase Activity**

The method described by Banowetz et al. [37] was used to measure the superoxide dismutase (SOD) activity. Briefly, reaction mixture (125 \( \mu \text{L} \)) contained 0.4 mM o-dianisidine, 0.5 mM DTPA, 26 \( \mu \text{M} \) riboflavin, 10 mM Piperazine-N,N′-bis (2-ethanesulfonic acid) (pH = 7.5), and 20 \( \mu \text{L} \) of enzyme extract. The absorbance at 450 nm was read initially and 30 min after the reaction had been irradiated with fluorescent light (120 W). The SOD activity was determined using a SOD standard calibration curve. The results were expressed as enzyme units (U) mg\(^{-1}\) of protein.

### 2.3. Soluble Sugars and Starch Content

Total soluble sugars (TSS) and starch content were determined using the anthrone method adopted by Ni et al. [38]. Approximately 40 mg of bud samples were previously ground to a fine powder in liquid nitrogen using a chilled mortar and pestle and homogenized with 2 mL pure acetone for 5 min. After centrifugation at 1100 rpm at 4 °C for 10 min, 80% ethanol was added to the pellets. The extract was incubated at 80 °C for 10 min in a water bath and again centrifuged at 1100 rpm at 4 °C for 10 min. The residues remaining in the tubes were stored for starch analysis. Regarding the total soluble sugar quantification, 100 \( \mu \text{L} \) supernatants were added to 500 \( \mu \text{L} \) of 0.2% anthrone reagent (Merck, EUA) and then placed at 100 °C for 10 min. The absorbance was measured at 630 nm, and total soluble sugar content was calculated using a glucose standard calibration curve and expressed as mg g\(^{-1}\) FW.

The starch content was determined on the solid fraction from the above extraction by heating the pellet in 1.1% HCl at 100 °C for 30 min. Afterwards, 100 \( \mu \text{L} \) of supernatant was added to 500 \( \mu \text{L} \) of 0.2% anthrone reagent and then placed at 100 °C for 10 min. The absorbance was measured at 630 nm; the starch content was calculated based on a glucose standard calibration curve, and the results were expressed as mg g\(^{-1}\) FW.

### 2.4. Gibberellic Acid Content

The endogenous gibberellic acid (GA\(_3\)) was assessed according to the method described by Ergün et al. [39] and Berrios [40]. Buds (60 mg) were previously ground to a fine powder in liquid nitrogen using a chilled mortar and pestle and then homogenized with 1 mL pure ethyl acetate and 200 \( \mu \text{L} \) pure HCl and stored at −20 °C for 24 h. The next day, the mixture was centrifuged at 5000 rpm for 15 min, and the supernatants were transferred to a clean tube. The same procedure was repeated for the pellet. The combined extract (1:1) was mixed with 3 mL of pure ethanol and 8 mL of HCl 3.75 M. The absorbance was measured at 254 nm, and the gibberellic acid was used as standard. The values were expressed as \( \mu \text{g} \) g\(^{-1}\) FW.

### 2.5. Statistical Analysis

Statistical analysis was performed using JMP version 7 statistical software (SAS Institute Inc., Cary, NC, USA). The effects of the site (vineyard), variety, and phenological stage on all biochemical parameters were evaluated using a three-way factorial ANOVA. A two-way ANOVA was also used to examine the effects of site and variety on the budburst percentage. All datasets satisfied the ANOVA assumptions—namely, homogeneity of variances and normality. When the differences were significant (\( p < 0.05 \)), Tukey’s post hoc test was carried out.

### 3. Results

#### 3.1. Percentage of Budburst

The results of the budburst percentage under forced conditions are presented in Table 1. Neither site (vineyard) nor variety had a significant effect (\( p > 0.05 \)) on budburst percentage.
However, the interaction effects of site and variety were significant \((p < 0.05)\) for this variable. The Fernão-Pires variety presented, in both vineyards, the higher number of budbursts, representing a budburst percentage of 70.0% and 69.4% in Celorico de Basto and Penafiel, respectively. In contrast, the percentage of budburst was significantly lower in the Loureiro variety, especially in Celorico de Basto, where it reached 46.7% of budburst.

