Chlorophyll Breakdown – How Chemistry Has Helped to Decipher a Striking Biological Enigma

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Dedicated to Professor Franz-Peter Monforts on the occasion of his 70th birthday.

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Abstract How the fall colors arise and how chlorophyll (Chl) breakdown occurs in higher plants has remained enigmatic until three decades ago. Fundamental insights into this fascinating puzzle have been gained, meanwhile, by basic contributions from plant biology and chemistry. This short review is a personal account of key advances from synthetic, mechanistic, and structural chemistry that led to the discovery of the bilin-type Chl catabolites and helped elucidate the metabolic processes that generated them from Chl.

1 Introduction

The appearance of the fall colors is an enchanting and puzzling phenomenon. The widespread disappearance of chlorophyll (Chl) in autumn and the re-greening of the vegetation in spring are probably the most visual signs of life, observable on Earth from outer space.1 Indeed, the annual apparent ‘recycling’ of the green pigment Chl is a massive biological process involving about 1000 million tons, worldwide, and occurring by seasonally alternating de novo Chl biosynthesis and Chl degradation.2 Strikingly, until about 30 years ago Chl seemed to disappear without leaving a trace, as non-green products of Chl breakdown remained elusive.2,3 In fact, all searches for genuine Chl catabolites were futile, since they concentrated on the detection of colored remains of the Chls. However, as is now well known, Chl breakdown products in higher plants accumulate as colorless linear tetrapyrroles, primarily.1,4–7 All the same, the bilin-type Chl catabolites, named phyllobilins (PBs),8 turn out to be remarkably related, structurally, to bilins,9 the colored products from heme breakdown.10

2 Discovery and Structure Elucidation of a First Non-Green Chl Catabolite

Chl breakdown has for a long time been considered to play a particular role in the recuperation of nitrogen.3 First traces of non-green Chl breakdown products were traced by Matile, Thomas and coworkers by comparing the pigment pattern of senescent leaves of wild type Festuca pratensis with those in ‘stay green’ mutants of this grass.11 Remarkably, colorless compounds could be spotted in the wild type that were absent in the green mutant, and which were presumed to represent Chl catabolites. These polar compounds were called ‘rusty pigments’, as they rapidly converted into rust-colored products.12 We learned to isolate one such ‘rusty pigment’ without degradation and color formation, and elucidated its chemical constitution in 1991 with the help of heteronuclear NMR spectroscopy and soft ionization mass spectrometry.4,13 It turned out to be fruitful, to solve
the desired structure directly by purely spectroscopic means,\textsuperscript{4} rather than submitting this polar Chl catabolite to chemical modifications, e.g., in order to decrease its polarity.

![Scheme 1](image1)

The colorless 'rusty pigment' from de-greened leaves of barley (*Hordeum vulgare*) was unambiguously revealed to be a bilane-type linear tetrapyrrole that carried structural hallmarks of the Chls, such as their characteristic ring C/E-portion.\textsuperscript{8} It was classified as a nonfluorescent Chl catabolite (NCC) and given the phenomenological name Hv-NCC-1 (1).\textsuperscript{14} More recently, it was classified semisystematically as a 1-formyl-19-oxo-16,19-dihydrophyllobilane,\textsuperscript{5,8} see Scheme 1. The structure of 1 indicated oxidative ring opening of the Chl macrocycle at the 'northern' meso position, between C4 and C5, thus, representing a 4,5-secophytoporphyrinoid.\textsuperscript{4,13} This cleavage site contrasted with all expectations from Chl chemistry,\textsuperscript{15} but turned out to be strikingly reminiscent of the site of ring opening in heme breakdown.\textsuperscript{16} However, in contrast to the typical colored bilins from heme breakdown,\textsuperscript{9,13} the meso-carbon C5 of Chl a was retained in the formyl group of the NCC 1, and its three remaining meso positions were saturated. Additional peripheral polar functionality of 1 pointed to puzzling catabolic reactions in its formation, raising further questions to its biochemical formation and to the general relevance of its structure for the general problem of Chl breakdown in higher plants.

