A set of data on green, ripening and senescent vanilla pod (Vanilla planifolia; Orchidaceae): anatomy, enzymes, phenolics and lipids

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Abstract — Introduction. Mature green vanilla pods accumulate 4-O-(3-methoxy-benzaldehyde)-β-D-glucoside (glucovanillin), which, upon hydrolysis by an endogenous β-glucosidase, liberates vanillin, the major aroma component of vanilla. Little is known on the spatial distribution of aroma-generating phenolics, and the enzymes responsible for their liberation (β-glucosidase) and oxidation (peroxidase). We report here quantitative data with respect to these three components in relation to the anatomy of the pod. Furthermore, the spatial progression of oxidation is shown. Materials and methods. Mature green vanilla pods were analyzed for their contents of phenolics (HPLC), and β-glucosidase and peroxidase activities by spectrophotometric techniques using p-nitrophenyl glucoside and vanillin as substrates, respectively. Lipids were examined under fluorescence microscopy after Nile red staining. Oxidation development was observed on transverse slices of pods. Results and discussion. Phenolics, and β-glucosidase and peroxidase activities showed gradients of increasing-decreasing concentrations from the stem to the blossom end of pods. The β-glucosidase activity is distributed in between the placenta, mesocarp, and trichomes in a [7 / 2 / 1] proportion while that of peroxidase shows a [38 / 1] ratio in the mesocarp and placenta, and was absent from trichomes. Oxidation begins from the blossom end in the placenta, progressively invading the mesocarp and moving towards the stem end. Conclusion. The green mature vanilla pod is spatially heterogeneous for its phenolics, and β-glucosidase and peroxidase activities, its placenta playing an important role in the liberation of vanillin and its subsequent oxidation.

Ensemble de résultats sur la gousse de vanille verte, murissante et sénescente (Vanilla planifolia; Orchidaceae) : anatomie, enzymes, composés phénoliques et lipides.


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1. Introduction

Vanilla pod, the fruit of *Vanilla* sp. vine, exhales an excellent aroma, making it a high added-value product. When it is green mature at the time of harvest (8–9 months after pollination), it does not smell; it is only after a lengthy treatment that it becomes a commercial product. Up until the present day, preparation of vanilla comprises several steps [1–4], “killing” in a hot water bath, “sweating” in a cloth-lined close wooden box, drying on racks in the morning sun, and conditioning in a wax paper-lined wooden trunk, the number and length of the various steps depending on the area of production. Basically, the aims and consequences of these steps are:

- “killing”, to stop the vegetative life of the pod to prevent dehiscence [3]; compartmentation of the inner volumes of the cells (vacuole, cytoplasm, etc.) [5], outset of hydrolysis of glucovanillin by the vanilla β-glucosidase [6] and of oxidation with production of brown pigments; no variation in the water content [3],
- “sweating”, to take benefit of the inner heat generated by the “killing” to go on with hydrolysis of glucovanillin by the vanilla β-glucosidase [7] and continue oxidation; no variation in the water content,
- “drying”, a lengthy step (several months) aiming at bringing pods to 30–50% dry matter content,
- conditioning, to further increase the dry matter content to 50–60%, depending on its duration.

Commercial pods will have variable vanillin contents depending on origin (country, areas of production, and, for the same origin, variability of the pod batches cured), agropedoclimatic conditions, cultural practices, curing conditions, etc. [1, 8–10].

Although a great amount of work has been done on vanilla for a century, there are still zones of uncertainty concerning the sequence of events leading to the development of the vanilla flavor. The enzymatic release of vanillin from its non-aromatic precursor, glucovanillin, has been established for a long time [1], but it is only recently that the vanilla enzyme responsible was characterized [6]; although Arana [1] demonstrated earlier that glucovanillin was unequally distributed along the length of the pod, no fine anatomical picture of its distribution in the different compartments of the pod (mesocarp, placenta, trichomes) was available until recently [11–13]; it is the same in the case of the β-glucosidase [11].

Oxidative alteration of vanillin has also been known for a long time [1] and is exemplified by drastic color changes occurring during the curing process, the green vanilla pod turning black in the end. Since then several authors have measured the peroxidase activity of vanilla [14–16], and a peroxidase was recently purified and characterized [17]. However, it is only recently that Gatfield *et al.* [18] showed the presence in some vanilla extracts of a taste-active compound, divanillin, an oxidation product of vanillin by a peroxidase, and no anatomical picture of the distribution of peroxidase activity in the vanilla pod is, to our knowledge, available today.

