



Article Acylated Anthocyanins from Red Cabbage and Purple Sweet Potato Can Bind Metal Ions and Produce Stable Blue Colors

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Abstract: Red cabbage (RC) and purple sweet potato (PSP) are naturally rich in acylated cyanidin glycosides that can bind metal ions and develop intramolecular π -stacking interactions between the cyanidin chromophore and the phenolic acyl residues. In this work, a large set of RC and PSP anthocyanins was investigated for its coloring properties in the presence of iron and aluminum ions. Although relatively modest, the structural differences between RC and PSP anthocyanins, i.e., the acylation site at the external glucose of the sophorosyl moiety (C2-OH for RC vs. C6-OH for PSP) and the presence of coordinating acyl groups (caffeoyl) in PSP anthocyanins only, made a large difference in the color expressed by their metal complexes. For instance, the Al³⁺-induced bathochromic shifts for RC anthocyanins reached ca. 50 nm at pH 6 and pH 7, vs. at best ca. 20 nm for PSP anthocyanins. With Fe^{2+} (quickly oxidized to Fe^{3+} in the complexes), the bathochromic shifts for RC anthocyanins were higher, i.e., up to ca. 90 nm at pH 7 and 110 nm at pH 5.7. A kinetic analysis at different metal/ligand molar ratios combined with an investigation by high-resolution mass spectrometry suggested the formation of metal-anthocyanin complexes of 1:1, 1:2, and 1:3 stoichiometries. Contrary to predictions based on steric hindrance, acylation by noncoordinating acyl residues favored metal binding and resulted in complexes having much higher molar absorption coefficients. Moreover, the competition between metal binding and water addition to the free ligands (leading to colorless forms) was less severe, although very dependent on the acylation site(s). Overall, anthocyanins from purple sweet potato, and even more from red cabbage, have a strong potential for development as food colorants expressing red to blue hues depending on pH and metal ion.

Keywords: anthocyanin; hydroxycinnamic acid; polyphenol; pigment; acylation; aluminum; iron; metal complexation

1. Introduction

The color of red cabbage (RC) and purple sweet potato (PSP) is due to closely related anthocyanins displaying a cyanidin or peonidin (3'-O-methylcyanidin) 3-O-sophoroside-5-O-glucoside structure [1,2]. Cyanidin derivatives (the major RC pigments) are especially interesting colorants owing to their ability to bind metal ions (via their catechol B-ring) in neutral or mildly acidic solution. Indeed, metal-induced cyanidin deprotonation leads to a quinonoid chromophore that can express intense purple to blue colors [1,3]. Another important consequence of metal–anthocyanin binding is increased color stability. Indeed, through the electrophilic flavylium ion (main colored form in acidic solution), anthocyanins, unlike their metal chelates, are vulnerable to water addition with the concomitant reversible formation of colorless forms (hemiketal and chalcones) [4]. A remarkable feature of RC anthocyanins is that the sophorosyl moiety is typically acylated by one or two residue(s) of *p*-hydroxycinnamic acid (HCA = *p*-coumaric, ferulic, caffeic, or sinapic acid). Phenolic



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Copyright: © 2021 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). acyl groups are known to favor folded conformations in which the anthocyanidin (chromophore) and the acyl residues develop π -stacking interactions. Like metal binding, this phenomenon, called intramolecular copigmentation, causes a bathochromic shift (BS) in the visible absorption band and protects the chromophore against water addition [5–7]. The combination of π -stacking interactions and metal binding is actually required to achieve maximal blue color stability with anthocyanins. Indeed, phenolic acyl groups stacked onto the cyanidin nucleus could either directly participate in metal binding (e.g., caffeic acid residues through their catechol ring) or at least strengthen metal binding by building a hydrophobic pocket around the metal-cyanidin complex. Recently, a remarkable RC anthocyanin (called pigment B or PB), displaying a single ideally located sinapoyl residue, was shown to form an aluminum(III) complex of 1:3 stoichiometry in which the three PB ligands in octahedral coordination to Al^{3+} adopt a chiral arrangement, causing an intense positive Cotton effect in the visible part of its circular dichroism spectrum [8]. The Al(PB)₃ complex is strongly stabilized by the π -stacking interactions taking place between each cyanidin chromophore and the sinapoyl residue of an adjacent ligand. Moreover, this original supramolecular structure imposes a large torsion angle around the bond connecting the B- and C-rings of the three cyanidin nuclei. This unique combination of structural characteristics results in an intense vibrant blue color of high stability, making PB and its metal complexes potential lead compounds for the replacement of artificial blue colorants by natural alternatives.

In this work, a selection of acylated cyanidin glycosides from red cabbage (including PB) and purple sweet potato is revisited for its affinity for aluminum and iron ions. In particular, the influence of the acyl groups on the binding kinetics, color stability, and rate of oxidative degradation of the complexes is systematically addressed. Indeed, the addition of iron ions was shown to accelerate the oxidative degradation of nonacylated cyanidin glycosides, while the presence of phenolic acyl groups tends to cancel this effect [9]. In this work, the conditions (pH, metal type, and concentration) permitting the optimal development of a blue color is also explored.

2. Results and Discussion

Metal–anthocyanin binding is of great importance for plants, not only because it is an efficient way to express blue colors to attract pollinating insects [6], but also as a detoxification mechanism against metal excess [10], which can operate with a variety of metal ions (e.g., Fe, Al, Pb, Cd, Mo, Mg, Ni, and V in corn roots). With Cu²⁺, the binding is followed by Cu²⁺ reduction and anthocyanin oxidation [11]. However, quantitative physicochemical investigations of metal–anthocyanin binding are scarce. Such approaches have to address the structural transformations of anthocyanins in aqueous solution, the kinetics of metal binding and its stoichiometry, the critical influence of pH, and the possible influence of phenolic acyl groups and even of the selected buffer (depending on its own affinity for metal ions). This is the specific focus of this work, based on a large series of diversely acylated cyanidin glycosides and two of the most important metals (Al, Fe) in terms of blue color development [6].

