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Phenolic distribution in apple epidermal and outer cortex tissue by multispectral deep-UV autofluorescence cryo-imaging

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Abstract

Phenolic compounds in fruit are involved in responses to biotic and abiotic stresses and are responsible for organoleptic properties. To establish the distribution of these secondary metabolites at the tissue and sub-cellular scales, mapping of fluorescence in apple epidermis and outer cortex tissue in cryogenic condition was performed after deep-UV excitation at 275 nm. Douce Moën and Guillevic cider apple varieties were sampled and frozen after harvest, after 30 days at 4 °C and after 20 days at room temperature. Image analysis of fluorescence emission images acquired between 300 and 650 nm allowed the assignment of fluorescence signals to phenolic compound families based on reference molecules. Emission attributed to monomeric and/or condensed flavanol was localized in whole tissue with major fluorescence in the cuticle region. Hydroxycinnamic acids were found predominantly in the outer cortex and appeared in the cell wall. Fluorescent pigments were mostly found in the epidermis. The distribution of flavanols in the sub-cuticle and phenolic acids in the outer cortex distinguished apple varieties. Storage conditions had no impact on phenolic distribution. The proposed fluorescent imaging and analysis approach enables studies on phenolic distribution in relation to fruit development, biotic/abiotic stress resistance and quality.

- Key words: Malus domestica, phenolic compounds, autofluorescence, multispectral imaging,
- 41 deep-UV synchrotron light, cryo-microscopy.

1. Introduction

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Phenolic compounds are common plant secondary metabolites that have been grouped into three classes: flavonoids, phenolic acids and stilbenes [1]. The flavonoids encompass flavanols that are also found as oligomers and polymers (i.e., proanthocyanidins and condensed tannins), anthocyanins and flavonols, while phenolic acids comprise hydroxybenzoic acids and hydroxycinnamic acids (Figure 1). Except for flavanols and hydroxybenzoic acids, which are mainly present in their free form, most phenolic compounds are glycosylated and/or acylated derivatives. Their ultraviolet absorption capacity limits lightinduced damage to tissues while they are part of plant defence mechanisms against external biotic stresses [2, 3]. In fleshy fruit, phenolic compounds are known for their antioxidant contribution to foods and organoleptic properties, such as colour, astringency or bitterness [4-6]. Various studies in apple and grape have shown important differences in phenolic content between varieties according to fruit development and tissue [7]. Phenolic compounds are commonly characterized by biochemical techniques on whole fruit, skin, flesh, seed or juice extracts. However, there are few reports on the distribution of specific phenolic compounds at the tissue and sub-cellular scales under in-vivo-like conditions. Flavanols and condensed tannins were determined to be localized in the epidermis, parenchyma and seeds by microscopy of fresh or chemically fixed apple [8], grape [9, 10] and other fruits [11-14]. Flavanols were localized by DMACA staining (4-dimethylaminocinnamaldehyde) [14-16] while flavonols were highlighted by fluorescence microscopy using the specific dye DPBA (diphenylboric acid-2-aminoethyl ester) [17, 18]. However, chemical preparation damage to fleshy fruit tissue integrity affects the cellular localization of diffusible compounds [19], and dyes are limited by their specificities.

To address these issues, mapping phenolic compounds through their autofluorescence characteristics appears to be a good alternative as no specific stain is required.

Hydroxycinnamic acids express a blue fluorescence when excited at the UV wavelength of approximately 350 nm [20, 21] while anthocyanins and flavonols are autofluorescent at visible wavelengths [22]. Monomeric and condensed tannin ((+)catechin, (-)epi-catechin, procyanidins) fluorescence requires deep-UV wavelength excitation at approximately 280 nm [23] [24]. The use of different bandpass filters to select specific ranges of fluorescence emissions offers the opportunity to localize different compounds within the same sample through a multispectral approach [25]. However, data on the fluorescence emission of specific phenolic compounds are scarce due to the high number of entities that constitute this metabolite family and because access to pure reference phenolic compounds is limited.