**Table 1.** Mean values of budburst percentage of Alvarinho (AL), Fernão-Pires (FP), and Loureiro (LO) varieties in Celorico de Basto and Penafiel vineyards obtained by forced bursting. The mean values followed by different letters are significantly different at \(p < 0.05\) by Tukey’s HSD test. Different letters in the same column indicate significant differences. Site (S), variety (V), and interaction effects (S × V).

| Site (S)       | Variety (V) | Budburst (%) |
|----------------|-------------|--------------|
| Celorico de Basto | AL          | 63.3 ± 9.7 ab|
|                 | FP          | 70.0 ± 10.7 a|
|                 | LO          | 46.7 ± 8.7 b |
| Penafiel        | AL          | 58.9 ± 16.9 b|
|                 | FP          | 69.4 ± 11.8 a|
|                 | LO          | 55.3 ± 11.3 b|

\(p\)-value

| Site (S) | Variety (V) | Budburst (%) |
|----------|-------------|--------------|
| S        | ns          |              |
| V        | ns          |              |
| S × V    | *           |              |

Symbols 'ns' indicates non-significant differences \((p > 0.05)\), and * \(p < 0.05\).

### 3.2. Malondialdehyde and Proline Contents

The effect of site \((p < 0.001)\), variety \((p < 0.001)\), and the phenological stage \((p < 0.001)\), as well as the interactions of variety × phenological stage \((p < 0.05)\) and site × variety × phenological stage \((p < 0.05)\), were statistically significant on the MDA content (Table 2). Between varieties, Loureiro and Alvarinho exhibited higher MDA content, with values ranging from 0.0018 to 0.0023 µg g\(^{-1}\) and from 0.0018 to 0.0022 µg g\(^{-1}\), respectively (Figure 2a). In all varieties, the MDA content increased from dormancy to budburst and was more substantial in the Loureiro variety from Penafiel, with values ranging from 0.0019 to 0.0026 µg g\(^{-1}\). In turn, the Fernão-Pires variety presented the lowest increase (0.0018 µg g\(^{-1}\)) when compared with Alvarinho (0.0021 µg g\(^{-1}\)) and Loureiro varieties (0.0021 µg g\(^{-1}\)).

**Table 2.** Significance of site (S), variety (V), phenological stage (P) and interaction effects \((S × V; S × P; V × P; S × V × P)\) on MDA, proline, antioxidant enzymatic activity \((APX, POX, CAT, SOD, GST)\), nonstructural carbohydrates \((TSS and starch)\), and \(GA_3\), according to repeated-measures ANOVA. The values are significantly different at \(p < 0.05\) by Tukey’s HSD test.

|                        | S     | V     | P     | S × V | S × P | V × P | S × V × P |
|------------------------|-------|-------|-------|-------|-------|-------|-----------|
| MDA (µg g\(^{-1}\))    | <0.001| <0.001| <0.001| 0.423 | 0.295 | 0.010 | 0.001     |
| Proline (mg g\(^{-1}\))| <0.001| <0.001| <0.001| <0.001| 0.651 | <0.001| 0.023     |
| APX (µg mg\(^{-1}\) of protein) | <0.001| <0.001| <0.001| <0.001| 0.021 | <0.001| <0.001     |
| POX (µg mg\(^{-1}\) of protein) | <0.001| 0.008 | <0.001| <0.001| 0.029 | <0.001| <0.001     |
| CAT (µg mg\(^{-1}\) of protein) | 0.704 | 0.280 | <0.001| 0.009 | 0.002 | 0.584 | <0.001     |
| SOD (µg mg\(^{-1}\) of protein) | <0.001| <0.001| <0.001| <0.001| 0.375 | <0.001| <0.001     |
| GST (µg mg\(^{-1}\) of protein) | <0.001| <0.001| <0.001| <0.001| 0.763 | 0.500 |           |
| TSS (mg g\(^{-1}\))   | <0.001| <0.001| <0.001| <0.001| 0.013 | <0.001| 0.039     |
| Starch (mg g\(^{-1}\)) | <0.001| <0.001| <0.001| 0.001 | 0.716 | 0.009 | <0.001     |
| \(GA_3\) (µg g\(^{-1}\) FW) | 0.136 | <0.001| <0.001| <0.001| 0.013 | <0.001| 0.039     |
With respect to phenology, APX, POX, and SOD activities were higher at dormancy, while were also found. In general, the activity of the enzymes was higher in Celorico de Basto APX \((0.37 \text{ mg} \text{ g}^{-1})\) when compared with Fernão-Pires (0.37 mg g\(^{-1}\)) and Alvarinho (0.27 mg g\(^{-1}\)). Table 2. In addition, significant interactions between site and variety were observed for Alvarinho, Fernão-Pires, and Loureiro. In variety, phenological stage, and the interaction among these factors (Table 2). The results presented significantly higher APX, POX activities, followed by Loureiro and Alvarinho. In Appendix D different letters indicate significant differences between phenological stages \((p < 0.05)\).

