# Natural hydroxyanthraquinoid pigments as potent food grade colorants: an overview

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Abstract: Natural pigments and colorants are widely used in the world in many industries such as textile dying, food processing or cosmetic manufacturing. Among the natural products of interest are various compounds belonging to carotenoids, anthocyanins, chlorophylls, melanins, betalains... The review emphasizes pigments with anthraquinoid skeleton and gives an overview on hydroxyanthraquinoids described in Nature, the first one ever published. Trends in consumption, production and regulation of natural food grade colorants are given, in the current global market. The second part focuses on the description of the chemical structures of the main anthraquinoid colouring compounds, their properties and their biosynthetic pathways. Main natural sources of such pigments are summarized, followed by discussion about toxicity and carcinogenicity observed in some cases. As a conclusion, current industrial applications of natural hydroxyanthraquinoids are described with two examples, carminic acid from an insect and Arpink red<sup>TM</sup> from a filamentous fungus.

Keywords: anthraquinone, hydroxyanthraquinone, natural colorant, food colorant, microbial pigment, biotechnology, mycotoxin contamination

#### Introduction

Food grade colorants can loosely be categorized as 'natural' or 'synthetic'. The term 'natural colorants' indicates that the source of the colorant is natural even if varying definitions and regulations exist according to the country in question. For example, under the United States (US) Food and Drug Administration (FDA) regulations, a colorant added to a food product cannot be considered as 'natural', no matter what the source is; unless the colorant is natural to the food product itself. The FDA regulates the natural and synthetic colorants of food applications in two classes. In general, the synthetic colorants (that do not exist in nature) are subjected to a certification requirement to assure that each batch of material manufactured meets the standard specifications, while natural colorants are "exempt from certification" and may be manufactured and marketed without certification of FDA (no US Food, Drug and Cosmetic Act (FD&C)-number). In contrast, E-numbers are used for all colorants for food applications in the European Union (EU). Colorants for food applications listed by both the FDA and the EU are tested for biosafety before their promotion and commercialization, and are further controlled by national legislation specifying those colorants that may be used, the type of food that may be coloured, the quantity that may be added and the limit of maximum daily intake.

For a very long time, the use of food colorants focused on synthetic ones. However, over the last few decades, synthetic colorants tend to be perceived as undesirable by consumers, due to the harmful effects of some synthetic pigments on human health, including allergic reactions, mutagenicity and carcinogenicity (e.g. skin cancer)<sup>1</sup>. manufacturers have considered replacing synthetic colorants in their food products with natural colouring alternatives in response to pressure from both customers and regulators<sup>2</sup>. Whereas 43 colorants were authorized in the EU as food additives in 1994, actually almost a hundred of food grade colorants are authorized in the EU and have been assigned by an 'E-number'; almost 40% of these were of natural origin<sup>1,3</sup> These natural colorants are usually applied in several industrial food processes for the same reasons as the synthetic counterparts: (i) to enhance the product's natural colour whose ingredients are unable to provide a sufficient colour; (ii) to standardize the colour and appearance of product like confectionery; (iii) to restore what has been lost during processing or (iv) to add a novel sensory aspect that attracts customers.

Among the natural pigments of interest are various compounds belonging to carotenoids, anthocyanins,

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chlorophylls, melanins, betalains, quinones... This review emphasizes pigments with anthraquinoid skeleton and gives an overview on hydroxyanthraquinoid pigments which are widely present in Nature and are gaining increasing interest by academics and people from the industry.

### 1 Trends in consumption, production and regulation of natural food grade colorants in the current global market

Currently, the natural food colouring industry market is growing 10%-15% annually. Natural varieties share of the global food colorant market increased from about 31% in 2005 to 36% in 2009. The current consumer preference for natural food grade colorants is associated with their image of being healthy and of good quality. According to a report from Leatherhead Food Research, Shaun Weston mentions that the global market for food grade colorants is expected to reach \$1.6 billion USD by 2015, up to 10% from its present levels and fuelled mainly by the growth in natural colorants and colouring foodstuffs (data from Leatherhead Food International LFI) (www.leatherheadfood.com). The main industrial technology used for the production of natural colorants for food applications depends on the extraction of coloured pigments from edible plants, fruits or vegetables. Table 1 shows the main natural food grade colorants authorized and currently available in the current global market. Common natural colorants include turmeric, curcumin, annatto, paprika, caramel and cochineal extract. Natural colorants are often commercially available in powder, oil-soluble emulsion, or water-soluble emulsion forms.

Table 1. Main natural food grade colorants authorized and currently available in the current global market

Color/shade	E-number*	Natural colorant (	Chemical category
From plants, fruit	or vegetables:		
Yellow	E100, E100 (i)	Curcumin	Curcuminoid
Yellow	E100 (ii)	Turmeric	Curcuminoid
Green	E140	Chlorophylls	Tetrapyrrole
Green	E141	Chlorophyllins	Tetrapyrrole
Brown	E150a-d	Caramel	Melanoidin
Orange-yellow	E160a (i)	<b>Mixed Carotenes</b>	Carotenoid
Orange-yellow	E160a (ii)	$\beta$ -carotene	Carotenoid
Yellow to orange	E160b	Annatto	Carotenoid
Yellow to orange	E160b (i)	Annato (Bixin)	Carotenoid
Yellow to orange	E160b (ii)	Annato (Norbixin)	Carotenoid
Red	E160c	Paprika (Capsanthir	n) Carotenoid
Yellow to red	E160d	Lycopene	Carotenoid
Yellow to red	E160e	Apocarotenal	Carotenoid
Orange-yellow	E161a	Flavoxanthin	Carotenoid
Orange-yellow	E161b	Lutein	Carotenoid
Orange-yellow	E161d	Rubixanthin	Carotenoid
Orange-yellow	E161e	Violaxanthin	Carotenoid
Orange-yellow	E161f	Rhodoxanthin	Carotenoid
Orange, Red	E161h	Zeaxanthin	Carotenoid
Red	E162	Red Beet Juice	Betalain
Red, Blue or Violet	E163a	Cyanidin	Anthocyanin
Red, Blue or Violet	E163e	Peonidin	Anthocyanin

From animal:			
Yellow	E101, E101a	Riboflavin	Flavin
Magenta-red	E120 (ii)	Carminic acid	Anthraquinone
		(Cochineal extract)	
Orange-yellow	E161c	Cryptoxanthin	Carotenoid
Orange, Red	E161g	Canthaxanthin	Carotenoid
From microorga	nisms:		
Yellow	E101 (iii)	Riboflavin (from Bacillus subtil. Other sources: Ashby Candida gulliermndi Clostridium and Debaryomyces su	va gossypii, i, acetobutylicum
Orange-yellow	E160a (ii)	<b>β-carotene</b> (from <i>Blakeslea trisp</i>	Carotenoid pora)
Orange-yellow	E160a (iv)	β-carotene (from Dunaliella sali Other sources: Dunal	
Yellow to red	E160d (iii)	<b>Lycopene</b> (from <i>Blakeslea trisp</i>	Carotenoid pora)
Yellow to red	E-161j	Astaxanthin (from Haematococcu Other sources: Haem lacustris, Xanthophy, dendrorhous	atococcus
Orange, Red	E161g	Canthaxanthin (from Haematococcu	Carotenoid as lacustris)
		Other sources: Brady	rhizobium sp.

\*E-number of the corresponding authorized food colorant in the European Union

Many scientific papers describe the extraction, characterization and properties of natural pigments from fruits, vegetables, lichens and marine life<sup>1,3</sup>. However, the potential of these renewable resources as sources for new commercial natural food grade colorants would still be limited both by the manufacturing costs and the availability of the raw material, which would need to be cultivated in sufficient quantities for industrial extraction. The microbial pigment production by biotechnology would have the advantage of producing higher yields. This kind of pigment production is not at all dependent on the availability and external supply of particular raw materials. In addition, microbial pigments are often more stable and water-soluble than those of plant sources<sup>4,5</sup>. The really first European success story in pigment production using a microorganism is  $\beta$ -carotene (additive E-160a(ii); orange-yellow pigment) from the fungus *Blakeslea trispora* by DSM<sup>TM</sup>. Among microalgae, some successful stories yield to efficient production of carotenoids using Dunaliella salina (e.g.,  $\beta$ -carotene, additive E-160a(iv)) or *Haematococcus* pluvialis (e.g., astaxanthin, additive E-161j; yellow to red pigment)<sup>6</sup> (Table 1). Nowadays, fermentative productions of natural food grade colorants are available in the global market<sup>7</sup>. This approval of microbial carotenoids as food colorants has strengthened the prospects for new natural colorants<sup>2</sup>.

However, some new microbial pigments might not be accepted if they were to be introduced into industrial food manufacturing today<sup>8</sup>. The commercially available *Monascus* pigments are a perfect example. These fungal pigments are natural azaphilone pigment mixtures. The red colorant obtained is produced commercially using strains of *Monascus* fungi in the Orient for centuries used as a food colorant for making red rice wine, red soybean cheese, meat and marine





products. However, Monascus pigments are still not allowed as a food additive in either the US or the EU; there have been controversial views presented over their safe use<sup>8</sup>. This could be essentially because of the presence of the mycotoxin citrinin (yellow compound) and some other potential toxic metabolites which may occur in some batches with Monascus fungi9-The production of citrinin limits the commercial use of Monascus fungi as producers of natural food grade colorants<sup>1</sup> Research continues on new azaphilone pigments produced from non-mycotoxigenic fungal strains, such as Epicoccum nigrum, Penicillium aculeatum or P. pinophilum—that are incapable of co-producing citrinin-in the prospects for new natural food grade colorants<sup>8,13</sup>. The case of the fungal Arpink red<sup>TM</sup> colorant, i.e. a natural food colorant manufactured by the Czech company, Ascolor Biotech s.r.o. is also atypical. This company has produced a chomophore of the anthraquinoid type as a natural food colorant, by fermentation and bioprocess engineering using the strain Penicillium oxalicum var. Armeniaca CCM 8242 obtained from soil (the variety was never formally described). The Arpink red<sup>TM</sup> colorant has received a two-year temporary approval by the EU for distribution as a food additive, exclusively in the Czech Republic from 2004 to 2006. The extraction, isolation and characterization of natural anthraquinoid pigments have been also reported from other filamentous fungi with different shades such as red, reddish brown, bronze and maroon.

# 2 Natural hydroxyanthraquinoid pigments: chemical structures of the main colouring components, their properties and their biosynthetic pathways

Anthraguinones are a class of compounds of the quinone family that consists of several hundreds of compounds that differ in the nature and positions of substituent groups. Anthraquinoid derivatives are derivatives of the basic structure 9,10-anthracenedione or also called 9,10-dioxoanthracene, i.e. a tricyclic aromatic organic compound with formula C<sub>14</sub>H<sub>8</sub>O<sub>2</sub> and whose ketone groups are on the central ring in position C-9 and C-10. Figure 1 shows the skeleton structure of anthraquinoid derivatives. In general, for each anthraquinoid derivative there are eight possible hydrogens that can be substituted. The term 'hydroxyanthraquinoid (HAQN)' derivatives usually refers to derivatives of 9,10-hydroxyanthraquinone, i.e. derivatives of 9,10-anthraquinone where any number n of hydrogen atoms have been replaced by nhydroxyl (-OH) groups. In this case the number n of hydroxyl group is indicated by a multiplier prefix (mono-, di-, tri-, up to octa-). The HAQN derivatives absorb visible light and are coloured, whereas strictly 9,10-anthraquinone derivatives are colorless like tectoquinone<sup>14</sup>.

