Search for Transcriptional and Metabolic Markers of Grape Pre-Ripening and Ripening and Insights into Specific Aroma Development in Three Portuguese Cultivars

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Abstract

Background: Grapes (Vitis species) are economically the most important fruit crop worldwide. However, the complexity of molecular and biochemical events that lead to ripening of berries as well as how aroma is developed are not fully understood.

Methodology/Principal Findings: In an attempt to identify the common mechanisms associated with the onset of ripening independently of the cultivar, grapes of Portuguese elite cultivars, Trincadeira, Aragónés, and Touriga Nacional, were studied. The mRNA expression profiles corresponding to veraison (EL35) and mature berries (EL36) were compared. Across the three varieties, 9.8% (2255) probesets corresponding to 1915 unigenes were robustly differentially expressed at EL 36 compared to EL 35. Eleven functional categories were represented in this differential gene set. Information on gene expression related to primary and secondary metabolism was verified by RT-qPCR analysis of selected candidate genes at four developmental stages (EL32, EL35, EL36 and EL 38). Gene expression data were integrated with metabolic profiling data from GC-EI-TOF/MS and headspace GC-EI-MS platforms.

Conclusions/Significance: Putative molecular and metabolic markers of grape pre-ripening and ripening related to primary and secondary metabolism were established and revealed a substantial developmental reprogramming of cellular metabolism. Altogether the results provide valuable new information on the main metabolic events leading to grape ripening. Furthermore, we provide first hints about how the development of a cultivar specific aroma is controlled at transcriptional level.

Introduction

Grapes (Vitis species) are economically the most important fruit crop worldwide due to their use in production of wine, grape juice, table grapes, among other. The consumption of grapes and wine presents numerous nutritional and health benefits for humans [1]. Grape seeds have significant content of phenolic compounds such as gallic acid, catechin and epicatechin, and a wide variety of proanthocyanidins which demonstrate significant cancer prevention potential [2]. Red wines contain more than 200 polyphenolic compounds that are thought to act as antioxidants. In particular, resveratrol exhibits cardioprotective effects and anticancer properties [2].

The Vitaceae family consists of approximately one thousand species grouped in seventeen genera. The most cultivated Vitis vinifera comprises up to 5000 true cultivars. In traditional vine areas, the production should present typicity that is dependent on grapevine variety among other factors. Therefore, vine improvement, specifically the regional development of typical wines, is greatly limited to the natural variability of available cultivars. In this respect, the less known Portuguese and Spanish cultivars offer plenty of choice to develop wines with different characteristics that

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may constitute a competitive advantage in a demanding global market.

The Portuguese wines are traditionally made by blending generally three or four varieties. Nevertheless, following a recent consumer trend the cultivars Touriga Nacional, Trincadeira and Aragónés have been used to produce high quality monovarietal red wines. Their exceptional characteristics such as the potential to produce highly aromatic wines have recently attracted the interest of vine growers and viticulturists abroad with the aim of introducing new competitive products into the market. Touriga Nacional is considered to be one of the most, if not the most important native variety in Portugal despite its tiny berries with a high skin to pulp ratio. Aragónés, also called Tinta Roriz in Northern Portugal or Tempranillo in Spain, offers high yields and is indispensable in the blend of a good Porto wine. Trincadeira is ideally suited to grow in hot, dry and very bright areas, but has varying (irregular) yield and is prone to infection with pathogenic moulds. However, in good years Trincadeira will produce exceptionally great wines.

The process of development and ripening of non-climacteric fruits such as grapes is not well known. Grape berry development consists of two successive sigmoidal growth periods separated by a lag phase; from anthesis to ripening grape development can be divided into three major phases [3] with more detailed descriptive designations, known as the modified E-L system, being used to define more precise growth stages over the entire grapevine lifecycle [4]. During the first growth period, the berry is formed, and biosynthesis of tannins and hydroxycinnamic acids occurs, as well as accumulation of organic acids, such as tartrate and malate. The onset of ripening, véraison, constitutes a transition phase during which growth declines and during which berries soften and color production is initiated, e.g. the anthocyanin accumulation in red grapes. Ripening, the final phase of cell expansion growth, is characterized by an increase in pH and accumulation of soluble sugars, anthocyanins and flavour-enhancing compounds.

Specific combinations of aromas, flavours, tannins, sugars and acids, create the unique “varietal character” of a ripe wine grape. The many chemical compounds contributing to flavour, i.e. the combination of taste and aroma, in wines are determined in the vineyard by interacting factors, such as the natural environment, vineyard management practices, and vine genotypes. Increased knowledge of the grape ripening process will contribute to maintain a sustainable production of high quality grapes in a changing environment, one major challenge for the viticulture in this century.

Recent reports characterized grape ripening of specific varieties, namely Shiraz [5], Cabernet Sauvignon [6], Pinot Noir [7], Barbera [8], Corvina [9], Muscat Hamberg [10] and Trincadeira [11]. More recently, a comparison was conducted between Cabernet Sauvignon and Norton grapes [12] regarding transcriptional regulation and flavonoid biosynthesis. These publications showed that activities of several metabolic pathways, including the tricarboxylic acid cycle, the shikimate pathway, and amino acid metabolism, shifted during fruit growth and ripening. Nevertheless, it is still difficult to ascertain which molecular and metabolic mechanisms are specific characteristics of a single variety and which can be considered general markers of ripening. It is particularly important to determine, how the nutritional quality for wine production purposes is achieved. In a previous NMR based study the quality of Trincadeira was associated with low phenolics content but higher sugars and organic acid levels while Touriga Nacional and Aragónés were characterized by high contents of phenolics and specific amino acids [13]. This NMR analysis enabled the identification of approximately thirty metabolites.

We now report the GC-MS based metabolome analysis of the same cultivars that complements previous results and importantly increased the coverage to a total of 121 volatile and soluble taste and aroma components. Furthermore, we integrated information from transcriptomic and from paired metabolomic analyses and now apply two essential functional genomics tools to characterize and to establish common mechanisms and candidate markers of grape ripening among the three most important Portuguese grapevine varieties. Next to the robust responses of all varieties we searched for the specific properties and, thus, gather insights into how a cultivar specific aroma is developed.

Results and Discussion

Phenotypic and metabolic characterization of grape berries

Grape berries were sampled at four developmental stages according to EL system [4] during 2008 growing season. The staging took into account the berry weight and the content of the main anthocyanins, organic acids and sugars (Fig.s 1, 2). The developmental stages investigated in this study were identified as EL 32 characterized by small hard green berries accumulating organic acids, EL 35 corresponding to véraison, EL 36 involving sugar and anthocyanin accumulation, and active growth due to cell enlargement and EL 38 corresponding to harvest time. The date of véraison was set at approximately 9 weeks post-flowering for each of the three varieties.

Metabolic profiling of grapes was carried out using a GC-TOF/MS platform that enabled the relative quantification of several classes of compounds such as fatty acids, phenylpropanoids and sugars among the polar compounds (Table 1, File S1). Additional metabolic profiling was achieved using a headspace GC-ELMS platform for the relative quantification of volatiles. In what concerns amino acids, few could be quantified since the conditions used favored the identification of secondary metabolites. Homoserine and pyrog glutamic acid which includes glutamine and glutamic acid were the amino acids identified (Table 1).

Specific chromatogram ion peaks were selected for relative quantification (based on peak height) of certain metabolites during ripening as shown in Fig. 2.

Regarding berry weight corresponding to ripe and harvest stages, Aragónés tends to present bigger berries, followed by Trincadeira and then Touriga Nacional. This implies that berries of Touriga Nacional contain a higher amount of skin per berry which may be related to the increased content in anthocyanins observed for this variety (Fig.1). The higher weight observed in Aragónés berries may also be partially responsible for higher content in fructose and glucose (Fig. 2) since these compounds accumulate preferentially in the pulp.

To confirm if these and other metabolites were present in significantly different amounts during ripening Kruskal-Wallis and Wilcoxon Rank sum tests were performed using ion peak responses (File S2). For the multiple comparisons the Benjamini & Hochberg correction was the most adequate. These tests showed that the majority of the metabolites (77, 69%) exhibited a different content either comparing among developmental stages or among varieties.

These ion peak responses were also used for Multivariate Data Analysis using the unsupervised method of Independent Principal Component analysis (ICA). A good discrimination was obtained between pre- véraison/ véraison and post- véraison stages for all the varieties and both analysis of volatiles and polar metabolites.
Figure 1. Fresh berry weight (g) and total anthocyanin content (absorbance at 530 nm per g of freeze dried material) of berries at developmental stages of EL 32, EL 35, EL 36 and EL 38. Bars represent standard deviation. doi:10.1371/journal.pone.0060422.g001

Figure 2. Relative quantification of malic acid, tartaric acid, fructose and glucose based on ion peak response. Malate and tartrate contents are higher at pre-véraison stages peaking at EL 32 whereas contents in fructose and glucose increase at post-véraison stages reaching maximal levels at EL 38. doi:10.1371/journal.pone.0060422.g002
Table 1. List of metabolites identified by GC-EI-TOF/MS and GC-EI-MS during ripening of the three varieties.

