Chlorophyll Breakdown as Seen in Bananas: Sign of Aging and Ripening – A Mini-Review

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Abstract
The ripening of bananas is seen by a characteristic change of their color from deep green to bright yellow. Likewise, their over-ripening and eventual rotting are accompanied by the appearance of an unappetizing brown. Chlorophyll breakdown is a major contributor to the visual signs of these processes in bananas.Outlined here are the basic structures of chlorophyll catabolites in higher plants, with particular reference to ripening and aging bananas. In these fruits, unique fluorescent chlorophyll catabolites accumulate and give rise to their fascinating blue luminescence.

Introduction
The seasonal appearance and disappearance of the green plant pigments in deciduous trees and in fruits belong to the most colorful and fascinating natural phenomena. Indeed, chlorophyll metabolism probably is the most visual sign of life on earth, and can even be observed from outer space [1]. However, only within the last 2 decades has chlorophyll breakdown in plants begun to reveal some of its mysteries (fig. 1) [1–6].
Hv-NCC-1 (1) (fig. 1) [1]. Hv-NCC-1 is a linear tetrapyrole, which is colorless due to deconjugation of the four pyrrole units. Its structure thus gave the first hints as to the changes that happen to chlorophylls during their breakdown in higher plants: it indicated oxygenolytic opening of the macrocyclic ring of chlorophyll a at the northern meso-position and loss of both the central magnesium ion and the lipophilic phytol group.

In the meantime, NCCs have been identified in senescent leaves from a variety of vascular plants and their basic structural pattern has been established: it involves four deconjugated pyrrolic units in a substituted formyl-bilane, whose formyl group is derived from the former meso-carbon of chlorophyll a. In solution, NCCs display a characteristic UV/Vis absorption maximum near 320 nm, which is due to the formyl pyrrole moiety [5, 12].

Blue Fluorescent Catabolites of Chlorophyll

In the course of the elucidation of the chlorophyll breakdown path, the eventual identification of the direct precursors of the NCCs as 'fluorescent' chlorophyll catabolites (FCCs) was a further major breakthrough [13]. Minute amounts of fluorescent compounds were observed early in senescent leaves, due to an easily detected blue emission (near 450 nm) [10]. However, they occurred only fleetingly during leaf senescence and were suggested to be intermediate chlorophyll breakdown products. The first structural characterization of such a fluorescent compound, of primary FCC (pFCC) (fig. 1), indeed confirmed it to be a chlorophyll catabolite [13]. At the same time the structure of pFCC suggested it to be the direct ('primary') product of an elusive precursor that would be colored red [1, 13]. The hypothetical red chlorophyll catabolite was prepared by chemical synthesis [14], and its intermediary role as the elusive red precursor of pFCC could be confirmed (fig. 1) [1, 3, 15].

Typical natural FCCs do not accumulate as they convert to NCCs rapidly under physiological conditions. When pFCC or its natural epimer, epi-pFCC [16], were exposed to slightly acidic conditions, these two FCCs were completely converted to a specific NCC each [17, 18]. This thermodynamically favored, stereoselective chemical isomerization depended critically on the presence of the free propionic acid function of the pFCCs and it was inhibited in the related (synthetic) methyl esters of the pFCCs [18]. The isomerization of natural FCCs to NCCs was thus suggested to occur spontaneously in the acidic milieu of the vacuoles, and without participation of an enzyme [3, 5, 17], as was, in fact, considered earlier, when the first structure of a pFCC was elucidated [13].

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**Fig. 1.** Outline of chlorophyll breakdown in senescent higher plants with reference to (suggested) relevant enzyme activities [1, 3–6]. Chlorophylls [Chl a (R = CH₃) or Chl b (R = CH = O)] are degraded by catabolic enzymes in the chloroplast (a, b = Chl b reductase, hydroxymethyl-Chl a reductase; c, d = magnesium dechelatase, pheophytinase) to pheophorbide a (Pheo a) and then (by e = Pheo a oxygenase) to 'red' chlorophyll catabolite (RCC), which is reduced (by f = RCC reductase) to the pFCC. FCCs are exported from the chloroplast into the cytosol, where they may be further modified enzymatically (g). Most FCCs are then imported into the vacuole, where (h) their isomerization to NCCs occurs, such as to Hv-NCC-1.
Common knowledge suggests the development of yellow and red colors during ripening of fruit as an indirect sign of the breakdown of chlorophyll (fig. 2). However, the remains of the green plant pigments in ripe fruit have until recently been unknown [9]. Analysis of fresh extracts of peels of ripe apples of the ‘Golden Delicious’ brand and of ‘Williams’ pears revealed the presence of two NCCs. The same two NCCs were also detected in senescent leaves of the two fruit trees (see formula shown in fig. 2) [9], suggesting a common biochemical path of chlorophyll breakdown in senescent leaves and in ripening fruit. NCCs were thus indicated to be the typical tetrapyrrolic remains of the chlorophylls in higher plants [4, 9].