Thus, the aim of this publication is to gather apparently dispersed data, all of them converging for a better comprehension of the formation of the vanilla aroma. More precisely, objectives were:

- to analyze the variability in populations of green vanilla pods with regard to glucovanillin and β-glucosidase contents,
- to examine in detail the existence of longitudinal and radial gradients in the pod for β-glucosidase, peroxidase, phenolics and sugars,
- to describe the anatomical distribution of β-glucosidase, peroxidase and phenolics in the pod,
- to describe the longitudinal development of phenol oxidation in the pod,

2. Materials and methods

2.1. Plant materials

Two large batches of green mature vanilla pods (~8–9 months after pollination) were from Réunion Island (France) and Madagascar; other minor batches were from Tamil...
Nadu (India) and Papantla (Vera Cruz, Mexico). Their mass ranges were 15–20 g. Their water content was determined by drying overnight at 105 °C.

2.2. Extraction and measurement of the β-glucosidase activity

Immediately after dissection [13], portions of mesocarp, placenta and trichomes (~30–50 mg each) were dilacerated with a MicroTurrax at ambient temperature in 10 mL of 0.1 M phosphate buffer (pH 7.0). After centrifugation (14 000 g, 5 min, 20 °C), the filtrate was diluted 10 × with the same buffer (20 °C).

Activity was measured as described by Odoux et al. [6]; briefly, an aliquot of enzyme extract was added to 200 µL of 4 mM p-nitrophenyl β-D-glucopyranoside in the above phosphate buffer and the total volume was made to 400 µL with phosphate buffer. After 20 min of incubation at 40 °C, 1 mL of 0.5 M NaOH was added and the optical density was read at λ = 400 nm. 1 nanokatal (nKat) is defined as the amount of enzyme hydrolyzing 1 nanomole of substrate per second in the above conditions.

Undissected representative portions of pods (~50–250 mg) were also submitted to the same extraction-determination protocol.

2.3. Extraction and measurement of the peroxidase activity

Immediately after dissection, portions of mesocarp, placenta and trichomes (~50–100 mg each) were dilacerated with a MicroTurrax at ambient temperature in 1.8 mL of 0.1 M acetate buffer (pH 4.5) containing 1 M NaCl, and the slurry was stirred for 30 min at ambient temperature. After centrifugation (14 000 g, 5 min, 20 °C), the supernatant (400 µL) was mixed with the same volume of buffer, and 200 µL of 40 mM vanillin in buffer was added, then 10 µL of 30.9 % hydrogen peroxide; the increase in absorbance was followed at λ = 480 nm between (1 and 3) min at 20 °C. A unit (U) of peroxidase activity was defined as the amount of enzyme giving a 0.001 absorbance unit increase per minute.

Estimation of “total” vanillin was achieved according to Odoux [3]; “total” vanillin content is that obtained after full hydrolysis of glucovanillin with almond β-glucosidase plus free vanillin, the latter being generally low in the total. Thus “total” vanillin is an acceptable approximation of the glucovanillin level.

2.4. Extraction and determination of glucovanillin and related phenolics

When mentioned, the word “phenolics” will include aglycons and their glycosylated forms. Analyses were performed according to Odoux and Brillouet [13]; briefly, portions of mesocarp, placenta and trichomes (~50–100 mg each) were extracted with [methanol:water] (50:50, v/v); the supernatant was then submitted to quantitative analysis by HPLC separation of phenolics and their glycoconjugates on a (250 × 4.6) mm Modulocart QS LiChrospher 5-µm ODS2 column operated at 0.5 mL⋅min⁻¹ and 30 °C. The mobile phase consisted of [water:formic acid] (98:2, v/v) (eluant A) and [water:acetonitrile:formic acid] (18:80:2, v/v/v) (eluant B). The elution program was as follows: 8–13% eluant B for 0–10 min; 13–20% eluant B for 10–30 min; 20–8% eluant B for 30–35 min. Duplicate samples were injected at a level of 20 µL. The column effluent was monitored at 280 nm. Triplicate samples were injected at a level of 10 µL. External standardization was achieved with pure glucosides, and aglycones. Data were expressed as % per fresh weight.

Undissected representative portions of pods (~50–250 mg) were also submitted to the same extraction-determination protocol.