A significant effect of site \((p < 0.001)\), variety \((p < 0.001)\), and phenological stage \((p < 0.001)\) on proline content was observed (Table 2). Regarding interactions, only the effect of site x phenological stage was not significant \((p > 0.05)\). The higher levels of this amino acid were recorded at budburst (Figure 2b). Except for Alvarinho, there was a significant increase in the proline content between dormancy and budburst, by 56–64\% in Fernão-Pires and 16–53\% in Loureiro, respectively (Figure 2). Between varieties, the proline content tended to be higher in Loureiro (0.51 mg g\(^{-1}\)) when compared with Fernão-Pires (0.37 mg g\(^{-1}\)) and Alvarinho (0.27 mg g\(^{-1}\)).

### 3.3. Enzyme Antioxidant Activity

The activities of almost all antioxidant enzymes were significantly influenced by site, variety, phenological stage, and the interaction among these factors (Table 2). The results revealed that the activities of APX, POX and SOD depend on phenological stage and variety (interaction effects of variety x phenological stage were all statistically significant, Table 2). In addition, significant interactions between site and variety were observed for APX \((p < 0.001)\), POX \((p < 0.001)\), SOD \((p < 0.001)\), GST \((p < 0.001)\), and CAT \((p < 0.001)\) activities. Furthermore, there were significant differences between varieties and phenological stages for APX \((p < 0.001)\), POX \((p < 0.001)\), and SOD \((p < 0.001)\) activities. Significant interactions among site, variety, and phenological stage \((p < 0.001)\) for enzyme activities were also found. In general, the activity of the enzymes was higher in Celorico de Basto than in Penafiel, except for CAT, whose activity remained similar in both sites. Fernão-Pires presented significantly higher APX, POX activities, followed by Loureiro and Alvarinho. In turn, Alvarinho showed the higher SOD activity, followed by Fernão-Pires and Loureiro. With respect to phenotype, APX, POX, and SOD activities were higher at dormancy, while CAT showed higher activity at budburst (Figure 3).

**Figure 2.** Malondialdehyde (MDA) content (a) and proline content (b) at dormancy and budburst for Alvarinho, Fernão-Pires, and Loureiro buds from Celorico de Basto and Penafiel vineyards. The columns represent the mean values and the bars the respective standard error of three replicates. Different letters indicate significant differences between phenological stages \((p < 0.05)\).
3.3. Enzyme Antioxidant Activity

The activities of almost all antioxidant enzymes were significantly influenced by site, variety (for site, variety, and interactions between these factors \( p < 0.001 \), phenological stage \( p < 0.001 \), and interactions between these factors \( p < 0.001 \), phenological stage \( p < 0.001 \), and interactions between these factors \( p < 0.001 \), respectively). The results also revealed that the CAT activity was close to zero in dormancy for all varieties and sites. The SOD activity was generally significantly higher in dormancy, compared with budburst. Through an analysis of Figure 3b, it is possible to verify that the Fernão-Pires variety from Celorico de Basto showed higher activity in dormant buds.

The results also revealed that the CAT activity was close to zero in dormancy for Alvarinho and Fernão-Pires (from Celorico de Basto and Penafiel sites). However, significantly higher enzyme activity was observed during budburst, particularly for Fernão-Pires from Celorico de Basto. The SOD activity was generally significantly higher in dormancy. However, Alvarinho from Celorico de Basto presented a higher activity of this enzyme at budburst. In addition, the GST activity was significantly higher at budburst, particularly for the Alvarinho and Loureiro varieties, both from Celorico de Basto (not shown), though with very low values (<0.001) for all varieties and sites.