Most HAQN colour compounds of natural origin have

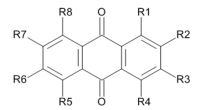


Figure 1. The skeleton structure of anthraquinoid derivatives





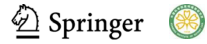
complex structures with several functional groups, which modify their absorption spectra. The chemical structures of natural HAQN pigments and their main physical and functional properties 15-52 are shown on Table 2. In the UV region, substituted 9,10-anthraquinone derivatives show intense benzenoid absorption bands fairly regularly within the ranges 240-260 and 320-330 nm. The quinonoid bands appear in a range from 260 to 290 nm and 9,10-hydroxyanthraquinone derivatives show an absorption band between 220 and 240 nm. The HAQN derivatives have attracted the attention of many researchers due to their large list of possible applications related to their interesting photoactivity and more particularly based on their chromatic properties. They possess good light-fastness properties, which makes metallization unnecessary. HAQN derivatives can form coordination complexes with several cations. They are relatively stable and the advantage of pigments of HAQN-type compared to azo pigments is their superior brightness. Moreover, ionization of a hydroxylic group results in a bathochromic shift. It appears that the colour of the HAON pigments depends on the position and number of the hydroxyl substituents in the different rings<sup>5</sup>

Natural HAQN pigments are produced by the secondary metabolism of organisms. One of the remarkable features of natural HAQN biosynthesis is that they are derived from a variety of different precursors and pathways. There are at least two biosynthetic pathways leading to HAQN pigments. On one hand, the most important is the polyketide pathway (acetate-malonate pathway) that includes suitable folding and condensation of an octaketide chain derived from acetate (acetyl-CoA) and malonate (malonyl-CoA) units<sup>3,32,54</sup>. The resultant polycarbonyl compounds serve as substrates for various cyclases that produce aromatic compounds that represent typical fungal metabolites. Natural HAQN pigments that are synthesized following this acetate-malonate pathway (see Figure 2) always show a characteristic substitution pattern, i.e. they show substitution on both aromatic rings and more particularly at least one hydroxyl group in position R1 and one hydroxyl or methoxyl (-OCH<sub>3</sub>) group in position R8: examples emodin, physcion, endocrocin, dermolutein, dermoglaucin, dermorubin and dermocybin. According to this polyketide pathway, the biosynthetic relationships show that the yellow compounds (e.g., emodin, physcion, endocrocin and dermolutein) exist in the beginning of the synthesis pathway whereas the red compounds like dermorubin and dermocybin are more complicated in structure and occur in the latter part of the biosynthesis pathway<sup>32</sup>. More recently, Bringmann et al.55 revealed that the pigment chrysophanol is shown to be formed, in an organism-specific way, by a third folding mode involving a remarkable cyclization of a bicyclic diketo precursor, thus establishing the first example of multiple convergence in polyketide biosynthesis.

On the other hand, HAQN pigments are formed via the shikimate or chorismate/o-succinylbenzoic acid pathway (Fig. 2). HAQN pigments that are synthesized via this pathway only have one of the rings unsubstituted and at least one hydroxyl group in position R1 on the ring C. The rings A and B are derived from chorismate and  $\alpha$ -ketoglutarate via o-succinylbenzoic acid (the HAQN biosynthesis branch at 1,4-dihydroxy-2-naphtoic acid), whereas ring C is formed from isopentenyl diphosphate either formed via the mevalonic acid pathway or the 2-C-methyl-D-erythritol 4-phosphate pathway<sup>54,56,57</sup>. The relevant colouring compounds are alizarin

Table 2. Chemical structures, physical and functional properties of natural hydroxyanthraquinoid (HAQN) pigments

HAQN dye (and it's abbreviation)	Chemical structure	Physical and functional properties	References
Monohydroxyanthraquinones (OH) <sub>1</sub>			
Damnacanthal 3-hydroxy-1-methoxy- anthraquinone-2-carboxaldehyde	O OCH₃ CHO OH	$C_{16}H_9O_4(OH)_1$ – Mw: 282 Mp: 210–211 °C Shade: pale yellow	(15, 16)
Lucidin primeveroside 3-O-primeverose-1-hydroxy-2-hydroxymethyl-anthraquinone (LuP)	O OH CH₂OH O-Pr	$C_{26}H_{27}O_{13}(OH)_1$ – Mw: 564 Mp: 210–212 °C Shade: red	(17)
Pachybasin 2-methyl-1-hydroxy-anthraquinone	O OH	C <sub>15</sub> H <sub>9</sub> O <sub>2</sub> (OH) <sub>1</sub> – Mw: 238 Mp: 176 °C; shade: yellow	
2-incluyi-1-nydioxy-anunaquinone	CH <sub>3</sub>	UV (EtOH) $\lambda_{max}$ 403, 281, 252, 224 nm; IR (KBr) $v_{max}$ 3083, 2967, 2933, 2862, 1677, 1642, 1593, 1578, 1403, 1380, 1338, 1311, 1290, 1235, 1148, 1068, 1039, 1002, 929, 870, 805, 793, 769, 712 cm <sup>-1</sup> ; <sup>1</sup> H NMR (CDCl <sub>3</sub> , 500 MHz) $\delta$ 12.67, 8.20, 7.74, 7.55, 7.17, 2.45.	(18)
Ruberythric acid (= alizarin primeveroside) 2-O-primeverose-1-hydroxy- anthraquinone	O OH O-Pr	$C_{25}H_{25}O_{12}(OH)_1$ – Mw: 534 Mp: 259–261 °C Shade: yellow Soluble in hot water	(17, 19)
(Rba)		Slightly soluble in alcohols and ether Insoluble in benzene	
- Dihydroxyanthraquinones (OH)2:			
2-(1-hydroxyethyl)-3,8-dihydroxy- 6-methoxy-anthraquinone	H <sub>3</sub> CO OH OH OH OH OH	$C_{17}H_{12}O_4(OH)_2 - Mw: 314$ Mp: $208-212$ °C; shade: orange UV (MeOH) $\lambda_{max}$ 466, 341, 305, 285, 219 nm; IR (KBr) $v_{max}$ 3420, 1667, 1630, 1580, 1480, 1440, 1390, 1360, 1305, 1270, 1240, 1215, 1160, 1080, 1030, 1005, 970, 930, 900, 875, 835, 800 cm <sup>-1</sup> ; <sup>1</sup> H NMR (CDCl <sub>3</sub> , 500 MHz) $\delta$ 12.96, 7.94, 7.76, 7.35, 6.70, 5.20, 3.93, 1.68.	(20, 21)
	OH O O	C <sub>17</sub> H <sub>10</sub> O <sub>4</sub> (OH) <sub>2</sub> – Mw: 312 Mp: 256–270 °C; shade: yellow	
2-acetyl-3,8-dihydroxy-6-methoxy- anthraquinone	H <sub>3</sub> CO OH	UV (MeOH) λ <sub>max</sub> 420, 348, 307, 280, 233 nm; <b>IR</b> (KBr) v <sub>max</sub> 1672, 1655, 1630, 1575, 1480, 1450, 1410, 1380, 1300, 1250, 1205, 1170, 1125, 1025, 970, 935, 905, 890, 865, 830, 805, 755, 745, 715, 695 cm <sup>-1</sup> ; <sup>1</sup> <b>H NMR</b> (CDCl <sub>3</sub> , 500 MHz) δ 12.91, 12.72, 8.76, 7.81, 7.39, 6.74, 2.81.	(20, 21)
Alizarin 1,2-dihydroxy-anthraquinone (Ali)	ООНОН	$C_{14}H_6O_2(OH)_2-Mw;\ 240$ Mp: $278{-}280^{\circ}C$ Shade : Yellow to red (acid); red to violet (alkaline)	(22, 23)
		Soluble in water, in NaOH 1M and in alcohols UV (MeOH) $\lambda_{max}$ 609, 567, 429, 422 nm.	
Aloe-emodin 3-hydroxymethyl-1,8-dihydroxy- anthraquinone (Ale)	OH O OH CH <sub>2</sub> OH	$C_{15}H_8O_3(OH)_2$ – Mw: 270 Mp: 223–224 °C Shade: orange-yellow	(24, 25, 26)
	0		





Anthraflavic acid (= anthraflavin) 2,6-dihydroxy-anthraquinone (Afv)	но	$C_{14}H_6O_3(OH)_2 - Mw$ : 240 Mp $\geq$ 320 °C Shade: yellow	(19)
Arpink red <sup>TM</sup>	H <sub>3</sub> C OH OH OH OH	$C_{22}H_{13}O_4(OH)_1$ Shade: red	(2, 5, 15)
Austrocortinin 2-methoxy-7-methyl-1,4-dihydroxy- anthraquinone	H <sub>3</sub> C OCH <sub>3</sub> OCH <sub>3</sub>	$C_{16}H_{10}O_3(OH)_2 - Mw$ : 284 Shade: red	(27)
Chrysophanol (=chrysophanic acid) 3-methyl-1,8-dihydroxy- anthraquinone (Chr)	OH O OH CH3	$C_{15}H_8O_2(OH)_2 - Mw$ : 254 Mp: 186 °C Shade: orange-yellow UV (EtOH) $\lambda_{max}$ 436, 288, 278, 256, 226 nm; <sup>1</sup> H NMR (DMSO, 500 MHz) $\delta$ 11.93, 7.81, 7.72, 7.56, 7.39, 7.23, 2.45.	(18, 28-31)
Danthron (= dantron or chrysazin) 1,8-dihydroxy-anthraquinone (Dan)	OH O OH	C <sub>14</sub> H <sub>6</sub> O <sub>2</sub> (OH) <sub>2</sub> – Mw: 240 Mp: 190-195 °C Shade: reddish to orange Very soluble in alkaline hydroxide solutions Insoluble in water, acetone, chloroform, diethyl ether and ethanol	(27)
Dermolutein 8-methoxy-3-methyl-1,6-dihydroxy- anthraquinone-2-carboxylic acid	OCH <sub>3</sub> O OH O OH OH OH	$C_{17}H_{10}O_{5}(OH)_{2} - Mw$ : 326 Shade: yellow	(32)
Fallacinal 3-formyl-6-methoxy-1,8-dihydroxy- anthraquinone	Н₃СО ОН	C <sub>16</sub> H <sub>8</sub> O <sub>4</sub> (OH) <sub>2</sub> – Mw: 298 Mp: 227–228 °C Shade: yellow <b>IR</b> (KBr) v <sub>max</sub> 2850, 2835, 2740, 1715, 1635, 1600 cm <sup>-1</sup> ; <sup>1</sup> <b>H NMR</b> (CDCl <sub>3</sub> , 500 MHz) δ 12.17, 12.19, 8.29, 7.76, 7.44, 6.74, 3.97.	(33)
Frangulin A (=franguloside) 3-O-primeverose-6-methyl-1,8- dihydroxy-anthraquinone (Fran)	H <sub>3</sub> C O-Rh	$C_{21}H_{18}O_7(OH)_2$ – Mw: 416 Shade: orange	(24)



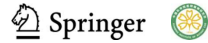


Lucidin (=henine) 2-hydroxymethy-1,3-dihydroxy-anthraquinone	O OH CH <sub>2</sub> OH	C <sub>15</sub> H <sub>8</sub> O <sub>3</sub> (OH) <sub>2</sub> – Mw: 270 Mp: 300 °C Shade: red	(19, 23, 34)
Munjistin 1,3-dihydroxy-anthraquinone-2- carboxylic acid (Mun)	ООНООН	$C_{15}H_6O_4(OH)_2 - Mw$ : 284 Shade: orange-red	(17)
Nordamnacanthal 2-formyl-1,3-dihydroxy -anthraquinone (Nor)	О ОН СНО	$C_{15}H_6O_3(OH)_2 - Mw$ : 268 Mp: 214–218 °C Shade: orange-yellow	(16, 17)
Phomarin 3-methyl-1,6-dihydroxy- anthraquinone	HO CH <sub>3</sub>	C <sub>15</sub> H <sub>8</sub> O <sub>2</sub> (OH) <sub>2</sub> – Mw: 254 Shade: yellow UV (MeOH) λ <sub>max</sub> 410, 293, 268, 219 nm; IR (KBr) ν <sub>max</sub> 3414, 3310, 2925, 2856, 1677, 1640, 1603, 1579, 1488, 1389, 1305, 1268, 1191, 1153, 1020, 945, 894, 854, 817, 795, 795 cm <sup>-1</sup> ; <sup>1</sup> H NMR (acetone, 500 MHz) δ 8.14, 7.65, 7.56, 7.28, 7.10.	(18)
Physcion (= parietin) 6-methoxy-3-methyl-1,8-dihydroxy- anthraquinone (Phy)	H <sub>3</sub> CO CH <sub>3</sub>	$C_{16}H_{10}O_3(OH)_2 - Mw$ : 284 Mp: 207 °C; shade: yellow UV (EtOH) $\lambda_{max}$ 435, 283, 265, 253, 226 nm; IR (CHCl <sub>3</sub> ) $v_{max}$ 2929, 2854, 1626, 1568, 1483, 1140 cm <sup>-1</sup> ; <sup>1</sup> H NMR (CDCl <sub>3</sub> , 500 MHz) $\delta$ 7.08, 7.62, 7.36, 6.69, 3.94, 2.45.	(35)
Questin 8-methoxy-3-methyl-1,6-dihydroxy- anthraquinone	HO CH <sub>3</sub>	$C_{16}H_{10}O_3(OH)_2-Mw:~284$ Mp: $301-303$ °C Shade: yellow to orange-brown Soluble in aqueous sodium carbonate	(36)
Quinizarin 1,4-dihydroxy-anthraquinone (Qza)	O OH	$C_{14}H_6O_2(OH)_2$ – Mw: 240 Mp: 198–199 °C Shade: Orange to red-brown Slightly soluble (but soluble in hot water) UV (MeOH) $\lambda_{max}$ 480 nm	(19, 34, 37, 38)
Rhein (=cassic acid) 1,8-dihydroxy-anthraquinone- 3-carboxylic acid (Rhe)	OH O OH	C <sub>15</sub> H <sub>6</sub> O <sub>4</sub> (OH) <sub>2</sub> – Mw: 284 Mp: 320–321 °C; shade: orange Insoluble in water <b>IR</b> (KBr) v <sub>max</sub> 3441, 3407, 1693, 1673, 1624, 1560, 1449 cm <sup>1</sup> ; <sup>1</sup> <b>H NMR</b> (CDCl <sub>3</sub> , 500 MHz) δ 11.87, 8.07, 7.81, 7.71, 7.69, 7.38.	(33)
Rubiadin 2-methyl-1,3-dihydroxy- anthraquinone	O OH CH <sub>3</sub>	$C_{15}H_8O_2(OH)_2$ – Mw: 254 Mp: 290 °C (in alcohols) and 302 °C (in glacial acetic acid) Shade: yellow UV (EtOH) $\lambda_{max}$ 415, 280, 246 nm.	(19, 39, 40)