To avoid artefacts related to small compound diffusion and loss during sample preparation, cryo-fixation, cryo-microtomy and fluorescence cryo-observation of biological samples have seen recent developments. Cryo-confocal observations at high resolutions were introduced with the realization of specific microscope stages and immersion objective lens prototypes [26, 27]. A simpler and readily accessible setup was developed to observe fleshy fruit tissue by cryo-laser scanning confocal microscopy (LSCM) [18]. Among the advantages of cryo-observations is the lower fluorescence bleaching compared to that during room temperature observations and thus the recovery of higher fluorescence intensity [26, 28].

Preliminary tests of fresh fruit auto-fluorescence responses after excitation at 375 nm were performed on apple and grape. The latter revealed promising autofluorescence signals, while a lower excitation wavelength was found necessary to assess flavanol localization. The objective of the present work was to map phenolic compounds in cryo-fixed outer tissues of apple. The mapping was performed at a large field of view by multispectral autofluorescence imaging after deep-UV excitation at 275 nm using the setup developed for fleshy fruit cryo-observation. Two cider apple varieties stored in three different conditions were studied to evaluate the genetic and physiological impacts on phenolic compound distribution.

- 92 Fluorescence excitation and emission characteristics of reference phenolic compounds were
- 93 recorded to help identify fruit tissue phenolics.

2. Materials and methods

95 *2.1. Materials*

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- 96 *2.1.1. Chemicals*
- Phenolic compounds used for reference fluorescence spectra were from the laboratory collections of ESA-GRAPPE (Angers, France) and BIA-PRP (Rennes, France). Formic acid was purchased from Sigma-Aldrich (UK) and methanol was purchased from CarloErba
- 100 (Italy).

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101 *2.1.2 Fruits*

- Apple (*Malus domestica*, Royal Gala) and grape (*Vitis vinifera*, Red Italian) for preliminary fluorescence analysis after 375 nm UV excitation were from a local retail store. These fruits were used without further storage. For the fluorescence mapping following excitation at 275 nm, two cider apple varieties, Douce Moën (*M. domestica*, Douce Moën) and Guillevic (*M. domestica*, Guillevic), were provided by IFPC (Le Rheu, France). Fruits were harvested on October 2017 in orchards at Surzur (Morbihan, France). Nine fruits of each variety were subjected to three different storage conditions: three were immediately sampled, three were stored at 4 °C for 30 days and three were stored at room temperature for 20 days before sampling.
- 111 2.2. *Methods*
- 112 2.2.1. Spectrofluorometry
- Phenolic references were dissolved at 2 mg mL⁻¹ in 2 mL of H₂O:MeOH (1:1 in volume) + 1% formic acid (in volume). Quartz cuvettes (10 x 4 mm, 114F-QS, Suprasil

HELLMA Analytics, France) were filled with 500 µL of standard solution and observed by spectrofluorimetry (F-4500, HITACHI, Japan). Emission and excitation spectra for each compound were recorded between 200 nm and 600 nm.

2.2.2. Laser scanning confocal microscopy (LSCM)

Cubes of $5 \times 5 \times 5$ mm³ were sampled from the epidermal region of equatorial fruit slices of commercial apple and grape. Sections of 100 μ m thickness were cut from fresh cubes with a vibrating blade microtome (HM 650V, MICROM, France) and immediately observed or dipped in standard buffer solution pH 10 (HI 70010C, NIST standard, HANNA INSTRUMENTS, Hungary) followed by H₂O milliQ wash prior to LSCM observations at room temperature. LSCM (A1 Eclipse Ti inverted microscope, NIKON Inc., Japan) was configured with laser diode excitation at 375 nm. Fluorescence emission was acquired at 20×10^{-1} magnification with a resolution of 0.63 μ m pixel⁻¹ and image size of 512 x 512 pixels² (16-bit). Four bandpass emission filters were used to map fluorescence: 400-450 nm, 500-530 nm, 560-600 nm and 630-700 nm. These form sets of multispectral images of four channels with spectral controlled excitation and emission parameters.