**Blue Luminescence of Ripening Bananas**

The sweet ‘dessert’ banana (cavendish cultivar of Musa acuminata) is one of the most important fruits worldwide. Degreening is a clearly visible indicator to distinguish between delicious sweet and unripe harsh stages of bananas as well as of many other fruits. Surprisingly, the biochemical pathway of chlorophyll degradation in the
yellowing peel of bananas differs from that in ripening apples and pears [7–9]. Instead of NCCs 'chemically stabilized' fluorescent catabolites ('persistent' FCCs) occur in a large variety and accumulate in the peels of freshly ripe (yellow) bananas. As depicted in figure 3, the accumulation of FCCs in the outer regions of the peel causes yellow bananas to glow blue, when observed under UV light (at 366 nm). The natural luminescence shows an emission maximum at 447 nm. Its intensity is highest for fresh ripe, bright yellow bananas and decreases during the further aging process of the fruit [see fig. 4 in ref. 7]. Intact bananas and extracts of yellow banana peels, as well as solutions of purified FCCs, all showed similar fluorescence behavior (fig. 4a), suggesting FCCs as the source of the blue glow [5, 7]. As strongly fluorescing compounds, FCCs are also easily spotted using chromatographic analysis in combination with fluorescence detection.

Spectroscopic structural analysis of FCCs in yellow banana peels revealed their unusual structures and the chemical basis for their 'stabilization' [7]. The most abundant FCC in bright yellow banana peels, Mc-FCC-64, was found to be esterified by a daucyl group (fig. 4, the daucic acid moiety is highlighted in red). Chlorophyll breakdown in bananas thus deviates from that revealed earlier in senescent leaves (fig. 5). Accumulation of FCCs in yellow banana peels can be rationalized by chemical arguments: the lack of a free propionic acid side chain hinders the easily occurring acid-catalyzed conversion of typical FCCs to NCCs, and the esterified FCCs were thus classified as 'persistent' FCCs [17, 18]. Clearly, this biosynthetic stabilization of FCCs against isomerization to NCCs and the resulting blue luminescence of the ripening banana fruit are striking new features of chlorophyll breakdown [7, 8].
The stunning results with banana fruit induced further studies with senescent leaves of bananas (M. acuminata). As could be observed under UV light, banana leaves also showed blue luminescence, which was again found to be due to accumulation of FCCs. Indeed, the FCCs from degreened banana leaves (Ma-FCCs) clearly differed from their relatives in the fruit peels (Mc-FCCs), although they were identified again as ‘persistent’ FCCs, i.e. FCCs with a (complex) esterification of their propionate function (fig. 5) [19]. These results now contrast with the earlier notion of a common path of chlorophyll breakdown in senescent leaves and ripening fruit [9].

Blue Halos of Cell Death in Aging Bananas

‘Senescence-associated’ dark spots appear on the peel of bananas during a short period of aging that follows upon the ripening and yellowing, as typical signs of post-harvest deterioration [20, 21] (fig. 5). The diagnostic senescence-associated spots appear on the peel accompanied by a fade-out of the natural blue luminescence [7]. At the same time remarkable intense, blue luminescent rings begin to develop around these dark spots, which are sections of the peel containing dead cells (fig. 6) [8]. Peels of bananas are very sensitive to physical damage of their surface, and the ripening process is also sensitive to lower temperatures. Only senescence-associated dark spots are surrounded by blue rings, and are thus diagnostic visual signs of banana overripening [8]. In contrast, dark features due to mechanical damage to the surface do not show the formation of fluorescent areas (fig. 6).