Soranjidiol 2-methyl-1,6-dihydroxy- anthraquinone	O OH CH <sub>3</sub>	$C_{15}H_8O_2(OH)_2 - Mw: 254$	(39, 40)
Teloschistin (= fallacinol or phallacinol) 3-hydroxymethyl-1,8-dihydroxy-6-methoxy-anthraquinone	H <sub>3</sub> CO	C <sub>16</sub> H <sub>10</sub> O <sub>4</sub> (OH) <sub>2</sub> – Mw: 300 Mp: 236–237 °C; Shade: yellow IR (KBr) v <sub>max</sub> 3450, 2840, 1670, 1630, 1625 cm <sup>-1</sup> ; <sup>1</sup> H NMR (CDCl <sub>3</sub> , 500 MHz) δ 12.30, 12.20, 7.97, 7.40, 7.39, 6.70, 3.98.	(33)
Xanthopurpurin 1,3-dihydroxy-anthraquinone (Xpu)	OH OH	$C_{14}H_6O_2(OH)_2-Mw$ : 240 Shade: red	(19, 23, 37)
- Trihydroxyanthraquinones (OH)3:			
Anthragallol (= alizarin brown) 1,2,3-trihydroxy-anthraquinone (Agl)	ОНОН	$C_{14}H_5O_2(OH)_3$ – Mw: 256 Mp: 312–313 °C Shade: orange Soluble in alcohol, ether and glacial acetic acid Slightly soluble in water and chloroform	(19)
Citreorosein 6-hydroxymethyl- 1,3,8-trihydroxy-anthraquinone	HOH₂C OH	$C_{15}H_7O_3(OH)_3$ – Mw: 286 Shade: yellow UV (EtOH) $\lambda_{max}$ 448, 435, 290, 268, 266, 253, 252, 221, 207 nm.	(41, 42)
<b>Dermoglaucin</b> 3-methyl-1,7,8-trihydroxy- 6-methoxy-anthraquinone	HO OH O OH CH <sub>3</sub>	C <sub>16</sub> H <sub>9</sub> O <sub>3</sub> (OH) <sub>3</sub> – Mw: 300 Shade: red	(32)
Dermorubin 3-methyl-1,4,6-trihydroxy- 8-methoxy-anthraquinone- 2-carboxylic acid	OCH <sub>3</sub> O OH O OH OH OH OH	$C_{17}H_9O_5(OH)_3 - Mw$ : 344 Shade: red	(32)
Emodin (= frangula emodin) 3-methyl-1,6,8-trihydroxy- anthraquinone (Emo)	HO O OH CH <sub>3</sub>	C <sub>13</sub> H <sub>7</sub> O <sub>2</sub> (OH) <sub>3</sub> – Mw: 270 Mp: 254–256 °C; Shade: orange Soluble in ethanol, in DMSO UV (EtOH) λ <sub>max</sub> 437, 289, 265, 252, 222 nm; <b>IR</b> (KBr) ν <sub>max</sub> 3353, 3061, 1730, 1670, 1624, 1558, 1475, 1451, 759 cm <sup>-1</sup> ; <sup>1</sup> <b>H NMR</b> (CDCl <sub>3</sub> , 500 MHz) δ 12.07, 11.99, 11.19, 7.45, 7.12, 7.09, 6.56, 2.40.	(33, 42, 43)
Endocrocin 3-methyl-1,6,8-trihydroxy- anthraquinone-2-carboxylic acid	OH O OH O OH OH OH OH	$C_{16}H_7O_4(OH)_3 - Mw: 314$ Shade: yellow	(32)





Erythroglaucin 3-methyl-1,4,8-trihydroxy- 6-methoxy-anthraquinone	OH O OH  CH <sub>3</sub>	$C_{16}H_9O_3(OH)_3 - Mw$ : 300 Shade: red	(4)
Flavokermesic acid (= Laccaic acid D) 1-methyl-3,6,8-trihydroxy- anthraquinone-2-carboxylic (Flk)	O OH  O CH <sub>3</sub> O OH  HO OH  O OH	$C_{16}H_7O_4(OH)_3$ – Mw: 314 Shade: yellow	(15, 44, 45)
Flavopurpurin (= alizarin Y) 1,2,6-trihydroxy-anthraquinone	но	$C_{15}H_5O_2(OH)_3 - Mw$ : 256 Shade: yellow	(19)
Helminthosporin 3-methyl-1,5,8-trihydroxy- anthraquinone	OH O OH CH <sub>3</sub>	$C_{15}H_7O_2(OH)_3 - Mw$ : 270 Shade: brown	(4)
Islandicin 2-methyl-1,4,5-trihydroxy- anthraquinone	O OH CH <sub>3</sub>	$\begin{split} &C_{15}H_7O_2(OH)_3-Mw:270\\ &Mp:217-219~^{\circ}C\\ &Shade:~red\\ &\textbf{UV}~(EtOH)~\lambda_{max}~550,~532,~510,~479,~294,\\ &253,~~234~~nm~;\\ &\textbf{IR}~(KBr)~v_{max}~1602~cm^{-1} \end{split}$	(46)
Morindone 6-methyl-1,2,5-trihydroxy- anthraquinone	O OH OH	C <sub>15</sub> H <sub>7</sub> O <sub>2</sub> (OH) <sub>3</sub> – Mw: 270 Shade: yellowish-red <sup>1</sup> H NMR (CDCl <sub>3</sub> , 500 MHz) δ 13.19, 12.95, 7.61, 7.52, 7.24, 6.99, 2.14.	(47)
Pseudopurpurin 1,3,4-trihydroxy-anthraquinone- 2-carboxylic acid (Psp)	ООНООНООН	$C_{15}H_5O_4(OH)_3 - Mw$ : 300 Shade: red	(15)
Purpurin 1,2,4-trihydroxy-anthraquinone (Pur)	ООНОН	$C_{14}H_5O_2(OH)_3$ – Mw: 256 Mp: 265–270 °C; shade : yellow to red (acid); red to violet (alkaline) Soluble in water and chloroform Insoluble in hexane UV (EtOH) $\lambda_{max}$ 521, 515, 480 nm.	(15, 23, 37, 38)
Rubrocristin 2-methyl-1,4,7-trihydroxy- 5-methoxy-anthraquinone	HO OCH <sub>3</sub> O OH CH <sub>3</sub>	$C_{16}H_9O_3(OH)_3 - Mw$ : 300 Shade: red	(36)





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Skyrin	HO OH O OH	$C_{30}H_{15}O_7(OH)_3 - Mw$ : 358 Shade : yellow to red	(48)
Tetrahydroxyanthraquinones (OH)			
Averythrin 2-(1-Hexenyl)-1,3,6,8-tetrahydroxy- anthraquinone (Avt)	HO OH OH CH3	$C_{20}H_{14}O_{2}(OH)_{4} - Mw: 354$ Shade : orange	(49)
Carminic acid 2-a-D-glucopyranosyl-8-methyl- 1,3,4,6-tetrahydroxy-anthraquinone- 7-carboxylic acid (Car)	CH <sub>2</sub> OH OH OH OH	$C_{22}H_{16}O_{9}(OH)_{4} - Mw$ : 492 Mp: 120 °C; shade: red to violet (alkaline); or orange (acid) Good solubility in water UV (EtOH) $\lambda_{max}$ 495, 491, 311, 278 nm.	(15, 22, 37, 50)
Catenarin 3-methyl-1,4,5,7-tetrahydroxy- anthraquinone	HO OH CH <sub>3</sub>	C <sub>15</sub> H <sub>6</sub> O <sub>2</sub> (OH) <sub>4</sub> – Mw: 286 Mp: 240 °C; shade: red UV (MeOH) λ <sub>max</sub> 525, 512, 490, 480, 463, 306, 278, 257, 230 nm; <sup>1</sup> H NMR (CDCl <sub>3</sub> , 500 MHz) δ 13.34, 12.42, 12.35, 7.32, 7.13, 6.66, 2.35.	(51)
Cynodontin 2-methyl-1,4,5,8-tetrahydroxy- anthraquinone (Cyn)	OH O OH CH <sub>3</sub>	$C_{15}H_6O_2(OH)_4 - Mw$ : 286 Shade: bronze	(4)
<b>Dermocybin</b> 3-methyl-1,5,7,8-tetrahydroxy-6-methoxy- anthraquinone	H <sub>3</sub> CO OH O CH <sub>3</sub>	$C_{16}H_8O_3(OH)_4$ – Mw: 316 Mp: 228–229 °C Shade: red UV (EtOH) $\lambda_{max}$ 521, 486, 459, 279, 262, 219 nm.	(52)
Kermesic acid; 8-methyl-1,3,4,6-tetrahydroxy- anthraquinone-7-carboxylic acid (Ker)	HO CH <sub>3</sub> O OH OH OH	$C_{16}H_6O_4(OH)_4$ – Mw: 330 Mp > 320 °C Shade: Dark red in acidic pH; Violet in aqueous NaOH UV (MeOH) $\lambda_{max}$ 545, 496 nm.	(37, 44, 45)
Laccaic acid A (LaA)	HO O OH OH OH	C <sub>26</sub> H <sub>15</sub> NO <sub>8</sub> (OH) <sub>4</sub> – Mw: 537 Shade: red UV (H <sub>2</sub> SO <sub>4</sub> ) λ <sub>max</sub> 558, 518, 361, 302 nm.	(37, 44, 45)





Laccaic acid B (LaB)	HO O OH OH	$C_{24}H_{12}O_8(OH)_4 - Mw$ : 496 Shade: red	(37, 44, 45)
Laccaic acid C (LaC)	HO OH OH	$C_{25}H_{13}NO_9(OH)_4 - Mw$ : 539 Shade: dark red	(37, 44, 45)
Laccaic acid E (LaE)	HO O OH OH OH	$C_{24}H_{13}NO_{7}(OH)_{4} - Mw$ : 495 Shade: red UV (H <sub>2</sub> SO <sub>4</sub> ) $\lambda_{max}$ 496 nm.	(37, 44, 45)
<b>Tritisporin</b> 3-hydroxymethyl-1,4,6,8- tetrahydroxy-anthraquinone	HO OH CH <sub>2</sub> OH	C <sub>15</sub> H <sub>7</sub> O <sub>3</sub> (OH) <sub>3</sub> – Mw: 286 Shade: brownish-red	(4)

\*Mw: molecular weight; Mp: Melting point; Pr: primeverose; Rh: rhamnose

(yellow(acid) to red(alkali)), pseudopurpurin (orange), purpurin (dark red) and lucidin (red). A practical HAQN classification, according to the respective biosynthetic pathway of the compound and the position of the functional groups added on the 9,10-anthraquinone skeleton, is shown on Table 3. This classification is partially based on that proposed by Rafaëlly et al. in 2008<sup>45</sup>. It appears that the natural HAQN pigments formed via the chorismate/o-succinylbenzoic acid pathway are all classified in the entire 'group E' of HAQN dyes because they have substitution only on one aromatic ring, like alizarin and purpurin. In contrast, the HAQN pigments that are synthesized via the polyketide pathway are classified in the 'group A<sub>1</sub>' (compounds show substitution on both aromatic rings and at least two hydroxyl groups in both R1 and R8 positions, like emodin, chrysophanol or physcion) or into the 'group A2' of HAQN dyes (compounds show substitution on both aromatic rings and at least two hydroxyl groups in R1 and R6 positions and one methoxyl group in R8 position, like dermolutein and dermorubin).

### 3 The main natural sources of hydroxyanthraquinoid pigments

The natural HAQN pigments are mainly found in plants like *Rubiaceae*, *Polygonaceae*, *Rhamnaceae*, *Fabaceae*, *Liliaceae*, *Bignoniaceae* and *Pedaliaceae*, in lichens and in the animal kingdom (insects).