3.4. Soluble Sugars and Starch

Soluble sugars (TSS) content was significantly influenced by site \( (p < 0.001) \), variety \( (p < 0.001) \), phenological stage \( (p < 0.001) \), and interactions between these factors \( (p < 0.001) \). In general, the Alvarinho variety exhibited a higher content of soluble sugars \( (6.18 \text{ mg g}^{-1} \text{ FW}) \), followed by Fernão-Pires \( (5.58 \text{ mg g}^{-1} \text{ FW}) \) and Loureiro \( (4.59 \text{ mg g}^{-1} \text{ FW}) \). In addition, the soluble sugars content was remarkably higher in Penafiel \( (5.83 \text{ mg g}^{-1} \text{ FW}) \) than in Celorico de Basto \( (5.07 \text{ mg g}^{-1} \text{ FW}) \), and at budburst \( (6.49 \text{ mg g}^{-1} \text{ FW}) \), compared with dormancy \( (4.42 \text{ mg g}^{-1} \text{ FW}) \) (Figure 4a). The only exceptions were Fernão-Pires (Celorico de Basto) and Alvarinho (Penafiel), which presented similar values in both phenological stages.

![Figure 3. Enzymatic activity of ascorbate peroxidase (APX) (a), peroxidase (POX) (b), catalase (CAT) (c), superoxide dismutase (SOD) (d) in two phenological stages (dormancy and budburst) in Alvarinho, Fernão-Pires, and Loureiro buds from Celorico de Basto and Penafiel sites. The columns represent the mean values and the bars the respective standard error of three replicates. Different letters indicate significant differences between phenological stages \( (p < 0.05) \).](image-url)
nho, Fernão-Pires, and Loureiro buds from Celorico de Basto and Penafiel sites. The columns represent the mean values and the bars the respective standard error of three replicates. Different letters indicate significant differences between phenological stages \((p < 0.05)\).

### 3.4. Soluble Sugars and Starch

Soluble sugars (TSS) content was significantly influenced by site \((p < 0.001)\), variety \((p < 0.001)\), phenological stage \((p < 0.001)\), and interactions between these factors (Table 2).

In general, the Alvarinho variety exhibited a higher content of soluble sugars \(6.18 \text{ mg g}^{-1} \text{FW}\), followed by Fernão-Pires \(5.58 \text{ mg g}^{-1} \text{FW}\) and Loureiro \(4.59 \text{ mg g}^{-1} \text{FW}\). In addition, the soluble sugars content was remarkably higher in Penafiel \(5.83 \text{ mg g}^{-1} \text{FW}\) than in Celorico de Basto \(5.07 \text{ mg g}^{-1} \text{FW}\), and at budburst \(6.49 \text{ mg g}^{-1} \text{FW}\), compared with dormancy \(4.42 \text{ mg g}^{-1} \text{FW}\) (Figure 4a). The only exceptions were Fernão-Pires (Celorico de Basto) and Alvarinho (Penafiel), which presented similar values in both phenological stages.

The starch content also varied between site \((p < 0.001)\), variety \((p < 0.001)\) and phenological stage \((p < 0.001)\) and interactions, with an exception for site \(\times\) phenological stage interaction (Table 2). The values recorded from Penafiel \(57.97 \text{ mg g}^{-1} \text{FW}\) were statistically higher, compared with Celorico de Basto \(37.83 \text{ mg g}^{-1} \text{FW}\), Figure 4b. Regarding varieties, the starch content was higher in Fernão-Pires \(58.6 \text{ mg g}^{-1} \text{FW}\) and lower in Alvarinho, with \(37.2 \text{ mg g}^{-1} \text{FW}\). There was a considerable decrease in the starch content between dormancy \(59.94 \text{ mg g}^{-1} \text{FW}\) and budburst \(35.86 \text{ mg g}^{-1} \text{FW}\), with the reduction (about 56\%) more pronounced for Fernão-Pires and Alvarinho varieties, both from Celorico de Basto.

