1 The evidence for anthocyanins in the betalain-pigmented genus Hylocereus is weak

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9 ABSTRACT

Here we respond to Zhou et al., 2020 "Combined Transcriptome and Metabolome analysis of Pitaya fruit 10 unveiled the mechanisms underlying Peel and pulp color formation" published in BMC Genomics. Given 11 12 the evolutionary conserved anthocyanin biosynthesis pathway in betalain-pigmented species, we are 13 open to the idea that species with both anthocyanins and betalains might exist. However, in absence of 14 LC-MS/MS spectra, apparent lack of biological replicates, and no comparison to authentic standards, the 15 findings of Zhou et al., 2020 are not a strong basis to propose the presence of anthocyanins in betalain-16 pigmented pitaya. In addition, our re-analysis of the datasets indicates the misidentification of 17 important genes and the omission of key anthocyanin synthesis genes ANS and DFR. Finally, our re-18 analysis of the RNA-Seq dataset reveals no correlation between anthocyanin biosynthesis gene 19 expression and pigment status.

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21 **KEYWORDS**

- 22 Anthocyanins, betalains, flavonoids, Caryophyllales, mutual pigment exclusion
- 23

24 MAIN TEXT

Betalain pigments are restricted to the Caryophyllales, where they replace the otherwise ubiquitous anthocyanins in several families [1]. However, not all families in the Caryophyllales produce betalains.

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27 The complex pigment distribution over evolutionary lineages can be explained by at least four

- 28 independent origins of the betalain biosynthesis [2]. Interestingly, anthocyanins have not been
- 29 observed in betalain-pigmented species [1, 3, 4]. Consequently, the theory of mutual exclusion of both
- 30 pigments was established and repeatedly supported by numerous studies in the last decades [3, 4].
- 31 Although co-occurrence of anthocyanins and betalains can be achieved through genetic engineering [5],

32 simultaneous accumulation of both pigments within a native species has previously never been reported

in nature. Zhou et al. 2020 [6] stated that "the anthocyanin coexistence with betalains is unneglectable"

in their publication about the pigmentation of pitayas. Here, we outline some reasons why we do not

think the study by Zhou et al.,2020 provides solid evidence for the presence of anthocyanins in betalain-

36 pigmented pitayas.

Patterns of proposed anthocyanin accumulation are weak basis for subsequent correlation with genes expression

39 Zhou et al., investigated metabolic differences between three pitaya cultivars: red peel/red pulp (RR), 40 yellow peel/white pulp (YW), and green peel/white pulp (GW). They claim that 70 different 41 anthocyanins are differentially accumulated in peels and pulps of these cultivars, but Table S9 lists only 42 14 anthocyanins. Unfortunately, these crucial metabolic analyses were apparently restricted to a single 43 sample per cultivar (Table S9) which prevents any solid conclusions about the quantity of pigments. 44 Nevertheless, we calculated the total amounts of detected anthocyanins per tissue and cultivar based 45 on the data they provide (**Table 1**). While the total anthocyanin amount in the red peel is substantially 46 higher than the amounts in yellow or green peel, there is only a very small difference between the 47 different pulp samples. The difference between the two white pulps is substantially higher than the 48 difference between one white and the red pulp. There are more anthocyanins reported in the yellow or 49 green peel than in the red pulp. The detection of delphinidin and malvidin glycosides (blue pigments) 50 would imply the presence of a functional flavonoid 3',5'-hydroxylase (F3'5'H) in all three cultivars. 51 However, Zhou et al., do not mention such an enzyme and our analyses revealed no evidence for the 52 presence of F3'5'H transcripts in pitaya. Zhou et al., report that "metabolites with similar fragment ions 53 were suggested to be the same compounds". But as we previously outlined this method does not follow 54 best practice [7]. We would expect at least to see the LC-MS/MS spectra and co-elution/fragmentation 55 of the pigments versus authentic reference compounds [7], especially when reporting the unexpected 56 occurrence of anthocyanins in a betalain-pigmented species.