#### 3.1 Hydroxyanthraquinoid pigments from plants

In plants, the dyestuff is often extracted from dried roots. HAQN pigments are mostly present as sugar derivatives—the glycosides—but the free form—the aglycones—are widely distributed as well. For example, the European madder roots contain 2%–3.5% of the dry weight of di- and tri-

hydroxyanthraquinone-glycosides and, in general for higher plants, the HAQN-based colorant content from the dry mass is often under 5%<sup>15</sup>. The anthraquinone glycosides are formed when one or more sugar molecules, mostly glucose or rhamnose, are bound to the aglycone by a  $\beta$ -glycoside linkage to hydroxyl group at position C-8 (in the case of glucose) or the one at C-6 (in the case of rhamnose)<sup>58</sup>. During storage, hydrolysis of the glycosides occurs, which is completed under acidic conditions. In the literature, a total of more than 35 anthraquinoid compounds have been reported to be extracted from roots of European madder (Rubia tinctorum Linn., i.e. the most important species of the plant family Rubiaceae), even if a part of the compounds is believed to be artefacts formed during extraction or drying<sup>15</sup>. The main HAQN colouring compounds of plants of the Rubiaceae family (e.g. Rubia spp., Galium spp., Morinda spp., Hypericum spp., Polygonum spp. and Cinchona spp.) 14,15,23,34,38-40,45,59-73 are alizarin (yellow to red, group E<sub>3</sub>), pseudopurpurin (orange, group E<sub>2</sub>), purpurin (dark red, group E2), lucidin-3-O-primeveroside (red, group acid (golden-yellow, group E<sub>3</sub>), ruberythric  $E_2$ ). nordamnacanthal (orange, group E<sub>3</sub>) and munjistin (orange-red, group E<sub>3</sub>) (see Table 4). These colouring compounds are all classified in the entire 'group E' of HAQN dyes and they are formed through the chorismate/o-succinylbenzoic acid pathway as mentioned above.

In contrast, in other higher plants such as the *Polygonaceae* (*Rheum spp.*, *Rumex spp.*)<sup>24,29,31,43,74-80</sup>, *Rhamnaceae* (*Rhamnus spp.*)<sup>25,81</sup>, *Fabaceae* (*Cassia spp.*)<sup>30,33,82</sup>, *Liliaceae* (*Aloes spp.*)<sup>83,84</sup> and *Pedaliaceae* (*Ceratotheca spp.*)<sup>85</sup> families, the most common naturally occurring HAQN pigments are synthesized *via* the polyketide pathway (Fig. 2). Relevant pigments are emodin (yellow), aloe-emodin (yellow), physcion (yellow), rhein (orange) and chrysophanol (orange-red) (see





Table 4). These HAQN pigments are all classified in the 'group  $A_1$ ' of HAQN because they show substitution on both

aromatic rings and at least two hydroxyl groups in both R1 and R8 positions.

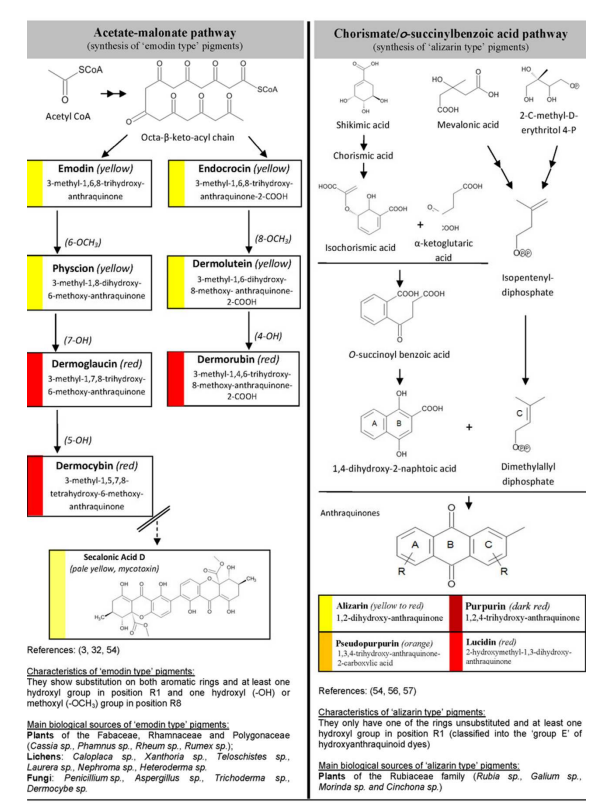


Figure 2. The two main biosynthetic pathways of hydroxyanthraquinoid (HAQN) pigments in organisms





Table~3.~Position~of~functional~groups~added~on~9, 10-anthraquinone~skeleton~in~natural~hydroxyanthraquinoid~(HAQN)~pigments~and~their~classification~into~several~groups

HAQN compound	R1	R2	R3	R4	R5	R6	R7	R8	shade
Group A1: at least two hydroxyl groups	in both R1 a	ınd R8 positi	ons:						
Aloe-emodin	OH	Н	$CH_2OH$	H	H	Н	H	OH	orange
Averythrin	OH	$C_6H_{11}H$	OH	H	H	OH	Н	OH	orange
Chrysophanol	OH	H	$CH_3$	H	H	H	H	ОН	orange-red
Citreorosein	OH	Н	OH	H	H	CH <sub>2</sub> OH	H	OH	yellow
Cynodontin	OH	H	CH <sub>3</sub>	OH	OH	Н	H	OH	bronze
Danthron (= chrysarin)	OH	H	H	H	H	H	H	OH	reddish to orange
Dermoglaucin	OH	H	CH <sub>3</sub>	H	Н	OCH <sub>3</sub>	OH	OH	red
Dermocybin Emadin	OH	H	CH <sub>3</sub>	H	OH	OCH <sub>3</sub>	OH	OH	red
Emodin Endocrocin	OH OH	H COOH	CH <sub>3</sub>	H H	H H	OH OH	H H	OH OH	orange
Erythroglaucin	OH OH	Н	CH <sub>3</sub>	н ОН	н Н		н Н	OH OH	yellow red
Fallacinal	ОH	Н	CH <sub>3</sub> CHO	И Н	п Н	OCH <sub>3</sub> OCH <sub>3</sub>	п Н	ОH	rea yellow
Frangulin A	OH	Н	O-Rh	Н	H	CH <sub>3</sub>	H	OH	•
Helminthosporin	OH	Н	CH <sub>3</sub>	H	OH	H	H	OH	orange maroon
Physcion (= parietin)	OH	Н	CH <sub>3</sub>	H	Н	OCH <sub>3</sub>	Н	OH	yellow
Rhein	OH	Н	COOH	H	H	Н	H	OH	orange
Teloschistin (= fallacinol)	OH	Н	CH <sub>2</sub> OH	Н	Н	OCH <sub>3</sub>	Н	OH	yellow
Tritisporin	OH	Н	CH <sub>2</sub> OH	OH	Н	OH OH	Н	OH	reddish brown
1			=				11	on	readish brown
Group A2: at least two hydroxyl groups							**	OCH	
Questin Dermolutein	OH	H	CH <sub>3</sub>	H	H	OH	H	OCH <sub>3</sub>	yellow to orange
	OH	COOH	CH <sub>3</sub>	Н	H	OH	H	OCH <sub>3</sub>	yellow
Dermorubin	ОН	COOH	$CH_3$	OH	Н	ОН	Н	OCH <sub>3</sub>	red
Group B: Four hydroxyl groups in R1, I	R3, R4 and I	R6 positions,		carboxyl g	roup in R7	position:			
Kermesic acid	ОН	H	OH	OH	H	ОН	COOH	$CH_3$	red
Carminic acid	ОН	Glc	OH	ОН	Н	ОН	СООН	$CH_3$	red
Laccaic acid A	ОН	$C_{10}H_{12}O_2N$		OH	H	ОН	СООН	COOH	red
Laccaic acid B	OH	$C_8H_9O_2$	ОН	OH	H	ОН	СООН	COOH	red
Laccaic acid C	OH	$C_9H_{10}O_3N$	ОН	OH	H	ОН	СООН	COOH	red
Laccaic acid E	ОН	$C_8H_{10}ON$	ОН	OH	Н	OH	СООН	COOH	red
Group C1: at least two hydroxyl groups	in R1 and R	4 positions, a	and at lea	st one fun	ctional gro	ир (-ОН, -С	H <sub>3</sub> ) in R7 p	osition:	
Austrocortinin	ОН	$OCH_3$	Н	OH	H	Н	$CH_3$	H	red
Catenarin	ОН	H	$CH_3$	OH	OH	H	OH	H	red
Rubrocristin	ОН	$CH_3$	Н	ОН	$OCH_3$	H	OH	H	red
Group C2: at least two hydroxyl groups	in R1 and R	6 positions, a	and positi	on R8 uns	substituted:				
Flavopurpurin	ОН	OH	н	Н	H	ОН	Н	H	yellow
Phomarin	ОН	Н	$CH_3$	Н	H	ОН	Н	H	yellow
Soranjidiol	ОН	$CH_3$	Н	Н	H	ОН	Н	H	yellowish-red
Group C3: at least two hydroxyl groups	in R1 and R	5 nositions	and nociti	on RR uns	uhetitutad.				
Islandicin	OH	CH <sub>3</sub>	mu posiii H	OH NO UNS	онозиниен. ОН	Н	Н	Н	red
Morindone	OH	OH	Н	Н	ОН	CH <sub>3</sub>	Н	Н	vellowish-red
									yellowish rea
Group D: no hydroxyl group in R1 posit									
Anthraflavic acid (= anthraflavin)	Н	OH	H	H	Н	OH	H	H	yellow
Skyrin	$C_{15}H_9O_5$	ОН	Н	OH	ОН	Н	CH <sub>3</sub>	Н	yellow to red
Compounds with functional groups only	on one arom	atic ring (gro	оир Е):						
Curren Els no hudrocon internationales	Landa midh	and and Go							
Group E1: no hydrogen intramolecular 3-MeO-hystazarin	bonas with	c <b>arbonyi jun</b> OH	Ction: OCH <sub>3</sub>	Н	Н	Н	Н	Н	
Damnacanthal	OCH <sub>3</sub>	СНО	OCn <sub>3</sub> OH	п Н	п Н	Н	п Н	п Н	nala vallav
		CHO	OH	11	11	11	11	11	pale yellow
Group E2: at least 2 hydroxyl groups in	-								
	R1 and R4								
Purpurin	R1 and R4 OH	OH	Н	ОН	Н	Н	Н	Н	dark red
Purpurin Pseudopurpurin	R1 and R4 OH OH	OH COOH	OH	OH	H	Н	H	Н	orange
Purpurin	R1 and R4 OH	OH							
Purpurin Pseudopurpurin	R1 and R4 OH OH OH	OH COOH H	OH H	OH OH	H H	Н	H	Н	orange
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a Alizarin	R1 and R4 OH OH OH	OH COOH H	OH H	OH OH	H H	Н	H	Н	orange
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a Alizarin Anthragallol	R1 and R4 OH OH OH Iways one in	OH COOH H position R1	OH H and none	OH OH e in positio	Н Н <b>on R4:</b>	H H	H H	H H	orange orange-red
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a Alizarin	R1 and R4 OH OH OH lways one in	OH COOH H position R1	OH H <b>and none</b> H	OH OH e in positio H	Н Н <b>en R4:</b> Н	Н Н	H H	Н Н	orange orange-red yellow to red
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a Alizarin Anthragallol	R1 and R4 OH OH OH lways one in OH	OH COOH H position R1 OH OH	OH H <b>and none</b> H OH	OH OH e in positio H H	Н Н <b>on R4:</b> Н Н	н н н н	H H H	Н Н Н	orange orange-red yellow to red orange
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a. Alizarin Anthragallol Lucidin-3-O-primeveroside Lucidin (= henine) Munjistin	R1 and R4 OH OH OH Ways one in OH OH	OH COOH H position R1 OH OH CH <sub>2</sub> OH	OH H and none H OH O-Pr	OH OH e in positio H H H	Н Н <b>on R4:</b> Н Н Н	Н Н Н Н	H H H H	Н Н Н Н	orange orange-red yellow to red orange red
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a Alizarin Anthragallol Lucidin-3-O-primeveroside Lucidin (= henine)	R1 and R4 OH OH OH Ways one in OH OH OH	OH COOH H position R1 OH OH CH <sub>2</sub> OH CH <sub>2</sub> OH	OH H and none H OH O-Pr OH	OH OH e in positio H H H H	Н Н <b>on R4:</b> Н Н Н	Н Н Н Н Н	Н Н Н Н Н	Н Н Н Н Н	orange orange-red  yellow to red orange red red
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a. Alizarin Anthragallol Lucidin-3-O-primeveroside Lucidin (= henine) Munjistin Nordamnacanthal Pachybasin	RI and R4 OH	OH COOH H position R1 OH OH CH <sub>2</sub> OH CH <sub>2</sub> OH COOH	OH H and none H OH O-Pr OH OH	OH OH e in positio H H H H	Н Н <i>on R4:</i> Н Н Н Н	Н Н Н Н Н Н	Н Н Н Н Н Н	Н Н Н Н Н Н	orange orange-red  yellow to red orange red red orange-red
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a. Alizarin Anthragallol Lucidin-3-O-primeveroside Lucidin (= henine) Munjistin Nordamnacanthal Pachybasin Ruberythric acid	RI and R4 OH	OH COOH H position R1 OH OH CH2OH CH2OH COOH	OH H and none H OH O-Pr OH OH OH	OH OH e in positio H H H H H	Н Н Н <b>R4:</b> Н Н Н Н Н Н	Н Н Н Н Н Н Н	н н н н н н	н н н н н н	orange orange-red  yellow to red orange red orange-red orange-red orange-yellow
Purpurin Pseudopurpurin Quinizarin  Group E3: 2 or 3 hydroxyl groups but a. Alizarin Anthragallol Lucidin-3-O-primeveroside Lucidin (= henine) Munjistin Nordamnacanthal Pachybasin	RI and R4 OH	OH COOH H  position R1 OH OH CH2OH CH2OH COOH CHO CH3 O-Pr CH3	OH H and none H OH O-Pr OH OH OH OH	OH OH e in positio H H H H H H	Н Н ОП R4: Н Н Н Н Н Н	H H H H H H H	н н н н н н н	н н н н н н н	orange orange-red  yellow to red orange red red orange-red orange-yellow yellow