2.5. Extraction and determination of sugars

Undissected representative portions of pods (~40–250 mg) were extracted by 1.5 mL of 0.01 N sulfuric acid for 2 h at ambient temperature, then centrifuged (14 000 g, 5 min). HPLC analyses were performed using a
Spectra-SERIES separation system P100 (Thermo Separation Products, USA) including a quaternary pump and controlled by Chemstation A.10.02 software. Separations were achieved using a (300 × 7.8 mm i.d.) AMINEX® Ion HPX-87H (7 µm, BioRad) column with a guard column operated at 25 °C. The mobile phase consisted of 0.01 N sulfuric acid at 0.6 mL·min⁻¹. Duplicate samples were injected at a level of 20 µL. The column effluent was monitored with a Shimadzu differential refractometer. Quantification was achieved by injection of solutions of known concentrations of sucrose, glucose and fructose; data were expressed as % per fresh weight.

2.6. Fluorescence microscopy

Fresh cross-sections (100 µm) were obtained from pods using a Microm HM650V vibratome, stained for few minutes with Nile red (0.5 mg·mL⁻¹ acetone, then 1/500 dilution in water), then observed with a Leica DM6000 epifluorescence microscope (filter cube A, excitation 450–500 nm, emission > 528 nm) (Leica Microsystems, Rueil-Malmaison, France).

2.7. Chemicals

Vanillin and p-hydroxybenzaldehyde were from Fluka (Buchs, Switzerland). Glucovanillin was from ChromaDex (Irvine, CA, USA), and glucoside of p-hydroxybenzaldehyde was synthesized according to Dignum et al. [19].

3. Results

3.1. Variability in the total vanillin content and β-glucosidase activity amongst populations of mature green vanilla pods

Two large batches of mature green vanilla pods were analyzed for their “total” vanillin content and β-glucosidase activity (figure 1). One must note that the mature green stage was chosen by local producers according to visual estimation of the pods. As seen, this population exhibits a wide variability from 0.7% to more than 5.2% of the pod dry weight, with the majority of pods having 4.5% of vanillin, i.e., 9.2% as glucovanillin. The same remark may be applied to measurement of the β-glucosidase activity: indeed, it varied in a second population from 110 nKat·g⁻¹ fresh weight to more than 1680 nKat·g⁻¹ fresh weight.

3.2. Longitudinal distribution of mass, water, β-glucosidase, peroxidase, sugars and phenolics in mature green vanilla pods

After removal of the corky lignified zones of attachment to the stem and residual blossom pieces (3 mm length), vanilla pods were cut into 16 slices of equal thicknesses from their stem to their blossom end and analyzed (figure 2).

Mass percents and contents of water and dry matter along the length of a mature green vanilla pod show that a pronounced water gradient develops along this axis from the highly hydrated stem end (slices n°1, 2, ~89.2%), descending to ~84.5% in the drier ~2/3-length zone (slice n°12); then, water
content increases up to the blossom end (slice n°16, ∼ 87.8%) (figure 2A). All calculations done, slice n°14, i.e., close to the blossom end, is the most representative of the entire pod water content. The mass distribution follows an almost symmetrical pattern: indeed, the pod shows a tronconic shape from slices n°1 to 6, then adopts a pseudo-cylindrical structure (n°7 to 15), and finally terminates as a cone (n°16).

The β-glucosidase activity exhibits a pattern of variation similar to that of the mass (figure 2B): actually, it is also somewhat superimposable on the variation in dry matter (figure 2A). Its distribution in nkatal by 1/3-length portion (slices n°1 to 6, n°7 to 11, n°12 to 16) was: 22 / 40 / 38. Peroxidase is differently distributed: from slices n°6 to 14, its level was constant while it increased gently up to the stem end; on the other end of the pod (blossom), it increased tremendously up to the terminal slice n°16. Its distribution in units by 1/3-length portion was 31 / 27 / 42. Sugars (i.e., sucrose, glucose and fructose in a constant weight ratio of 0.47 / 0.26 / 0.27) decreased from the stem end to slice n°13, then they increased moderately up to the floral end (figure 2B).