![Scheme 2](image2)

3 Structure Elucidation of Fleetingly Existent Blue-Fluorescent Chl Catabolites

The repeated observation of fleetingly existent blue-fluorescent compounds alongside of the NCCs in extracts of senescent leaves suggested their relevance in Chl breakdown.\textsuperscript{17} In vitro experiments by Hörtensteiner and Matile showed the requirement for molecular oxygen and identified the Mg-free and dephytylated Chl derivative phophorlbide a (Pheo a, 6) as a precursor for the presumed Chl catabolites.\textsuperscript{18} From a preparation with Pheo a as the substrate by using an enzyme active extract of senescent leaves of oil seed rape (*Brassica napus*), a rather instable blue-fluorescent compound became available in small quantities, believed to be a Chl catabolite. By spectroscopic means its structure was, again, revealed as a linear tetrapyrrole,\textsuperscript{19} closely related to 1, and to three polar *Brassica napus* NCCs that had meanwhile been found in senescent cotyledons of oil seed rape\textsuperscript{20} (typical UV/Vis absorption spectra are collected in the reviews\textsuperscript{5,8}). The fluorescent compound was, therefore, phenomenologically named *Bn*-FCC-2 (5) and classified as a fluorescent Chl catabolite (FCC) (see Scheme 2).\textsuperscript{16} The chemical constitution of 5 supported its hypothetical role as a biological precursor of the *Bn*-NCCs 2–4. The molecular formula of the FCC 5 also indicated a close relationship to 6, involving the mere formal incorporation of two oxygen atoms and four hydrogen atoms in the course of the presumed formation of 5 from 6. Hence, the moderately polar FCC 5 was deduced to represent a straight forward

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Biographical Sketch

Bernhard Kräutler studied chemistry at the ETH in Zürich, where he received his PhD working with Prof. Albert Eschenmoser. After postdoctoral studies with Prof. Allen J. Bard (University of Texas, Austin) and with Prof. Nicholas J. Turro (Columbia University, New York City) he returned to the ETH to start his own research group. In the fall of 1985 he was a visiting Professor at the Roger Adams Labs of the University of Illinois. In 1991 he became Full Professor of Organic Chemistry at the University of Innsbruck, where he has been Professor Emeritus since October 2015. His current research interests include the chemistry and chemical biology of chlorophyll and vitamin B12.
(‘primary’) product of the breakdown of 6, and was assigned the specific role as ‘primary’ FCC (pFCC).19 As the relative configuration of the three stereocenters of pFCC (5) at C8, C12, and C13 was the same as that in Pheo a (6), their absolute configuration was also inferred to be retained. Hence, pFCC (5) is a 8S,12S,13S,16Z,19-oxo-12,13,16-tetrahydrophyllobilene-b.5,8

However, in extracts of senescent leaves of sweet pepper (Capsicum annuum) a different fluorescent Chl catabolite (Ca-FCC-2) was observed, which was identified as an isomer of pFCC (5). Ca-FCC-2 was deduced to display the same (relative) stereochemistry at C8, C12, and C13, but to differ from pFCC (5) by the configuration at C16. Hence, Ca-FCC-2 was named epi-pFCC (epi-5), with the semisynthetic name 8S,12S,13S,16Z-1-formyl-19-oxo-12,13,16-tetrahydrophyllobilene-b.5,21 The absolute configuration of C16 of pFCC (5) or of epi-pFCC (epi-5) is not established, so that it is classified provisionally as ‘normal = n’ in 5, or as ‘epimeric = epi’ in epi-5, and in their respective catabolic descendants.5

The structural analysis suggested the formation of ‘primary’ FCCs from Pheo a (6) to involve more than one enzyme and to require the existence of an intermediate, probably less saturated at its ‘western’ meso position than 5 or epi-5.19 In analogy to the red excretion products of Chl breakdown of the alga Auxenochlorella protothecoides, characterized structurally in Gossauer’s group,22 related red linear tetrapyrroles were now taken into consideration as so far elusive intermediates of the Chl breakdown in higher plants (see Scheme 3).19,23