2.1. The Color and Spectral Properties of the Metal Complexes

Anthocyanins under their flavylium form (AH⁺) are typically diacids undergoing a first proton loss from C7-OH ($pK_{a1} \approx 4$), followed by a second one from C4'-OH around neutrality [12]. Thus, at pH 7, cyanidin glycosides from RC or PSP are a mixture of neutral (A₇) and anionic (A_{4'7}) bases ($pK_{a2} = 7.0-7.3$ [7,13]), in agreement with the broad absorption band observed (Figure 1). From the experimental spectra at pH 5–7, the pK_a values, and the spectrum of the pure flavylium ion (pH 1), the spectra of the pure neutral and anionic bases can be calculated [13]. Compared to the neutral base, the anionic base not only has a much higher λ_{max} , but also a higher molar absorption coefficient at λ_{max} (Figure 1).

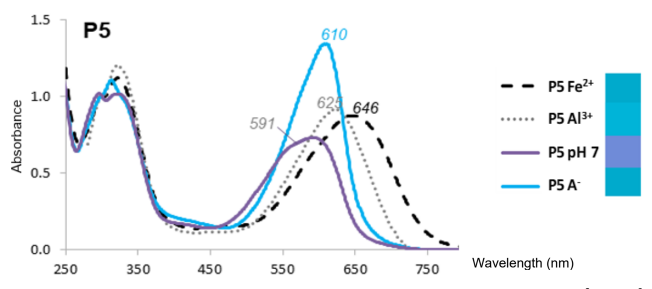


Figure 1. UV-VIS spectra of Pigment 5 (50 μ M) at pH 7, its pure anionic base (A⁻, calculated), and its Fe²⁺ and Al³⁺ complexes (1 equiv.). Right: color patches from the L*a*b* coordinates calculated from the visible spectra.

The major PSP anthocyanins are acylated peonidin glycosides, which do not bind metal ions through their chromophore. However, cyanidin glycosides are also present in PSP. Unlike the RC anthocyanins, the HCA residues of the PSP anthocyanins are only located at the primary C6-OH positions of the sophorosyl moiety (Scheme 1). In particular, the acyl residue borne by Glc-2 (R₃) is expected to be more mobile than its homolog in red cabbage (R₂ = sinapoyl). This happens to make a large difference in terms of color variation: the Al³⁺-induced bathochromic shifts for PB (red cabbage) are ca. 50 nm at pH 6 and pH 7 vs. at best ca. 20 nm for P4' (R₃ = feruloyl, Table 1). The flexible acyl residue of P4' is probably much less apt to develop π -stacking interactions with the cyanidin nucleus than the more rigid sinapoyl residue of PB. Consistently, it has been demonstrated that RC anthocyanins having a single HCA residue at R₁ (P1–P3) are much more susceptible to water addition than PB [8] as a consequence of the latter adopting folded conformations in which the cyanidin and HCA moieties are in molecular contact. However, PSP pigments have a specific advantage over RC pigments: the presence of caffeoyl residues at Glc-1 and/or Glc-2, which themselves can bind metal ions.

Table 1. Spectral characteristics of the metal complexes of red cabbage and purple sweet potato anthocyanins (1 equiv. metal ion). $\Delta \lambda_{\text{max}} = \lambda_{\text{max}}(+\text{metal}) - \lambda_{\text{max}}(\text{no metal})$. $\Delta A = A(+\text{metal}) - A(\text{no metal})$.

			Fe ²⁺	F			Al ³⁺		
Red Cabbage Pigments		PA	P2	P5	РВ	PA	P2	P5	PB
	λ_{max} (+metal) (nm)	559	569	640	656		571	587	629
pH 6	$\Delta\lambda_{\max}$ (nm)	21	25	87	76	no binding	27	34	49
	$\Delta A/A$ (no metal) at λ_{\max} (%)	-25	-15	3	65		-15	-9	16
pH 7	λ_{max} (+metal) (nm)	596	592	646	656	583	585	625	637
	$\Delta\lambda_{\rm max}$ (nm)	14	-4	55	71	4	5	34	52
	$\Delta A/A$ (no metal) at λ_{max} (%)	-7	-3	19	43	3	-1	17	25
Purple Sweet Potato Pigments						P4′	P6′	P7′	P9b′
pH 6	λ_{max} (+metal) (nm)	/	/	/	/	563	574	575	581
	$\Delta\lambda_{\max}$ (nm)	/	/	/	/	16	26	23	29
pH 7	λ_{max} (+metal) (nm)	/	/	/	/	580	591	598	598
	$\Delta\lambda_{\max}$ (nm)	/	/	/	/	8	21	22	16

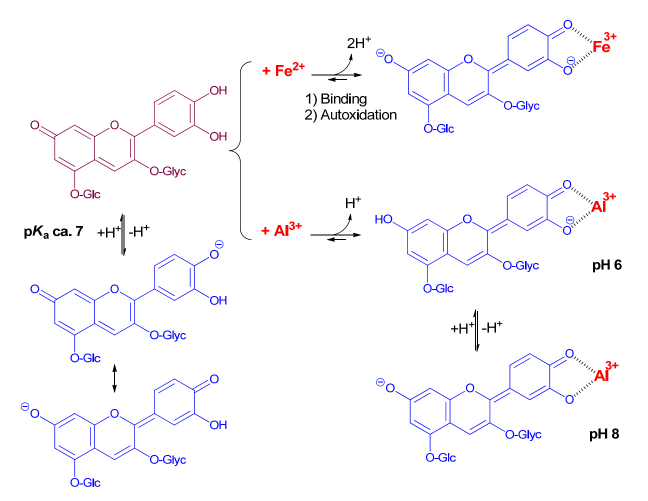
HO		В	,OH O—R₃ /_OH			
Ĺ	A C		Сон	HCA Residue	R' 1	R' 2
_OH _	5 4		Glc-2	pC	Н	Н
HOLIO	,ò	R ₂		Cf	ОН	Н
OF	- -			Fl	OMe	Η
ج <mark>R</mark> '			4	Sp	OMe	OMe
O HCA residue	-OH 2	О́́́́́́о́н R ₁				
-	Pigment	R 1	R ₂	R3		
	PA a)	Н	Н	Н		
	P1 a)	pC	Н	Н		
	P2 a)	F1	Н	Н		
-	P3 a)	Sp	Н	Н		
	PB a)	Н	Sp	Н		
	P4 a)	pC	Sp	Н		
	P5 a)	F1	Sp	Н		
-	P6 a)	Sp	Sp	H		
	P4' b)	Н	H	Fl		
	P6′b)	Cf	H	Н		
	P7′b)	Cf	H	Cf		
-	P9b' b)	Cf	Н	Fl		

Scheme 1. Structure of the RC and PSP anthocyanins studied. When present, hydroxycinnamoyl residues R₁, R₂, and R₃ are *p*-coumaroyl (pC), feruloyl (Fl), caffeoyl (Cf), and/or sinapoyl (Sp). Note: a) from red cabbage; b) from purple sweet potato.