| Class         | Metabolite                                      |
|---------------|-------------------------------------------------|
| volatile      | (E)-2-Heptenal                                  |
| volatile      | (E)-2-Hexenal                                   |
| volatile      | (E)-2-Pentenal                                  |
| volatile      | (E,E)-2,4-Hexadien-1-al                         |
| volatile      | (Z)-3-Octen-1-ol                                |
| volatile      | [3-Hydroxymandelic acid, ethyl ester, di-TMS]   |
| volatile      | 2-Ethylfuran                                    |
| volatile      | 2-Pentylfuran                                   |
| volatile      | 3-Methylbutanal                                 |
| volatile      | 3-Methylbutanol                                 |
| volatile      | 6-Methyl-5-hepten-2-one                         |
| volatile      | Benzaldehyde                                    |
| volatile      | Decanal                                         |
| volatile      | Ethanol                                         |
| volatile      | Hexanal                                         |
| volatile      | Nonanal                                         |
| volatile      | Octanal                                         |
| volatile      | Ethyl Acetate                                   |
| volatile      | Heptanal                                        |
| volatile      | Limonene                                        |
| volatile      | Undecane                                        |
| acid          | Aconitic acid, cis-*                            |
| acid          | Benzoic acid                                    |
| acid          | Benzoic acid, 3,4-dihydroxy-                    |
| acid          | Butanoic acid, 2,4-dihydroxy-                  |
| acid          | Citric acid                                     |
| acid          | Fumaric acid                                    |
| acid          | Gallic acid                                     |
| acid          | Gluconic acid, 2-oxo-                          |
| acid          | Glutaric acid, 2-hydroxy-                       |
| acid          | Glutaric acid, 2-oxo-                          |
| acid          | Glutaric acid, 3-hydroxy-3-methyl-              |
| acid          | Glycolic acid                                   |
| acid          | Hexanoic acid, 2-ethyl-                         |
| acid          | Lactic acid                                     |
| acid          | Malic acid                                      |
| acid          | Malic acid, 2-methyl-                           |
| acid          | Pyruvic acid                                    |
| acid          | Shikimic acid                                   |
| acid          | Succinic acid                                   |
| acid          | Tartaric acid                                   |
| acid          | Vanillic acid                                   |
| amino acid    | Homoserine                                      |
| amino acid    | Pyroglutamic acid                               |
| fatty acid    | Docosanoic acid                                 |
| fatty acid    | Eicosanoic acid                                 |
| fatty acid    | Hexacosanoic acid                               |
| fatty acid    | Hexadecanoic acid                               |
| fatty acid    | Hexanoic acid                                   |
| fatty acid    | Nonanoic acid                                   |
| fatty acid    | Octacosanoic acid                               |
| fatty acid    | Octadecanenoic acid, n-                        |
| fatty acid    | Octadecanoic acid                               |
| fatty acid    | Octadecenoic acid, 9-(Z)-                      |
| fatty acid    | Octadecenoic acid, 9-(Z)-                      |
| fatty acid    | Tetraconanoic acid                              |
| fatty acid    | Tetraconanoic acid                              |
| fatty acid    | Triaconanoic acid                               |
| lipid         | Ampelopsin                                      |
| lipid         | Campesterol                                     |
| lipid         | Phytol                                          |
| lipid         | Sitosteryl, beta-                               |
| lipid         | Stigmaster-3-ol, (3-beta)-                     |
| lipid         | Tocopherol, alpha-                              |
| lipid         | Tocopherol, beta-                               |
| N-compound    | Ethanolamine                                     |
| N-compound    | Pyridine, 2-hydroxy-                            |
| N-compound    | Pyridine, 3-hydroxy-                            |
| N-compound    | Triethanolamine                                 |
| phenylpropanoid| Quercetin                                       |
| phenylpropanoid| Resveratrol, cis-                              |
| phenylpropanoid| Caffeic acid, trans-                           |
| phenylpropanoid| Catechin                                        |
| phenylpropanoid| Epigallocatechin                               |
| phenylpropanoid| Cinnamic acid, 4-hydroxy-, trans-              |
| phosphate     | Ethanolaminophosphate                           |
| phosphate     | Glycerol-3-phosphate                            |
| phosphate     | Phosphoric acid                                 |
| polyhydroxy acid| Dehydroascorbic acid dimer                    |
| polyhydroxy acid| Gluconic acid-1,5-lactone                    |
| polyhydroxy acid| Glyceric acid                                  |
| polyhydroxy acid| Lyxonic acid                                   |
| polyhydroxy acid| Threonic acid                                  |
| polyhydroxy acid| Threonic acid-1,4-lactone                     |
| polyol        | Erythritol                                      |
| polyol        | Glycerol                                        |
| sugar         | Fructose                                        |
| sugar         | Glucose                                         |
| sugar         | Sedoheptulose, 2,7-anhydro-, beta-             |
| sugar         | Sucrose                                         |

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(Fig. 3). However, the developmental stages of EL 32 and EL 35 (véraison) were only separated according to the polar metabolite fractions while samples corresponding to EL 36 and EL 38 clustered together in both types of analysis (though some degree of discrimination could be observed for polar metabolites). Regarding volatiles stages EL 35 and EL 32 were separated from EL 36 and EL 38 by two independent components accounting for 41.99% of variance. In what concerns polar metabolites the same
discrimination accounted for 58.94% of variance. Trincadeira tends to cluster more independently of the other varieties as it has been previously observed using a NMR platform for metabolic profiling [13].

The heat map showed that the metabolites have a differential accumulation during berry development and ripening (Fig. 4). Some alcohols and aldehydes such as ethanol, (E)-2-hexenal and heptanal reached the highest levels in ripe and harvest stage of fruits, and contributed to the formation of aroma. The same holds true for sugars and many fatty acids. Organic acids tend to decrease during ripening in all varieties, though exhibiting a high degree of variance among them. Phosphoric acid was an exception since it increased at harvest stage in all varieties. Phenylpropanoids and flavonoids may also decrease or increase during ripening.

**Microarray, functional categorization of Unigenes and RT-qPCR**

The mRNA expression profiles of two time points (EL35 and EL36) during 2008 season were compared using the Affymetrix GrapeGen® genome array containing 23096 probesets corresponding to 18711 unique sequences. Testing was performed using biological triplicates for each time point. After performing a Bayes t-statistics from the linear models for microarray data (limma) for differential expression analysis, P-values were corrected for multiple-testing using the Benjamini-Hochberg’s method [14]. The total number of probesets that were differentially expressed at least in one variety (fold change ≥ 1.5 and FDR < 0.05 or fold change ≤ −1.5 and FDR < 0.05) was 8696 corresponding to 37.65% of the total probesets represented in the chip (File S3). From these, a common gene set of 2255 probesets were differentially expressed in all the three studied varieties, whereas...
Figure 4. Heat map showing metabolite levels analyzed during berry ripening. Each square represents the average of the biological replicates.
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Analysis of gene expression and metabolite content during grape berry ripening

Carbohydrate metabolism. Berries start to accumulate after véraison the carbohydrates produced during photosynthesis and imported from the leaves.

Sucrose concentrations remained relatively constant with a slight increase towards harvest stage. This is in agreement with transcript abundance of a gene encoding an enzyme involved in sucrose biosynthesis that was higher at EL 36 when compared with EL35, namely sucrose phosphatase (VVTU21174_s_at, VIT_08s0032g00840) (Table 2).

A gene coding for a myb domain protein R1 transcription factor (VVTU2631_at, VIT_11s0001g09850) was up-regulated at EL 36 vs EL 35 in all varieties (File S3). This gene presents homology to a gene coding for an enzyme involved in sucrose catabolism of sucrose to fructose and glucose, decreased during ripening (Table 2, File S3) as observed for Pinot Noir [7] and Syrah [18].

Furthermore, the transcript abundance of two vacuolar invertases, Gin1 and Gin2 (VVTU3710_at, VIT_02s0154g00969, VVTU13187_at, VIT_16s0022g00670), which catalyze the catabolism of sucrose to fructose and glucose, decreased during ripening (Table 2, File S3) as observed for Pinot Noir [7] and Syrah [18].

Plastid glycolysis seems to be inhibited at the onset and following véraison as we suggested previously [11]. On the other hand, cytoplasmic glycolysis seems to be activated. In fact, genes coding for pyruvate kinase, cytosolic isozyme (VVTU18887_s_at, VIT_08s0007g04170) are up-regulated at EL 36 vs EL 35 and a gene coding for a cytosolic fructose-1,6-bisphosphatase is down-regulated (VVTU12977_s_at, VIT_18s0072g00770). Although different berry tissues may have different trends of glycolysis [8], cellular compartmentation should be taken into account [11]. This subject deserves further confirmation as a marker of ripening.

Nevertheless, the whole rate of glycolysis seems to decrease in ripe berries as suggested by the decrease in glyceric acid (File S2), a precursor of important biochemical intermediates in that process.

During ripening there is a major decrease in malate content as it was confirmed here by using GC-MS. The study of catabolism of this acid is very important for wine improvement due to its involvement in TCA cycle, gluconeogenesis, ethanol fermentation and for production of secondary metabolites (reviewed by [23]).

In the cytoplasm, malate can be produced from phosphoenolpyruvate produced in glycolysis through the activities of phosphoenolpyruvate carboxylase (PEPC) and malate dehydrogenase (MDH). One gene coding for a PEPC was down-regulated during ripening in Trincadeira berries at EL 35 and EL 36 [11]. However, genes putatively involved in synthesis of starch such as coding for starch synthase 1, chloroplast precursor (VVTU3341_at, VIT_16s0090g01780) and starch synthase (VVTU13257_s_at, VIT_02s0025g02790) were up-regulated at EL 36 vs EL 35 in all varieties. On the other hand, a gene coding for an isomaltase-type starch-debranching enzyme 1 (VVTU13947_s_at, VIT_11s0078g00310) is down-regulated. This enzyme is assumed to play a role in biosynthesis of starch [19].

The control of activity of starch synthase and starch degradation enzymes is complex in storage organs such as fruits. Different starch degradation pathways may be specific to early development and not active in late development [20].

We have observed a high up-regulation of a gene coding for galactosyl synthase (VVTU3450_at, VIT_05s0020g00330) at EL 36 vs EL 35 (Table 2). The biosynthesis of raffinose and stachyose occurs via sequential transfers of galactosyl units to sucrose. The galactosyl donor is galactosyl synthesized from UDP-galactose and myo-inositol, in a reaction catalyzed by galactosyl synthase. The biosynthesis of raffinose and stachyose was reported during ripening of olive, also a non-climacteric fruit [21]. On the other hand, we observed up-regulation at EL 36 vs EL 35 of a gene coding for 1-phosphatidylinositol-4-phosphate 5-kinase (VVTU9324_at, VIT_18s0001g03920), an enzyme that participates in inositol phosphate metabolism and phosphatidylinositol signalling. It should be noted the considerable higher fold change displayed by the gene coding for galactosyl synthase (VVTU3450_at, VIT_05s0020g00330) in Touriga Nacional (~105× compared to 40.81× in Aragones and 6.64× in Trincadeira). This may eventually imply a different composition in raffinose and stachyose for the three varieties. Raffinose is a minor carbohydrate in grape tissues but it has been reported that accumulation of raffinose in leaves of Vitis vinifera might be an early step of cold acclimation [22].