4 The Red Chl Catabolite – Key Ring-Opened Tetrapyrrole Accessed by Partial Synthesis

Since a red Chl catabolite was unknown in higher plants, we set out to prepare the likely candidate by partial synthesis from the methyl ester of Pheo a (6), in order to help test the suggested intermediacy of such a red bilin-type Chl catabolite in higher plants. Our synthesis of the presumed red Chl catabolite (RCC, 7) roughly followed a methodology, developed by Iturraspe and Gossauer, for the preparation of the corresponding analogs from the green alga.24 For this purpose, 6 was converted into the green Cd(ll)-methyl-photophorbidate 8, which was subsequently subjected to photo-oxidation at –40 °C, furnishing the corresponding 4,5-secoporphyrinoid Cd complex 9 in about 32% yield (see Scheme 4).23 Reduction of the latter with sodium borohydride at room temperature and work-up of the reaction mixture with dilute hydrochloric acid furnished the red 4,5-secoporphyrinoid Me-7 (as a 3:1 mixture of 8S-epimers) in 72% total yield. Under these conditions, (i) the electrophilic meso position between the ‘eastern’ rings of the 4,5-secoporphyrinoid Cd complex 9 was reduced by the hydridre reagent selectively, and (ii) the subsequent acid treatment removed the Cd ion quantitatively. The main isomer Me-7 was isolated by semipreparative HPLC and shown to exhibit the natural β-configuration at C8, as deduced from NOE investigations.23 The semisynthetic dimethyl ester Me-723 was also identified with a dimethyl-ester isolate of a red Chl catabolite from Auxenochlorella protothecoides.25 By using pig liver esterase, Me-7 was hydrolyzed highly regioselectively to the remarkably stable RCC (7),23 the (then) elusive key red catabolite and precursor of pFCC (5) and epi-pFCC (epi-5).

Samples of semisynthetic RCC (7, 8S,12S,13S,16Z,19-oxo-12,13,16,19-tetrahydrophyllobiladiene-b,c)5 were tested for their presumed central role in Chl breakdown in leaves. Indeed, an extract from senescent barley leaves containing active stroma enzymes converted 7 stereoselectively into pFCC (5). Predominant conversion of 7 to epi-5 was, likewise, observed with a corresponding
preparation obtained from sweet pepper.26 Furthermore, traces of RCC (7) were now detected in an aerated incubation of Pheo a with chloroplast membranes from senescent sweet pepper cotyledons washed free of stroma components. These in vitro experiments confirmed the relevance of two enzymatic steps involved in the conversion of Pheo a to pFCC (5), with RCC (7) representing the original enzyme-bound intermediate with a 4,5-seco-porphyrinoid bilin-type structure.5,26 Meanwhile, the two inferred enzymes have been identified by Hörtensteiner and coworkers as the non-heme iron-dependent enzyme Pheo a oxygenase (PAO)27 and the cofactor-free ferredoxin-dependent RCC reductase (RCCR).6,28 In fact, two plant-specific lines of RCCRs have been identified by Hörtensteiner and coworkers as the PAO/phyllobilin pathway.5,31,32

Samples of semisynthetic red Chl catabolite (RCC, 7) and its methyl ester Me-7 were also used for exploring their chemical conversion into the corresponding 'primary' FCCs or their methyl esters. Electrochemical reduction of semisynthetic Me-7 in a deoxygenated methanolic solution at –1.3 V vs. a 0.1 N calomel electrode produced two major FCC methyl ester fractions, besides yellow regioisomers.33 The separated fractions of the two pure semisynthetic FCC methyl esters were assigned the structures of methyl esters of pFCC (5) and epi-pFCC (epi-5), on the basis of NMR-spectroscopic analyses. The free catabolite RCC (7) was, likewise, reduced electrochemically at –1.3 V vs. 0.1 N calomel electrode in a deoxygenated methanolic solution containing phenol, which produced roughly equal amounts of the two ‘primary’ FCCs.5,34 with a chromophore structure reminiscent of functionality present in some heme-derived bilins9 (see Scheme 6). Because of their tendency to isomerize in weakly acidic solution (see below), the epimeric FCCs needed to be isolated and stored with due precautions. These electrochemical experiments not only opened up a short preparative route to FCCs (or phyllolumobilins, PluBs), but they also represented first model reactions for the puzzling enzymatic conversion of an RCC (7) to a ‘primary’ FCC by the cofactor-free RCCRs. Indeed, our studies suggested a mode of action of RCCRs requiring an external supply with electrons and protons, as deduced meanwhile in a range of bilin reductases,25 including RCCR from Arabidopsis thaliana.36