Pr: primeverose; Glc: glucose; Rh: rhamnose





Table 4. Natural occ	urrence of hydroxyanthraquinoid (HAQN) pigments in plants	
Latin name	Main colouring components	References
Rubiaceae		
Rubia tinctorum L. (= European madder)	alizarin; purpurin; pseudopurpurin; lucidin; rubiadin; xanthopurpurin; munjinstin; anthraflavin; quinizarin; danthron; anthragallol; nordamnacanthal; ruberythric acid; lucidin primeveroside, alizarin-2-methyl ether; lucidin-ω -ethyl-ether; munjistin ethyl ether	(23, 34, 38, 59)
Rubia cordifolia L.	rubiadin; alizarin; purpurin; pseudopurpurin; lucidin; munjinstin; xanthopurpurin; tectoquinone	(14, 15, 23, 60)
Rubia akane	purpurin; ruberythric acid	(15, 45)
Rubia peregrina L.	pseudopurpurin	(15)
Galium aparine L.	nordamnacanthal; xanthopurpurin; rubiadin	(61)
Galium sinaicum	7-methyl-anthragallol-1,3-dimethyl ether; 7-methyl-anthragallol-2-methyl ether; 6-methyl-anthragallol-3-methyl ether; 8-hydroxy-anthragallol-2,3-dimethyl ether; 7-formyl-anthragallol-1,3-dimethylether; 6-hydroxy-xanthopurpurin; 6-methoxy-lucidin- $\omega$ -ethyl ether; copareolatin; copareolatin-6,7-dimethyl ether; copareolatin-5,7-dimethyl ether	(62)
Galium verum L. (= Lady's bedstraw)	alizarin; 1,3-dihydroxy-2-methoxymethyl; 1,3-dimethoxy-2-hydroxy,1,3-dihydroxy-2-acetoxy; 1-hydroxy-2-hydroxymethyl; 1,3-dihydroxy-2-methyl; 1-methoxy-2-hydroxyanthraquinones; 1,3-dihydroxy-2-hydroxymethyl-6-methoxy anthraquinones	(15, 63)
Galium mollugo L.	pseudopurpurin	(15)
Galium spurium	8-hydroxy-3-methoxy-7-methyl-1,2-methylenedioxy-anthraquinone;	(64)
	2,8-dihydroxy-1,3-dimethoxy-7-methyl-anthraquinone	
Morinda officinalis	alizarin; purpurin; pseudopurpurin; lucidin; rubiadin; 2-hydroxy-1-methoxy-anthraquinone; 1,3,8-trihydroxy-2-methyl-anthraquinone	(40)
Morinda elliptica	alizarin; purpurin; pseudopurpurin; lucidin; rubiadin; moridone; soranjidol; nordamnacanthal; alizarin-1-methylether; lucidin-ω-methylether	(39)
Morinda citrifolia	damnacanthal; morindone; morindin; alizarin; physcion; morenone; morenone; ruberythric acid; rubiadin; lucidin	(65)
Cinchona ledgeriana	purpurin; rubiadin; anthragallol-1,2-dimethylether; anthragallol-1,3-dimethylether; 1-hydroxy-2-hydroxymethylanthraquinone; 1-hydroxy-2-methylanthraquinone; morindone-5-methylether (or 1,7-dihydroxy-8-methoxy-2-methylanthraquinone)	(66, 67)
Cinchona pubescens	purpurin; alizarin-2-methylether; anthragallol-1,2-dimethylether; purpurin-1-methylether; 1-hydroxy-2-hydroxymethyl-anthraquinone; 2-hydroxy-1,3,4-trimethoxy-anthraquinone	(68)
Cinchona succirubra	emodin; anthrapurpurin; quinizarin; 2,6-dihydroxyanthraquinone; 1,8-dihydroxyanthraquinone	(69)
Cinchona robusta	robustaquinones (A-H); 1,3,8-trihydroxy-2-methyl anthraquinone; copareolatin 6-methylether	(70)
Asperula tinctoria L.	alizarin; rubiadin	(15, 45)
Asperula arvensis L.	alizarin	(45)
Oldenlandia umbellata L.	alizarin;1,2,3-trimethoxyanthraquinone; 3-MeO-hystazarin, ruberythric acid; 1,3-dimethoxy-2-hydroxyanthraquinone; 1,2-dimethoxyanthraquinone; 1-methoxy-2-hydroxyanthraquinone; 1,2-dihydroxyanthraquinone	(45, 71, 72)
Hedyotis auricularia L	alizarin	(45)
Crucianella maritima L.	alizarin; 3-formyl-1-hydroxy-2-methoxy anthraquinone; alizarin-1-methyl ether; 1,4-dihydroxy-2-methoxy-anthraquinone	(45, 73)
Coprosma lucida	anthragallol; lucidin; rubiadin	(45)
Hymenodictyon excelsum	anthragallol	(45)
Polygonaceae		
Rheum officinale	emodin; chrysophanol; rhein	(43, 74)
Rheum palmatum	chrysophanol; aloe-emodin; rhein; physcion; citreorosein	(24, 75)
Rheum emodi	emodin; chrysophanol; aloe-emodin; rhein; physcion	(29, 76)
Rheum rhabarbarum	emodin; chrysophanol; aloe-emodin; rhein; physcion	(31)
Rumex dentatus	chrysophanol; physcion	(24)
Rumex crispus	chrysophanol; parietin	(77, 78)
Rumex acetosa	chrysophanol; physcion; emodin; emodin-8- $O$ - $\beta$ -D-glucopyranoside	(79)
Rumex obtusifolius	aloe-emodin; chrysophanol; emodin	(80)
Rumex spp. (19 spp.)	emodin; chrysophanol; physcion; aloe-emodin; rhein	(80)
Rhamnaceae		
Rhamnus saxatilis	emodin; chrysophanol; aloe-emodin; rhein; physcion	(25)
Rhamnus alpinus L.	aloe-emodin; rhein; emodin; chrysophanol; physcion	(81)
Fabaceae		
Cassia occidentalis L.	emodin, chrysophanol, aloe-emodin, rhein, physcion	(30)
Cassia tora	emodin; rhein; physcion	(33)
Senna alata	aloe-emodin; emodin; rhein; chrysophanol	(82)
Liliaceae		
Aloes spp. (32 spp.)	chrysophanol; asphodelin; chrysophanol-8-methyl ether; aloechrysone; helminthosporin; aloesaponols; aloesaponarins	(83, 84)
Bignoniaceae		
Tecoma ipes	tectoquinone	(45)
Pedaliaceae		
Ceratotheca triloba (Bernh.	1) 1-hydroxy-4-methylanthraquinone	(85)





#### 3.2 Hydroxyanthraquinoid pigments from lichens

HAQN pigments found in some lichens are synthesized *via* the polyketide pathway. For example, the main colouring compounds in the lichens of the family *Teloschistaceae* (*Caloplaca sp., Xanthoria* sp. or *Teloschistes sp.*)<sup>15,48,86–90</sup> and the family *Trypetheliaceae* (*Laurera benguelensis*)<sup>87,88</sup> are emodin, physcion, teloschistin (yellow, group A<sub>1</sub>) and fallacinal (yellow, group A<sub>1</sub>) (see Table 5). The lichens *Nephroma laevigatum* and *Heteroderma obscurata* also contain emodin<sup>91</sup> whereas skyrin (*i.e.* a yellow to red pigment classified in the 'group D' of HAQN which show substitution on both aromatic rings; see Table 3) is the main component of *Cladonia* species<sup>48</sup>.

#### 3.3 Hydroxyanthraquinoid pigments from insects

In animals, HAQN-type pigments are known to be present only in a few insect species (see Table 5). Concerning the red carminic acid, kermesic acid and laccaic acid obtained from cochineal (*Dactylopius coccus*)<sup>15,22,37,50</sup> kermes (*Kermes vermilio*)<sup>15,44</sup> and lac (*Kerria lacca*)<sup>15,44,45</sup>, respectively, they contain functional groups on both aromatic rings and particularly four hydroxyl groups in R1, R3, R4 and R6 positions, and one carboxyl group in R7 position. So these animal anthraquinoid glycosides are all classified in the 'group B' of HAQN (see Table 3). In both cochineal and kermes the pigments were obtained from the body and eggs of the female insect. Although the various species of the genus Porphyrophora, e.g. Armenian cochineal (P. hamelli) and Polish cochineal (P. polonica), also contain carminic acid, dried specimens of Dactylopius coccus have a much higher content (15%-20%) of carminic acid, compared with only 0.8% and 0.6% for the Armenian and Polish ones, respectively<sup>15,44,45</sup>. Lac insects of the *Kerria* family (e.g.

Kerria lacca and K. chinensis) contain mainly laccaic acids like laccaic acid A.

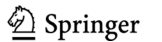
#### 3.4 Hydroxyanthraquinoid pigments from fungi

HAQN pigments are widespread in nature and have been also found abundantly in microorganisms, particularly in filamentous fungi belonging to Penicillium spp. and Aspergillus spp., with different shades (see Table 6). For example, the pigment emodin was isolated from strains of Penicillium citrinum and P. islandicum<sup>5,92</sup>. The natural food colorant Arpink red<sup>TM</sup> manufactured by the Ascolor Biotech Czech company was claimed to be produced by fermentation and bioprocess engineering using the strain Penicillium oxalicum var. Armeniaca CCM 8242 obtained from soil<sup>5,92</sup>. On the second day of cultivation of this fungus in liquid broth containing carbohydrates, zinc sulfate and magnesium sulfate, a red colorant is released in the medium, increasing up to 1.5-2.0 g/L of broth after 3–4 days<sup>2</sup>. After biosynthesis of the red colorant, the liquid is separated from the biomass by centrifugation or filtration. The liquid is then acidified to pH 3.0-2.5 to precipitate the colorant. The precipitate is dissolved in ethyl alcohol and filtered. Following removal of alcohol, the colorant is obtained in the crystalline form as a dark red powder. In strains of *Penicillium purpurogenum*<sup>93</sup>, a red pigment of HAQN-type (none completely characterized) was also observed

Some strains of Aspergillus (A. glaucus, A. cristatus and A. repens)<sup>4,5,35,36,92–94</sup> were found to produce known yellow and red HAQN compounds such as emodin (yellow, group  $A_1$ ), physcion (yellow, group  $A_1$ ), questin (yellow to orange-brown, group  $A_2$ ), erythroglaucin (red, group  $A_1$ ), catenarin (red, group  $C_1$ ; see Table 3) and rubrocristin (red, group  $C_1$ ; see Table 3). However, by using Penicillium or Aspergillus strains,

Table 5. Natural occurrence of hydroxyanthraquinoid (HAQN) pigments in lichens and in insects