Interestingly, the sugar alcohol erythritol is more present in Aragones cultivar at harvest stage, and may contribute to the final wine taste (File S2).
### Table 2. Selection of genes differentially expressed during ripening of the three grape varieties (considering a fold change $\geq 1.5$ and FDR < 0.05 or fold change $\leq -1.5$ and FDR < 0.05).

| ProbeID          | Trincadeira 2008 | Touriga Nacional 2008 | Aragonés 2008 | Unique gene 12XID | Annotation                                                                                                                                 |
|------------------|------------------|-----------------------|--------------|------------------|------------------------------------------------------------------------------------------------------------------------------------------|
| VVTU1208_at      | −3.56            | −6.02                 | −3.1         | VIT_01s0011g02740 | Phosphoenolpyruvate carboxylase                                                                                                        |
| VVTU12977_s_at   | −2.59            | −2.13                 | −2.26        | VIT_18s0072g00770 | fructose-1,6-bisphosphatase, cytosolic                                                                                                    |
| VVTU13187_at     | −22.87           | −26.8                 | −33.59       | VIT_16s0022g00670 | vacuolar invertases, GIN2                                                                                                               |
| VVTU13257_s_at   | 5.18             | 2.54                  | 2.57         | VIT_02s0025g02790 | starch synthase                                                                                                                         |
| VVTU13333_s_at   | 1.83             | 1.55                  | 1.56         | VIT_16s0098g01200 | 6-phosphogluconolactonase                                                                                                               |
| VVTU13825_at     | −2               | 2.05                  | 2.26         | VIT_02s0025g01410 | Acyl-CoA synthetase long-chain member 2                                                                                                  |
| VVTU1391_at      | 2.9              | 4.14                  | 1.9          | VIT_00s0391g00070 | 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase                                                                                     |
| VVTU13947_s_at   | −3.9             | −2.6                  | −2.19        | VIT_11s0078g00310 | isoamylase-type starch-debranching enzyme 1                                                                                               |
| VVTU13950_s_at   | 9.22             | −20.03                | −5.66        | VIT_08s0007g07400 | Glutamate dehydrogenase 1                                                                                                               |
| VVTU18887_s_at   | 1.67             | 1.68                  | 2.09         | VIT_09s0002g00990 | Triacylglycerol lipase                                                                                                                   |
| VVTU1903_at      | −1.82            | −1.89                 | −1.67        | VIT_13s0019g05250 | Maltase dehydrogenase [NADP], chloroplast precursor (NADP-MDH)                                                                          |
| VVTU211174_s_at  | 1.58             | 2.01                  | 2.43         | VIT_08s0032g00840 | Sucrose-phosphatase                                                                                                                      |
| VVTU22296_s_at   | −2.5             | −2.34                 | −1.77        | VIT_08s0040g03320 | glutamate N-acetyltransferase                                                                                                             |
| VVTU24552_at     | −2.12            | −3.73                 | −2.06        | VIT_11s0016g00130 | omega-6 fatty acid desaturase, chloroplast (FAD6) (FADC)                                                                               |
| VVTU25722_at     | 2.84             | 2.15                  | 2.45         | VIT_18s0001g11910 | 1-acyl-sn-glycerol-3-phosphate acyltransferase 4                                                                                          |
| VVTU3450_at      | 6.64             | 104.84                | 40.81        | VIT_05s0020g00330 | galactinol synthase                                                                                                                     |
| VVTU3496_at      | −2.49            | −2.7                  | −3.21        | VIT_01s0146g00070 | 11-beta-hydroxysteroid dehydrogenase                                                                                                     |
| VVTU35404_at     | −1.89            | −2.21                 | −2.04        | VIT_03s0038g00450 | Squalene synthase                                                                                                                       |
| VVTU35129_s_at   | 10.64            | 11.55                 | 10.6         | VIT_16s0001g15460 | stearyl acyl carrier protein desaturase                                                                                                   |
| VVTU3541_at      | 2.45             | 2.62                  | 2.18         | VIT_16s0098g01780 | soluble starch synthase 1, chloroplast precursor                                                                                         |
| VVTU35625_s_at   | −2.5             | −1.98                 | −2.01        | VIT_19s0085g00880 | succinate-semialdehyde dehydrogenase (SSADH1)                                                                                             |
| VVTU3709_at      | 3.04             | 1.79                  | 3.29         | VIT_09s0002g06970 | Palmitoyl-monoacyl-sn-glycerol delta-7 desaturase, chloroplast                                                                         |
| VVTU3710_at      | −3.03            | −2.03                 | −5.65        | VIT_02s0154g00090 | vacuolar invertases, GIN1                                                                                                                |
| VVTU37457_s_at   | 2.36             | 3.2                   | 1.68         | VIT_01s0127g00260 | ATP-citrate synthase                                                                                                                     |
| VVTU3838_at      | 1.82             | 2.35                  | 1.61         | VIT_09s0018g01940 | pyruvate dehydrogenase E1 component alpha subunit                                                                                         |
| VVTU40835_s_at   | −2               | 2.58                  | 1.65         | VIT_05s0020g03080 | Acyl-CoA synthetase long-chain member 6                                                                                                   |
| VVTU4095_at      | −2.55            | −4.9                  | −2.4         | VIT_03s0088g01190 | malate dehydrogenase, glyoxysomal precursor                                                                                               |
| VVTU4210_at      | 1.83             | 6.23                  | 4.23         | VIT_14s0068g01760 | Alcohol dehydrogenase                                                                                                                    |
| VVTU5073_at      | −2.14            | −2.79                 | −3.29        | VIT_08s0105g00430 | omega-3 fatty acid desaturase, chloroplast precursor                                                                                       |
| VVTU5906_at      | 1.62             | 3.65                  | 3.15         | VIT_07s0031g02840 | Diacylglycerol kinase 2                                                                                                                  |
| VVTU6948_at      | 1.53             | 6.69                  | 3.19         | VIT_04s0008g06570 | chorismate mutase, cytosolic (CM2)                                                                                                        |
| VVTU8069_at      | −2.01            | −2.32                 | −2.18        | VIT_08s0007g05710 | L-Galactono-1,4-lactone dehydrogenase                                                                                                     |
| VVTU8356_at      | 5.11             | 10.04                 | 4.14         | VIT_07s0005g01240 | triacylglycerol lipase                                                                                                                   |
| VVTU9324_at      | 1.52             | 1.52                  | 2.35         | VIT_18s0001g03290 | 1-phosphatidylinositol-4-phosphate 5-kinase                                                                                               |
| VVTU9635_s_at    | −3.06            | −4                    | −2.9         | VIT_03s0038g02510 | glyoxylate reductase                                                                                                                     |
| VVTU9987_s_at    | −3.41            | −2.23                 | −2.84        | VIT_05s0020g04510 | GDP-mannose 3,5-epimerase 1                                                                                                              |
| **FLAVONOID AND STILBENE METABOLISMS AND AROMA DEVELOPMENT** |                 |                       |              |                  |                                                                                                                                          |
| VVTU11634_at     | 2.72             | 3.83                  | 3.03         | VIT_09s0002g06420 | Lactoylglutathione lyase                                                                                                                 |
| VVTU11927_at     | 2.7              | 10.66                 | 2.9          | VIT_05s0062g00980 | aldo/keto reductase AKR                                                                                                                  |
| VVTU12032_s_at   | 6.62             | 7.34                  | 2.89         | VIT_06s0004g08150 | trans-cinnamate 4-monooxygenase                                                                                                          |
| VVTU13018_s_at   | −2               | −1                    | 1.57         | VIT_15s0046g02300 | Beta-cyanoalanine synthase                                                                                                               |
Two of these genes (VVTU12208_at, VIT_01s0011g02740, VVTU4095_at, VIT_03s0088g01190) were more down-regulated in Touriga Nacional (Table 2) which may be related to an enhanced decrease in malic acid content of this variety at EL 38 (Fig. 2).

It is not surprising that a gene coding for a TONOPLAST DICARBOXYLATE TRANSPORTER with homology to a sodium-dicarboxylate cotransporter (VVTU17663_s_at, VIT_00s2188g00010), which is putatively involved in the transport of malate in the vacuole, was down-regulated at EL 36 vs EL 35 (File S3).

Organic acids such as malic and tartaric acids are well known for their contribution to wine taste. At veraison large amounts of malate released from the vacuole may be directly transported to the mitochondria and fed into an increased TCA cycle flux. The mitochondrial pyruvate dehydrogenase complex provides acetyl-CoA for the TCA cycle and NADH for oxidative phosphorylation. The alpha (VVTU3838_at, Table 2.