5 Synthesis of ‘Primary’ Fluorescent Chl Catabolites by Reduction of Red Chl Catabolite

Comparison of the core structures of FCCs and of natural nonfluorescent Chl catabolites (NCCs or phyllobilanes), indicated them to represent pairs of constitutional isomers and suggested the existence of an isomerization path from

| Scheme 5 | Enzymatic conversion of Pheo a in the presence of 18,18O2 by the monooxygenase PAO and by the RCC reductase of type RCCR-1 produced 18O-pFCC. |

| Scheme 6 | Electrochemical reduction of the red Chl catabolite (RCC) in MeOH furnished a roughly (1:1) mixture of the colorless PBs pFCC (5) and epi-pFCC (epi-5), and a yellow constitutional isomer, as a relevant side product. |
FCCs to the corresponding NCCs. This hypothesis was first tested with a sample of authentic 5. Indeed, in aqueous solutions the ‘primary’ FCC 5 isomerized stereoselectively to the moderately polar NCC 5, named 8,10R,16epi-1-formyl-19-oxo-16,19-dihydrophyllobilane (see Scheme 7). The phyllobilane 5 was identified with a natural NCC isolated in small amounts from senescent leaves of the deciduous tree Cercidiphyllum japonicum, and named phenomenologically as Cj-NCC-2.37

The crucial stereochemical outcome of the FCC to NCC isomerization, which installs the newly saturated C10 with spontaneous chemical process to account for NCC formation under physiological conditions in the acidic vacuoles of the plant cell.37 A sequence of tautomerization and epimerization steps was proposed to account for this reaction, which is driven thermodynamically by the isomerization of the characteristic FCC chromophore into the more stable one of an NCC.37 Interestingly, the saturated meso position C10 in natural NCCs appears to conform to a common R-configuration (as derived from their CD spectra), supporting the general relevance of the isomerization path activated by the propionic acid function of typical natural FCCs.5 Indeed, in agreement with their tendency to convert spontaneously to their (more stable) NCC isomers, typical natural FCCs are only fleetingly existent in senescent leaves as intermediates of Chl breakdown.6

In contrast to pFCC and epi-pFCC, their FCC methyl esters Me-5 and Me-epi-5 were persistent and required strong acid (trifluoroacetic acid), in order to promote their rather stereounselective isomerization to NCC methyl esters. The theoretically complete set of four stereoisomers was obtained, i.e. Me-10 and Me-epi-10, and their C10 epimers Me-ent-10 and Me-ent-epi-10. Thus, the methyl esters of the so far unknown type of enantiomeric NCCs (Me-ent-10 and Me-ent-epi-10) was obtained, as well as the methyl ester (Me-epi-10) of the natural NCC named Cj-NCC-2 (see Scheme 8).34 As observed elsewhere, the chiroptical properties of the NCCs were dominated by the effect of the absolute configuration at C10.34

On the basis of the list of roughly thirty structurally different natural NCCs, known at present,5 a corresponding number of natural FCCs would, hence, be expected to occur in senescent plants as short-lived intermediates of Chl breakdown. So far only six of the rather unstable natural FCCs have been characterized structurally that carry the critical free propionic acid function.5 However, the existence of two ‘epimeric’ ‘primary’ FCCs, 5 and epi-5, opens up a path to the two corresponding epimeric lines of the natural NCCs.5 The stereochemical classification of a range of NCCs and of other downstream phyllobilins (PBs) as ‘n’ or ‘epi’ was achieved by the identification of key NCCs, which were first screened by mass spectrometry38 and UV/Vis spectroscopy39 with known NCC reference compounds (further NMR-spectral and HPLC-based comparison). Thus, the assignment of the structure, e.g., of NCCs from spinach leaves as belonging to the ‘epi’-series was secured by identifying the polar So-NCC-2 (epi-1) as C16-epimer of 1 (Hv-

Scheme 7 In weakly acidic solution pFCC (5) and epi-pFCC (epi-5) isomerize stereoselectively to the NCCs 10 and epi-10, respectively.
NCC-1). For this purpose, an additional stereounselective synthesis of a pair of reference compounds with the common 1,2-dihydroxyethyl side chain of the two NCCs was carried out. It was achieved by regioselective OsO₄ oxidation of the exocyclic vinyl group of Cj-NCC-1 (Scheme 9).42,43

In the three NCCs from senescent oil seed rape leaves (the Bn-NCCs 2–4) a β-keto-carboxylic acid function (from eventual hydrolysis of the original β-keto methyl ester function of pFCC) turned out to be surprisingly stable and spontaneous decarboxylation did not occur with any one of them.40 Furthermore, from enzymatic hydrolysis of the methyl ester function of epi-11 the semisynthetic carboxylic acid analogue epi-12 was obtained, also found in senescent leaves of spinach as So-NCC-3.40 Decarboxylation of the β-keto-carboxylate epi-12 was remarkably slow and required heating with dilute sulfuric acid to give the corresponding ‘pyro’-NCC epi-13 (see Scheme 10), a 10R,16-epi-3'-hydroxy-8'-decarboxamethoxy-1-formyl-19-oxo-16,19-dihydrophyllobilane.44