2.2.3. Synchrotron deep-UV wide-field fluorescence microscopy (DUV)

One 5 x 5 x 5 mm³ cube was randomly sampled from the epidermal region of a cider apple fruit equatorial slice. The cube was frozen in liquid nitrogen-cooled isopentane and stored at -20 °C prior to analysis. Sections (60 µm thick) were cut from frozen cubes using a cryostat microtome (HM 500 OM, MICROM, France) at -20 °C. The section was placed between quartz lamella (R52-5000, ESCO Optics, USA) glued together by frozen water droplets. Synchrotron UV microscopy was performed at the DISCO (Dichroism, Imaging, mass Spectrometry for Chemistry and biOlogy) beamline at the SOLEIL (Source Optimisée de Lumière à Energie Intermédiaire du LURE (Laboratoire à Utilisation du Rayonnement

Electromagnétique)) synchrotron radiation facility [29] (Gif-sur-Yvette, Saint-Aubin, France) on the inverted epi-fluorescence TELEMOS microscope [30]. Observation was realized under cryogenic conditions using a handmade setup composed of a Peltier stage (PE100, LINKAM Scientifics, UK) cooled by dry ice as described in [18]. The wide-field microscope TELEMOS modified by the DISCO beamline staff was a Zeiss Axio Observer Z-1 designed to observe samples after deep-UV (DUV) excitation. The microscope was equipped with a motorized sample plate (MS-2000 XY, Applied Scientific Instrument, USA) to move the sample along the X and Y axes. Sample movement along the Z axis was controlled by the inverted microscope. The setup used the 10x magnification Zeiss Ultrafluar lens (N.A 0.2, WD 7.4) and a sharp 300 nm dichroic mirror "DM" (Omega Optical), which reflected the 280 nm excitation and transmitted emission light above 300 nm. Multispectral fluorescence emission was recorded using six different emission bandpass filters: 300-306 nm, 327-353 nm, 352-388 nm, 412-438 nm, 420-480 nm and 600-650 nm (Semrock, Rochester, USA). In the text, filters are referred to by their low wavelength limit. A back-illuminated CCD camera (Pixis BUV, Princeton Instrument, USA) recorded the images (1024 x 1024 pixels; 1 x 1 µm² per pixel; grey level coded in 16 bit). Acquisition times were 12.5 s for the 300 nm filter, 5 s for the 327, 352, 412 and 420 nm filters, and 7.5 s for the 600 nm filter. Acquisition was performed from the sample cuticle to approximately one mm inside the tissue. With three biological repetitions and three different storage modalities, a total of nine images were registered per variety.

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The beamline intensity, which was focused at the centre of the observation field, led to raw images that required corrections to homogenize illumination. In addition, the camera provided a non-null background without illumination. Both inhomogeneous illumination and background corrections were realized according to the following formula:

 $Ic = \frac{I - Dark}{White - Dark}$

where Ic is the corrected image; I is the raw image; Dark is the background and White

is the flat field

For this purpose, nine different "dark" images were recorded without sample and without any illumination, one for each of the six emission filters. The "White" images were acquired using a frame $(1.0 \text{ x } 1.0 \text{ cm}^2, \text{AB-0576}, \text{Gene Frame, THERMOSCIENTIFIC, UK})$ displaying a homogeneous autofluorescent signal. Thresholding was also required to define the sample

image from the camera noise. Image pretreatments were performed using MATLAB® 2017b.

The corrected images were cropped to 300 x 400 µm² size to fit within a pertinent sample image area. To take into account out-of-plane sample sections, several acquisitions were recorded for the same sample area with a maximum of six focal planes. As this method was not confocal imaging, a clear image was reconstructed by combining the different focal plane views with the extended focus software HeliconFocus6® (version 6.8.0) using the pyramidal method.

2.2.3.1. Multispectral analysis of DUV fluorescence images

DUV fluorescence emission was valued based on the grey scale intensity of pixels in cropped images. Principal component analysis was applied separately on each multispectral image, and fluorescence profiles were extracted from selected emission filters.

2.2.3.2. Image Analysis

Individual emission images were unfolded to form one vector [31]. The image vectors were combined into one table per sample containing six columns corresponding to the six filters and n rows corresponding to the n pixels of the image. Principal component analysis (PCA) was applied to the resulting data table, and correlation circles were drawn from the

PCA loadings. Principal component images were obtained by refolding scores [31]. Component images were filtered to keep 99% of pixels closer to the median value and to remove eventual aberrant pixel values prior to normalization of grey levels between 0 and 255. PCA reconstructed colour images were created from components 1, 2 and 3 identified in the blue, red and green channels, respectively.