Our recent work [13] suggests that such pigments, in which the cyanidin and caffeoyl units tend to stack onto each other, can actually sequester Fe^{2+} and Al^{3+} by the simultaneous involvement of both catechol rings, thereby largely increasing the metal-induced bathochromic shift. This was confirmed in the current work: the bathochromic shifts induced by Al^{3+} (1 equiv.) at pH 7 were 8, 21, and 22 nm for P4' (R₃ = feruloyl), P6' (R₁ = caffeoyl), and P7 (R₁ = R₃ = caffeoyl), respectively (Figure S1A, Table 1). It was also clear that the caffeoyl residue at R₁ was critical to promote the bluing effect, while the other one at R₃ was not. By contrast, P9b (R₁ = caffeoyl, R₃ = feruloyl) was clearly less efficient (BS = 16 nm), possibly pointing to a less-favorable binding because of the relatively bulky feruloyl residue.

The pH dependence of the λ_{max} values in the pH range 6–8 for the free forms (Table S1) was consistent with the conversion of the neutral base (A₇) to the anionic base (A_{4'7}, proton loss from C4'-OH), in full agreement with the corresponding pK_{a2} value of the RC and PSP anthocyanins [7,13]. As Al³⁺ also triggers proton loss from C4'-OH, the bathochromic shift (BS) induced in the free forms by increasing the pH from 6 to 8 should be close to the one accompanying Al³⁺ binding at pH 6. Additionally, only small BS were expected upon Al³⁺ binding at pH 8. While the latter prediction was well-verified experimentally for all PSP pigments (Al³⁺-induced BS at pH 8 < 10 nm, Table S1), the former was not: indeed, the pH-induced BS (*ca.* 50 nm) were much larger than the Al³⁺-induced BS at pH 6 (from 16 nm for P4' to 29 nm for P9b'). Moreover, bathochromism was clearly observed in the complexes' visible band when the pH was increased from 6 to 8 (BS = 25–33 nm, Table S1). Hence, it can be proposed that C7-OH in the complexes was not dissociated at pH 6, and lost its proton when the pH is raised to 8 (Scheme 2). This proposal is consistent with the theoretical visible spectrum calculated on the Al(PB)₃ complex, which, while involving 3 ligands undissociated at C7-OH, were found fully consistent with the experimental spectrum [8].

Moreover, comparing the neutral base of the 7-O- β -D-glucosyloxy-4'-hydroxyflavylium ion (proton loss from C4'-OH) with that of its 4'-O- β -D-glucosyloxy-7-hydroxyflavylium regioisomer (proton loss from C7-OH) also showed that the former displayed a λ_{max} value that was 20 nm higher [14] and a molar absorption coefficient almost 3 times as large. Finally, DFT calculations on pyranoanthocyanins [15] confirmed that the neutral base formed by proton loss from C4'-OH (A_{4'}, a very minor species) displayed an intense absorption band at higher wavelengths than the major tautomer (proton loss from C7-OH). Overall, turning A₇ into A_{4'} upon metal binding is actually expected to promote both bathochromism and hyperchromism.



Scheme 2. Metal–anthocyanin binding.

Red cabbage anthocyanins with R_2 = sinapoyl, whether mono- or diacylated, were distinct from the PSP anthocyanins and the other RC pigments in that the Al³⁺-induced BSs were larger, especially at pH 7 (36 and 52 nm for P5 and PB, respectively, vs. barely 5 and 8 nm for P2 and P4', respectively) (Table 1, Figures 1 and S1B). The λ_{max} values of the complexes were even higher than that of the anionic base. Thus, even if proton loss from C7-OH was not complete at pH 7 for the Al³⁺ complexes, the strong π -stacking interactions occurring between the cyanidin nucleus and R_2 = sinapoyl and the concomitant torsion imposed between the B- and C-rings [8] effectively turned the color to an intense cyan hue (Figure 1) close to the one expressed by the major synthetic blue food colorants Brilliant Blue and indigotine.

With Fe^{2+} , the BSs were even larger: at pH 7, 87 and 76 nm for P5 and PB, respectively, vs. 25 nm for P2 (Table 1, Figures 1 and S1B). As already reported for other Fe^{2+} -polyphenol complexes [16,17], bound Fe^{2+} was rapidly autoxidized to Fe^{3+} . This reaction was: (a) promoted by the higher affinity of catechols for Fe^{3+} (vs. Fe^{2+}); (b) confirmed by the

similarity of the final spectra, whether Fe²⁺ or Fe³⁺ was added [13]; and (c) consistent with the broad absorption band of the iron complexes and their high λ_{max} , which both suggest ligand-to-Fe³⁺ charge transfer. However, iron autoxidation clearly followed metal binding. Indeed, despite the higher intrinsic affinity of catechols for Fe³⁺, anthocyanins bound Fe²⁺ much more rapidly in our model (Figures 2 and S2) [13], as competition between anthocyanins and the phosphate anions for the metal was much less severe with Fe²⁺.