7 Persistent Fluorescent Chl Catabolites and Blue- Luminescent Bananas

In the peels of ripening bananas a variety of strikingly persistent natural FCCs accumulates and give the ripe banana a blue glow.45,46,47 Spectroscopic analysis of such banana FCCs revealed the presence of propionate esters as their new structural feature, as, e.g., in Ma-FCC-56 (epi-14) (see Scheme 9).42,43

Indeed, this type of natural ester modification deactivates FCCs against their isomerization to NCCs and makes such ‘hypermodified’ FCCs (hmFCCs) persistent, so that they may accumulate in ripening fruit.45,48 as well as in senescent leaves of some evergreens.49,50 A new basic variant of the hmFCCs was discovered recently in Vv-FCC-55 (epi-15) isolated in senescent leaves of grapevine (Vitis vinifera). It exhibited two-fold attachment of a sugar unit furnishing a ‘bicyclo’-FCC (bcFCC) with a β-glycopyranosyl linker bridging the propionate side chain extending from C12 and the hydroxethyl substituent extending from C3 of the phyllobilin core.51 Formation of hmFCCs and bcFCCs indicates biosynthetic investments in ripening fruit and in senescent leaves that suggests still unknown biological roles of such PBs in the plants.51,52

8 Discovery, Structure Elucidation, and Biological Formation of Dioxobilin-Type Chl Catabolites

The absorption of colorless PBs at around 320 nm, which is due to the characteristic formyl-pyrrole unit of the PBs identified until about 2008,53 allowed on-line detection of PBs by routine HPLC analyses of extracts of senescent leaves,13,20,40,41,54 of vegetables,40,55 and ripe fruit.56,57 Fluorescence at 450 nm was also used for on-line identification in more recent analyses.45,46 The analytical setup was extended to detection at 250 nm, when Losey and Engel reported on two epimeric nonfluorescent ‘urobilinogenoid’ Chl catabolites (or DNCCs, see below) from senescent leaves
of barley that absorbed little at 320 nm.\textsuperscript{58} Likewise, in extracts of senescent leaves of Norway maple, NCCs were absent and the colorless and nonfluorescent dioxobilin-type NCC (DNCC) \textbf{16} was found instead.\textsuperscript{59} Interestingly, the maple DNCC \textbf{16} was a chirioptically distinct stereoisomer (possibly the enantiomer) of one of the earlier described ‘urobilinogenoidic’ Chl catabolites from barley leaves (see Figure 2).\textsuperscript{58} This curious finding contrasted with the presumed formation of DNCCs from NCCs,\textsuperscript{58} and raised the question of the biological formation of dioxobilin-type PBs.

In collaboration with the Hörtensteiner lab, dioxobilin-type NCCs (DNCCs) were established as the dominant colorless PBs in senescent leaves of \textit{A. thaliana}, and the cytochrome-P450 enzyme CYP89A9 was shown to be crucial for their formation.\textsuperscript{60} CYP89A9 is a new (heme-dependent) P450-type oxygenase that deformylates the ring A

\begin{equation}
\text{DFCC} \rightarrow \text{Me-DFCC} \rightarrow \text{DFCC} \rightarrow \text{Me-DFCC} \rightarrow \text{DFCC}
\end{equation}

of the PAO/phyllobilin pathway of Chl breakdown of higher plants.\textsuperscript{8} The oxidative deformylation by CYP89A9 was first observed in extracts of senescent leaves of \textit{A. thaliana} (and were first synthesized at the 32-position, such as \textbf{5}) itself and resulted in a mixture of type-I PxBs and type-II PBs and have been detected in recent years in a range of senescent leaves,\textsuperscript{51,59,60,65,66} in some vegetables,\textsuperscript{55} and also in fruit.\textsuperscript{57} DNCCs more directly derived from pFCC (\textbf{5}) by oxidative deformylation and their hydroxymethylated analogs (such as the 4HM-DNCC \textbf{18}) are much less abundant in senescent leaves.\textsuperscript{61,68}

\section{Occurrence, Partial Synthesis, and Structure of Phyllochromobilins, the Colored Bilin-Type Chl Catabolites}