57 **Table 1**: Total suggested anthocyanin amount reported by Zhou et al., 2020 in three pitaya cultivars.

Cultivar/tissue	Total amount of suggested anthocyanins
RR-peel	6,669,221
GW-peel	786,331
YW-peel	797,863
RR-pulp	631,184
GW-pulp	479,385
YW-pulp	155,051

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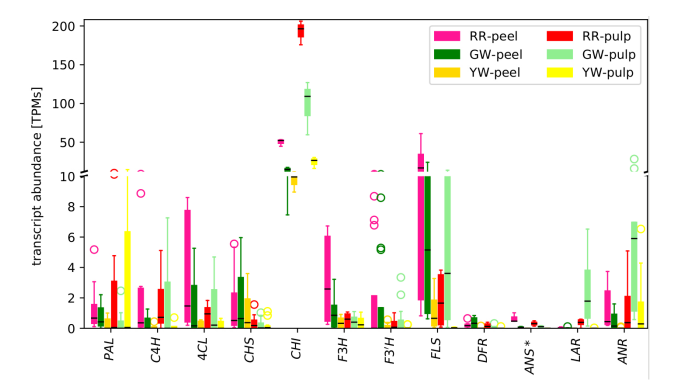
59 Transcriptomic analysis indicates likely block in anthocyanidin biosynthesis at the level of DFR and

60 ANS genes

51 Zhou et al., claim "Our results demonstrated that anthocyanin biosynthesis was one of the significantly

62 enriched pathways". We do not see any evidence for this statement, because the analysis presented in

63 their Fig. 5 covers only the general phenylpropanoid pathway and selected steps of the flavonoid biosynthesis. Our re-evaluation based on the construction of gene trees for all steps in the pathway 64 (Additional file 1, Additional file 2) indicates several cases of misidentification and missed gene copies, 65 66 although their annotation is difficult to evaluate as multiple transcripts are reported for each gene class, 67 and clear orthology is not established in absence of a phylogenetic analysis. Important genes of the 68 anthocyanin biosynthesis like DFR and ANS are missing. For example, DFR (Cluster-16519.0) and ANS 69 (Cluster-7001.0) are both present in the transcriptome assembly, but were not presented by Zhou *et al.*. 70 Zhou et al., write "The comparative pitaya transcriptome showed the differential regulation of the 71 anthocyanin pathway and genes controlling almost every single step in the pathway were differentially 72 regulated". However, two very important anthocyanin biosynthesis genes, DFR and ANS, are not 73 presented in Fig. 5 of Zhou et al.. Since expression of DFR and ANS would be crucial for the formation of 74 anthocyanins, we performed a re-analysis of the flavonoid biosynthesis including genes for the reactions 75 leading to anthocyanidins (Fig. 1). Our analysis is based on a transcriptome assembly of RR (red fruits) 76 and indicates a block in the anthocyanin biosynthesis at DFR and ANS (Fig. 1). We would expect to see 77 high transcript abundance of all genes necessary for anthocyanin formation if anthocyanins would be 78 substantially contributing to the red colour, especially at the intensity they report. We included LAR and 79 ANR in our analysis because the enzymes encoded by these genes are responsible for proanthocyanidin 80 biosynthesis. Anthocyanins and proanthocyanins have shared precursors and could be considered as 81 competing pathways. The presence of ANR transcript at high levels indicates that the lowly abundant 82 DFR and ANS transcripts might be involved in the proanthocyanidin biosynthesis rather than the 83 anthocyanin biosynthesis. Proanthocyanidin production could be an explanation for the identification of DFR, ANS, LAR, and ANR transcripts. However, it is important to emphasize that the transcript 84 85 abundances of DFR and ANS are in any case extremely low (average TPM < 1), and inconsistent with high 86 levels of anthocyanins. These low transcript abundances for DFR and ANS align well with previous 87 reports about the absence of anthocyanin in betalain-pigmented species [8–10].