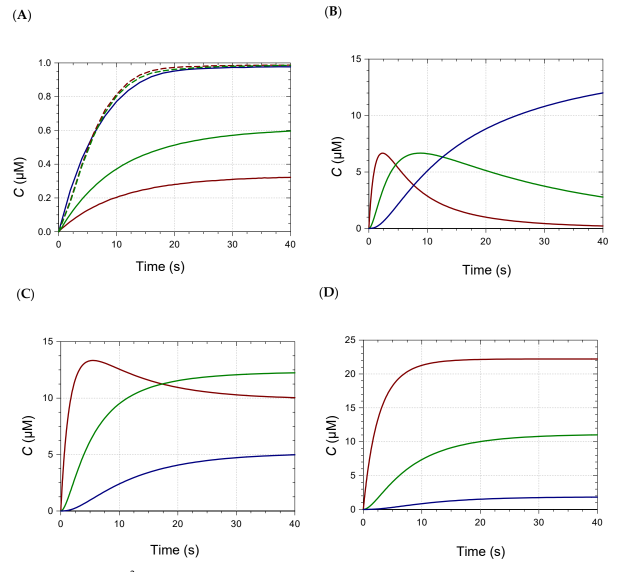


Figure 2. (A) Kinetics of Fe²⁺–Pigment B binding (pH 7, 25 °C). Pigment concentration = 50 μ M, metal/pigment molar ratio = 0.05 (_), 0.1 (_), 0.3 (_), 0.6 (__), 1 (__). (B–D) Speciation diagrams of Fe²⁺–Pigment B complexes. (B) 0.3 equiv. Fe²⁺. (C) 0.6 equiv. Fe²⁺. (D) 1 equiv. Fe²⁺. ML (_), ML₂ (_), ML₃ (_). ML_n: metal–ligand complex having a 1:n metal/ligand stoichiometry.

While the visible band of the Al^{3+} complexes of the RC anthocyanins showed the same pH dependence as for the PSP anthocyanins, the visible band of their Fe³⁺ complexes was remarkably insensitive to pH in the subgroup with R₂ = sinapoyl (Figure S1B). In particular, the BS featuring iron–PB binding at pH 5.68 hit a record high of ca. 110 nm, i.e., twice as much as with Al^{3+} at the same pH. Hence, it can be proposed that C7-OH in the iron complexes is dissociated even at low pH (Scheme 2).

The colorimetric data of the PSP cyanidin glycosides and their Al³⁺ complexes (Table S2) provide additional evidence of the bluing effect induced by raising the pH from 6 to 8, or adding increasing Al³⁺ concentrations at a given pH. For comparison, a hue

angle of 207.3 was recorded for the PB-Al³⁺ complex at pH 7, i.e., a close match for that of the synthetic triarylcarbonium colorant Brilliant Blue ($h^0 = 209.4$), regarded as a reference for a vibrant cyan hue in the confectionary industry [8]. No such match was observed with the PSP anthocyanins and the best result recorded, i.e., the P6'-Al³⁺ complex at pH 8 ($h^0 = 221.7$) remained off target. However, it confirmed that a single caffeoyl residue at R₁ was sufficient to promote a strong bluing effect.

2.2. Kinetic Analysis and Stoichiometry of Metal Binding

When small volumes of concentrated Fe²⁺ or Al³⁺ aqueous solutions were added immediately after diluting the RC pigments into pH 6–8 phosphate buffers, a relatively fast metal–anthocyanin binding occurred (Figures 2 and S2). Contrary to predictions based on steric hindrance, the observed trend was that acylation by noncoordinating HCA residues favored metal binding. For instance, nonacylated PA weakly bound Al³⁺ at pH 7 and 8 (weak spectral changes preventing the kinetic analysis), and strong Al³⁺–P2 binding only occurred at pH 8. By contrast, P5 strongly bound Al³⁺ at both pHs.

The minimal metal/ligand molar ratio to reach full binding (saturation of the visible band of the complex) is an indicator of the complex's stoichiometry. This ratio lay between 1/3 and 2/3 for PA, PB, P2, and P5 (Figures 2 and S3), as already observed with iron [9]. Thus, mixtures of 1:1, 1:2, and 1:3 complexes were expected in variable proportions according to the metal/ligand molar ratio. Consistently, in our recent work, the $Al^{3+}(PB)_3$ and $Al^{3+}(P6)_3$ complexes were evidenced by high-resolution mass spectrometry in a dilute ammonium acetate buffer, but not the Al³⁺(P3)₃ homolog [8]. This was confirmed in the present work (Table S3). Under the same conditions, 1:1 and 1:2 iron-anthocyanin complexes were detected with P5 and PB (Table 2, Figure S4). With monoacylated P2, only the 1:1 complex was detected, whatever the M/L molar ratio between 1/6 and 1. The detection of 1:3 complexes is more challenging, as the corresponding ions must bear at least 3 charges for the m/z ratio to fall below 1500, the upper limit of detection. Satisfying agreements between experimental and theoretical m/z values were observed for the main ions and their major isotopes. Moreover, HRMS data were consistent with iron having a +3 oxidation degree in the complexes. For instance, the FeP5 monocation (exp. m/z 1208.2307, 1209.2352, 1210.2398) is proposed to be $[P5 - 3H^+ + Fe^{3+}]^+$ (P5 referring to the flavylium cation, theoretical *m*/*z* 1208.2302, 1209.2335, 1210.2362). The corresponding complex involving Fe^{2+} would be $[P5 - 2H^+ + Fe^{2+}]^+$ (*m*/*z* 1209.2381, 1210.2414, 1211.2440). Despite the possible match of the experimental spectrum with isotopes of the Fe²⁺ complex having one or two 13 C-atoms, the intense signal at m/z 1208.2307 (Figure S4) clearly required an Fe^{3+} ion. Unexpectedly, varying the M/L molar ratio between 1/6 and 1 did not strongly impact the signal intensity of the 1:1 complex relative to free ligand (Table S4). Moreover, the signal intensities of the 1:1; 1:2, and 1:3 complexes could not be compared due to the charge-specific ion sensitivity of the MS detector.

A simple model assuming stepwise 1:1, 1:2, and 1:3 binding was tested to account for the kinetics of metal binding as a function of the metal concentration. To keep the number of adjustable parameters to a minimum, the rate constants of the first, second, and third steps were assumed to be 3k (3 available binding sites), 2k (2 available binding sites) and k(1 available binding site), respectively. Moreover, the molar absorption coefficients of the ML, ML₂, and ML₃ complexes (M = metal, L = ligand) were assumed to be ε , 2ε , and 3ε , respectively. Satisfactory curve-fittings of the A(670 nm) vs. time curves for different M/L molar ratios were thus obtained, leading to optimized values for parameters k and ε , and permitting the plotting of the time dependence of the concentrations of the 3 complexes (Table 3, Figure S5).