### Table 2. Cont.

| ProbeID       | Trincadeira 2008 | Touriga Nacional 2008 | Aragónés 2008 | Unique gene 12XID | Annotation                                                                 |
|---------------|------------------|-----------------------|---------------|-------------------|-----------------------------------------------------------------------------|
| VVTU13618_x_at| 2,1              | 2,31                  | 2,03          | VIT_16s0050g01580  | UDP-glucose: anthocyanidin 5,3-O-glucosyltransferase                       |
| VVTU13738_at  | -5,78            | -6,69                 | -6,79         | VIT_16s0050g0190   | Beta-carotene hydroxylase                                                  |
| VVTU13791_at  | -                | 2,57                  | -             | VIT_02s0025g00240  | Beta-carotene hydroxylase                                                  |
| VVTU13955_at  | 1,69             | 2,64                  | -             | VIT_14s0128g00780  | lipoxigenase                                                               |
| VVTU14620_at  | 1,79             | 6,38                  | 3,66          | VIT_18s0041g00740  | UDP-glucose: anthocyanidin 5,3-O-glucosyltransferase                       |
| VVTU14878_s_at| -                | 3,53                  | -             | VIT_04s0023g02200  | S-adenosyl-L-methionine:salicylic acid carboxyl methyltransferase          |
| VVTU15353_at  | 4,04             | 12,57                 | 2,7           | VIT_11s0065g00350  | trans-cinnamate 4-monooxygenase                                             |
| VVTU15628_s_at| 6,01             | 29,77                 | 4,29          | VIT_16s0100g00950  | Stilbene synthase 3                                                        |
| VVTU16103_at  | 1,54             | 3,68                  | 2,03          | VIT_18s0001g03470  | Flavonol synthase                                                          |
| VVTU16622_at  | 258,59           | 46,4                  | 16,14         | VIT_06s0225g00230  | allin lyase precursor                                                      |
| VVTU17111_s_at| -1,63            | -1,97                 | -1,71         | VIT_03s0180g00200  | Limonoid UDP-glucosyltransferase                                           |
| VVTU17555_s_at| -4,68            | -                     | -             | VIT_10s0035g03750  | 9-cis-epoxycarotenoid dioxygenase 2                                       |
| VVTU21329_at  | 9,32             | 3,43                  | 1,99          | VIT_16s0039g00880  | CYP89H3                                                                    |
| VVTU21725_at  | -                | 3,24                  | 2,79          | VIT_15s0046g03600  | (+)-neomenthol dehydrogenase                                               |
| VVTU2626_at   | -                | 2,57                  | -             | VIT_17s0000g05580  | isopiperitenol dehydrogenase                                               |
| VVTU34553_s_at| 6,36             | 33,68                 | 4,42          | VIT_16s0100g01030  | stilbene synthase [Vitis quinquangularis]                                   |
| VVTU34913_at  | 3,25             | 8,38                  | 1,81          | VIT_16s0100g00810  | stilbene synthase [Vitis vinifera]                                         |
| VVTU3533_s_at | 11,83            | 5,84                  | 2,04          | VIT_05s0077g01300  | Aldo-keto reductase                                                        |
| VVTU35884_at  | 2,73             | -                     | -             | VIT_06s0095g03010  | flavonoid 3',5'-hydroxylase [Vitis vinifera]                               |
| VVTU36515_at  | -4,79            | -                     | -2,73         | VIT_09s0271g00030  | myrcene synthase                                                          |
| VVTU36927_x_at| -14.1            | -3,01                 | -4,28         | VIT_12s0059g01790  | Caffeic acid O-methyltransferase                                            |
| VVTU37595_s_at| -                | -                     | -1,82         | VIT_12s0059g01060  | hydroperoxide lyase (HPL1)                                                 |
| VVTU39097_at  | -2.47            | -3,28                 | -3,79         | VIT_04s0079g00680  | phytorethene synthase, chloroplast precursor                               |
| VVTU4228_at   | -                | 1.7                   | -             | VIT_14s0030g01740  | Zeta-carotene desaturase ZDS1                                               |
| VVTU4697_at   | -6,77            | -10,64                | -9,48         | VIT_08s0007g01040  | aldo-keto reductase                                                        |
| VVTU52782_at  | 2,86             | 7,98                  | 6,35          | VIT_09s0018g01490  | Anthraniloyl-CoA: methanol anthraniloyl transferase                        |
| VVTU5372_s_at | -3,83            | -2,78                 | -2,53         | VIT_04s0023g02280  | S-adenosyl-L-methionine:salicylic acid carboxyl methyltransferase          |
| VVTU6355_at   | -3,24            | -4,38                 | -3,08         | VIT_07s0031g00620  | zeaxanthin epoxidase (ZEP) (ABA1)                                           |
| VVTU7133_s_at | -5,94            | -9,97                 | -4,21         | VIT_15s0046g03570  | (+)-neomenthol dehydrogenase                                               |
| VVTU8254_at   | -                | 2,79                  | -             | VIT_02s0087g00930  | 9-cis-epoxycarotenoid dioxygenase                                          |
| VVTU8862_at   | 7,64             | -                     | 7,85          | VIT_01s0111g05920  | S-adenosyl-L-methionine:salicylic acid carboxyl methyltransferase          |
| VVTU9837_s_at | -2,05            | -2,04                 | -2,4          | VIT_04s0008g03560  | lactoylglutathione lyase                                                   |
| VVTU9888_at   | -2,28            | -3,28                 | -2,1          | VIT_05s0049g01130  | aldo-keto reductase                                                        |

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VIT_09s0018g01940 subunit of the pyruvate dehydrogenase complex was induced after véraison, though this enzyme is known to be highly regulated post-translationally.

We observed an increase in the expression of a gene coding for ATP-citrate synthase (VVTU37457_s_at, VIT_01s0127g00260) at EL 36 vs EL 35 in all varieties though the highest level of expression was observed in Touriga Nacional. At EL 32 and EL 35 this variety tends to present more citric acid than Aragãoês and Trincadeira. Aragãoês showed elevated concentration of succinate whereas Trincadeira of fumarate (File S2).

While malic and tartaric acids showed the highest content in green berries in all varieties, citrate, succinate and fumarate showed the highest content in different developmental stages (green or véraison) depending on the variety. During ripening a transcript encoding a Succinic semialdehyde dehydrogenase (SSADH1; VVTU35625_s_at, VIT_19s0085g00880) putatively involved in succinate synthesis is down-regulated in all varieties as obtained by both microarray and RT qPCR analysis (Fig. 6). It was also noticed that several genes coding for enzymes of the TCA cycle such as aconitate hydratase and malate dehydrogenase showed different expression among the three varieties, being some genes up or down-regulated in one variety but not in other (File S3).

The increase in the rate of glycolysis due to an excess of sugars leads to an increase in pyruvate that may trigger aerobic fermentative metabolism [24].

Figure 5. PCA plot showing transcriptional discrimination of developmental stages and cultivars. The first (PC1) and the second (PC2) principal components are represented.
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Figure 6. Functional categories distribution in the common gene set of the 2255 modulated probesets and in the entire GrapeGen Chip®.
doi:10.1371/journal.pone.0060422.g006
In fact, the production of ethanol by pyruvate decarboxylase and alcohol dehydrogenase may occur in ripening fruit [23]. Here it was observed an increase in ethanol during ripening in all varieties though pyruvate levels showed no significant change. It was observed up-regulation at EL 36 vs EL 35 of genes coding for alcohol dehydrogenase and aldehyde dehydrogenase [7] which may be indicative of a shift to an aerobic fermentative metabolism during ripening [25].

We observed that a gene coding for an Alcohol dehydrogenase (VVTU4210_at, VIT_14s0068g01760) was up-regulated at EL 36 vs EL 35. This data may indicate that aerobic fermentation is occurring during ripening of all varieties. Other genes coding for alcohol dehydrogenases were up- or down-regulated in only one or two varieties (File S3). Interestingly, acetic acid, a product of ethanol metabolism which is known for giving a sour, pungent fatty and sweaty aroma to wines was previously detected in higher amounts at harvest stage (EL 38) in Trincadeira comparing to the other varieties Aragonês and Touriga Nacional [13].

There is a general tendency for an increase in dehydroascorbic acid dimer (an oxidized form of ascorbic acid) at EL 36 and EL 38. This is not surprising since ripening is characterized by increased oxidative stress [7,11]. Moreover, ascorbic acid present in abundant amounts in fruits undergoes rapid oxidation to dehydroascorbic acid [26] yielding reduced glutathione. Previously, the content in glutathione was shown to increase during grape ripening with 90% being reduced [27] in agreement with these results. Moreover, ascorbate was shown to decrease during ripening [11] together as indicated here with genes coding for GDP-mannose 3,5-epimerase 1 (VVTU9987_s_at, VIT_05s0020g04510), VTC2 (VITAMIN C DEFECTIVE), L-Galactono-1,4-lactone dehydrogenase (VVTU8069_at, VIT_08s0007g05710) enzymes putatively involved in ascorbate biosynthesis. At the same time there is a decrease in threonic acid (File S2) which is a polyhydroxy acid derived from L-ascorbic acid [28].

The pentose phosphate pathway is a process that generates NADPH and pentoses used in reductive biosynthesis reactions within cells (e.g. fatty acid synthesis). An increase in expression of a gene coding for 6-phosphogluconolactonase (VVTU13533_s_at, VIT_16s0098g01200) at EL 36 vs EL 35 in all varieties from the pentose phosphate pathway may be related to increase in fatty acids content as it will be discussed in paragraph on lipid

Figure 7. Real time qPCR validation of the expression profiles of ten genes in the three varieties under analysis. Data are reported as means ± SE of three technical and two-three biological replicates. Transcript levels were calculated using the standard curve method and normalized against grapevine actin gene (VVTU17999_s_at) used as reference control. Flavonol synthase (VVTU16103_at); Palmitoyl-monogalactosyldiacylglycerol delta-7 desaturase, chloroplast (VVTU3709_at); Succinate semialdehyde dehydrogenase (VVTU35625_s_at); Tocopherol cyclase (VVTU22626_s_at); L-Galactono-1,4-lactone dehydrogenase (VVTU8069_at); Limonoid UDP-glycosyltransferase (VVTU17111_s_at); Terpene synthase (VVTPS34); Cytochrome 450 monooxygenase CYP89H3 (VVTU1329_at) and Caffeic acid O-methyltransferase (VVTU36927_x_at). doi:10.1371/journal.pone.0060422.g007
metabolism. Interestingly, there is a big increase in β-sedoheptulose at EL 38 in agreement with major fatty acid accumulation.

In conclusion, the analysis of expression of genes involved in carbohydrate metabolism and contents in metabolites in all varieties indicates that véraison is a stage of active carbohydrate metabolism and transport. Ripening is characterized by a complex mechanism of starch breakdown with little sucrose accumulation but major accumulation of fructose and glucose. Raffinose and stachyose may play an important role in ripening of grapes. On the other hand, the pentose phosphate pathway seems to be involved in grape ripening together with aerobic fermentative metabolism. The TCA cycle may operate differently among the varieties since they were accumulating different contents in organic acids in particular in pre-véraison and véraison stages.