The original classification of \textit{Hv}-NCC-1 (\textbf{1}) as a ‘rusty pigment’ referred to the readily occurring transformations of NCCs to colored compounds (typical UV/Vis absorption spectra are collected in the reviews\textsuperscript{13}). Chemical oxidation of \textit{Cj}-NCC-1 (\textbf{epi-11}) by DDQ at low temperature and work-up at room temperature with a short treatment with trifluoroacetic acid produced the yellow bilin-type tetrapyrrole \textbf{21Z} as main product, characterized as \textit{8S,10R,15Z}-3\textsuperscript{-}hydroxy-1-formyl-19-oxo-19,N24-dihydrophyllobilene-\textit{c}, a type-I phylloxyanthobilin (PxB) commonly classified as a yellow Chl catabolite (YCC).\textsuperscript{69} The yellow pigment \textbf{21Z}, and its more polar \textit{15E}-isomer \textbf{21E}, have also been observed in extracts of senescent leaves of \textit{C. japonicum} (and were first given the phenomenological name \textit{Cj-YCC-2} and \textit{Cj-YCC-1}, respectively).\textsuperscript{70} These two PxBs have a chromophore identical to the one present in bilirubin.\textsuperscript{67,71} In methanolic solution they interconvert by \textit{Z/E}-isomerization upon irradiation with day light (see Scheme 12).\textsuperscript{70}
However, 21Z and its methyl ester (Me-21Z) exhibit a remarkably medium-responsive photochemistry, which is due to the tendency of these two type-I 15Z-PxBs to associate to dimers (in a ‘hand shake motif’) in less polar media, e.g., when dissolved in CHCl₃ or in membrane-mimetic micellar detergent solutions.⁷² Photoysis of a solution of 21Z in chloroform led to selective formation of a C₅-symmetric, unstable [2+2]-cycloadduct that reverted to 21Z cleanly at room temperature.⁷² In dimethylsulfoxide solution 21Z was monomeric and exhibited a weak emission at 493 nm. It bound Zn ions cleanly in a 2:1 complex, which exhibited a strong red-shifted fluorescence at 538 nm, about 100 times more intense than that of 21Z itself.⁷³

Fortunately, the methyl ester Me-21Z furnished single crystals, suitable for X-ray analysis. The crystal analysis confirmed the NMR-based structure determination and showed Me-21Z to associate into H-bonded and π-stacked dimers in the crystal.⁷² It also allowed for the unambiguous determination of the absolute configuration of the asymmetric carbons C₈ and C₁₀ as S and R, respectively,⁷² confirming the tentative stereochemical assignment of carbons C₈ and C₁₀ in the prevailing epimers of natural NCCs,⁴³ which was derived earlier in the course of the studies of the acid-induced isomerization of epi-5 to epi-10.⁷⁷

The repeated observation of YCCs in extracts of senescent leaves and ripened fruit⁶⁹,⁷⁰,⁷⁴,⁷⁵ has been intriguing. Indeed, aerated aqueous or methanolic extracts of several senescent leaves converted epi-11 (and its C₁₆-epimer 11) cleanly to polar NCCs at ambient temperature, which were oxidized at C₁₅ stereo- and regioselectively. These oxidized NCCs eliminated water (or methanol) under weakly acidic conditions to furnish the YCC 21Z selectively and in good yield.⁷⁴ This finding led us to use a variety of leaf extracts (especially of the evergreen Spatiphyllum wallisii) to prepare YCCs from corresponding NCCs by a type of ‘green synthesis’.⁴⁴ The identity of the ‘oxidative activity’ in leaves is still puzzling.⁷⁴ However, the ‘green synthesis’ method with leaves of Sp. wallisii was also used for preparative oxidation of the semisynthetic ‘pyro’-NCC epi-13 to the corresponding optically active ‘pyro’-YCC 22Z (8⁵S,10R,15Z-3⁻-hydroxy-8⁻-decarboxymethyl-1-formyl-19-oxo-19,N₂₄-dihydroxyphylllobilene-c) (see Scheme 13). The methyl ester form Me-22Z crystallized readily and exhibited a dimer structure virtually superimposable to that of the YCC methyl ester Me-21Z.⁴⁴