Variations in the concentrations of glucovanillin and vanillin (% per FW) show profiles somewhat symmetrical to the water content but slightly shifted (figure 2C). Their levels began from almost nil in the stem end, then progressively climbed up to slice n°14, and finally dramatically fell down to the blossom end; this is also true on a dry matter basis, glucovanillin varying from 0.08 (slice n°1) to a maximum of 9.43 (slice n°12) % (DW), and vanillin from 0.01 to 0.85 % (DW). The same trends were observed for \( p \)-hydroxy-benzaldehyde glucoside and its aglycon; however, it must be noted that, while glucovanillin is more or less stable from slices n°8 to 15, \( p \)-hydroxy-benzaldehyde glucoside goes on increasing from slices n°10 to 14. Weight concentration ratios between glucosides and their aglycones were stable at: \([\text{glucovanillin} / \text{vanillin}] = 11.2, \ [\text{p-hydroxybenzaldehyde glucoside} / \text{p-hydroxybenzaldehyde}] = 5.6\). Distribution of total phenolics (FW) by 1/3-length portion (slices n°1 to 6, n°7 to 11, n°12 to 16) was: 17 / 43 / 40.

Similar variations were observed on several other green pods of different origins.

### 3.3. Radial distribution of β-glucosidase and peroxidase in the different compartments of mature green vanilla pods

Transverse slices (n°14) of vanilla pods were finely dissected under a stereomicroscope, yielding chlorophyllous mesocarp, placenta and trichomes. The mesocarp portion was further dissected into five concentric coronae [(I) to (V)] and placentae were divided into three layers [(VI) to (VIII)] from their base to their top (figure 3).
**Figure 3.**
Top: transverse slice of a mature green vanilla pod (n°14) showing the diverse compartments analyzed: (I) epidermis, (II)–(III) outer mesocarp, (IV)–(V) inner mesocarp, (VI) placental base, (VII) lobes of the *placentae*, (VIII) funicles, (IX) trichomes. Bottom: anatomical distributions of the β-glucosidase (A), and peroxidase (B) activities, and glucovanillin (C) (from ref. [13]). Intensities of colors (clear to deep) are roughly indicative of numerical values given in table I; white indicates nil.

**Table I.**
Distribution of the β-glucosidase and peroxidase activities, and glucovanillin in the different compartments of a transverse slice (n°14) of a mature vanilla pod (n = 10).

<table>
<thead>
<tr>
<th>Compartment</th>
<th>Compartments</th>
<th>N°</th>
<th>β-Glucosidase activity (nKat·g⁻¹ FW)</th>
<th>Peroxidase activity (U·g⁻¹ FW)</th>
<th>Glucovanillin² (% FW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mesocarp</td>
<td>Epidermis (I)</td>
<td>268 ± 119</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Outer mesocarp (II)</td>
<td>311 ± 192</td>
<td></td>
<td>7124 ± 183²</td>
<td>0.03 ± 0.02²</td>
</tr>
<tr>
<td></td>
<td>Outer mesocarp (III)</td>
<td>294 ± 105</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Inner mesocarp (IV)</td>
<td>326 ± 177</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Inner mesocarp (V)</td>
<td>210 ± 123</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placentae</td>
<td>Placental base (VI)</td>
<td>1047 ± 305</td>
<td>9.63 ± 1.99</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lobes of the <em>placentae</em> (VII)</td>
<td>1377 ± 112</td>
<td>186 ± 9²</td>
<td>4.98 ± 1.27²</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Funicles (VIII)</td>
<td>1210 ± 152</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Trichomes</td>
<td>Trichomes (IX)</td>
<td>200 ± 89</td>
<td>Not detected</td>
<td>3.93 ± 1.28</td>
<td></td>
</tr>
</tbody>
</table>

1 Measured without sub-dissections of mesocarp and *placentae*.
2 From Odoux and Brillouet [13]; (VII)–(VIII) are the placental lobes in this ref.
The β-glucosidase activity was measured on all these fractions including trichomes, while the peroxidase activity was determined on whole mesocarp, *placentae* and trichomes without sub-dissection (*table I, figure 3*). The β-glucosidase was mainly and rather uniformly located in the *placentae*; mesocarp was 3-4× poorer and no radial gradient was observed from its periphery to its inner portion. Trichomes were the poorest compartment (1.5× lower than mesocarp).

The peroxidase activity was differently distributed; it was absent from the trichomes and showed a [38 / 1] repartition between mesocarp and *placentae* (FW).