Compound	Formula	Th. <i>m/z</i> (rel. abundance) ^(a)	Exp. <i>m/z</i> (rel. abundance) ^(b)	δ (ppm)	
P5+	$C_{54}H_{59}O_{28}$	1155.3187	1155.3198	1.0	
		1208.2302 (100)	1208.2307 (100)	0.4	
$[P5 - 3H^+ + Fe^{3+}]^+$	C ₅₄ H ₅₆ O ₂₈ Fe	1209.2335 (62.4)	1209.2352 (98)	1.4	
		1210.2362 (25.2)	1210.2397 (46)	2.9	
		604.6188 (100)	604.6189 (100)	0.2	
$[P5 - 2H^+ + Fe^{3+}]^{2+}$	C ₅₄ H ₅₇ O ₂₈ Fe	605.1204 (62.0)	605.1211 (83)	1.2	
		605.6217 (24.9)	605.6227 (32)	1.7	
		788.1856 (84.1)	788.1836 (87)	2.5	
$[2P5 - 2H^+ + Fe^{3+}]^{3+}$	C ₁₀₈ H ₁₁₆ O ₅₆ Fe	788.5199 (100)	788.5183 (100)	2.0	
		788.8542 (69.8)	788.8537 (35)	0.6	
PB^+	$C_{44}H_{51}O_{25}$	979.2714	979.2722	0.8	
		1032.1829 (100)	1032.1836 (100)	0.7	
$[PB - 3H^+ + Fe^{3+}]^+$	$C_{44}H_{48}O_{25}Fe$	1033.1862 (51.4)	1033.1888 (71)	2.5	
		1034.1887 (18.4)	1034.1921 (25)	3.3	
		516.5951 (100)	516.5952 (100)	0.2	
$[PB - 2H^+ + Fe^{3+}]^{2+}$	C ₄₄ H ₄₉ O ₂₅ Fe	517.0967 (51.1)	517.0969 (53)	0.4	
		517.5980 (18.2)	517.5982 (16)	0.4	
		1005.7273 (100)	1005.7277 (63)	0.4	
$[2PB - 3H^{+} + Fe^{3+}]^{2+}$	C ₈₈ H ₉₉ O ₅₀ Fe	1006.2289 (98.5)	1006.2304 (100)	1.5	
[210 - 511 + 10]		1006.7303 (59.0)	1006.7320 (80)	1.7	
		1007.2317 (26.2)	1007.2343 (45)	2.6	
$[PB - 2H^+]^-$	$C_{44}H_{49}O_{25}$	977.2563	977.2570	0.7	
		1030.1683 (100)	1030.1684 (100)	0.1	
$[PB - 5H^+ + Fe^{3+}]^-$	$C_{44}H_{46}O_{25}Fe$	1031.1716 (51.4)	1031.1723 (53)	0.7	
		1032.1741 (18.4)	1032.1752 (16)	1.1	
		514.5806 (100)	514.5807 (100)	0.2	
$[PB - 6H^+ + Fe^{3+}]^{2-}$	C ₄₄ H ₄₅ O ₂₅ Fe	515.0822 (51.1)	515.0822 (47)	0.0	
		515.5834 (18.2)	515.5837 (12)	0.6	
		1003.7128 (100)	1003.7124 (100)	0.4	
$[2PB - 7H^+ + Fe^{3+}]^{2-}$	C ₈₈ H ₉₅ O ₅₀ Fe	1004.2143 (98.5)	1004.2153 (93)	1.0	
		1004.7157 (58.9)	1004.7166 (65)	0.9	
		1005.2171 (26.2)	1005.2180 (40)	0.9	

Table 2. Ions detected for the iron complexes of pigments P5 = Cya-3-(Fl)Glc-2-(Sp)Glc-3-Glc and PB = Cya-3-Glc-2-(Sp)Glc-3-Glc from red cabbage (metal/ligand molar ratio = 1).

(a) Relative abundances calculated on the website https://www.chemcalc.org (accessed on 2 March 2021).
 (b) Relative abundances estimated with the Freestyle software (Thermo Fisher Scientific, Waltham, MA, USA).

For given pigment and metal ion, the optimized ε values were fairly stable when the metal concentration was varied, which was satisfactory. This was less true for the optimized *k* values, which suggests that the model of independent binding steps was too simple. With the P5–Al³⁺ pair at pH7, much better curve-fittings were actually obtained by implementing 3 optimizable rate constants in the model. Despite its crude approximations, the model offered a simple way to quantitatively assess the influence of pH, metal type, and acylation pattern on the rate of metal–anthocyanin binding. Overall, binding was faster at pH 8 than at pH 7, and faster with Fe²⁺ than with Al³⁺. Overall, under our conditions, acylation did not impede metal binding. Most importantly, pigments having R2 = sinapoyl (PB, P5) formed complexes with much higher molar absorption coefficients (typically, a factor 2–3), which was an obvious advantage for color development. **Table 3.** Kinetic analyses of metal–anthocyanin binding according to the stepwise formation of ML, ML₂, and ML₃ complexes (25 °C, monitoring at 670 nm). ML_n: metal–ligand complex having a 1:n metal/ligand stoichiometry. For simplicity, the successive binding steps are described by rate constants 3k, 2k, and k, respectively, and each bound ligand by the same molar absorption coefficient ε . Some repetitions are shown. Values between brackets are standard deviations for the curve-fitting procedure.