Along these lines, yellow PBs (phyllloxanthobilins, PxBs) have been observed in extracts of senescent leaves of grapevine⁵¹ and of an A. thaliana MES16-mutant,⁶⁸ and were provisionally classified as type-II PxBs, proposed to represent oxidation products of the major isomeric DNCCs in each of these leaves. The major DNCC epi-23 of the C₁₆-‘epi’ series from senescent leaves of grapevine (named Vv-
DNCC-5151 and the C16-‘n’-epimer 23 of the A. thaliana mutant62 were used as substrates for selective oxidation by the ‘green synthesis’ method using leaves of Sp. wallisii. This furnished the corresponding dioxobilin-type YCCs (named DYCCVv and DYCCAt) in roughly 70% yield (see Scheme 14).76 The oxidation in the aerated leaf extract occurred again at the ‘western’ meso position, with remarkable regioselectivity, as observed earlier in the corresponding NCC-oxidation processes,74 thus, desaturating the stereo-differentiating asymmetric C16. In fact, the oxidation of both, the ‘n’- and ‘epi’-DNCC, apparently furnished a single DYCC 24 (8S,10R,15Z-3α-hydroxy-1,19-dioxo-1,4,19,N24-tetrahydrophyllobilene-c), suggesting the asymmetric C4 to have the same absolute configuration in both DNCCs and their oxidation products. Hence, in the course of the oxidative deformaylation of the corresponding epimeric FCC precursors by CYP89A9, C4 would be installed with the same absolute configuration in both DNCCs and their oxidation products.77

This furnished the corresponding dioxobilin-type YCCs (such as in the YCCs 21Z and 22Z) to be a crucial factor in stabilizing their H-bonded and π-stacked dimers.44,72

The blue-light-absorbing chromophore of YCCs is prone to further oxidation in the presence of air to pink-colored Chl catabolites (PiCCs), or type-I phylloroseobilins (PrBs), and PiCC 25 is a side product of the formation of the YCCs 21Z and 21E by low-temperature DDQ or light-induced oxidation of NCCs.69,70 Traces of 25 are also found in extracts of senescent leaves of C. japonicum.70 The most useful preparative method for the synthesis of 25 from 21Z turned out to be a two-step procedure encompassing (i) complexation of the YCC 21Z with Zn(II) ions in dimethylformamide solution,73 which, upon exposure to air, led to the clean formation of the blue Zn(II) complex 26 of the PiCC 25.77 followed (ii) by removal of the Zn ion of 26 by treatment of its solution in acetonitrile with 20 mM aqueous phosphate buffer (pH 4.7), furnishing 25 in an overall yield of 91%.77 The structure of 25 was determined as a 10Z,15E-phylloroseobiladiene-β-c (see Scheme 15), on the basis of its NMR-spectral NOE data. A crystal structure analysis of 25 (as the potassium salt) confirmed this assignment of the molecular structure completely and showed 25 to crystallize as H-bonded and π-stacked pairs of enantiomers (25 is a racemate from equilibration at its acidified asymmetric 8β-position).77

Complexation of the effectively tridentate 25 by divalent transition-metal ions [such as Zn(II), Ni(II), Cu(II), Cd(II)] occurs readily and induces the isomerization of the effective double bond C10=C11 to the Z-configura- tion in the blue complexes M(II)-26 with the four divergent metal ions M(II).77,78 Solutions of the blue complexes Zn-26 and Cd-26 with the closed shell ions Zn(II) and Cd(II), respectively, exhibit an intense red luminescence, allowing for the quantitative analysis of Zn(II) and Cd(II) ions down to the nM range77 (typical UV/Vis absorption spectra are collected in the reviews5,8).

In exploratory experiments, the dioxobilin-type PxBl (DYCC 24) was likewise revealed to have a pronounced tendency to oxidize in the presence of Zn(II) ions to the blue Zn(II) complex 27, from which the chiral and optically active type-II PrB (DPiCC) 28 was set free with dilute phos-

![Scheme 14] DNCCs 23 and epi-23 are oxidized by chemical oxidants and by a puzzling ‘oxidative activity’ in leaves (e.g.) of Sp. wallisii at the ‘western’ moiety to the apparently common DYCC 24.

![Scheme 15] YCC 21Z slowly oxidizes with air in methanolic solution to PiCC 25. In the presence of air and Zn(II) ions 21Z is rapidly complexed and oxidized to the blue Zn complex Zn-26, from which Zn ions are removed by dilute phosphoric acid, furnishing the pink PiCC 25 in high yield.

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phosphate buffer (pH 4.7) in 92% yield (see Scheme 16). The weakly luminescent 28 bound Zn(II) ions with high affinity and furnished the red-luminescent Zn(II) complex 27 in practically quantitative yield.