Lipid metabolism. The glyoxylate cycle plays a role in metabolism of fatty acids during seed germination; but was relevant to our analyses even though seeds have been removed from our samples. At EL 36 vs EL 35 there is down-regulation of a gene coding for glyoxylate reductase (VVTU9635_s_at, VIT_03s0038g02510) in all varieties in agreement with a decrease in glycolic acid (Tables 1, 2). The net result of the glyoxylate cycle is the production of glucose from fatty acids. However, fatty acid oxidation does not seem to be favored during ripening in pulp and skin but instead fatty acid biosynthesis appeared to be enhanced. Indeed, we have observed at EL 36 and EL 38 an increase in content in several saturated fatty acids (Table 1, File S2) as previously reported for another Vitis vinifera variety [29]. Moreover, genes coding for key enzymes of glyoxylate cycle such as malate synthase and isocitrate lyase were down-regulated though not in all varieties (File S3).

It was observed up-regulation at EL 36 vs EL 35 of two genes coding for Acyl-CoA synthetase long-chain member 2 and 6 (VVTU13825_at, VIT_02s0025g01410, VVTU40835_s_at, VIT_05s0020g03080) in Touriga Nacional and Aragones. The skin of berries from these varieties is thicker than from Trincadeira (Agudelo-Romero et al., unpublished). Fatty acids can be found in cells in activated form performed by enzymes acyl-CoA synthetases. The resultant acyl-CoAs are used in the biosynthesis of many cellular products such wax and cutin which are components of berry skins. After véraison grape berries continue to enlarge due to cell expansion driven by the increase of sugar content in the cell vacuole and a continuous flow of water; therefore a thick cuticle is necessary to avoid evapotranspiration.

The surface wax of the grape berry is chemically a mixture of long-chain alcohols, esters of such alcohols and fatty acids, free fatty acids, long-chain aldehydes, and hydrocarbons [30]. The GC-MS platform enabled the identification of several saturated fatty acids which in general increase at EL 36 and/or EL 38, namely octacosanoic, hexacosanoic and triacontanoic acids (Table 1, File S2). This increase was much more noticed in Touriga Nacional berries whereas docosanoic and eicosanoic acids showed increased levels at EL 38 in Aragones cultivar. This increase is not surprising since saturated long-chain fatty acids are major constituents of grape waxes that accumulate in ripe berries. The hexacosanoic acid or cerotic acid is a component of carnauba wax, an extremely good wax presenting several commercial applications. It is interesting to note that this fatty acid increased significantly in Touriga Nacional samples (File S2). The composition of their skin e.g wax density may influence the resistance of this variety to fungal attack [31] comparing to other varieties such as Trincadeira.

Lipid components in fruits though occurring in low amounts are important due to their nutritional value and due to their contribution to aroma and flavor. Fatty acids as n-hexadecanoic acid and 9,12-(Z, Z)-octadecadienoic acid (linoleic acid) have an important influence on the flavour of strawberry fruit [32].

A gene coding for a chloroplastidial omega 6-fatty acid desaturase (VVTU24552_at, VIT_11s0016g00130) was down-regulated at EL 36 vs EL 35 in all the varieties. This enzyme also known as FAD6 is the primary route of polysaturated fatty acids production in plants and converts oleic acid (18:1, a monounsaturated fatty acid) to linoleic acid [33] which provides a potential herbaceous aroma in ripe grapes [29]. In fact, we did not detect a significant decrease in linoleic acid. Also a gene coding for a chloroplastidial omega-3 fatty acid desaturase (VVTU5073_at, VIT_06s0105g00430) putatively involved in ω-3-linoleic acid biosynthesis is down-regulated at EL 36 vs EL 35 in all varieties.

A gene coding for a stearyl acyl carrier protein desaturase was up-regulated at EL 36 vs EL 35 in all varieties (VVTU35129_s_at, VIT_18s0001g15460). Stearoyl ACP-desaturase, commonly known as fatty acid biosynthesis 2 (FAB2), is a key enzyme that catalyzes the conversion of stearic (18:0) into oleic acid (18:1) [34]. Nevertheless we did not observe a significant decrease in stearic acid also known as octadecanoic acid. Other fatty acids such as octadecenoic and hexadecanoic also showed no significant differences throughout ripening (File S2). However, in 'Pedro Ximinez’ Vitis vinifera grapes hexadecanoic acid (palmitic acid) increased during ripening [29] highlighting metabolic specificities among varieties.

Regarding steroid biosynthesis we have observed down-regulation of genes coding for squalene synthase (VVTU35040_s_at, VIT_08s0038g00450), and 11β-hydroxysteroid dehydrogenase (VVTU3496_at, VIT_01s0146g00070) suggesting decreased steroid synthesis at EL 36. In fact, a transcript encoding 11β-hydroxysteroid dehydrogenase has been recently considered a biomarker of pre-veraison and veraison stages [9]. Campesterol decreased during ripening in all the varieties while stigmasteran-3-ol showed no significant variation in content.

In tomato, among all sterol lipids, the ratio of stigmastanol to β-sitosterol the two major sterols in tomato, increased 2.3-fold during ripening [35]. It has been reported an increase in the total sterol content of pericarp tissue during ripening of tomato fruit from mature green to red ripe stages [36]. In berries of Vitis vinifera at least for the sterols evaluated this does not seem to be the case. In addition, a gene involved in brassinosteroids biosynthesis from campesterol and coding for a steroid 5 alpha reductase DET2 (VVTU6606_at, VIT_08s0007g01760) was down-regulated at EL 36 vs EL 35 in all the varieties (File S3). This class of hormones plays an important role in the onset of grape ripening [37].

We observed an increase in α-tocopherol and β-tocopherol in all varieties during ripening but in particular in Touriga Nacional. The increase in tocopherol has been reported also for tomato and recently shown to be regulated by APETAL2a (AP2a) transcription factor [30]. It is interesting to note that a gene coding for AP2a (VVTU2494_at, VIT_07s0013g00220) was up-regulated at EL 36 vs EL 35 in all the varieties (File S3). It remains to be established whether this transcription factor is a major regulator of grape ripening as occurs in tomato [30].

The increase was mostly noticed for α-tocopherol which presents the highest anti-oxidant activity. Previously, we reported an increase in glutathione during ripening of Trincadeira grapes and suggested to be related to increased oxidative stress at the onset of ripening and during ripening as suggested by [7]. Interestingly, we observed up-regulation in Touriga Nacional and Trincadeira at EL 36 vs EL 35 of a gene coding for tocopherol cyclase (VVTU22626_s_at, VIT_04s0009g03600, File S3) which catalyzes the penultimate step of tocopherol synthesis. The
precursor of tocopherol, phytol also increases at EL 36 and EL 38 in Touriga Nacional berries.

Derivatives of fatty acids may be structural components of cell membranes, and/or used in triacylglycerol production [39]. Two genes coding for triacylglycerol lipases (VVTU15611_at, VIT_06s0002g00990, VVTU2356_at, VIT_07s0005g01240) were up-regulated at EL 36 vs EL 35, and may be involved in the release of fatty acids. The resultant product of glycerol-3-phosphate which decreases during ripening (File S2) may be metabolized for synthesis of esters of fatty acids for surface waxes [40,41]. It was noticed an up-regulation of a gene coding for chloroplastidial palmitoyl-monogalactosydialcyclyglycerol delta-7 desaturase in all the varieties up to EL 38 as revealed by RT qPCR data (VVTU3709_at, VIT_09s0002g06970, Fig. 7) putatively involved in unsaturated fatty acid biosynthesis for that purpose. Interestingly, glycerol which is also important for wines taste presented the highest concentration at EL 38 in Aragone’s (File S2).

On the other hand, it was noticed the up-regulation of a gene coding for a diacylglycerol kinase 2 (VVTU5906_at, VIT_07s0031g02840) which functions in the recycling of diacylglycerol, yielding phosphatidic acid, a compound that has emerged as an important lipid second messenger in plants, being involved in a wide range of biotic (e.g. pathogens) and abiotic (e.g. osmotic, temperature) stress responses [42]. These stresses are known to increase during grape ripening [11].

It was observed the up-regulation at EL 36 vs EL 35 in all varieties of a gene coding for a 1-acyl-sn-glycerol-3-phosphate acyltransferase (VVTU25722_at, VIT_18s0001g11910) which intervenes in the phospholipid biosynthetic process. This suggests that this class of lipids may increase during ripening as it occurs with Capsicum annuum, another non-climaticer fruit [43]. Indeed, the most abundant fraction of grape lipids has been reported to be the phospholipids [44].

The metabolism of lipids is very active during grape ripening due to the breakdown of plastid thylakoids, and differentiation of chloroplasts into non-photosynthetic chromoplasts. In these plastids fatty acid biosynthesis is enhanced for wax biosynthesis, as well as for aroma production which is due also to a specific fatty acid and correspondent esters combination. Steroid composition also varies among varieties but tends to decrease during ripening. On the contrary, tocopherol increases probably to cope with increased stress occurring during grape ripening.

**Amino acid metabolism.** Homoserine and pyroglutamatic acid which when monitored by the chosen GC-MS profiling approach also includes glutamine and glutamic acid were the amino acids identified (Table 1). In this context, the use of an NMR- based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). In this context, the use of an NMR-based platform enabled the quantification of many amino acids identified (Table 1). However, it is interesting to note that genes related to glutamate metabolism were down-regulated at EL 36 vs EL 35 in all the varieties and may eventually be considered as negative markers of ripening such as glutamate N-acetyltransferase (VVTU22296_s_at, VIT_08s0040g03320) and glutamate dehydrogenase 1 (VVTU13950_s_at, VIT_08s0007g07400).

**L-Homoserine is a precursor of methionine biosynthesis.** Interestingly, it has been shown that homoserine accumulation in the chloroplast triggers a novel form of downy mildew resistance that is independent of known defense signaling pathways [45]. Though the levels of this amino acid have remained the same throughout ripening in all varieties we have detected increased methionine levels towards harvest especially in Trincadeira [13].

S-adenosyl-L-methionine is a precursor of both ethylene and polynylene biosynthesis. Recently, we showed that during ripening there is an increase in polynylene catabolism in the three varieties [46].