Localization of phenolic compounds in samples was also analysed by computing fluorescence intensity profiles from the cuticle to the inner tissue. This "image distance" analysis was realized using FIJI/ImageJ® software [32] and the mathematical morphology plugin process/erode (3D) [33]. Thresholding was first applied to create a binary mask of the sample. Distance from the cuticle was obtained by the following procedure: one-pixel erosion (pixel size = 1 x 1 µm²) was applied to the sample mask and the resulting binary image was subtracted from the initial sample binary image. This step was iteratively applied until the last image column was reached. The mean profile was smoothed to reduce intensity variations due to the presence of cell lumens with low fluorescence and cell walls with high fluorescence. This was realized by applying local regression (LOESS) with a degree of smoothing of 0.4. Normalization between 0 and 1 was realized on the maximum smoothed mean value to allow profiles comparison between samples. Examples of raw and smoothed profiles are shown for Douce Moën at harvest in Supplementary Figures 1 and 2. The smoothed profiles from the three biological replicates per storage modality were averaged and the mean profile per modality was plotted with its 95% confident interval (Supplementary Figure 3).

2.2.3.3. Statistical analyses

The effect of storage was evaluated by Student's t-tests of the smoothed mean pixel values of three biological replicates per pixel distance. As no significant differences were found between storage modalities, Student's t-test for variety effect was performed on the nine replicates.

Principal component analysis, LOESS smoothing and Student's t-tests were performed in R [34].

3. Results

3.1. Fluorescence properties of reference phenolic compound

Excitation and emission fluorescence spectra were registered from standard phenolic compounds (Supplementary figure 4) and the maximum wavelength responses recorded (Table 1). If associating specific compounds with peculiar emission wavelengths is difficult due to the width of spectra with large overlaps, distinguishing families of compounds, such as flavanols (including condensed tannins), phenolic acids, anthocyanins and flavonols remains possible. For example, phenolic acids, such as p-coumaric acid, ferulic acid and caffeic acid have a maximum emission at approximately 420 nm, while flavanols such as (+) catechin, (-) epi-catechin and procyanidin share maximum emission at approximately 320 nm. In fact, due to their large excitation spectra, most phenolic compounds will fluoresce under excitation at 275 nm. Among the tested reference compounds, only phloridzin, rutin and arbutin will not fluoresce after deep-UV excitation.

3.2. Preliminary analysis of autofluorescence observed after excitation at 375 nm

In a preliminary study, autofluorescence observations of fleshy fruit were realized with a laboratory confocal laser scanning microscope using a UV diode laser at 375 nm. This test was performed to check the pertinence of autofluorescence analysis on two types of fleshy fruits: Royal Gala apple and Italian Red grape. In all cases, the cuticle presented bright and intense fluorescence as previously observed [35]. Red spots were observed within the cells of both fruit but particularly in grape, while light yellow fluorescence was observed in vacuole-like compartments only in apple. Red fluorescence could correspond to chlorophyll residues while yellow emission could be related to compounds of the flavonol class, such as quercetin

derivatives [36] [37]. A weak blue emission was also observed in the cell wall at native pH and was found to increase at pH 10, as reported for ferulic or coumaric acids [20]. These emission variations after UV excitation at 375 nm provided support for the mapping of phenolic compounds using multispectral autofluorescence imaging. However, a lower excitation wavelength was sought to access compounds in the flavanol family.

3.3. Autofluorescence after excitation at 275 nm

The autofluorescence of flavanols in fleshy fruit tissue requires excitation at a low UV wavelength, which is yet inaccessible with a common microscopy laser beam. The synchrotron beamline at DISCO allows for exciting samples in the 200-350 nm range. To limit the diffusion of phenolic compounds, apple tissue was first cryo-fixed prior to cryo-observation of multispectral fluorescence from 300 nm to 650 nm (Figure 3). Outer-cortex parenchyma fluorescence was distinguished from that of epidermal tissue, which encompasses cuticle and sub-cuticular cell layers (Figure 3). Between 327 and 388 nm, sub-cuticular cell layer fluorescence was more intense and disappeared at higher bandpass wavelength filters except, for the 600 nm filter. Intense fluorescent spots were observed inside the cell mainly with 300 and 327 nm filters, but in all cases, cells walls showed a more intense signal than cells.