![Figure 4. Total soluble sugar (TSS) (a) and starch content (b) at dormancy and budburst for Alvarinho, Fernão-Pires, and Loureiro buds from Celorico de Basto and Penafiel sites. The columns represent the mean values and the bars the respective standard error of three replicates. Different letters indicate significant differences between phenological stages \((p < 0.05)\).](image)

### 3.5. Gibberellic Acid Content

Significant differences between varieties \((p < 0.001)\) and phenological stages \((p < 0.001)\) were found in the gibberellic acid \((\text{GA}_3)\) concentration (Table 2). \(\text{GA}_3\) was significantly higher in Alvarinho \(1.14 \text{ mg g}^{-1} \text{FW}\) and lower in Fernão-Pires variety \(0.84 \text{ mg g}^{-1} \text{FW}\) (Figure 5). Significant differences between phenological stages \((p < 0.001)\) were also found, with the most significant increase in the \(\text{GA}_3\) content at budburst. A higher increment in \(\text{GA}_3\) content from dormancy to budburst was observed in the Loureiro variety from Celorico de Basto and Alvarinho and Loureiro from Penafiel, with an increase of 12\%, 11\%, and 17\%, respectively. There was, however, no significant difference in this hormone between the phenological stages for Alvarinho from Celorico de Basto.
Significant differences between varieties (p < 0.001) and phenological stages (p < 0.05). The three varieties showed different responses in terms of enzymatic activity regarding different sites and phenological stages (Table 2). Thus, APX, POX, and SOD activities were significantly higher during dormancy and subsequently decreased at the budburst, with different levels between varieties and vineyards. On the other hand, CAT activity was lower in the dormancy period and significantly increased at the budburst. These results are in agreement with studies developed in dormant buds of apple [47] and fig trees [15], where higher APX and POD activities and lower CAT activity during dormancy until bud break were observed. An increase in CAT activity occurred in the absence of SOD, indicating that SOD is not the primary enzyme for $\text{H}_2\text{O}_2$ formation, as noted by Abassi et al. [47].
A few studies have shown that oxidative stress can play an important role in bud dormancy break, as increased \( \text{H}_2\text{O}_2 \) levels in buds seem to act as a signalling molecule at this stage [10,14]. The increase in detoxification pathways and antioxidant defence—namely, CAT, SOD, APX enzymes, and proteins of the peroxidase family, in breaking dormancy attests that oxidative stress is an important part of the dormancy process [10]. Studies of dormancy-breaking agents’ application have provided more insight into how oxidative stress induces dormancy and budburst. It has been proposed that the application of these compounds inhibits CAT activity, which induces an increase in \( \text{H}_2\text{O}_2 \) content, consequently causing oxidative stress [2,44,48–50].

The results showed that despite the apparent inactivity of buds in the dormant stage, the enzymatic activity detected may indicate that, internally, a dormancy break mechanism may be occurring since buds were previously submitted to a period of low temperatures, to accumulate chilling hours. To our knowledge, there are no studies reporting values of enzyme antioxidant activity in buds of different grapevine varieties.

The lipid peroxidation of the membrane, evaluated by the formation of MDA (a secondary product of lipid peroxidation), can be considered an indicator of the cell damage caused by ROS [16,51]. In all varieties, lipid peroxidation, as indicated by MDA content, was higher at budburst, compared with dormancy. Alvarinho and Loureiro were the varieties with higher MDA values, particularly from the Penafiel site (Figure 2a).

Proline plays an important role in response to stress in plants since it contributes to cellular osmotic adjustment and removal of ROS [24]. A significant increase in proline content was observed during the budburst (Figure 2b). As noted, the effect of this increase was different between vineyards and varieties, with higher levels observed in the buds of the Loureiro and Fernão-Pires varieties from Penafiel. An increase in proline during break dormancy has been observed in perennial species, such as apple [44,52,53], Japanese apricot [27], and grapevine [2,54], treated with dormancy-breaking agents. These findings highlight the role of this amino acid, which may act as a compatible osmolyte and may have protected cellular structures against ROS; in addition, as a signalling molecule, it may have induced the events leading to break dormancy.

An increase in endogenous levels of \( \text{GA}_3 \) in budburst for three varieties was found (Figure 5). Previous studies report that gibberellins are essential regulators of dormancy and bud break, meeting downregulated at the induction of dormancy and upregulated during dormancy break and budburst [24]. For this reason, the increase observed may be related to vegetative growth processes, once a promoter of cell growth and elongation. Chengguo et al. [55] reported a rise in \( \text{GA}_3 \) content during dormancy break and budburst in sweet cherry buds, which may be related to the decrease in ABA content in buds, with environmental temperature as a regulatory factor. However, Zheng et al. [26] reported that gibberellins have distinct functions in grapevine buds, during and after the dormancy break. According to these authors, despite the positive role of gibberellins in natural dormancy break, an exogenous application of this hormone can have inhibitory effects on the budburst.