### 3.4. Longitudinal and radial developments of oxidation in ripening-senescent vanilla pods

A ripening-senescent pod clearly shows oxidation at the blossom end progressively invading up to one-fourth of the length where green tissues are still present (*figure 4*); oxidation is also visible at the stem end (~10% of pod length). Transverse slices were obtained from different regions, starting from the green region and finishing at the blossom end, from the green area (n°1) to greenish-brown (n°2), to brownish (n°3), to blackish (n°4), to deep black (n°5) ones. On the fresh green slice, one may see the chlorophyllous mesocarp, the diffuse white placental zone, the glassy white V- and J-shaped trichome corners, and, on the frozen equivalent slice, a white area clearly demarcating the *placentae* (see also ref. [11]); the first signs of oxidation resulting from the transverse cutting are visible in the fresh mesocarp portion as two brownish circles, vascular bundles. On fresh slice n°2, one sees oxidation superimposed on the placental zone; on the frozen ring, three stages are observed: an intact upper left white *placenta*, an upper right slightly brown one and below a deep brown one and there, oxidation invaded the mesocarp. On fresh slice n°3, a typical clear reddish-brown urchin-like shape radiating from the *placentae* outwards half the mesocarp thickness; one must note that no oxidation is visible in between the mesocarpic vascular bundles. In this slice, one may see residual whitish placental zones and the trichomes stay white (see also *figure 5*). When moving towards the blossom end, slice n°4 shows a more or less regular annulus uniformly stained reddish while trichomes and funicles appear white; half of the mesocarp is still greenish. Finally, on slice n°5, one sees that the entire slice, except trichomes and funicles, turned deep chocolate brown.

On the enlarged view of slice n°3 (*figure 5*), one may see that the ends of rays from the radiating shape are actually vascular bundles: three per triangle side, and a triplet at each corner in the outer mesocarp area made of twin little bundles and a larger one (see also *figure 4*, slice n°2, and refs. [11, 20]). There, three small bundles are also seen in the whitish upper right *placenta*. At this stage, one may see that oxidation draws upside-down funnel-like shapes, their bottom being centered on the mesocarpic vascular bundles; areas in between bundles remain yellowish-green.

### 3.5. Anatomical distribution of lipids in mature green vanilla pods

In a partial transverse section of a native mature green vanilla pod after lipid staining with Nile red (*figure 6A*), neutral lipid droplets intensely fluorescing in sulfur yellow were seen in the single layer of endocarp cells lining the locule in between the trichomes and bordering the placental *laminae* (or blades); they were also seen in a single cell layer beneath the trichomes. Whether these two single layers are of the same ontogenic origin remains uncertain, and there is no apparent discontinuity in their junction area; the lipid droplets in the subtrichomal layer, however, appear smaller than in the properly so-called endocarpic layer. The trichomes also contain numerous small lipid droplets. Differently from the two above compartments (endocarp and trichomes), deposits of an intensely golden orange fluorescing substance, not spherically structured in droplets, were also seen in the terminal region of the placental blades, *i.e.*, the funicles. The autofluorescence of the upper part of the placental *laminae* (lobes) is apparently restricted to the funicular ends (*figure 6B*).
4. Discussion

Arana [1, 2] measured the glucovanillin content in three portions of green mature pods (stem region, middle and blossom region; presumably, 1/3 of the length each), and found that glucovanillin mass distribution in these three regions, taking into account their weight proportions, was, in the pod, 20/40/40 of the total. We confirm more precisely
that pronounced gradients exist all along the pods for water, glucovanillin, vanillin, p-hydroxy-benzaldehyde glucoside, and its aglycon; the shapes of these gradients much resemble those of mass and dry matter distributions, i.e., negligible levels of phenolics at the stem end, progressively increasing up to pod slice n°14, then decreasing strongly until the floral end. It is noteworthy that sugars followed a symmetrical trend. Coming back like Arana [1] to a 1/3 distribution of the three regions (i.e., stem end: fractions n°1 to 6; middle fractions: n°7 to 11; and blossom end: fractions n°12 to 16; figure 2), we found, all calculations done, a 17/43/40 distribution for the total phenolics (glucovanillin being largely dominant), a ratio almost identical to Arana’s; this means that the two-thirds of the pod from its apex contain more than 80% of the total vanillin precursor. It is noteworthy that the 22/40/38 distribution for the β-glucosidase activity is quite similar to the phenolics one.