Dye source	Latin name	Main colouring components	References
Lichens:			
Teloschistaceae	Xanthoria spp.	physcion; emodin; parietin; fallacinal; teloschistin; citreorosein; erythroglaucin; fallacinol	(15, 48, 86, 87)
	Xanthoria parietina L.	physcion; fallacinol; fallacinal; emodin; parietic acid	(88)
	Xanthoria mandschurica	erythroglaucin; parietin	(89)
	Xanthoria fallax	fallacinal; emodin; fallacinol; erythroglaucin; parietin	(89)
	Caloplaca spp.	2-chloroemodin; citreorosein; emodin; fallacinal; parietin; physcion; teloschistin	(48, 87)
	Caloplaca cerina	emodin; fallacinal; physcion; teloschistin	(33)
	Caloplaca erythrantha	emodin; 7-chloroemodin	(86)
	Teloschistes exilis	parietin; teloschistin	(86)
	Teloschistes spp.(29 spp.)	parietin; emodin; teloschistin; fallacinal; parietic acid;erythroglaucin	(90)
Trypetheliaceae	Laurera benguelensis	parietin; physcion; citreorosein; emodin; fallacinal; teloschistin	(87, 88)
Nephromataceae	Nephroma laevigatum	emodin; 7-chloroemodin; 7-chloro-1- <i>O</i> -methylemodin;5-chloro-ω-hydroxyemodin 7-chloro-1- <i>O</i> -methyl-ω-hydroxyemodin; 5-chloroemodin;5-chloro-1- <i>O</i> -methylemodin; 5-chloro-1- <i>O</i> -methyl-ω-hydroxyemodin;	(91)
Physciaceae	Heteroderma obscurata	emodin	(91)
Cladoniaceae	Cladonia spp.	skyrin	(48)
Insects:			
Dactylopius	Dactylopius coccus Costa (cochineal)	carminic acid [food additive E120(ii)]	(15, 22, 37, 50)
Porphyrophora	Porphyrophora hameli B. & Porphyrophora polonica L.	carminic acid; flavokermesic acid (LaE); kermesic acid	(15, 44, 45)
Kermes	Kermes vermilio Planchon (=Kermococcus vermilio)	kermesic acid	(15, 44)
Kerria	Kerria lacca Kerr	laccaic acids (A, B, C, E)	(15, 44, 45)





several known mycotoxins were coproduced in the medium. e.g. secalonic acid D, oxaline, citrinin, tanzawaic acid A, cyclochlorotine, islanditoxin, luteoskyrin, erythroskyrin, rugulosin or aspergiolide A (Table 6). Many of these mycotoxins are pigmented, that is, naphtoquinones from Aspergillus and Penicillium. All these fungal secondary metabolites (on one hand, the yellow and red HAQN pigments that show substitution on both aromatic rings and, on the other hand, the naphtoquinone-type mycotoxins) arise biosynthetically by the same polyketide pathway. The cytotoxic activity of naphtoquinones, and of mycotoxins in general, against mouse leukemia and HeLa cells has been mainly reported in the literature. Moreover, along with the antibiotic and toxic activities, naphtoquinones revealed mutagenic and carcinogenic properties. The results suggested that these fungal strains could not be used to provide safe fungal hydroxyanthraquinoid pigments as potent natural food grade colorants

Species of *Eurotium* spp. (*E. amstelodami, E. chevalieri* and *E. herbariorum*)<sup>5,92</sup> were found to produce the yellow pigment physcion and the red pigment erythroglaucin (group A<sub>1</sub>), however they produce in addition the mycotoxin echinulin and two benzaldehyde coulouring compounds: flavoglaucin (yellow) and auroglaucin (red) (see Table 6). In the same way, it has been demonstrated that a coproduction of red hydroxyanthraquinoid pigments (with no hydroxyl substituents at the positions R1 and R4) and mycotoxins such as fusaric acid, nectriafurone, monoliformin and gibepyrones, occurs by using strains of *Fusarium oxysporum* isolated from roots of diseased citrus trees <sup>95–97</sup> (Table 6).

Apart from those mycotoxigenic fungi, there are other filamentous fungi that have the ability to produce known HAQN pigments that arise biosynthetically by the polyketide pathway more particularly, without coproduction of mycotoxins. A strain of *Dermocybe sanguinea* (= *Cortinarius sanguineus*) has been identified as producing the red HAQN glycoside dermocybin-1- $\beta$ -D-glycopyranoside giving the typical red colour of the fruiting bodies and the spores, in addition with both emodin and physcion pigments<sup>3,15,18,52</sup>. In the fresh fungi as much as 90% of the pigments exist as glycosides. The detection of emodin-glycosides and physcion-glycosides was also pointed from a strain of *Dermocytes spp*. 98

Strains of Trichoderma aureoviride<sup>99</sup> and T. harzianum<sup>29,100</sup> were found to produce yellow pigment pachybasin (group E<sub>3</sub>) and also the orange-red pigment chrysophanol (group  $A_1$ ). Both species of *Trichoderma polysporum* and *T. viride*  $^{99,101,102}$ can also produce pachybasin in addition to the emodin and chrysophanol pigments. Several HAQN-type pigments have been isolated from cultures of *Curvularia lunata*<sup>3-5</sup>. The main pigments characterized were erythroglaucin (red, group A<sub>1</sub>), catenarin (red, group C<sub>1</sub>), chrysophanol (orange-red, group A<sub>1</sub>), helminthosporin (maroon, group A<sub>1</sub>) and cynodontin (bronze, group  $A_1$ ). Cynodontin extracted from the biomass of C. lunata has been converted successfully to two anthraquinone biodyes (Disperse blue 7 and Acid Green 28). The properties of these biodyes applied to knitted polyamides were compared with those of conventional dyes and found to be identical to all-important aspects<sup>3</sup>. Several species of *Drechslera* (e.g. D. teres, D. graminea, D. tritici-repentis, D. phlei, D. dictyoides and D. avenae) give HAQN pigments like catenarin (red, group C<sub>1</sub>), helminthosporin (maroon, group A<sub>1</sub>), cynodontin

(bronze, group  $A_1$ ), tritisporin (reddish brown, group  $A_1$ ) and erythroglaucin (red, group  $A_1$ ), without coproduction of mycotoxins<sup>3,103</sup>. Other HAQN pigments like averythrin (orange, group  $A_1$ ) and averythrin-6-monomethyl ether were isolated and identified from a culture of *Herpotrichia rhodosticta* without coproduction of known mycotoxins<sup>104</sup>. More recently, a red pigment produced by a strain of *Isaria farinosa* was recently elucidated as a chromophore of the anthraquinone type<sup>105</sup>. Similarly, the red pigment produced by *Paecilomyces sinclairii*, which was beforehand discovered but uncharacterized<sup>106</sup>, is certainly of an identical chemical nature, *i.e.* an amino group linked to an anthraquinone structure<sup>105</sup>.

#### 4 Toxicity and carcinogenicity of some natural hydroxyanthraquinoid pigments

Anthraquinoid derivatives, including natural HAQN pigments, possess a broad spectrum of biological activities, including anti-inflammatory, anti-cancer, anti-viral, antifungal, anti-bacterial, astringent and purgative. In general, natural HAQN pigments and their intermediates have not been reported as strongly toxic substances, even if it is known that some anthraquinoid dyes are toxic or mutagenic 107,108. Due to its use as a food colorant in Japan, the safety of European madder extracts has been studied in the literature. For example, in an extensive study the European madder roots were extracted using different solvents and extracts were fractionated by chromatography. Several colour components extracted from madder roots were positive to mutagenicity tests as the yellow rubiadin pigment (group E<sub>3</sub>; see Table 3) and the red lucidin pigment (group E<sub>3</sub>) aglycones, which are metabolites of lucidin-3-O-primeveroside. From structure mutagenicity studies it was concluded that 1,3-dihydroxyanthraguinones that bear a methyl (-CH<sub>3</sub>) or hydroxymethyl (-CH<sub>2</sub>OH) group in position R2, e.g. rubiadin or lucidin, respectively, are mutagenic. For direct mutagenicity an oxygenated state of the benzylic carbon-2 is required. Mutagenic studies about lucidin more particularly showed that a reactive compound is formed from the metabolism of the pigment, which then reacts with DNA and possibly other macromolecules to form covalent adducts 109,110. Other 1,3dihydroxyanthraquinones that do not possess a methyl or hydroxymethyl group in position R2, such as the orange pigment nordamnacanthal (group E<sub>3</sub>) and the orange-red munjistin pigment (group E<sub>3</sub>), are not found to be mutagenic, since the dehydration to the exomethylenic compound is not possible under physiological conditions<sup>15</sup>. In a 13-week repeated oral dose toxicity study of madder colour, which was performed using F344 rats, the animals were fed a diet containing 0, 0.6, 1.2, 2.5 or 5.0% of colouring compounds extracted from madder roots. The results suggested that madder colour exerts mild toxicity, targeting liver, kidneys and possibly red blood cells and white blood cells, some renal changes being evident from 0.6% madder colour in diet. This is considered to be the lowest-observed adverse effect level (305.8-309.2 mg/kg of body weight per day)<sup>111</sup>. Data are in agreement with another study performed in the same year in a medium-term multiorgan carcinogenesis bioassay in male F344 rats, which reported that madder colour demonstrated significant tumour-promoting effects in the liver and kidneys<sup>112</sup>. More recently, an additional two-year carcinogenicity study conducted on male and female F344





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Table 6. Natural occurrence of hydroxyanthraquinoid (HAQN) pigments in microorganisms