Regarding the contents of total soluble sugars and starch, significant changes between the dormant bud and budburst of the three varieties were found. There was an increase in the total soluble sugar concentration and a concomitant decrease in starch concentration between the two phenological stages, in all varieties. In dormant buds, starch concentration was higher than that of soluble sugars, a tendency that was manifestly reversed in the budburst (Figure 4, Table 2). Studies on apple [13,52], walnut [53], and fig trees [15] observed a reduction in the starch content and a simultaneous enhancement in the total soluble sugar during endodormancy. These findings are related to the carbohydrate metabolism that occurs between dormancy, dormancy break, and budburst, as a result of cold hardiness.

Carbohydrate reserves are the main source of energy for metabolic changes that occur during dormancy and budburst until the leaves acquire photosynthetic capacity [56–59]. In addition to these functions, starch and soluble sugars availability serve as freezing protectants, leading to an enhanced freezing tolerance during winter [53,60]. According to
Mohamed [41,54], during dormancy, part of the starch reserves is hydrolysed into soluble sugar, by increasing the activity of \( \alpha \)-amylase, an enzyme responsible for degrading starch and induced by low temperatures. Thus, a rise in the starch hydrolysis induces a soluble sugars content increase at this stage.

Low total soluble sugars and starch concentrations in the bud tissues are also one of the causes indicated for the incidence of bud necrosis and low budburst [61,62]. In these situations, the carbohydrate content may be insufficient to meet the energy needs required for metabolic and respiratory activities necessary to break dormancy and support plant vegetative growth [62,63].

5. Conclusions

The buds’ biochemical changes addressed in the present study are considered very important to understand the responses of grapevines, as regards the physiological and biochemical mechanisms that lead to dormancy and budburst. The comprehension of these mechanisms may contribute to improved vineyard management and reduction in wine production fluctuations over the years.

The present research indicated that MDA, proline, soluble sugars, and endogenous \( \text{GA}_3 \) contents were higher during budburst, compared with dormancy. The enzymatic activity, MDA, proline, and carbohydrate content also varied between varieties and sites, which may be related to the antioxidant metabolism of each variety and climatic conditions.

The results highlight the complexity of biochemical processes/mechanisms involved in the transition from dormancy to budburst. Though the study was mainly performed under controlled environmental conditions, the methodologies and results may be applied to different varieties and wine regions (particularly Iberian regions).

In future studies, enzymatic activities in buds of different varieties should be monitored throughout dormancy until budburst under field conditions, for understanding the effects of environmental factors on this mechanism. Additionally, there is a need to identify potential biochemical markers or indicators of oxidative stress in buds at different phenological stages.

Author Contributions: Conceptualisation, A.I.M., A.C.M., and E.A.B.; data curation, A.I.M. and A.A.; funding acquisition, A.C.M.; investigation, all authors.; methodology, all authors; project administration, A.C.M.; resources, all authors; supervision, A.C.M. and E.A.B.; writing—original draft preparation, A.I.M.; writing—review and editing, all authors. All authors have read and agreed to the published version of the manuscript.

Funding: This study was funded by the INTERAC project- “Integrated Research Environmental, Agro-Chain and Technology”, no. NORTE-01-0145-FEDER-000017, in its line of research entitled VitalityWine, co-financed by European Regional Development Fund (ERDF) through NORTE 2020. The study was also funded by the VITISHIDRI project—“Estratégias para a gestão do stress hídrico da vinha no Douro Superior” financially supported by the European Agricultural Fund for Rural Development and the Rural Development Programme 2020.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Acknowledgments: This work is supported by National Funds by FCT-Portuguese Foundation for Science and Technology, under the project UIDB/04033/2020. The authors acknowledge Aveleda S.A. for the collaboration in making available their vineyards for the study.

Conflicts of Interest: The authors declare no conflict of interest. The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, or in the decision to publish the results.
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