If one considers the glucovanillin levels expressed as mM in the water phase of the pod (i.e., 100 mM, dry matter basis), one may calculate that its concentration from slice n°8 to slice n°15 (blossom end) equals ~50 mM, i.e., $10 \times K_m$ of the vanilla β-glucosidase in dilute solutions [6], or $2.2 \times K_{0.5}$ of the same enzyme working in heat-treated pods [7]; thus, in the first comparison, this
enzyme would work for a while at its $V_{\text{max}}$, while, in the second case, hydrolysis would also proceed at $V_{\text{max}}$, but for a shorter period of time. As in addition to that, the level of the $\beta$-glucosidase activity shows a gradient similar to that of glucovanillin, and although most of this activity is lost upon “killing” [7, 21, 22], the residual enzyme would, at the beginning, work at its maximum speed in the middle-blossom end regions; then, its reaction rate would decrease progressively downwards to the stem end, as its activity level and glucovanillin concentration do.

Both the above observations are thus to be related to the preferential occurrence of vanillin crystals on the epidermis of the middle region and blossom end of cured vanilla pods [1, 2]. It is also noticeable that a black blossom end-split overripe pod was invaded by mold at the stem end only where vanillin, presumably toxic to some fungal species [23], is present at trace level [pers. com.].

Considering the peroxidase and $\beta$-glucosidase gradient shapes, one understands now why, as long as sufficient vanillin is liberated by the $\beta$-glucosidase at the blossom end where peroxidase is very high, oxidation starts in this portion by forming blossom-end-yellow pods (figure 4) [1, 10]. The determinism of this preferential oxidation at the floral end must be searched for in the structure and anatomy of the pod: vascular bundles, comprising xylem vessels, run its whole length from the stem to the floral pieces. They are present in a constant number of 27 (figures 4, 5 and ref. [20]) distributed as follows: in the mesocarp – 3×3 (medium diameter; sides of the triangular section) plus 3 (large diameter; corners) and 3 twin (small diameter; corners); in the placental bases – 3×3 (small diameter; sides of the triangular section). Since the volume occupied by these elements is constant along the pod, and because both ends of the pod have much more reduced diameters than the rest of it, then the volume (weight) proportions of xylem elements in these tissues increase there to the detriment of parenchyma (number and/or size of cells); conversely, in the pseudo-cylindrical portion (slices n°7 to 5), the ratio (xylem/parenchyma) is constant, and so is the peroxidase activity. Since cell wall-bound [24] and soluble peroxidases [25] are known to play a role in the lignification of secondary walls of the xylem by converting the monolignols (coumaryl, coniferyl and synapyl alcohols) into free radicals which spontaneously polymerize to give lignin, one may hypothesize that xylem peroxidase(s) is somewhat involved in vanillin oxidation; however, the existence of other non-vascular peroxidases cannot be ruled out (see below). Since glucovanillin, and thus vanillin and $\beta$-glucosidase are very low in the very stem end, and although peroxidase is, as at the blossom end, higher than in the central core, oxidation does not develop there.
The progressive longitudinal invasion of vanillin oxidation from the blossom to the stem ends (see below) is also well explained by these gradients: indeed, glucovanillin is not prone to oxidation due to engagement of its vanillin phenoxy group in the \( \beta \)-glucosidic linkage, while free vanillin is; thus, action of the \( \beta \)-glucosidase is a prerequisite to oxidation, and since its level decreases continuously from slice \( \#14 \) downwards to the stem end, the amount of vanillin liberated per unit of time will also decrease, and, thus, the browning front will progressively move to the stem end, i.e., the length of time to get from green to yellowish to chocolate brown will increase.