Paceins   HAVA Negiments composited stander   Pacein texts   HAVA Negiments   Contracts   HAVA Negiments   Contracts   HAVA Negiments   Contracts   Pacein texts   Pacein	(					ş
Potent taxic HAQN pigments   Potent taxic HAQN pigments   Potent taxic HAQN pigments   Conditions	Cenus	Species	HAON colouring compounds (shade)		Othertoxic compounds	Keterences
P. consistone         Arpink red <sup>31</sup> (red)         Arpink red <sup>31</sup> (red)         Sectionic soid D and coaline           P. consistone         Skyrin (vellow to red)         emodin (yellow)         cerchonic soid D and coaline           P. bistonicum         skyrin (vellow to red)         emodin (yellow)         cyclebalooute, sind and an and and and and an an and an an an and an			Non-toxic HAQN pigments	Potent toxic HAQN pigments	(mycotoxins, color or colorless)	
P. clintum         - cutinum         - cutinum         cutinum         cutinum           P. clintum         P. clintum         4. clintum         cyclochtronic scal A         cyclochtronic scal A           P. clintum         crintum         cythroglaucit (cell, cythroglaucit (cell), cythroglauci	Penicillium	P. oxalicum	Arpink red <sup>TM</sup> (red)	1	secalonic acid D and oxaline	(5, 92)
P. Isturalecum         Skyrin (yellow to red.)         skyrin (yellow to red.)         Protechtorine, allanduron.           P. Instrument         explanaçam         explanaçamin (ved.) explanaçati (red.)		P. citrinum		emodin (yellow)	citrinin and tanzawaic acid A	(5, 92)
P purpuragemum         rod HAQN pignent (conc completely charactrized)         - emodin & physicion (yellow)         - emodin & physicion (yellow)           A. gámicas         everphenypatra (taraca), trideparti (edelás brown)         - emodin & physicion (yellow)         - emodin & physicion (yellow)           A. repers         crythreglaucin (red), rubbocristin (red), physicion (yellow)         - physicion (yellow)         - physicion (yellow)           A. repers         crythreglaucin (red), rubbocristin (red), symadoutin (red), physicion (yellow)         - physicion (yellow)         - physicion (yellow)           Beaving and category (red), remaining portin (unicon), trideparti (red), germentin (red), setheropeani (red), dermocribin (red), dermocribin (red), dermocribin (red)         - chapter (red), physicion (yellow)         - chapter (red), physicion (yellow)           B. suggiung         dermocribin (yellow)         - chapter (red), dermocribin (red), chryosphanol (red)         - chapter (red), remain (red), chryosphanol (red)         - chapter (red), chryosphanol (red)         - chapter (red)		P. islandicum	skyrin (yellow to red)	emodin (yellow)	cyclochlorotine, islanditoxin, luteoskyrin, erythroskyrin and rugulosin	(5, 92)
A glanecas         certaintediate (red., certainti (red.), certainti		P. purpurogenum	red HAQN pigment (none completely characterized)	1		(93)
A. crisianes         centerain (cell), evitroglaudia (red), erthroglaudia (red),	Aspergillus	A. glaucus	erythroglaucin (red), catenarin (red), cynodontin (bronze), helminthosporin (maroon), tritisporin (reddish brown)	emodin & physcion (yellow)	aspergiolide A	(4, 5, 92)
A. repens         erythrogaucin (red)         physicion (yellow)         -           Euronium, spp         catenaria (red), erythrogaucin (red), cynodoutin (bronzo)         physicion (yellow)         -           Fusanium spp         catenaria (red), erythrogaucin (reds) brown)         chrysophanol (red)         -           Fusanium spp         catenaria (red), erythrogaucin (reds) brown)         chrysophanol (red)         -           F. oxysporum         2-devely1-38-diby droxy-6-methoxy-anthraquianon (red)         -         -           D. surguinea         dermocybin-1-fb-Dytveopyranoside (red), dermorphin (red)         -         -           D. surguinea         dermocybin-1-fb-Dytveopyranoside (red), dermorphin (red)         -         -           P. candidam         pachybasin (yellow)         chronophanol (red)         -           P. candidam         pachybasin (yellow)         chrysophanol (red)         -           I. duraning         pachybasin (yellow)         chrysophanol (red)         -           I. hurizan         catenarin (red)		A. cristatus	catenarin (red), erythroglaucin (red), rubrocristin (red), questin (yellow)	emodin & physcion (yellow)		(36)
Enraction, spp         catenaria (red), evildodutii (bronze), tendodutii (bronze), tendodutii (bronze), tendorii, cyandodutii (cod), cyandodutiii (cod), cyandodutii (cod), cyandodutiii (cod), cyandodutii (cod), cyandodutiii (c		A. repens	erythroglaucin (red)	physcion (yellow)	1	(35, 94)
Fixarium sp. cateuntin (red.) evrythraglaucia (red.) evrotoantin (bronze), helminthosparia (red.) evrythraglaucia (red.) evrotoantin (red.) evrythraglaucia (red.) evrotoantin (red.) evrythraglaucia (red.) evrythraglaucia (red.) evrythraglaucia (red.) eractosy-anthraguinone (red.) emodin & physicion-glycosides  D. sanguinea dermocyhin-1-fb-D-glycopyranoside (red.) dermorubin (red.) emodin & physicion-glycosides  D. sanguinea dermocyhin-1-fb-D-glycopyranoside (red.) dermorubin (red.) emodin & physicion-glycosides  D. sanguinea pachybasia (yellow) pachybasia (yellow) pachybasia (yellow) pachybasia (yellow) pachybasia (yellow) pachybasia (yellow)  T. saneoriride pachybasia (yellow)  D. sanguinea cateuaria (red.) erythraglaucia (red.) emodin (yellow), chrysophanol (red.)  D. sanguinea cateuaria (red.) erythraglaucia (red.) emodin (yellow), chrysophanol (red.)  D. sanguinea cateuaria (red.)	Eurotium	Eurotium. spp.	catenarin (red), erythroglaucin (red), cynodontin (bronze), helminthosporin (maroon), tritisporin (reddish brown)	physcion (yellow)	echinulin	(5, 92)
F. oxysporum         2-(1-bydroxy-c-methory-anthraquinone (red)         modit of physicion-glycosides         fisaric acid, nectrialitrone, monoiliformin and glycpyroues           D. sp. WAT22963         dermocybin-1-d-D-glycopyr-anoside (red), dermorubin (red), dermoglaucin (red), 5-chlorodermorubin         amodin & physcion-glycosides         monoiliformin and glycpyroues           D. sanguinea         dermolutein (yellow), phomarin (yellow)         dermocybin-1-d-D-glycopyranoside (red), dermoglaucin (red), 5-chlorodermorubin         chrysophanol (red)         -           P. cargian         pachybasin (yellow)         pachybasin (yellow)         dermolutein (yellow)         dermolutein (yellow)           P. cargian         pachybasin (yellow)         pachybasin (yellow)         dermolutein (yellow)         dermolutein (yellow)           T. harzianum         pachybasin (yellow)         dermolutein (yellow)         dermolutein (yellow)         dermolutein (yellow)           T. harzianum         pachybasin (yellow)         dermolutein (yellow)         dermolutein (yellow)         dermolutein (yellow)           T. harzianum         pachybasin (yellow)         dermolutein (yellow)         dermolutein (yellow)         dermolutein (yellow)           T. harzianum         pachybasin (yellow)         dermolutein (yellow)         dermolutein (yellow)         dermolutein (yellow)           D. teres         catenarin (red)         dermolutein (yellow)	Fusarium	Fusarium spp.	catenarin (red), erythroglaucin (red), cynodontin (bronze), helminthosporin (maroon), tritisporin (reddish brown)	chrysophanol (red)		(5)
D. sanguinea dermocybin-1-f-D-glycopyranoside (red.) dermorubin (red.)  P. candidum pachybasin (yellow), dermoglaucin (red.) 5-chlorodermorubin (red.)  P. candidum pachybasin (yellow), phomarin (yellow) phomarin (yellow) phomarin (yellow) phomarin (yellow) pachybasin (yellow) yellow) pachybasin (yellow) pachybasin (yellow) pachybasin (yellow) pachybasin (yellow) pachybasin (yellow) pachybasin (yellow) y		F. oxysporum	2-(1-hydroxyethyl)-3,8-dihydroxy-6-methoxy-anthraquinone 2-acetyl-3,8-dihydroxy-6-methoxy-anthraquinone (red)		fusaric acid, nectriafurone, monoliformin and gibepyrones	(92, 95-97)
D. sanguinea dermocybin-1-f-D-glycopyranoside (red.), dermocrubin	Dermocytes	D. sp. WAT22963		emodin & physcion-glycosides	ı	(86)
P. candidum         pachybasin (yellow)         chrysophanol (red)         -           P. exigua         pachybasin (yellow), phomarin (yellow)         chrysophanol (red)         -           T. aureovirde         pachybasin (yellow)         chrysophanol (red)         -           T. harzianum         pachybasin (yellow)         chrysophanol (red)         -           T. viride         pachybasin (yellow)         chrysophanol (red)         -           T. viride         catenarin (red)         cynodontin (bronze)         chrysophanol (red)         -           D. leves         catenarin (red)         -         -         -           D. phei         catenarin (red)         -         -           D. phlei         catenarin (red)         -         -           D. dictyoides         catenarin (red)         -         -           D. dictyoides         cynodontin (bronze), averythrin (ora	Dermocybe	D. sanguinea	dermocybin-1-β-10-glycopyranoside (red.), dermorubin (red.), dermolutein (yellow), dermoglaucin (red.), 5-chlorodermorubin	emodin & physcion-glycosides		(3, 15, 18, 52)
P. exigua         pactybasin (yellow), phomarin (yellow)         emodin Gellow), chrysophanol (red)         -           T. aureoviride         pactybasin (yellow)         chrysophanol (red)         -           T. harzianum         pactybasin (yellow)         chrysophanol (red)         -           T. pobysporum         pactybasin (yellow)         13,6,8-tetraHAQN, 2,4,5,7-tetraHAQN         emodin (yellow), chrysophanol (red)         -           T. viride         catenarin (red), erythroglaucin (red), erynodontin (bronze), helminthosporin (red)         -         -         -           D. teres         catenarin (red)         -         -         -         -         -           D. tritici-repentis         catenarin (red)         -         -         -         -         -         -           D. phiei         catenarin (red)         -	Pachybasium		pachybasin (yellow)	chrysophanol (red)	•	(18)
T. aureoviride         pachybasin (yellow)         chrysophanol (red)         chrysophanol (red)           T. harzianum         pachybasin (yellow)         chrysophanol (red)         chrysophanol (red)           T. viride         pachybasin (yellow)         13.6,8-tetraHAQN         emodin (yellow), chrysophanol (red)         -           T. viride         catenarin (red)         erydrogatein (red) serythrogancin (red) shown)         chrysophanol (red)         -           D. teres         catenarin (red)         -         -         -           D. teres         catenarin (red)         -         -           D. tritici-repentis         catenarin (red)         -         -           D. phlei         catenarin (red)         -         -           D. phlei         -         -         -           D. dicyoides         catenarin (red)         -         -           D. avenae         cynodontin (bronze), helminthosporin (maroon)         -         -           D. avenae         cynodontin (red)         -         -           D. farinosa         red HAQN pigment (none completely characterized)         -         -           D. farinosa         -         -         -           D. farinosa         -         -         - </td <td>Phoma</td> <td>P. exigua</td> <td>pachybasin (yellow), phomarin (yellow)</td> <td>emodin (yellow), chrysophanol (red)</td> <td></td> <td>(18)</td>	Phoma	P. exigua	pachybasin (yellow), phomarin (yellow)	emodin (yellow), chrysophanol (red)		(18)
T. harzianum         pachybasin (yellow)         chrysophanol (red)         -           T. pobysporum         pachybasin (yellow)         chrysophanol (red)         -           T. viride         pachybasin (yellow)         1,3,6,8-tetraHAQN         2,4,5,7-tetraHAQN         -           C. lunata         pachybasin (yellow)         1,3,6,8-tetraHAQN         chrysophanol (red)         -           C. lunata         pendin thosporin (red), erythroganin (red), erythroganin (red), erythroganin (red)         -         -           D. teres         catenarin (red)         -         -         -           D. tritici-repentis         catenarin (red)         -         -           D. phlei         -         -         -           D. phlei         -         -         -           D. dicryoides         catenarin (red)         -         -           D. dicryoides         cynodontin (bronze), helminthosporin (maroon)         -         -           D. avenae         cynodontin (roange), averythrin-6-monomethyl ether         -         -           I. Jarinosa         red HAQN pigment (none completely characterized)         -         -           I. Jarinosa         austrocortinin (red)         -         -	Trichoderma		pachybasin (yellow)	chrysophanol (red)	ı	(66)
T. polysporum pachybasin (yellow)  T. viride pachybasin (yellow), 1,3,6,8-tetraHAQN, 2,4,5,7-tetraHAQN emodin (yellow), chrysophanol (red) - catenarin (red), erythroglaucin (red), erythroglaucin (red), erythroglaucin (red), cynodontin (bronze), chrysophanol (red) - catenarin (red) emodin (pellow), chrysophanol (red) - catenarin (red)		T. harzianum	pachybasin (yellow)	chrysophanol (red)	1	(29, 100)
T. viride pachybasin (yellow), 1,3,6,8-tetraHAQN, 2,4,5,7-tetraHAQN emodin (yellow), chrysophanol (red) - catenarin (red), erythroglaucin (red), erythroglaucin (red), erythroglaucin (red), erythroglaucin (red) catenarin (red) -		T. polysporum	pachybasin (yellow)	emodin (yellow), chrysophanol (red)	ı	(99, 101)
C. Iumata         catenarin (red), erythroglaucin (red), cynodontin (bronze), helminthosporin (maroon), tritisporin (reddish brown)         chrysophanol (red)         -           D. teres         catenarin (red)         -         -         -           D. tritici-repentis         catenarin (red)         -         -         -           D. phlei         -         -         -         -           D. aletyoides         catenarin (red)         -         -         -           D. aletyoides         catenarin (red)         -         -         -           D. avenae         cynodontin (bronze), helminthosporin (maroon)         -         -         -           H. rhodosticta         averythrin (orange), averythrin (orange), averythrin (orange), averythrin (orange), averythrin (orange), averythrin (cange), averythrin (cange)         -         -           I. farinosa         red HAQN pigment (none completely characterized)         -         -         -		T. viride	pachybasin (yellow), 1,3,6,8-tetraHAQN, 2,4,5,7-tetraHAQN	emodin (yellow), chrysophanol (red)	•	(99, 102)
D. teres         catenarin (red)         -         -           D. graminea,         catenarin (red)         -         -           D. tritici-repentis         catenarin (red)         -         -           D. phlei         -         -         -           D. dictyoides         catenarin (red)         -         -           D. avenae         cynodontin (bronze), helminthosporin (maroon)         -         -           H. rhodosticta         averythrin (orange), averythrin (orange), averythrin (orange), averythrin (orange), averythrin (orange), averythrin (orange), averythrin (crange)         -         -           I. farinosa         red HAQN pigment (none completely characterized)         -         -           Austrocortinin (red)         -         -	Curvularia	C. lunata	catenarin (red., erythroglaucin (red.), cynodontin (bronze), helminthosporin (maroon), tritisporin (reddish brown)	chrysophanol (red)	•	(3-5)
D. graminea,         catenarin (red)         -         -           D. phiei         -         -         -           D. phiei         -         -         -           D. dictyoides         catenarin (red)         -         -           D. avenae         cynodontin (bronze), helminthosporin (maroon)         -         -           H. rhodosticta         averythrin (orange), averythrin (orange), averythrin-6-monomethyl ether         -         -           I. farinosa         red HAQN pigment (none completely characterized)         -         -           austrocortinin (red)         -         -	Drechslera	D. teres	catenarin (red)	ı	ı	(103)
D. trittei-repentis catenarin (red)		D. graminea,	catenarin (red)	1		(103)
D. phlei catenarin (red)		D. tritici-repentis	catenarin (red)		•	(103)
D. dictyoides catenarin (red)		D. phlei	catenarin (red)	1		(103)
D. avenae cynodontin (bronze), helminthosporin (maroon)		D. dictyoides	catenarin (red)		•	(103)
H. rhodosticta averythrin (orange), averythrin-6-monomethyl ether - I. farinosa red HAQN pigment (none completely characterized) - austrocortinin (red) -		D. avenae	cynodontin (bronze), helminthosporin (maroon)			(103)
L. farinosa red HAQN pigment (none completely characterized) austrocortinin (red) -	Herpotrichia		averythrin (orange), averythrin-6-monomethyl ether		•	(104)
austrocortinin (red)	Isaria	I. farinosa	red HAQN pigment (none completely characterized)			(105)
	Fungi K_BK5	2	austrocortinin (red)		•	(27)



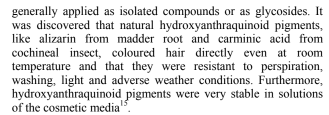
which were fed a diet containing 0, 2.5 or 5.0% of colouring compounds extracted from madder roots clearly indicate that this dyestuff—rich in alizarin, lucidin-3-*O*-primeveroside and ruberythric acid, all classified in the 'group E<sub>3</sub>' of HAQN and synthesized *via* the chorismate/*o*-succinylbenzoic acid pathway as mentioned above—exerts a carcinogenic potential in both the kidney and the liver, even with the lower dose of the study<sup>113</sup>. These studies support data in other previous studies<sup>114,115</sup> and provide clear evidence that madder colour exerts unequivocal carcinogenicity against renal tubule cells and hepatocytes in rats. Therefore, the authors<sup>113</sup> concluded that further studies on these individual HAQN components should be performed to clarify which anthraquinone is responsible for carcinogenicity.