Regarding the synthesis of aromatic amino acids such as phenylalanine which is a precursor of the phenolic compounds that accumulate during ripening, it was observed the up-regulation at EL 36 vs EL 35 of genes coding for a 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase (VVTU1391_at, VIT_06s0031g00070) and a cytosolic chorismate mutase (VVTU6948_at, VIT_04s0008g06570) putatively involved in shikimate metabolism and phenylalanine biosynthesis. Therefore, it is not surprising that shikimate content is decreased in berries at ripe and harvest stages (File S2) though it was substantially higher at EL 32 and EL 35 in Aragone’s and Touriga Nacional cultivars than in Trincadeira. Those varieties are known to accumulate more phenolic compounds [13]. Also different genes coding for phenylalanine ammonia lyase were up-regulated at EL 36 vs EL 35 though not in all varieties (File S3).

From the above it can be suggested that the metabolites of glutamate, methionine and phenylalanine play a major role in grape ripening.

**Flavonoid and stilbene metabolism.** Genes coding for enzymes acting on flavonoids, stilbenes, and anthocyanins synthesis were noticed to be induced during grape ripening [7,11]. Our results confirm these for all the varieties and establish these genes as candidate markers of grape ripening. In addition, many genes coding for transcriptions factors of the MYB family were up-regulated at EL 36 vs EL 35 in all the varieties and may be involved in the regulation of the flavonoid pathway as other previously characterized MYB transcription factors [12]. These are the case of genes coding for myb domains proteins 73, 14, R1 and 94 (File S3).

A gene coding for a flavonol synthase (VVTU16103_at, VIT_16s0001g03470) was up-regulated at EL 36 vs EL 35. This enzyme is responsible for the conversion of dihydroflavonol to flavonols which are important co-pigments that stabilize anthocyanins in wine. In the skin of Shiraz berries, VvFLS1 expression is highest between flowering and fruit set, then Declines to veraison, and increases again during ripening coincident with accumulation of flavonols per berry [47]. The same seems to be the case of this gene coding for flavonol synthase at least in Touriga Nacional (Fig. 7).

However, in this work it was detected a decrease in quercetin following EL 32 in all varieties including Touriga Nacional (Fig. 0) in agreement with a previous identified decrease in quercetin glycoside using NMR [13]. Ampelopsin or dihydromyricetin is a flavanone presenting anti-cancer properties and derivatives from the flavonol myricetin [48]. Interestingly, it was detected at higher concentrations in ripe berries (EL 36) (Fig. 9) and mainly in Aragone’s. This fact highlights the use of GC-MS as a platform to investigate the medicinal potential of samples.

A reduction in tannin concentration from pre-veraison to harvest has been well documented [49]. The tannin catechin decreases during ripening in all varieties as shown in Fig. 8 for Touriga Nacional. However, it should be noted that Trincadeira presents much lower levels of catechin at EL 32 and EL 35 than the other varieties. Interestingly, epigallocatechin was higher in Aragone’s at harvest stage and may contribute to higher astringency of the wine. This compounds is a precursor of epigallocatechin-3-gallate which is known for the anti-oxidant properties [50]. The observed
decrease in gallic acid during ripening is expected since this compound makes part of tannins such as epigallocatechin.

It is clear the overall flavonoid profile of grapes changes significantly over time [51] and includes a significant reduction in catechin and gallic acid.

One gene coding for an UDP-glucose: anthocyanidin 5,3-O-glucosyltransferase (VVTU14620_at, VIT_18s0041g00740) was up-regulated at EL36 vs EL 35. Another gene coding for this enzyme was down-regulated (VVTU13618_x_at, VIT_16s0050g01580) at EL 36 when compared to EL 35 but was up-regulated at EL 35 when compared to EL 32 at least for Trincaideira cultivar [11]. This suggests that the pattern of anthocyanins may change during berry ripening as recently observed for Corvina berries [51].

These authors showed a rapid accumulation of nonacylated anthocyanins and flavones during veraison, concomitant with a sharp decline in the levels of aromatic organic acids and procyanidins. This could be due to the use of the organic acid pool to provide precursors for the synthesis of other molecules. For example, both coumaric acid and caffeic acid could in principle be used (as coumaryl-CoA and caffeoyl-CoA, respectively) as substrates for the production of chalcones by chalcone synthase. The resulting chalcones could then be used for the synthesis of flavones and anthocyanins [51]. Interestingly, we have detected a decrease in caffeic acid in ripe berries of all the varieties being no longer detected at harvest stage. Touriga Nacional was the variety accumulating more caffeic acid at EL 32 and shown to present more anthocyanins at EL 36 and EL 38 (Fig. 1). The same holds true for trans-4-hydroxycinnamic acid or p-coumaric acid (Fig. 8), which is a precursor of flavonoids, stilbenes and coumarins [52].

Two genes coding for trans-cinnamate 4-monooxygenase (VVTU15353_at, VIT_11s0065g00350, VVTU12032_s_at, VIT_06s0004g08150) were more expressed at EL 36 vs EL 35 in Touriga Nacional (Fig. 8). Previously, and through an NMR platform it was established that the green and veraison stages (EL 32, EL 35) of Touriga Nacional presented more hydroxycinnamic acid derivatives such as caffic acid than Aragoneš and Trincaideira [13] and that was recently related to increased anti-tumoral properties of that cultivar [53].

One gene coding for flavonoid 3,5-hydroxylase (VVTU35884_at, VIT_06s0009g03010) was up-regulated at EL 36 vs EL 35 but only in Trincaideira (Table 2, File S3). This enzyme catalyzes the hydroxylation of the B-ring of flavonoids and can act on flavonols, anthocyanins, and proanthocyanidins. Bogs and co-workers [54] showed that in berry skin, expression of these genes was low at the onset of ripening but increased after veraison.
concomitant with the accumulation of 3′- and 3′,5′-hydroxylated anthocyanins. The hydroxylation pattern of the B-ring is an important determinant of flavonoids coloration, stability, and antioxidant capacity [55].

Regarding stilbenes, their accumulation during ripening of grape berries is well known [2]. It is interesting to note, however, that resveratrol or trans-5, 4′,5′-trihydroxystilbene is present at higher concentrations in Touriga Nacional berries at ripe (Fig. 8) and harvest stages in agreement with a higher fold change detected at EL 36 vs EL 35 in this variety for several genes coding for stilbenes synthases (VVTU34913_at, VIT_16s0100g00810, VVTU15628_s_at, VIT_16s0100g00950, VVTU34553_s_at, VIT_16s0100g01090).

Phenylpropanoid metabolism plays a major role in grape ripening changing the content in hydroxycinnamic acid derivatives, flavonoids, tannins, anthocyanins, and stilbenes from green to harvest stage of grapes and presenting cultivar specific variation with a tendency of Touriga Nacional to present more phenolics.

### Aroma development

Monoterpenes, C15-norisoprenoids, and some benzenoid compounds are the most important grape aroma substances present in the pulp and skin of berries in both free and glycoside forms. Their specific profiles depend mainly on the variety though environmental cues may have impact (Flamini and Traldi, 2010).

The largest class of plant volatiles is derived from isoprenoid pathways [56]. The mevalonate pathway is a source of isoprene units for the biosynthesis of cytokeins and brassinosteroids. In plants, both the cytosolic mevalonate and the plastidic mevalonate isopentenyl pyrophosphate and its isomer dimethylallyl pyrophosphate. These two precursors give rise to volatile and non-volatile terpenoid compounds such as monoterpenes (C10), sesquiterpenes (C15) diterpenes (C20), triterpenes (C30), carotenoids, sterols and phytols. In this respect, it should be noted that Touriga Nacional presented a higher quantity of phytol at EL 36 and EL 38 than Aragoneś and Trincadeira.

Variety, terroir, sunlight exposure, soil water retention capacity as well as ripening stage affect carotenoid concentration in the grapes which is known to decrease following veraison [57]. So it is not surprising that genes coding for a chloroplast precursor of phytoene synthase (VVTU39097_at, VIT_04s0079g00680), for a zeaxanthin epoxidase (VVTU6355_at, VIT_07s0031g00620) and for a Beta-carotene hydroxylase (VVTU13738_at, VIT_16s0050g01090) are down-regulated at EL 36 vs EL 35 in all varieties. However, it is interesting to note that in Touriga Nacional we found up-regulation at EL 36 vs EL 35 of genes coding for a Zeta-carotene desaturase ZDS1 (VVTU4228_at, VIT_14s0030g01740) and a Beta-carotene hydroxylase (VVTU13791_at, VIT_02s0025g00240) involved in carotenoid biosynthesis as well as a gene coding for phytoene synthase (VVTU9210_a, VIT_13s0019g01270) that was up-regulated in Trincadeira.

Volatile terpenes may derive from the cleavage of carotenoids by carotenoid cleavage dioxygenases [56]. A gene coding for a 9-cis-epoxy-carotenoid dioxygenase 2 (isoenzyme carotenoid cleavage dioxygenase 1; VVTU8254_at, VIT_02s0067g00930) was up-regulated at EL36 vs EL 35 in Touriga Nacional whereas another (VVTU17555_s_at, VIT_10s0003g03750) was down-regulated in Trincadeira. Some flavor volatiles peak at pre-veraison or at the ripening stages and others at ripe or harvest stage and this may be also dependent on the variety. The volatiles (E)-2-Pentenal, (Z)-3-Octen-1-ol were higher at EL32 while (E)-2-Heptenal was more present at both pre-veraison or veraison stages. The volatiles ethanol, hexanal, decanal and nonanal increased their content at ripe and harvest stages while (E)-2-Hexenal peaked at EL 36. The content in the putatively identified undecane peaked at EL 35 or EL 36 depending on the variety while heptanal at EL 36 and limonene at EL 38.

Carotenoid cleavage dioxygenases have been reported to generate 6-Methyl-5-hepten-2-one from lycopene [58]. This compound is an important flavor volatile in tomato [59] and tends to be higher in Trincadeira at EL 38 comparing to the other varieties. Different aroma profiles may also be related to the fact that a gene coding for myrcene synthase (VVTU36515_at, VIT_06s0271g00030) was down-regulated at EL 36 vs EL 35 in Trincadeira and Aragoneś but not in Touriga Nacional.