To better distinguish differences in sample fluorescence, principal component analysis was applied to the collection of images obtained with the different bandpass filters. All principal component analyses performed on the three modalities presented similar loadings. For example, the loading and component images are shown in Figure 4 for representative samples of Douce Moën and Guillevic apple varieties at harvest. Fluorescence through 300, 327, 352, 412, 420 and 600 nm filters contributed mainly to components 1 and 2 and fluorescence at 300, 327 and 600 nm contributed more weakly to components 3 and 4. Most filters contributed to component 1, which corresponded to the general variations of

fluorescence intensity and explained 65.8% and 61.7% of the total variance for Douce Moën and Guillevic, respectively. The component images showed that the highest overall fluorescence intensity was mainly localized in the walls of outer-cortical cells. The second component, expressing the contribution of the 300 and 327 nm filters (positive side) in opposition to the emission filters 412 and 420 nm (negative side), contrasted the sub-cuticular cell layers. Filters 352 and 600 nm were inversely related to this second component for the two varieties. The third component pointed out the cuticle region with a relatively higher fluorescence measured using the 327 nm emission filter (in red) compared to the 300 and 600 nm filters (in blue), showing the rest of the sample. The fourth component showed a higher level of noise and a relatively higher fluorescence level associated with the 300 nm filter (in blue) in the epidermal region as well as in the cell walls of sub-cuticular cells and outer cortical cells. Thus, fluorescence distinguished three areas: the cuticle region with a relatively higher fluorescence emission using the 327 nm filter, the sub-cuticular cell layers with the 300 nm and 600 nm filters, and the walls of outer-cortical cells with the 352, 412 and 420 nm filters. The three first component images of the representative samples were combined in an RGB image to better visualize the tissue regions according to fluorescence (Figure 4). To objectivize differences observed in fluorescence distribution, normalized intensity profiles for all images of the two varieties at the different storage conditions were measured.

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Intensity profiles from the cuticular to the outer cortical tissue were obtained for the four bandpass filters discriminating the most apple tissues: 300, 327, 420 and 600 nm. As cell lumen and cell walls showed marked differences in fluorescence intensities, smoothing of each profile was realized by local regression (LOESS) at the expense of fine structures contributing to fluorescence (cells, cell walls, sub-cellular structures). The smoothed and normalized curves allowed comparison of trends in fluorescence variations along the first 400 µm under the cuticle. Examples of raw and smoothed profiles are shown for the Douce Moën

variety at harvest in Supplementary Figures 1 and 2. Considering close fruit calibres within varieties, Student's t-test at each distance point per filter showed that storage modalities had no effect on fluorescence profiles (p-value > 0.05; Supplementary Figure 3). Thus, all biological replicates (n= 9) were used to evaluate the effect of apple varieties on fluorescence intensity profiles. Significant differences were particularly noted with the 327 nm and 420 nm filters (Figure 5). For the 327 filter, an inverse gradient was observed between the two varieties with a rupture point at approximately 100 µm from the cuticle. Douce Moën showed higher fluorescence intensity than Guillevic, particularly in the first 10 µm under the cuticle but was lower than that of Guillevic 60 µm away from the cuticle. The 420 nm filter also presented two significantly different areas, first in the epidermis where the Guillevic signal was higher than that of Douce Moën and opposite to the 327 nm filter profile and second in the outer cortex (300 to the 400 µm limit of the images) where Guillevic fluorescence was lower. A maximum was observed at approximately 100 µm for Guillevic, while for Douce Moën, it was at approximately 330 µm from the cuticle. Weak significant differences were observed between Guillevic and Douce Moën in the sub-cuticular cell layers for the 300 nm filter and in the outer cortex for the 600 nm filter. These two profiles also showed similar ruptures at approximately 75-100 µm from the cuticle. The mean smoothed plot profiles (Figure 6) showed good agreement with the fluorescence distribution observed in the representative PCA reconstructed images (Figure 4). The three regions observed may be approximately delimited from these results (Figure 6) for Douce Moën to range from approximately 0 to 50 µm, 50 to 200 µm and 200 µm to 400 µm while for Guillevic, to range from approximately 0 to 60 µm, 60 to 150 µm and 150 to 400 µm. These regions may correspond to the cuticle and the sub-cuticle forming the epidermis and the outer cortex as depicted in Figure 3.