In vivo fluorescence analysis of these halo-like rings (fig. 6) suggested the presence of fluorescing chlorophyll catabolites due to a typical fluorescence emission maximum at 447 nm, similar to that of isolated FCCs. Quantification using HPLC analysis verified this unexpected accumulation of fluorescent chlorophyll degradation products. Elevated levels of FCCs were found within the bright fluorescent areas, whereas considerably lower FCC levels were detected in dull yellow areas. In the dark spots the levels of FCCs were still smaller, all consistent with quantitative fluorescence measurements with intact bananas [7]. Structure elucidation revealed the main contributor to this remarkable blue luminescence to be another ‘persistent’ FCC, called Mc-FCC-49 (fig. 6, inset). Mc-FCC-49 is the product of a further enzyme-catalyzed ‘hypermodification’ of Mc-FCC-56, the major fluorescent catabolite from the yellow banana peel (fig. 5, 6).

Dark senescence-associated spots are known to originate from near the stomata on the ripening banana peel [22], and are presumed to arise from spontaneous oxidative processes [20]. Detailed microscopic analyses confirmed the presence of the cellular remains of stomata in the center of the ‘dark’ spots and revealed the blue rings to mark an area on the peel surface in which the cells were still intact and ‘alive’, but to be considered ‘senescent’ [8].

Emergence and disappearance of blue halos could be studied noninvasively, and in time- and 2-dimensional space-resolved fashion by in vivo fluorescence measurements (fig. 7) [8]. Indeed, the selective accumulation of specifically ‘hypermodified’ FCCs suggests a physiological benefit from the prolonged persistence of these linear tetrapyrroles and calls for further investigations of their fate. On the one hand, the phenomenon might be of relevance for helping to inhibit the decline of vital functions of the fruit [23]. Hence it may prove to be a helpful non-invasive, molecular tool to study underlying cellular aging processes. On the other hand, fruit-eating animals could have learned through survival pressure [24] to see the blue bioluminescence of bananas as a noticeable signal of fruit ripeness.

Fig. 6. Typical signs of postharvest deterioration of a banana peel observed under 366-nm UV light (top) and white light (bottom). Senescence-associated dark spots exhibit typical ‘blue luminescent halos’. In contrast, mechanical damage causes dark surface areas that have no ‘halos’. The chemical formula depicts the ‘hypermodified’ FCC Mc-FCC-49, which accumulates in ‘blue luminescent halos’. The propionate ester group is marked in red, while the glucoside moiety on ring B is highlighted in yellow.
Chlorophyll Catabolites as Antioxidants

Chlorophyll degradation was, first of all, interpreted as a detoxification process [6], a part of a recycling strategy of higher plants to recover essential minerals (such as magnesium and reduced forms of nitrogen). Moreover, a recent investigation of chlorophyll catabolites in fruit [9] revealed remarkable antioxidant properties of NCCs. One of the ‘fruit’ NCCs found in ripening apples and pears [9] was tested in a standard auto-oxidation experiment designed for the analysis of bilirubin [25]: the rate of formation of hydroperoxides of linoleic acid was significantly reduced in the presence of the NCC. The (antioxidative) peroxo-radical scavenging effect of the tested NCC (for the formula see fig. 2) was only slightly inferior to that of bilirubin [25]. These results were of particular interest to us, because NCCs are structurally related to the tetapyrrolic heme breakdown product bilirubin and other heme-derived natural linear tetrapyrroles (fig. 8) [26]. Bilirubin was shown to be an antioxidant [25] and a cytoprotective component, relevant in the reduction of coronary heart diseases, retinal damage and cancer mortality [27]. The availability of the NCCs in plant-derived nutrition, as now documented for apples and pears, calls for their consideration as being of physiological interest in humans (and higher animals), and not as mere degradation products of the phototoxic green plant pigments.
These findings may even give a new twist to the meaning of the old saying ‘an apple a day keeps the doctor away’ [4, 5].

Conclusions

Chlorophyll breakdown provides a visually pleasing sign of ripening and of senescence in bananas [7, 8]. It has been apostrophized as a necessary ‘detoxification process’ of the green plant pigment during senescence [6]. This worldwide controlled degradation of the motor of the plant’s basic metabolism is an intriguing phenomenon. One may specifically wonder how metabolism in the ripening fruit may continue to function (nearly) without chlorophyll. Of course, the soft fruit eventually undergoes a further transition to senescence and cell death. However, the degreened tetrapyrrolic remains of the chlorophylls are distantly related to the heme-derived bilins, and chlorophyll catabolites may have relevant physiological roles in bananas and other fruits. Their presence in the nutrition of humans and frugivorous animals may also call for interest in their broader (physiological) effects. In addition, FCCs are luminescent molecular markers of ripening and cell death that may be useful for non-invasive in situ analysis of (fruit) ripening and senescence in higher plants.

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