Terpenes which are precursors for important aroma compounds start to accumulate at veraison [60]. A gene coding for a (E)-isopentenyl dehydrogenase (VVTU2626_at, VIT_17s0000g05580) putatively involved in monoterpene biosynthesis was up-regulated at EL 36 vs EL 35 but only in Touriga Nacional. On the other hand, a gene coding for (E)-isoterminal dehydrogenase (VVTU7133_s_at, VIT_15s0046g03570) putatively involved in menthol biosynthesis, a monoterpene, was down-regulated in all the varieties but another one (VVTU21725_at, VIT_15s0046g03600) was up-regulated at EL 36 vs EL 35 only Touriga Nacional and Aragoneś. From the above it is not surprising that wines made from Touriga Nacional berries are recognized as being highly aromatic.

Furthermore, the putatively identified cyclic terpene limonene that provides a lemon-like fresh aroma reached the highest concentrations at harvest stage (EL 38) in Trincadeira and in particular in Touriga Nacional. In fact, Touriga Nacional wines are know to have an aroma of orange tree flower. Higher-quality Touriga Nacional wines are characterized by a fruity-citric aroma described as sweet and fresh citrus evoking the bergamot fruit (Citrus bergamia) [61]. In accordance, we have noticed in Touriga Nacional and by real time qPCR a very significant increase at EL 38 (Fig.7) in the expression of a gene coding for a terpene synthase which previous functional characterization showed that produces mainly (E)-α-bergamotene (VvTPS10; [62]). On the other hand, a terpene synthase that produces mainly (E)-b-bocimene (VvTPS34,[62]) was up-regulated at EL 38 in all the varieties but mainly in Aragoneś highlighting aroma cultivar specificities.

We have observed down-regulation at EL 36 vs EL 35 of a gene coding for a limonoid UDP-glucosyltransferase in all varieties (VVTU17111_s_at, VIT_03s0180g00200 Table 2) putatively involved in limonoid glucoside synthesis from limonin [63]. In particular in Aragoneś a considerable decrease in expression was noticed from EL 32 to EL 35 as shown by RT qPCR (Fig. 7). Limonoids are highly oxygenated triterpenes and one of the components responsible for the bitterness in citrus fruit [63]. The lower synthesis of their glucosides may be related to increased citric aroma. This gene was also shown to be induced during early withering in grape [64].

One gene coding for a lactoylglutathione lyase (VVTU18134_at, VIT_09s0002g06420) was up-regulated whereas another one (VVTU9937_s_at, VIT_04s0008g03560) was down-regulated at EL 36 vs EL 35 in all varieties. The same holds true for four genes coding for aldo-keto reductase (VVTU4697_at, VIT_08s0079g01040, VVTU98888_at, VIT_05s0049g01130, VVTU11927_at, VIT_05s0062g00980, VVTU3533_s_at, VIT_05s0077g01300). Lactoylglutathione lyase or glyoxalase I is involved in the detoxification of methyglyoxal together with aldo-keto reductase. Dicarbonyl compounds, like methyglyoxal, are
Genes coding for carboxylesterases were up or down-regulated at EL 36 vs EL 35 in only some varieties (File S3). These enzymes are involved in hydrolysis of esters into acids and alcohols and may contribute to flavor development.

Interestingly, anthraniloyl-CoA: methanol anthraniloyl transferase putatively involved in the biosynthesis of fruit esters responsible for aroma was up-regulated from EL 35 to EL 36 in various varieties (VVTU53632_at, VIT_09b01060g01400). It is difficult to establish the activity of the enzyme coded by this gene since it shows strong homology with other acyltransferases responsible for synthesis of flavor compounds but it may be involved in the synthesis of the volatile methyl anthranilate [77].

S-adenosyl-L-methionine-dependent methyltransferases catalyze methylation of several molecules, and are key enzymes in phenylpropanoid, flavonoid and many other metabolic pathways. Enzymatic methylation of hydroxyl and carboxyl moieties is catalyzed by O-methyltransferases which include SAM: salicylic acid carboxy methyltransferases, SAM: jasmonic acid carboxyl methyltransferases and SAM: benzoic acid carboxyl methyltransferases, among others [78]. A gene coding for a S-adenosyl-L-methionine: salicylic acid carboxyl methyltransferase (VVTU5372_s_at, VIT_04s0029g02280) was down-regulated at EL36 vs EL 35, and may play an important role in specific aroma production at earlier stages of berry development [78]. Other two genes, however, were expressed either in Touriga Nacional (VVTU14878_s_at, VIT_04s0023g02900) or in Aragone ˆs and Trincadeira (VVTU8892_at, VIT_01s0011g05920). A gene coding for a caffeic acid methyltransferase (VVTU36927_x_at, VIT_12s0059g01790) putatively involved in aroma production was also down-regulated at EL 36 vs EL 35 as it was already referred for a decrease in the compound caffeic acid. RT qPCR data showed that this gene presented a major decrease from EL 32 to EL 35 in particular in Aragone ˆs cultivar (Fig. 7).

Other methyltransferases such as O-methyltransferases may be involved in the synthesis of vanillin [79] which precursor is vanillic acid. The presence of this compound was only identified in grapes at EL 36 and EL 38 stages of ripening and shown to be present at higher amounts in Touriga Nacional, highlighting the known aromatic potential of these grapes. In fact, ethyl vanilllate was shown to be present in clonal red wines from Touriga Nacional (Falco da Costa, personal communication). It should be pointed out that methyltransferases constitute a protein family in which little correlation is observed between the level of sequence similarity among enzymes and the structural similarity of their substrates so their activities should be discussed with caution [56].

Cytochrome P450 are also involved in the formation of plant volatiles and in a vast array of biosynthetic pathways in secondary and primary metabolism [56]. A gene coding a CYP69H3 (VVTU21329_at, VIT_16s0039g00880) was up-regulated at EL 36 vs EL 35 in all varieties in particular in Trincadeira while the expression values at EL 38 were similar among the varieties (Fig. 7). Other genes coding for CYPs were up-regulated in one or two varieties and may eventually be involved in aroma development (File S3).

It was noticed the up-regulation of a gene coding for alliin lyase precursor (VVTU16622_at, VIT_08b0025g00230) in all varieties, especially in Trincadeira. Alliin lyase (alliinase) gives rise to volatile sulphur compounds from S-alk(en)yl cysteine sulphoxide flavor precursors derived from cysteine and glutathione [80]. This latter compound was shown to increase during ripening of Trincadeira grapes [11]. Moreover, glutathione and trans-2-hexenal intervene in the biosynthesis of S-(3-hexan-1-ol)-glutathione which is a flavor precursor in wines [81]. In fact, most of the volatile sulphur compounds that contribute to varietal character are released
during must fermentation from specific aroma/flavor precursors present in the grapes [82]

The varietal aroma is built on different classes of compounds that may be more present in one variety independently of the terroir. Nevertheless at veraison aromas and flavors begin to develop in all varieties and increase their complexity during ripening. For all the varieties studied pre-veraison and veraison stages seem to be enriched in carotenoids, benzaldehyde and caffeic acid and post veraison stages in volatile terpenes and sulfur compounds, C6 alcohols and aldehydes, and vannilic acid.

Conclusions

This work performed a comprehensive analysis and interpretation of the transcriptome and metabolome of pre-ripe and ripe grapes of three Portuguese cultivars. The combined analysis of transcripts and metabolites contributed to the establishment of putative markers of pre-ripening and ripening involved in carbohydrate, lipid, amino acid and phenylpropanoid metabolisms (Fig. 9). The decrease in metabolites of the TCA cycle during ripening may be due to their use for increased amino acid biosynthesis as previously revealed by NMR studies with the exception being glutamate [11]. On the other hand, shikimate synthesized from the precursor phosphoenolpyruvate may be consumed for the synthesis of phenolics whereas pyruvate is likely to be used for fatty acid biosynthesis, branched chain aminoacids and isoprenoids.

Good correlations were found for the content of campesterol and the genes coding for enzymes involved in steroid biosynthesis. The same holds true for sucrose and a gene coding for sucrose phosphatase; for ascorbate and a gene coding for L-Galactonol-1,4-lactone dehydrogenase; for glycolic acid and a gene coding for glyoxalate reductase; for resveratrol and genes coding for stilbenes synthase, among others. We have also validated a previous identified biomarker of pre-ripe berries which is a gene coding for 11β-hydroxysteroid dehydrogenase. These biomarkers can be further validated in the future for other cultivars and over different seasons to exclude any potential climate or terroir effect. The candidate molecular biomarkers presented here were already validated for Trincadeira during 2007 season (GEO accession number GSE28779).

On the other hand, the use of approaches enabling both gene expression and metabolites content gave strong insights into how aroma is developed during ripening, and also in to those features which are cultivar specific. The differential accumulation of organic acids among the three varieties may result in a different pool of phenolic compounds and amino acids. In addition, several genes coding for enzymes of either the phenylpropanoid or the terpenoid pathway were differently expressed among the three varieties and may reflect the different content in volatiles. On the other hand, this differential gene expression may lead to the synthesis of glycosides with aromatic potential that were not addressed in this work but certainly deserve attention.

Altogether the data discussed here contributed for a better understanding of the metabolic pathways involved in berry ripening through establishment of candidate molecular and metabolic markers that may define key switches in cellular reprogramming. It was also a contribution for the understanding of the complex process of flavor development that characterizes a specific variety.

Materials and Methods

Sample collection

Four biological replicates (each including 80-100 berries from 8-10 plants) were collected around 10 a.m. in 2008 at Plansels vines located in Montemor-o-Novo (Southern Portugal). The sampling was carried out with permission from the owner of Plansel, a private company. Samples from three important Portuguese cultivars (Aragoûes, Touriga Nacional and Trincadeira, Figure S1) corresponding to the developmental stages of EL 32, 35, 36, and 38 (EL refers to the modified Eichhorn and Lorenz
developmental scale as described by [4]) were immediately frozen in liquid nitrogen, transported to the lab in dry ice and kept at -80°C. Each biological replicate contained berries from a single row of plants, and from the sunny and shady sides of the plants that were pooled to form mixed samples. Rows distant 3 to 10 m were used. Plants from the three varieties were growing in the vineyard 15–30 m apart (Figure S2). The experiments were conducted on healthy grapes presenting no visible signs of fungal attack or other microbiological alteration. Meteorological conditions are shown in File S4.