10 Conclusion and Outlook

Chl breakdown has for a long time been considered to play a particular role in the recuperation of scarce plant nutrient, such as bioavailable forms of nitrogen. However, the earlier assumption that the plant would go after the four nitrogen atoms of the Chl molecules has received no support from the structures of the known PBs. Once Chl is broken down to the colorless PBs found in extracts of senescent leaves do not account for the amount of Chl present in green leaves (see e.g. 5) and known PBs do not accumulate for longer times in the tissue of senescent leaves and ripened fruit. The deduced in vivo disappearance of PBs may reflect their further degradation, physiological use involving further metabolic transformations, or extracellular transport processes. Typical PBs are excellent antioxidants (such as NCCs and YCCs) that are prone to undergo oxidation reactions. Their oxidation products, like the phyllo-Dieckmann reaction opening the -keto-ester function under-went a retro-Dieckmann reaction at the Chl-derived ring E moiety readily, furnishing the yellow 8,8′-seco-PB 29.

Because of the availability of the chemical structures of key bilin-type Chl catabolites several of the critical enzymes directly required in the breakdown process have been identified, establishing the common PAO/phyllobilin pathway of Chl breakdown in the so far studied angiosperms,5,32,82 The regulation of these important biochemical processes in the plants has attracted much interest and receives broad attention, currently, in order to obtain further insights into the plant’s way of adjusting the cellular availability of Chl to the presence and absence of light, to biotic and biological external and internal stress, etc.31,32 Interestingly, in such studies, the occasional earlier bioinformatics-based mis-assignments of enzymes have been corrected, such as the one of the P FCC-hydroxylase TIC55,81 However, the ubiquitous enzymes active in the downstream part of the pathway, involved in attaching sugar units once or twice by glycosylation or esterification, are still unknown.51

Most recent studies in an ongoing collaboration with Thomas Müller have turned to investigating the structures of phyllobilins in Innsbruck, that occur in de-greened tissue of some gymnosperms, ferns, and other ‘exotic’ higher plants, such as Ginkgo biloba.83 Interestingly, we have been finding structural evidence from detailed spectroscopic work for unprecedented and intriguing deviations of Chl breakdown from the PAO/phyllobilin pathway in these families of higher plants. Bioinformatics methods have identified genes coding for activities related to those of PAO in green alga,82,84 and to those of PAO and RCCR in ferns and gymnosperms.84 The results of the structural and spectro-
scopic studies will, again, invite follow-up investigations dealing with elucidating the corresponding new biochemical processes and their key regulatory processes.

An ever intriguing question in the field of Chl breakdown has concerned the plant biological ‘why’. Earlier views on Chl breakdown considered the degradation of the potentially phototoxic Chl, and its removal from the plant cells, to represent a mere detoxification. In contrast, phyllobilins have become a topic of interest in their own right, lately, because of their possible hypothetical physiological roles in plant cells. Indeed, the chemical features of phyllobilins suggest a wide range of specific biological roles: (i) persistent fluorescent Chl catabolites, such as hmtFCCs, bcFCCs, and bcDFCCs, act as optical brighteners in fruit and leaves, making them blue fluorescent; (ii) FCCs and YCCs are very effective amphiphilic antioxidants; (vi) bicyclo-PBs that extend the repertoire of plant cells in their control and optical brighteners in fruit and leaves, making them blue fluorescent; (ii) FCCs and YCCs are very effective amphiphilic antioxidants, adding to the list of pigments in plant cells; (v) phyllochromobilins and their metal complexes are intensely colored compounds, extending the repertoire of plant cells in their control and neutralization of reactive oxygen species; (vi) bicyclo-PBs (bcPBs) and other ‘persistent’ hmtPBs are heterocyclic natural products with structures that may make them attractive as anti-infective agents in plants and in other applications. Furthermore, phyllobilimobilins feature structures strikingly related to those of some (heme-derived) natural bilins, specifically when taking into account some colored type-II or dioxobilin-type PBs. Hence, phyllobilimobilins may play crucial roles as inhibitors of bilin-dependent enzymes, e.g., in photoregulation, or as effectors, in their own right, in such enzymes. Clearly, the discovery of the presumed, but still elusive plant-biological roles of phyllobilins is an exciting scientific quest, as is the possible use of phyllobilins in pharmaceutical applications.

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