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The fluorescence emission of reference phenolic compounds allowed us to propose the possible nature of chemicals in the different tissues (Table 1). Fluorescence observed with the 300 nm filter was more likely due to proteins as this range of emission is related to the aminoacids tyrosine and phenylalanine [38]. Fluorescence through the 327 nm filter may have corresponded to the tryptophan and most likely to catechins that are predominant in apple fruit tissue. Since these compounds are flavanol monomers and oligomers, by extension such fluorescence would also reveal condensed tannins. Likewise, fluorescence at 420 nm would represent the phenolic acid family, such as hydroxycinnamic acids, while the fluorescence at 600 nm would be pigments, such as anthocyanin or chlorophyll. According to these propositions, pigments would be particularly localized in the sub-cuticular cell layers of both varieties, while flavanols including condensed tannins would be present in the epidermis and outer cortex of both varieties, though with higher intensity in the sub-cuticular cell layers of Guillevic. The distribution of phenolic acids differed markedly between the two varieties. In Douce Moën, they were present mainly in the outer cortex, while in Guillevic they were also present in the sub-cuticular cell layers. Storage conditions did not appear to affect these distributions.

4. Discussion

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The results showed that multispectral observation of auto-fluorescence provides a convenient alternative for localizing phenolic compound families in fleshy fruit tissue with limited sample preparation and artefact introduction compared to classical resin embedding and staining methods. Adaptation of the cryogenic procedures developed for cryo-LSCM [18] to the wide-field TELEMOS microscope of the DISCO beamline at the SOLEIL synchrotron facility allowed observation of fluorescence from 275 nm, which excited flavanols and condensed tannins together with other phenolic compounds in well-preserved apple tissue with limited metabolite diffusion. The obtained images confirmed the efficiency of the

cryogenic process as intracellular content appeared filled, though of lower fluorescence intensity compared to cell walls. Cryo-observation also improved fluorescence intensity [26], likely by reducing fluorescent bleaching due to the temperature dependency of quencher diffusion [28]. This phenomenon was particularly helpful in establishing the qualitative distribution of fluorescent compounds in fruit tissue.

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Fluorescence tended to indicate phenolic compounds within cell walls. However, most phenolics are localized within cell organelles, vacuoles or the cytoplasm [39, 40] with fewer in cell walls. In apple, phenolic compounds have not been described as part of the cell wall polysaccharide composition [41] [42]. Some authors [43] suggested that tannin accumulation takes place in the vacuole, but their polymerization appeared near the cell walls. As apple fruit cells are highly hydrated, ice crystals formed during cryo-fixation of the waterfilled vacuoles [44] may have pushed fluorescent organelles and cytoplasmic materials close to the cell walls. As a consequence, part of the cell wall fluorescence observed with the 10x magnification lens (resolution: 1.24 µm / pixel) may have resulted from nearby intracellular fluorescent compounds/organelles. However, the blue emission observed by LSCM in the cell wall of fresh apple and grape tissue and its increased fluorescence at basic pH supported the presence of hydroxycinnamic acids in cell walls. This observation was realized at higher magnification (20x, resolution of 0.63 µm / pixel) than that of the TELEMOS setup and excluded the potential of ice crystal effects on cell wall fluorescence. Hydroxycinnamic acids have already been reported in apple fruit mostly in vacuoles due to their characterization in the juice [45] [46, 47], but they are also known to be present in plant cell walls [48]. Their low concentration in apple cell walls likely prevented their detection by classic analytical techniques. However, further work is needed to assess the nature of these compounds and whether they are esters of cell wall polysaccharides, as reported for other plants [20] [49, 50].