M/Pigment	$k~(imes 10^3)~({ m M}^{-1}~{ m s}^{-1})$	$arepsilon$ ($ imes 10^3$) (M $^{-1}$ cm $^{-1}$)		
	Pigment A + Fe ²⁺ , pH 7			
0.3	4.1 (0.1)	9.5 (0.1)		
0.6	4.0 (0.3), 3.2 (0.2), 3.7 (0.2)	8.6 (0.2), 8.8 (0.2), 9.0 (0.1)		
1.0	3.1 (0.3), 4.0 (0.4), 3.7 (0.3)	9.9 (0.4), 8.4 (0.2), 9.1 (0.2)		
	Pigment B + Fe ²⁺ , pH 7			
0.3	2.03 (0.15), 4.16 (0.36)	38.3 (1.4), 26.1 (0.4)		
0.6	6.17 (0.15), 2.32 (0.13)	22.6 (0.1), 21.7 (0.2)		
1.0	0.90 (0.02), 1.18 (0.04), 1.03 (0.04)	23.6 (0.1), 23.1 (0.1), 23.2 (0.1)		
	Pigment A + Al ³⁺ , pH 7 Weak bi	inding		
	Pigment B + Al ³⁺ , pH 7			
0.3	0.81 (0.01)	15.4 (0.1)		
0.6	0.92 (0.01), 0.47 (0.01)	15.0 (0.1), 15.6 (0.1)		
1	0.42 (0.01), 0.51 (0.01), 0.39 (0.01)	14.5 (0.1), 16.0 (0.1), 14.6 (0.1)		
	P5 + Fe ²⁺ , pH 7			
0.3	2.7 (0.1)	24.5 (0.1)		
0.6	2.5 (0.1)	22.1 (0.1)		
1	3.1 (0.1)	24.9 (0.1)		
	P5 + Fe ²⁺ , pH 8			
0.3	13.6 (0.3)	13.5 (0.1)		
0.6	6.4 (0.1)	12.9 (0.1)		
1	6.6 (0.3)	12.8 (0.1)		
	P2 + Fe ²⁺ , pH 7			
0.3	16.6 (0.8)	6.7 (0.1)		
0.6	4.4 (0.1)	8.7 (0.1)		
1	2.5 (0.1)	10.0 (0.1)		
	P2 + Fe ²⁺ , pH 8			
0.3	14.1 (1.5)	12.4 (0.4)		
0.6	9.1 (0.7)	14.0 (0.3)		
1	5.5 (0.4)	13.6 (0.2)		
	P5 + Al ³⁺ , pH 7 (a)			
0.3	3.7 (0.1), 1.3 (0.1), 0.39 (0.05)	16.7 (0.2)		
0.6	0.58 (0.03), 0.14 (0.06) (b)	17.6 (1.1)		
1	0.50 (0.01) (c)	14.0 (0.1)		
	P5 + Al ³⁺ , pH 8			
0.3	5.5 (0.1)	10.3 (0.1)		
0.5	6.1 (0.1)	8.9 (0.1)		
1	1.7 (0.1)	7.4 (0.1)		
	P2 + Al ³⁺ , pH 7 Weak bindin	18		
	P2 + Al ³⁺ , pH 8			
0.3	4.1 (0.1)	3.4 (0.1)		
0.6	3.1 (0.1)	3.7 (0.1)		
1	1.2 (0.1)	3.8 (0.1)		

Note: (a) 3 optimizable rate constants; (b) $k_3 = 0$; (c) $k_2 = k_3 = 0$.

2.3. Competition between Metal Binding and Water Addition

Metal binding is accompanied by the removal of the B-ring's phenolic protons. In other words, protons and metal ions compete for the O-atoms of the B-ring. Hence, lowering the pH gradually destabilizes the complexes and a minimal pH for the onset of metal binding is expected. Also, a weaker stability for the complexes could mean reversibility in their formation and thus competition with water addition to the free form (flavylium ion), which leads to color loss.

The influence of pH on iron binding was investigated with the two isomers PB and P3. For a given pH, the UV-VIS spectra were recorded immediately after pigment addition to buffer in the absence of Fe^{2+} and after maximal binding in the presence of Fe^{2+} . A rise in visible absorbance in the range 600–700 nm, which is typical of metal binding, could be perceived with both pigments at pH \geq 3 (data not shown). Although P3 and PB could not be clearly distinguished by the pH for the onset of iron binding, it was obvious that the hyperchromic and bathochromic shifts in mildly acidic solution (pH 4.24) were much more spectacular with PB than with P3 (Figure 3A,C). Part of the interpretation is rooted in the higher susceptibility of P3 to color loss by reversible water addition, which was both faster and more complete than for PB (Figure 3B). Indeed, when the pigment and Fe²⁺ (1 equiv.) were added to the acetate buffer, a sharp drop of visible absorbance at 530 nm (free pigment) was observed with P3 over the first minute, whereas the increase of visible absorbance at 670 nm (iron complex) was negligible (Figure 3A). In a second phase, the onset of iron binding occurred, and A(670 nm) slowly increased. In this case, the first step corresponding to the flavylium hydration (fast) was clearly decoupled from the second step (slow) of iron binding. By contrast, with PB, the drop of A(530 nm) over the first minute was limited and accompanied by a rise of A(670 nm), which was then amplified along the second phase. This is evidence that hydration and metal binding now compete from the beginning. It is interesting to note that if hydration was faster than metal binding in the case of P3, the higher affinity of the colored forms (vs. colorless forms) for the metal ion eventually permitted the reversal of the hydration equilibrium and the slow development of metal binding. However, even with PB, the intensity of the complex's band in the pH range 3-6 remained much lower than in neutral solution, where it reached saturation even in the presence of substoichiometric iron concentrations. As the λ_{max} of the iron–PB complex's visible band was pH-independent, the spectrum at pH 7 (Figure S1B) could be used to calculate the percentage of iron complex at equilibrium at pH 4.24: 47%. A similar calculation with P3 gave only 24%.

Attempts to fit the spectral changes observed at 530 nm (free form) and 670 nm (iron complex) to a simple kinetic model, assuming competition with water addition and reversible iron binding, failed. However, a more sophisticated scheme assuming reversible Fe^{2+} binding by the colored forms, followed by irreversible autoxidation of bound Fe^{2+} with concomitant formation of a Fe^{3+} complex (in equilibrium with the free species), provided perfect curve-fittings and acceptable rate constants for the different steps (Table S5). Overall, PB bound Fe^{2+} twice as rapidly as P3 did, and the Fe^{2+} –PB complex seemed less susceptible to autoxidation than the Fe^{2+} –P3 complex. Finally, the blue color development was much more intense with PB, as the molar absorption coefficient of the PB–Fe³⁺ complex in the blue domain was ca. twice as large than that of the P3–Fe³⁺ complex. These remarkable improvements only reflected the shift of the single sinapoyl residue from the 6 position of Glc-1 to the 2 position of Glc-2. This is a spectacular example of the crucial importance of strong anthocyanidin–hydroxycinnamoyl π -stacking interactions in the development of vibrant blue colors.