The anatomical distributions of glucovanillin and \( \beta \)-glucosidase were first shown by Odoux \textit{et al.} \[11\], the same authors later gave a more detailed analysis of the repartition of phenolics in the green mature pods \[13\]. We give here an even more complete anatomical picture and a quantitative analysis of the distributions of the \( \beta \)-glucosidase and peroxidase activities in a representative slice, the peroxidase repartition having, to our knowledge, never been published (figure 3, table 1). \( \beta \)-glucosidase, as formerly shown \[13\], is high in the placentae whatever the portion considered, and lower in the mesocarp and trichomes; again, one must point out the uselessness of this enzyme, at least at the beginning of hydrolysis, in the green glucovanillin-free mesocarp, contrary to the assertion of Arana \[1\]. Peroxidase is far higher in the mesocarp than in placentae, and was not detected in the trichome strips \[13\], according to the fresh matter distribution of the two compartments \[13\], it is thus essentially found in the mesocarp. This result is not surprising since, as mentioned above, the mesocarp contains most of the pod vascular bundles including xylem vessels. Ironically, the mesocarp does not contain oxidizable phenols (e.g., vanillin), but likely only those serving as substrates for lignin synthesis (e.g., \( p \)-coumaric acid) which are encountered in xylem vessels \[26\]. Placentae also include vascular bundles of small diameter and, per placental pair, two of them are located at the base of each placental lamina, one of them in between (figures 4, 5 and ref. \[20\]). However, and since oxidation seems homogeneous in placental laminae (figure 4, frozen slice \( \#2 \), upper right), i.e., there is no preferential oxidation surrounding the bundles, one may imagine that placental peroxidase(s) would not be only restricted to the xylem, but would also be of parenchymatous origin. Indeed, plant peroxidases have been, for instance, also encountered in vacuoles \[27\]. Dignum \textit{et al.} studied the stability of vanilla peroxidase in crude extracts \[16\], and Márquez \textit{et al.} recently purified from whole pods with yellow apices a cell wall-bound peroxidase from vanilla pod \[17\]. However, the fine localization of peroxidases in the pod and their status (cell wall-bound, and possibly soluble forms) is not yet known.

When examining the progression of oxidation along the longitudinal axis of pods, one may define successive steps:

- in the green parts, enzymatic hydrolysis of glucovanillin did not begin, and only some mesocarpic xylem vessels show, after section, signs of oxidation, likely of the phenyl-propanoids by the xylem peroxidase(s),
- when considering the blossom end, one may see that oxidation, in addition to the above observation, concerns the whole placental laminae only, i.e., the mesocarpic parenchyma is not involved. Indeed, at this stage, glucovanillin only present in the placentae and trichomes undergoes progressive hydrolysis with liberation of vanillin itself undergoing enzymatic oxidation in the placentae; at that time, the incompleteness of hydrolysis is visible in residual whitish placental areas,
- later on, oxidation progressively invades the inner mesocarp, drawing radiating shapes of coalescence with the mesocarpic xylem oxidation zones,
- oxidation develops further into a concentric annulus comprising half of the mesocarp surface, and the coloration intensity increases, which indicates the completeness of glucovanillin hydrolysis and further subsequent oxidation of vanillin,
- finally, oxidation products spread all over the pod section, and the color turns from reddish to chocolate brown.
Thus, it is now possible to draw a hypothetical scheme of events leading to the brown-blackish vanilla pod (figure 7):

- when ripening on the vine or during the traditional curing process, a tissular decompartmentation occurs [5], and phenolic β-glucosides and β-glucosidase become present,
- hydrolysis of glucovanillin and other glucoconjugates by the β-glucosidase begins in the placenta, and the presence of this enzyme in the mesocarp serves, at that time, no use; vanillin and other aglycons are released, and the whitish zone slowly disappears,
- rapidly or simultaneously, oxidation of these phenolics develops only in the placenta where substrates and peroxidase(s) are present; the mesocarp is not concerned, except its xylem vessels,
- reddish then brown oxidation products diffuse outwards into the inner mesocarp; however, the possibility that glucovanillin and/or vanillin migrate into the mesocarp and also undergo hydrolysis and oxidation there cannot be ruled out; nevertheless, one must recall that placenta have by far enough β-glucosidase for glucovanillin to be hydrolyzed in situ without diffusing into another compartment [11, 13, this work]. At that moment, it is important to recall the comment of Arana [1] on the prerequisite for glucovanillin to diffuse from the inner placental parts of the pods outwards to the mesocarp for being hydrolyzed by the β-glucosidase: actually, we show here that this sequence of events is not compulsory but may or may not happen. It is presumably the oxidation products accompanied or not by glucovanillin and vanillin which diffuse into the mesocarp. Indeed, the possibility that vanillin would also be oxidized by the mesocarp xylem peroxidase(s) cannot be ruled out. Until peroxidase activity is precisely localized in the mesocarp (vascular bundles and parenchyma?), this question remains open,
- finally, the whole pod turns deep brown, then blackens upon drying.