The cuticle was the most fluorescent zone of apple and grape fruit tissues observed by LSCM (Figure 2). The observation of cuticle autofluorescence was already reported in several fruits, including apples and grapes and may be related to high concentrations of proteins and phenolic acids [35] [51]. Though anthocyanins, flavonols, flavanols (including condensed tannins), hydroxycinnamic acids and proteins were essentially reported in the epidermis region [52-54], they were more rarely localized in sub-cuticular cell layers [8, 10, 14]. The high concentration of phenolic compounds in the sub-cuticular cell layers is probably related to defence against pathogens [55] or light damage to tissues. The firmer mechanical resistance of grape and apple skins compared to their flesh [56, 57] may also be associated to the important presence of phenolic compounds in the skin. Indeed, phenolic compounds and, in particular, flavonoids were proposed to increase the rigidity of the cutin matrix in the cuticle of ripe tomato [58]. Comparatively, excitation with DUV at 275 nm revealed a lower cuticle autofluorescence. Although flavonoids and phenolic acids, the main cuticle phenolic compounds, have large excitation and emission spectra (Supplementary Figure 4), 275 nm excitation was not optimal to yield high autofluorescence intensities. Though such a wavelength was beneficial for exciting flavanols and proteins fluorescence, it appears that flavanols are in low concentration - if any- within the cuticle but are present in the first cell layers underneath (Figure 3).

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The red-skinned Douce Moën and the yellow-skinned Guillevic cider apples were used as model fruits to help distinguish phenolic compound locations by their autofluorescence. Douce Moën is known to be richer in hydroxycinnamic acids and in monomeric tannins, such as (+) catechin, (-) epicatechin and procyanidin with lower tannin degrees of polymerization compared to Guillevic [59]. Anthocyanins were not characterized in the two varieties, but the more colourful Douce Moën skin evidenced higher anthocyanin and/or pigment contents than that of Guillevic. Based on distinct fluorescent emissions of

reference compounds after 275 nm excitation, autofluorescence in Douce Moën and Guillevic cryo-sections allowed the detection of pigments in the sub-cuticular cell layers in agreement with the known flavanols, anthocyanins and chlorophyll concentrations in apple skin [60, 61]. The main distribution of tannins in the sub-cuticular area together with that of phenolic acids in the outer cortex parenchyma may be related to the ratio reported for procyanidin (PC) and hydroxycinnamic acid (HC) concentrations in the fresh peel (HC/PC: 0.68) and parenchyma (HC/PC: 1.29)[62]. This different distribution was not reported in the literature for grape in which flavanols were mainly localized in the epidermis and in the seeds although hydroxycinnamic acid content was higher in skin than in pulp [16, 54, 63].

Changes in phenolic compounds distribution were not observed with storage modalities that were aimed at impacting apple ripening. This result is at odds with studies showing phenolic compound variations during ripening of fruit [7], such as apple [52, 64] and grape [10, 65]. The lack of autofluorescence discrimination between ripening stages may be related to the small tissue area observed. Instead, the present study evidenced a variation in phenolic distribution between apple varieties. This may be related to different tissue thicknesses as for both varieties, different rupture points in phenolic distributions were observed along distance profiles from the cuticle. These ruptures could correspond to different cell layers in the epidermis and the outer cortex. In particular, a rupture was discernible in fluorescence profiles at an approximately 100 µm depth from the skin samples. Apple histology studies have shown changes from smaller cells in the epidermis to larger cells in the outer cortex parenchyma tissue at an approximately 100-µm depth from the surface of mature fruit [66-68]. Thus, differences in phenolic compound distribution between the two varieties could reveal genetically distinct thicknesses of epidermal cell layers as well as different secondary metabolism in these tissues.

5. Conclusion

The cryo-method used here could benefit from further developments in sample preparation and means of cryo-multispectral fluorescence observations. Applying microwaves during cryo-fixation [69] to avoid ice crystal growth combined with the use of confocal microscope stages and lenses for high-resolution cryo-observations [26, 27] is expected to improve the localization of phenolic compounds in the cell and cell wall. Nevertheless, multispectral fluorescence cryo-imaging at 275 nm excitation allowed the distinction of three major classes of phenolic compounds in tissue under *in-vivo-like* conditions without any specific dyes. The detection of hydroxycinnamic acids in apple and grape cell walls requires their identification and linkage in the walls. Coupling multispectral analysis based on LSCM observations in the UV-visible wavelength and synchrotron deep-UV analysis could provide a convenient way to extend the range of accessible autofluorescent phenolic compound distributions in the same sample tissue section. These observations will enable other studies on the mechanical behaviour resulting from the presence of these compounds in fruit tissues in the context of fruit development, quality and processing.