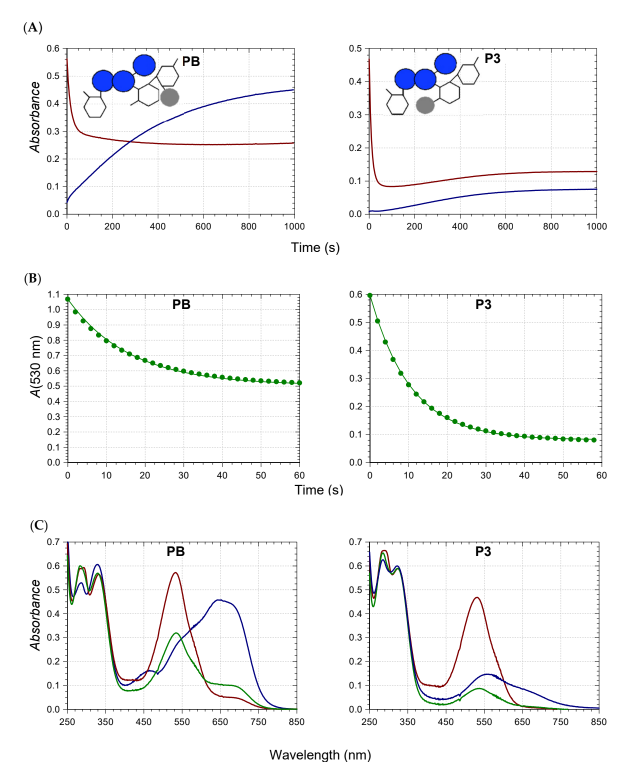


Figure 3. Cont.

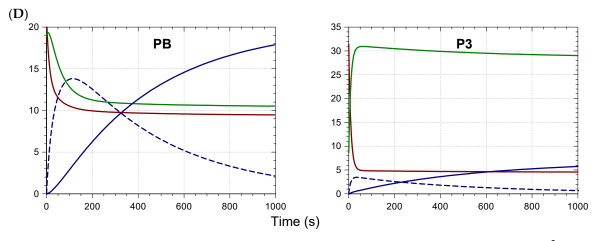


Figure 3. Iron–anthocyanin binding in a pH 4.24 acetate buffer (successive addition of pigment (40 μ M) and Fe²⁺ (1 equiv.)), Left: Pigment B; right: Pigment 3. Simplified schemes are provided as a reminder of the key structural differences. (A) Spectral monitoring at 530 nm (free pigment, —) and at 670 nm (iron complex, —). (B) Control in the absence of iron (solid lines are the result of the curve-fitting procedure). (C) Spectra at time zero (—), after ca. 1 min (—) and at the end of the kinetic run (—). (D) Time dependence of the species distribution: free colored forms (—); free colorless forms (—); Fe²⁺ complex (—).

2.4. Long-Term Stability of the Metal Complexes

Acylation promoted a moderate increase in color stability upon heating at 50 °C, again with an advantage conferred on pigments with R2 = sinapoyl (Table S6, Figure S6). This trend was hugely emphasized after Fe^{2+} addition (0.6 equiv.), a clear indication that the sinapoyl residue on Glc-2 was very efficient for the long-term stabilization of the metal complexes. Again, the comparison between the P3 and PB isomers was striking: a 25% color loss of the Fe²⁺-P3 complex was reached in 15–20 min vs. 4.5 h for the Fe²⁺-PB complex. As already observed [9], the spectroscopic titration of the residual pigment (after acidification to pH 1 for total dissociation of the iron complex and total conversion of the colorless forms into the flavylium ion) provided a more contrasted picture: in the absence of added iron, acylation offered to protect, and this so-called "acylation paradox" [9] could be interpreted by assuming that the colorless forms (hemiketal and chalcones) were much less susceptible to oxidative degradation than the electron-rich anionic base. In other words, being more vulnerable to reversible water addition, nonacylated PA and anthocyanins having a single HCA residue at R1 (P1, P3) were protected against autoxidation at the cost of losing their color. However, after Fe²⁺ addition, autoxidation was strongly accelerated for PA because the corresponding iron complex was not stable enough, and the protection offered by iron binding was significant for P4 and PB only. In our recent work, adding Fe²⁺ also was shown to increase the yield in some major degradation products of PA and P1 [18]. In summary, strong π -stacking interactions within the complexes ensured an efficient iron sequestration, thus preventing the pro-oxidant activity of loosely bound iron ions [16]. The tight metal binding also explained the total inhibition of intramolecular acyl transfer, normally occurring when solutions of P4 or PB are heated [18].

3. Materials and Methods

Red cabbage and purple sweet potato anthocyanins were isolated by preparatory LC according to already-published procedures [1,2]. Their structures are presented in Scheme 1.

3.1. Metal-Binding Experiments

3.1.1. Red Cabbage Anthocyanins

Metal–anthocyanin binding experiments were carried out according to previously reported procedures [13]. Fresh 5 mM solutions of Fe^{2+} and Al^{3+} were respectively pre-

pared from FeSO₄–7H₂O and AlCl₃–6H₂O (Sigma-Aldrich, St-Quentin Fallavier, France) in 1 mM aqueous HCl. Concentrated stock solutions of pigment (5 mM) were prepared in 50 mM aqueous HCl. Absorption spectra were recorded on an Agilent 8453 diode-array spectrometer in thermostated and magnetically stirred quartz cuvettes (pathlength = 1 cm). The following solutions were directly added to the cuvette in this order: 2 mL of 10 mM phosphate buffer (pH 7 or 8), 20 µL of anthocyanin stock solution and, after a few seconds (negligible formation of colorless forms), a small volume of the 5 mM Fe²⁺ or Al³⁺ solution (final iron/anthocyanin molar ratio = 1 or 2). The full UV-VIS spectra were recorded in kinetic mode for 1 to 2 min. For an optimal sensitivity, the detection in the visible range was set at 550 or 610 nm for Al³⁺ (close to the complex's λ_{max}) and at 670 nm for Fe²⁺ (charge-transfer contribution of the Fe³⁺ complexes). The hyperchromic and bathochromic shifts were calculated from the initial (free ligand) and final (metal complex) spectra as ($A_{max,f} - A_{max,0}$)/ $A_{max,0}$ and $\lambda_{max,f} - \lambda_{max,0}$, respectively.