89 Fig. 1: Transcript abundance of flavonoid biosynthesis genes in differently coloured pitaya cultivars.

90 This transcript abundance analysis is based on our Trinity assembly of all RR datasets. No bona fide ANS 91

was detected in the RR assembly, but the gene was represented with the closest homolog. RR=red

92 peel/red pulp, GW=green peel/white pulp, YW= yellow peel/white pulp

93

Conclusion 94

95 The evidence of anthocyanins lacks appropriate standards. Furthermore, the investigation of core anthocyanin biosynthesis genes via RNA-Seq does not provide insights into the accumulation of 96 97 anthocyanins, because there is no clear difference in bulk anthocyanin content between differently 98 pigmented pitaya varieties e.g. red vs. white. Consequently, there is little correlation between the 99 transcription of anthocyanin synthesis genes, and proposed levels of anthocyanins. There is clear 100 evidence of highly reduced DFR and ANS expression, which is not consistent with meaningful level so 101 anthocyanins. Altogether, we suggest that the evidence of anthocyanins in pitaya remains weak.

102

103 Methods

104 The applied methods are almost identical to our previous analysis of a very similar data set [7].

105 Transcriptome assembly

106 RNAseg datasets of different cultivars were retrieved from the Sequence Read Archive via fastg-dump 107 [11]. Trimming and adapter removal based on a set of all available Illumina adapters were performed via 108 Trimmomatic v0.39 [12] using SLIDINGWINDOW:4:15 LEADING:5 TRAILING:5 MINLEN:50 TOPHRED33. A 109 customized Python script was used to rename the surviving read pairs prior to the transcriptome 110 assembly. Clean read pairs were subjected to Trinity v2.4.0 [13] for *de novo* transcriptome assembly 111 using a k-mer size of 25. Short contigs below 200 bp were discarded. Previously described Python scripts [14] and BUSCO v3 [15] were applied for the calculation of assembly statistics for evaluation. Assembly 112 113 quality was assessed based on continuity and completeness. Although assemblies were generated for all 114 three species, the assembly generated on the basis of the data sets of Hylocereus undatus 115 (SRR11603186-SRR11603191) was used for all down-stream analyses.

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117 Transcriptome annotation

Prediction of encoded peptides was performed using a previously described approach to identify and retain the longest predicted peptide per contig [14]. Genes involved in the flavonoid biosynthesis were identified via KIPEs [16] using the peptide mode. Phylogenetic trees with pitaya candidate sequences and previously characterized sequences [16] were constructed with FastTree v2 [17] (WAG+CAT model) based on alignments constructed via MAFFT v7 [18] and cleaned with pxclsq [19] to achieve a minimal occupancy of 0.1 for all alignment columns.

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125 Transcript abundance quantification

126 Quantification of transcript abundance was performed with kallisto v0.44.0 [20] using the RNAseq reads

127 and our *Hylocereus undatus* transcriptome assembly [21]. Customized Python scripts were applied to

summarize individual count tables and to compare expression values [7].

129

130 ABBREVIATIONS

131 Not applicable

132

ADDITIONAL FILES

134 Additional file 1: Phylogenetic trees of genes in the flavonoid biosynthesis. Identified candidate

135 sequences are highlighted in red.

- 136 Additional file 2: Mapping table connecting the pitaya sequence names displayed in Additional file 1 to
- 137 the original sequence IDs assigned during the assembly process.

138

139 **DECLARATIONS**

- 140 Ethics approval and consent to participate
- 141 Not applicable
- 142
- 143 **Consent for publication**
- 144 Not applicable
- 145

146 Availability of data and materials

147 The datasets generated and/or analysed during the current study are available via PUB: 148 https://doi.org/10.4119/unibi/2956788. The Python scripts are available via github: 149 https://github.com/bpucker/pitaya.

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- 151 **Competing interests**
- 152 The authors declare that they have no competing interests.

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159 Authors' contributions

BP performed all analyses and prepared figures. BP and SFB wrote the manuscript. All authors reviewedthe manuscript.

162

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