RNA extraction

RNA extraction from pulp and skin of berries was carried out as previously described [11]. A DNase treatment was performed according to the suppliers’ instructions (Invitrogen, San Diego, CA, USA). Samples were then extracted in phenol/chloriform/isoamylalcohol (75:24:1,v/v/v), precipitated with sodium acetate and ethanol, washed in 70% ethanol and dissolved in water. RNA was further purified using RNAeasy Plant Mini kit (Qiagen, Valencia, CA, USA).

Target preparation and hybridization of oligo arrays

RNA quality was checked using the Agilent 2100 Bioanalyzer (Agilent technologies, Palo Alto, CA). cDNA was synthesized from 4 μg of total RNA using One-cycle target labeling and control reagents (Affymetrix, Santa Clara, CA) to produce biotin labeled cRNA which was then fragmented at 94°C for 35 min into 35–200 bases in length.

Three biological replicates were independently hybridized to the GrapeGena 520510F array (Affymetrix, Santa Clara, CA). Each sample was added to a hybridization solution containing 100 mM 2-(N-morpholino) ethanesulfonic acid, 1 M NaCl, and 200 bases in length.

Each microarray was washed and stained with streptavidin-phycoerythrin in a Fluidics station 450 (Affymetrix) and scanned at 7G System (Affymetrix).

Data and sequences analysis and gene annotation

Robust Multi-array Analysis (RNA) algorithm was used for background correction, normalization and expression levels summarization [83]. Next, differential expression analysis was performed with the Bayes t-statistics from the linear models for Microarray data (limma), included in the affylimGUI package. P-values were corrected for multiple-testing using the Benjamin-Hochberg’s method [14]. Data obtained from hybridization of GrapeGen chips was filtered considering a fold change >1.5 and corrected p value <0.05 or fold change ≤-1.5 and a corrected p value <0.05.

The probeset sequences were blasted against the genes predicted from the genome (blastn, e-value <e-20, minimum of 100 bp alignment) available at the NCBI website. Gene annotation was performed by updating the annotation performed in [84] following the same protocol as described by the authors to the new genes from the 12x coverage release of the genome assembly.

The microarray data were submitted to Gene Expression Omnibus (GEO), and are accessible through GEO accession number GSE35172.

Anthocyanins quantification

Fifty-one hundred milligrams of lyophilized material and 50% methanol were mixed to get a final suspension of 50 mg/mL (v/v). Two volumes of acidic methanol (1% HCl in 50% MeOH) were added and absorption read at 530 nm using quartz microcuvettes. Total relative anthocyanin concentration was expressed as the absorbance value at 530 nm/g of freeze-dried weight.

Quantitative RT-PCR

Complementary DNA was synthesized from 1.5 μg RNA using a RevertAid™ H Minus M-MuLV Reverse Transcriptase (Fermentas, Burlington, Canada) according to the manufacturer's instructions. Primers sequences (Table S1) were selected using Primer express software3.0 (Applied Biosystems, Foster City, CA).

Real-time PCR reactions were prepared using Maxima™ SYBR Green qPCR Master Mix (2x) (Fermentas, Burlington, Canada) and performed using the StepOne™ Real-Time PCR System (Applied Biosystems, Foster City, CA). Cycling conditions were 95°C for 2 min, then 40 cycles of 95°C for 1 min and 60°C for 20 min. Expression was determined for duplicate biological replicates and triplicate technical replicates using a serial dilution cDNA standard curve per gene. Data were calculated from the calibration curve and normalized using the expression curve of actin gene (VVTU17999_s_at) that presented absolutely no differential expression in the microarray analysis.

Soluble Metabolites

The profiling of soluble metabolites was performed as detailed previously [85,86]; by gas chromatography coupled to electron impact ionization/time-of-flight mass spectrometry (GC-EI/TOF-MS) using an Agilent 6890N24 gas chromatograph (Agilent Technologies, Boeblingen, Germany; http://www.agilent.com) with splitless injection onto a FactorFour VF-5 ms capillary column, 30m length, 0.25-mm inner diameter, 0.25-μm film thickness (Varian-Agilent Technologies), which was connected to a Pegasus III time-of-flight mass spectrometer (LECO Instrument GmbH, Monchengladbach, Germany; http://www.leco.de).

Soluble metabolites were extracted from 300 mg (fresh weight ±10% tolerance) deep frozen powder by 1 mL ethylacetate with 2 h agitation at 30°C. Aliquots of 300 μL from the ethylacetate fraction were dried by vacuum concentration and stored dry under inert gas at -20°C until further processing. Metabolites were methoxamminated and trimethylsilylated manually prior to GC-EI/TOF-MS analysis [86]. Retention indices were calibrated by addition of a C10, C12, C15, C18, C19, C22, C25, and C36 n-alkane mixture to each sample [87].

GC-EI/TOF-MS chromatograms were acquired, visually controlled, baseline corrected and exported in NetCDF file format using ChromaTOF software (Version 4.22; LECO, St. Joseph, USA). GC-MS data processing into a standardized numerical data matrix and compound identification were performed using the TagFinder software [88,89]. Compounds were identified by mass spectral and retention time index matching to the reference collection of the Golm metabolome database (GMD, http://gmd.mpimp-golm.mpg.de/; [90,91]. Guidelines for manually supervised metabolite identification were the presence of at least 3 specific mass fragments per compound and a retention index deviation < 1.0% [87]. All mass features of an experiment were normalized by sample fresh weight, internal standard (C22) and maximum scaled.

Volatile Metabolites

The profiling of volatile metabolites was performed using 1 g (fresh weight ±10% tolerance) of deep frozen powder which was suspended in 1 mL H2O and 0.2 g NaCl using 20 mL head-space vials (Gerstel, Mulheim an der Ruhr, Germany). Analyses were performed by solid phase micro extraction (SPME) and gas
chromatography coupled to electron impact ionization/quadrupole mass spectrometry (GC-EI-MS) using an Agilent 6890N/24 gas chromatograph (Agilent Technologies, Böblingen, Germany; http://www.agilent.com) and a StableFlex™ SPME-fiber with 65 µm polydimethylsiloxane/divinylbenzene (PDMS-DVB) coating (Supelco, Bellefonte, USA). SPME samples were taken from the headspace with 10 min incubation at 250°C onto a DB-624 capillary column with 60-m length, 0.25-mm inner diameter, 1.40-µm film thickness (Agilent Technologies, Böblingen, Germany). The GC temperature programming was 2 min isothermal at 40°C followed by a 10°C/min ramping to 260°C final temperature which was held constant for 10 min. The Agilent 5975B VL GC-MSD system was operated with a constant flow of helium at 1.0 mL/min. Desorption from the SPME fiber was at 16.6 psi with an initial 0.1 min pulsed-pressure at 25 psi. The subsequent purge was 1 min at a purge flow of 12.4 mL/min. System stability was controlled and the sample sequence randomized. GC-EI-MS chromatograms were acquired with mass range set to 30–300 m/z and a 20 Hz scan rate. Chromatography data files were visually controlled, exported in NetCDF file format using Agilent ChemStation-Software and baseline-corrected with MetAlign software [92]. GC-MS data processing into a standardized numerical data matrix and compound identification were performed as was described above using the TagFinder software [88,89]. Compounds were identified by mass spectral and retention time matching to the reference collection of the Gom Library for volatile compounds. Guidelines for manually supervised metabolite identification were the presence of at least 3 specific mass fragments per compound and a retention time deviation <3.0%. All mass features of an experiment were normalized by maximum scaling.

Relative Quantification of metabolite concentrations

For quantification purposes all mass features were evaluated for best specific, selective and quantitative representation of observed analytes. Laboratory and reagent contaminations were evaluated by non-sample control experiments. Metabolites were routinely assessed by relative changes expressed as response ratios, i.e. x-fold factors in comparison to a control condition or in comparison to the overall median of each metabolite measure.

Statistical analyses and data visualization

Statistical testing was performed using \( \log_{10} \)-transformed response ratios. Statistical assessments and heat map were performed with the multi-experiment viewer software, MeV (Version 4.6.2; http://www.tm4.org/mev/; [93]) or the Microsoft-Excel 2007 program. Independent component analysis (ICA) of the first 5 principal components were performed using \( \log_{10} \)-transformed response ratios via the MetaGeneAlyse web application (Version 1.7.1; http://metagenealyse.mpimp-golm.mpg.de) with missing value substitution, \( \log_{10} = 0 \). Excel files containing ion peak responses were used for Kruskal-Wallis and Wilcoxon rank sum tests in order to determine which samples have significantly different amounts of certain metabolites. For multiple comparisons the Benjamini & Hochberg correction was elected which defines a sequential p-value procedure that controls the expected proportion of falsely rejected hypotheses - the false discovery rate (FDR). The FDR is a less stringent condition than the family-wise error rate (FWER), so this method is among the most powerful procedures for multiple testing.

Supporting Information

Figure S1 Samples of berries at EL 38 stage of development. A Aragonè's B Touriga Nacional C Trincadeira. (PPTX)

Figure S2 Sketch of the sampling layout. (PPTX)

Table S1 List of primers used in real time reverse transcription-polymerase chain reaction. (DOCX)

File S1 List of volatiles and polar metabolites analyzed during grape ripening of the three cultivars under study. (XLSX)

File S2 Kruskal-Wallis and Wilcoxon Rank sum statistics applied to metabolomics data. For selected multiple comparisons the Benjamini & Hochberg correction was elected among other tests that are presented. (XLSX)

File S3 List of entire gene set (8696 probesets) and common gene set (2253 probesets) of modulated genes during grape ripening. Information concerning fold change values, annotation and functional category is provided. (XLSX)

File S4 Weather conditions from April to September in 2008 season. (XLSX)

Author Contributions

Conceived and designed the experiments: AMF JK. Performed the experiments: PAR AE AMF. Analyzed the data: PAR AE LS AMF. Contributed reagents/materials/analysis tools: MSP JK AMF. Wrote the paper: AMF JK.

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