Binding experiments were also carried out in the pH range 2–6 (50 mM acetate buffer) to determine the pH for the onset of metal binding and investigate the competition with water addition to the flavylium ion.

3.1.2. Purple Sweet Potato Anthocyanins

The anthocyanin isolates were diluted to a 50 μ M concentration in buffers of pH 6 (0.1 M sodium acetate), 7, and 8 (0.25 M TRIS). Small volumes of concentrated Al₂(SO₄)₃ solutions were then added to reach metal/anthocyanin ratios of 0.5, 1, and 5. Samples were equilibrated for 30 min at room temperature in the dark prior to analysis (in triplicates). UV-VIS spectra were collected from 380 to 700 nm using 300 μ L samples in poly-D-lysine-coated polystyrene 96-well plates with a SpectraMax 190 Microplate Reader (Molecular Devices, Sunnyvale, CA, USA).

3.2. Kinetic Analyses

The kinetic curves were analyzed with the Scientist software (Micromath, St. Louis, MO, USA). Sets of differential equations characteristic of the different kinetic processes (metal binding, water addition to the flavylium ion) were implemented in the models, as well as initial concentrations. Optimized values for the adjustable parameters (rate constants, molar absorption coefficients) and their standard deviations are reported.

3.3. Colorimetric Data

Color characteristics were expressed in the L*a*b* coordinates. L* corresponds to the light intensity, varying from 0 (no light) to 100. Parameters a* and b* quantify the contribution of four colors: green $(-a^*)$, red $(+a^*)$, blue $(-b^*)$ and yellow $(+b^*)$. With RC anthocyanins, the method to generate the L*a*b* coordinates and the corresponding color patches was as described in our recent work [13]. With PSP anthocyanins, the colorimetric data were generated as previously reported [2].

3.4. High-Resolution Mass Spectrometry (HRMS)

Stock solutions of anthocyanins P2, PB, and P5 were diluted to 0.1 mM in a 50 mM ammonium acetate buffer at pH 7. A 0.5 μ L volume was injected in the same solvent (flow injection analysis) over 1 min into an Orbitrap Exploris 480 mass spectrometer Exploris 480 (Thermo Fisher Scientific, Waltham, MA, USA) equipped with an H-ESI source. A static spray voltage of –3.5 kV in positive mode and of 2.5 kV in negative mode was applied with an ion-transfer tube temperature of 280 °C, a vaporizer temperature of 300 °C, and a N₂ sheath gas pressure of 40 psi. The full scan was recorded at a resolution of 24 × 10⁴ and corrected with an internal mass calibrant. Ions were searched between *m*/*z* 400 and 1600. The elution lasted ca. 0.3 min and an average spectrum was determined with the Freestyle 1.6 software based on the 20–30 spectra of intensities higher than 10%. A targeted detection of the Fe²⁺, Fe³⁺, and Al³⁺ complexes of stoichiometries 1:1, 1:2, and 1:3 and with charges ranging from –3 to +4 was carried out. Based on proposed raw formulae, the theoretical

isotopic patterns were calculated with the https://www.chemcalc.org/ web tool (accessed on 2 March 2021).

3.5. Thermal Degradation

Thermal degradation was performed at pH 7 and 50 °C in a thermostated water bath. The pigments were diluted to 50 μ M in the phosphate buffer at 50 °C and UV-VIS spectra were recorded over 24 h. The residual fraction of color species at pH 7 (a mixture of neutral and anionic bases) was determined at λ_{max} as % Color = 100 × $A_{\lambda max}(t)/A_{\lambda max}(t = 0)$. Aliquots of 1.5 mL were taken up at time zero, at regular time intervals over 8 h, and finally at t = 24 h. They were cooled down, acidified to pH 1, and stabilized at room temperature for 15 h (nonacylated anthocyanins) to 48 h (diacylated anthocyanins) (to ensure complete regeneration of the flavylium ion from the colorless forms). The absorption spectra were then recorded and the residual fraction of flavylium ion was calculated as % AH⁺ = 100 × $A_{\lambda max}(t)/A_{\lambda max}(t = 0)$ and plotted as a function of time. The time periods for a 25% color loss, or a 25% pigment loss (noted t_{25}), were determined from the theoretical curves generated by exponential curve-fitting.

4. Conclusions

Anthocyanins acylated by hydroxycinnamic acid residues are prone to develop interand intramolecular π -stacking interactions that make the flavylium nucleus less vulnerable to water addition. However, the acylation site plays a critical role, and among the red cabbage anthocyanins, it is clear that the sinapoyl residue at C2-OH of Glc-2 is by itself sufficient to provide resistance to water addition. Enhanced π -stacking interactions result from the ideal positioning of the sinapoyl residue (external Glc + C2 position), which probably reaches an optimal compromise between rigidity and flexibility.

The present work shows that optimal cyaniding–hydroxycinnamoyl π -stacking interactions can also exert a critical and favorable influence on metal binding. Indeed, only the metal complexes having R₂ = sinapoyl (pigments B, 4, 5, and 6) are deeply blue and thermally stable over long periods of time. Not only the sinapoyl residue (although noncoordinating by itself) does not compromise metal binding by steric hindrance, it also shifts the visible band to the blue domain and increases its intensity, and stabilizes the metal complexes and prevents the release of free pro-oxidant iron in the medium.

Given the diversity of metal ions possibly binding anthocyanins in plants, other metal complexes of acylated cyanidin glycosides might also be tested for their coloring properties. In summary, anthocyanins having a 3',4'-dihydroxysubstitution (cyanidin, delphinidin and petunidin glycosides) with adequately located hydroxycinnamoyl residues offer bright prospects for further development as natural food colorants.

Supplementary Materials: Supplementary materials can be found at https://www.mdpi.com/article/10.3390/ijms22094551/s1.

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