A particular point must be emphasized: throughout the ripening-ageing period when pods turn from green to black, the trichomes and funicles, although sources of glucovanillin and β-glucosidase (table I and

**Figure 7.**
Sequence of some consecutive enzymatic and chemical reactions occurring at the vanillin level in the vanilla pod.
Data on ripening vanilla pod (figure 3 and refs.[13, 28]); as formerly mentioned [13], the absence of catechin-HCl staining of vanillin-containing trichomes [12] was likely due to the presence of high quantities of lipid reserves in this compartment. The constitution of trichome lipids is now partly defined (γ-pyranones) [13], while the funicular ones remain unknown.

Finally, we also found that the "total" vanillin content, actually close to the glucovanillin concentration [3], of green vanilla pods was quite variable amongst a population: thus, to characterize a batch or follow maturation kinetics, one has to take into account this variability and carry out measurements on a sufficient number of pods. The same observation was true for the β-glucosidase activity, one of the key enzymes in the development of the vanilla aroma: it was variable amongst a population of pods of the same physiological age and origin. This is also likely true for a given set of vines from one year to another, depending on the climatic conditions of culture. Again, one must be cautious and analyze numerous pods before giving an amount of activity per unit of fresh weight.

5. Conclusion

Green mature vanilla pods are heterogeneous along their longitudinal axis, exhibiting pronounced gradients of decreasing-increasing concentrations of water, and increasing-decreasing ones of β-glucosidase and phenolics from the stem to the floral end; peroxidase activity is differently distributed, stable in a major portion of the pod and increasing at its two ends. These elements are also heterogeneously distributed in the vanilla pod tissues, phenolics and β-glucosidase being preponderant in the placenta while peroxidase predominates in the mesocarp. Finally, oxidation of phenols develops from the placental region of the blossom end towards the stem end; then the oxidation products (brown pigments) progressively invade the mesocarp and, in the end, the whole pod.

Green mature vanilla pods are variable for their glucovanillin and β-glucosidase contents, which must be taken into account in studies devoted to these components.

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References


Conjunto de resultados sobre la vaina de vainilla verde, en proceso de maduración y sensescente (Vanilla planifolia; Orchidaceae): anatomía, encimas, compuestos fenólicos y lípidos.

**Resumen — Introducción.** Las vainas de vainilla verdes maduras acumulan 4-O-(3-metoxibenzaldehído)-β-D-glucósido (glucovanilla) que, bajo el efecto de una hidrolisis mediante una β-glucosidasa endógena, libera vainillina, el principal compuesto de aroma de la vainilla. Las reparticiones espaciales de los compuestos fenólicos generadores de aroma y de las encimas responsables de su liberación (β-glucosidasa) y oxidación (peroxidasa) son poco conocidas. A continuación ofrecemos resultados cuantitativos de estos tres compuestos en relación con la anatomía de la vaina. Asimismo, se ilustrará la progresión espacial de la oxidación.

**Material y métodos.** Se analizaron vainas de vainilla verdes maduras para su contenido en compuestos fenólicos y actividades β-glucosidasa y peroxidasa mediante técnicas espectrofotométricas, gracias al empleo respectivo de glucósido de p-nitrofenilo y de vainilla como sustratos. Los lípidos se examinaron en microscopía fluorescente tras coloración roja de Nil. El desarrollo de la oxidación se observó en cortes transversales de vainas.

**Resultados y discusión.** Los compuestos fenólicos, así como las actividades β-glucosidasa y peroxidasa muestran gradientes de concentraciones crecientes-decrecientes de la extremidad peduncular hacia la extremidad floral de la vaina. La actividad β-glucosidasa se distribuye entre placentas, mesocarpio y tricomas con una relación [7 / 2 / 1], mientras que la actividad peroxidasa se reparte en proporciones [38 / 1] entre el mesocarpio y las placentas; está ausente de tricomas. La oxidación empieza en las placentas de la parte floral invadiendo progresivamente el mesocarpio y se desplaza hacia la zona peduncular. **Conclusión.** La vaina de vainilla verde madura es heterogénea desde un punto de vista espacial en lo que se refiere a sus compuestos fenólicos y actividades β-glucosidasa y peroxidasa, las placentas desempeñan un papel importante en la liberación de la vainillina y en su oxidación consecutiva.

**Francia / Vanilla planifolia / Orchidaceae / vainilla / β-glucosidasa / peroxidasa / glucovanillina / vainillina / lípidos / oxidación / mesocarpio / placentas / tricomas / xilema**