Other recent studies indicate that the dark red purpurin pigment extracted from Indian madder (*Rubia cordifolia*)—classified in the 'group E<sub>3</sub>' of HAQN dyes and synthesized *via* the chorismate/o-succinylbenzoic acid pathway—has an antimutagenic effect on the Ames *Salmonella* bacterial mutagenicity assay. The antigenotoxic effect was observed in *Drosophila melanogaster* against a range of environmental carcinogens. Inhibition of the formation of hepatic DNA adducts in male C57bl6 mice after a single dose of the heterocyclic amine dietary carcinogen Trp-P-2 (30 mg/kg) was observed by short-term dietary supplementation with purpurin<sup>116</sup>. In another study, purpurin was found to show inhibition of mutagenicity of a number of heterocyclic amines in the Ames mutagenicity test. The inhibition effect of purpurin was dependent upon pH, being better in neutral than acidic conditions<sup>15</sup>.

Concerning anthraquinoid pigments synthesized via the polyketide pathway, the orange pigment aloe-emodin (group  $A_1$ ) induced micronucleus frequencies in the *in vitro* micronucleus test in mouse lymphoma L5178Y cells<sup>117</sup>. The emodin pigment (group A<sub>1</sub>) has toxic and gene mutagenic properties. The activation mechanism of emodin into a direct mutagen to Salmonella typhimurium TA1537 was investigated by using the S9 and microsomes of rat livers. Emodin exhibited mutagenicity in the presence of NADPH or NADH<sup>118</sup>. Another study mentioned that emodin was clearly genotoxic in mouse lymphoma cells<sup>119</sup>. Emodin of fungal origin has been classified as diarrheagenic and genotoxic mycotoxin<sup>58</sup>. Similarly, as fungal chrysophanol and physcion are hypothesized to exert genotoxicity, they are also considered as mycotoxins today Thus the detection of emodin, physcion and/or chrysophanol from some strains of Aspergillus spp., Penicillium spp., Eurotium spp., Dermocybe sanguinea, Dermocytes spp., Trichoderma spp. and Curvularia lunata (see Table 6) suggests that some of these fungi are potent mycotoxigenic.

#### 5 Current industrial applications of natural hydroxyanthraquinoid dyestuffs

Traditionally, relevant hydroxyanthraquinoid dyestuffs such as the famous kermes parasite insect, the cochineal insect and the European madder root are essentially used to dye textiles. They are also used for other non-food applications, e.g. printing, cosmetics (hair colorants...) and pharmaceutical applications across the globe. They provide the most important red pigments used in artistic paintings. Hydroxyanthraquinoid red dyes were among the reds that dominated the dye markets of Europe. Natural hydroxyanthraquinoid pigments were



Colour compounds extracted from the roots of European madder have been used as mentioned above in Japan as colorants for food, e.g. confectionery, boiled fish and soft drinks, but they are not allowed as a food additive in either the US or the EU. Only the natural red colorant 'cochineal extract' (additive E-120(ii)) which is an extract of the dried bodies of the female cochineal insect, with around 20% carminic acid content, is allowed and widely used as a colouring agent in food processes in the EU (at dosage levels from 50 to 500 mg/kg) and in the US (only up to 5 mg/kg)<sup>7,15,22,38</sup>. Cochineal is commonly cultivated from the wild prickly pear cactus that grows thickly on the mountainsides in central Peru. Cochineal produces the pigment as a deterrent against other insects. The pigment can be obtained from the body and eggs of the insect. The few countries that produce commercial cochineal extracts are Peru, Mexico, the Canary Islands and, more recently, Chile and Bolivia. Only in Peru, the commercial production of cochineal extract is 200 ton/year, whereas in the Canary Islands production is only about 20 ton/year<sup>7</sup>. France is believed to be the world's largest importer of cochineal extract, but Italy and Japan come next. The insects are killed by immersion in hot water or hot ethanol or by exposure to sunlight, steam, or by oven heat. Approximately 130,000 insects or 2 kg dry insects are required to produce 1 kg of cochineal extract and approximately 200 kg of dried insects are produced weekly at the largest cochineal farm'. Cochineal extract is very soluble in water and exhibits shade changes with changes in pH. At pH 4 and below, it is orange; it turns from violet to red by increasing pH from 5 to 7. Traditionally, cochineal extract is extracted with water or aqueous alcohol at 90 °C to 100 °C by batch or continuous process. It is one of the few natural and water-soluble colorants that resist degradation with time. It has a good stability to heat, chemical oxidation, light and oxygen<sup>15</sup>. Often it is more stable than some synthetic food grade colorants but instable at low pH. A water insoluble form of cochineal extract is commonly used to colour several food products, e.g. sausage products, bakery and dairy products, confectionery, and often competes with red root beet (betanin) and anthocyanins in food colouring. The water-soluble form of cochineal extract is currently used in beverages, soft and alcoholic drinks such as aperitifs (e.g. Campari<sup>TM</sup>). Cochineal extract is not kosher and is not vegetarian. Its main limitation in food application is it's insolubility at low pH as mentioned above 15. Carmine, i.e. the additive E-120(i), is a complex of carminic acid with various metals: an aluminium lake of carminic acid is currently being used in the commercial preparation of carmine<sup>7,15,22</sup>. Variation in the ratio of carminic acid to aluminium produces a range of colours from pale strawberry to near black currant. Carmine is commonly traded as powder with a carminic acid content of 40% to 60% and liquid aqueous alkaline forms of carmine (and spray-dried derivatives) are also available with a carminic acid content of 2% to 7%. Cochineal extract and carmine are neither toxic nor known to be carcinogenic. Carmine is widely consumed in foods and beverages and has been rarely





implicated in adverse reactions. It can induce an anaphylacticshock reaction in a small number of people, due to impurities in the preparation, not due to the pigment itself. In fact, colouring compounds in natural food grade colorants are small molecular weight, non-protein chemicals that cannot be expected to give true food allergies, either immunoglobulin E (IgE)-mediated or cell-mediated allergy. However, natural colorants, for example carmine, are often extracted from biological materials that may contain many other compounds, including proteins in addition to the colouring compounds. In 1998, it was reported that IgE-mediated allergy might be caused by the consumption of carmine, due to the presence of protein residues<sup>120</sup>. Once IgE sensitization to these carmine proteins occurs, the level of exposure to these residual proteins through carmine-containing foods and beverages may be sufficient to elicit allergic reactions. For example, an anaphylactic reaction has been reported in a 34-year-old female atopic patient after ingestion of an orange beverage containing carmine. Symptoms like urticaria, rhinitis, nausea, vomiting, asthma, chills and diarrhea were observed. Skin prick tests carried out on the orange beverage, carmine and cosmetics containing the pigment were positive. In 1995, a reaction to carmine occurred in a 35-year-old woman after she ingested yoghurt that contained mixed fruits. Approximately 2 h after consumption she experienced symptoms of anaphylaxis including generalized urticaria, angioedema (localized swelling) and asthma<sup>121</sup>. In 1997, four adverse reactions following consumption of an alcoholic beverage containing carmine were reported in women ranging from 25 to 43 years old, with urticaria and angioedema. A skin prick test was performed and was found positive for carmine contained in the alcoholic beverage. Four instances of acute allergic reactions in a 28-year-old female after ingestion of orange beverage, strawberry milk and a red coloured cocktail containing carmine were also mentioned. An anaphylactic reaction in a 27-year-old woman has been reported after the consumption of a Popsicle coloured with carmine 122. Carmine and cochineal extract are different from the azo pigment 'cochineal red A' (additive E-124) which is a synthetic colorant.

Concerning the natural food colorant Arpink red<sup>TM</sup>, many toxicological data are also available: acute oral toxicity in mice of the pigment, 90-day subchronical toxicological study, acute dermal irritation/corrosion, acute eye irritation/corrosion, anti-tumour effectiveness, micronucleus test in mice, AMES test (Salmonella typhimurium reverse mutation assay), estimation of antibiotic activity, results of estimation of five mycotoxins. The fungal colorant gives a raspberry-red colour in an aqueous solution, stable at pH over 3.5. Neutral solutions are stable even after 30 min of boiling and colour shade does not change in relation with pH<sup>2</sup>. After evaluating all the materials provided by the Ascolor Biotech s.r.o company, the Codex Alimentarius Commission (Rotterdam meeting, March 11–15, 2002) made the following statement: "there will not be any objections to use the red colouring matter Arpink red<sup>TM</sup>"; The Arpink red<sup>TM</sup> colorant use was recommended as 100 mg/kg in meat products and in non-alcoholic drinks, 200 mg/kg in alcoholic drinks, 150 mg/kg in milk products including ice creams and 300 mg/kg in confectionery products<sup>2</sup>. After the first approval by the Codex Alimentarius, the Arpink red<sup>TM</sup> safety assessment was discussed during the 63rd meeting of Joint FAO/WHO Expert Committee on Food Additives (JECFA) in Geneva, June 8-17, 2004. The red colorant received a two-year temporary approval by the EU for

distribution as a food additive, exclusively in the Czech Republic from 2004 to 2006<sup>2</sup>. The file was still under progress at the European Food Safety Authority (EFSA) for some years. The situation now is not clear as Ascolor Biotech s.r.o. did not send data to authorities later on and seems to have closed its activities. Thus, there is no particular information on potential mycotoxin production and pathogenicity towards humans, despite the fact that the production of secalonic acid D, i.e. a pale yellow teratogenic mycotoxin, is well known from the fungus Penicillium oxalicum<sup>5</sup>. It has been shown that the biosynthesis of secalonic acid D (see Fig. 2 for the chemical structure) was dependent on the biosynthesis of the pigment emodin *via* the acetate-malonate pathway in a study conducted on the lichen *Laurera benguelensis*<sup>41</sup>. Diverse biological activities of secalonic acid D have been reported, such as a mycotoxin towards chicken and mice embryo, an inhibitor of various isozymes of protein kinase C and protein kinase A in murine secondary palate development<sup>123</sup>, as well as mouse and human cleft palatal inducing agent<sup>124</sup>.

In conclusion, this review provides relevant information regarding the properties of hydroxyanthraquinoid pigments, their biosynthetic pathway, their toxicity and carcinogenicity in recent decades. The collective information summarized in the review will act as an important segment for development of 'niche' fungal dyestuffs rich in hydroxyanthraquinoid pigments. These conclusions indicate that, even if the toxicological investigations of a new additive are not financially negligible, non-mycotoxigenic filamentous fungi such as strains of Drechslera spp., Herpotrichia spp., Paecilomyces spp. and Isaria spp. could be used for the production of dyestuffs rich in hydroxyanthraquinoid pigments as potent natural food grade colorants, with different shades according to the biomass composition: such as red (for main components catenarin & erythroglaucin), reddish brown (for tritisporin), bronze (for cynodontin), maroon (for helminthosporin) and orange-yellow (for pachybasin & averythrin). However, further studies should be performed on these fungal HAQN pigments to evaluate their potent carcinogenicity in humans from the food safety perspective. Current data on this topic are therefore insufficient.

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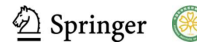
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