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Appendix A: Supplementary data

Supplementary data associated with this article can be found, in the online version.

Legend to Figures

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431 Figure 1: Generic phenolic compound structures. -R may correspond to -H, -OH or -OCH₃ 432 and –X may correspond to –H in monomeric form or other flavanol units in polymeric form (Vermerris and Nicholson 2006). 433 434 Figure 2: Local retail store fresh apple and grape autofluorescence at natural pH and at 435 pH=10.1 after excitation at 375 nm. The fluorescence emissions recorded are 400-450 nm (blue), 500-530 nm (green), 560-600 nm (yellow), and 630-700 nm (red). C: cuticle, CW: cell 436 437 wall, Chl: chlorophyll residues, Flv: flavonols. Scale bar: 20 µm. 438 Figure 3: Frozen apple autofluorescence for one replicate of Douce Moën and Guillevic at 439 harvest after excitation at 275 nm. Tissue regions are drawn on the right image resulting from 440 a combination of whole Guillevic filter emission images. C: cuticle, CW: cell wall, Scale bar: 40 μm. 441 442 Figure 4: Principal component analysis on the six emission filter images for one replicate of Douce Moën (blue character and arrows) and Guillevic (black character and arrows) at 443 444 harvest and the PCA images of the four components. On the component images, blue and red 445 pixels represent negative and positive contributions of the emission filters to the component. RGB reconstructed images from PCA components were built from dimension 1 (blue), 446 447 dimension 2 (red) and dimension 3 (green). C: cuticle, CW: cell wall 448 **Figure 5**: Smoothed (LOESS method: $\alpha = 0.4$) and normalized average plot profiles of Douce 449 Moën (DM) and Guillevic (GU) signal acquisition for emission filters 300 nm, 327 nm, 420 450 nm and 600 nm after excitation at 275 nm. The dotted lines represent the 95% confidence 451 interval. The coloured areas correspond to significant differences between DM and GU with a p-value < 0.01 by Student's test (n = 9). 452 **Figure 6**: Superposed smoothed (LOESS method: $\alpha = 0.4$) and normalized average plot 453

profiles of Douce Moën and Guillevic signal acquisition for emission filters 300 nm (grey

line), 327 nm (green line), 420 nm (blue line) and 600 nm (red line) after excitation at 275
nm. The profiles were plotted along the sample section distance from the outer cuticle to 400
µm deep inside the tissue.

Table 1: Fluorescence excitation/emission maxima of phenolic compounds registered by spectrofluorometry: regular case: literature, **bold case**: this study.

Compounds	λ _{max excitation} (nm)	λ _{max emission} (nm)	Compounds	λ _{max excitation} (nm)	λ _{max emission} (nm)
Emission range (<350 nm)			Emission range (400-500nm)		
Protein group			Phenolic acids group		
Phenylalanin[70]	258	285	Ferulic acid	310/370	420
Tyrosin [70]	276	302	p-Coumaric acid	350	420
Emission range (300-400 nm)			Caffeic acid	270/360	440
Flavanols group			Chlorogenic acid	360	460
(-) Epicatechin	290	320	Others compounds		
(+) Catechin	290	320	Kaempferol[21]	268	422
Procyanidin B1	285	320	Rutin	300	460
Procyanidin B2	295	320	Emission range (500-600nm)		
Epigallocatechin	275	320	Flavonols group		
Others compounds			Anthocyanins[22]	300/410	360/420/530
Tryptophan [22]	280	357	Cyanidin hydrate	278	550
2,5- Dihydroxybenzaldehyde[21]	278	360	Quercetin-3-O-glucoside	260/410	510
Syringic acid	310	360	Quercetin	420	520
Vanillic acid[21]	278	360	Quercetin galactoside	295	600
Gallic acid	320	370	Others compounds		1
Myricetin[21]	268	370	Arbutin	420	530
Resveratrol	310/360	390	Phloridzin	250/330	540
			Chlorophyll/chloroplast[37]	360	680

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Hydroxybenzoic acids

Flavanols

Phenolics acids

Stilbenes

Hydroxycinnamic acids

HO

HC

Resveratrol

Flavonoids

Flavonols









