Duplicated flavonoid 3’-hydroxylase and flavonoid 3’, 5’-hydroxylase genes in barley genome

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ABSTRACT

Background. Anthocyanin compounds playing multiple biological functions can be synthesized in different parts of barley (Hordeum vulgare L.) plant. The diversity of anthocyanin molecules is related with branching the pathway to alternative ways in which dihydroflavonols may be modified either with the help of flavonoid 3’-hydroxylase (F3’H) or flavonoid 3’,5’-hydroxylase (F3’5’H)—the cytochrome P450-dependent monooxygenases. The F3’H and F3’5’H gene families are among the least studied anthocyanin biosynthesis structural genes in barley. The aim of this study was to identify and characterise duplicated copies of the F3’H and F3’5’H genes in the barley genome.

Results. Four copies of the F3’5’H gene (on chromosomes 4HL, 6HL, 6HS and 7HS) and two copies of the F3’H gene (on chromosomes 1HL and 6HS) were identified in barley genome. These copies have either one or two introns. Amino acid sequences analysis demonstrated the presence of the flavonoid hydroxylase-featured conserved motifs in all copies of the F3’H and F3’5’H genes with the exception of F3’5’H-3 carrying a loss-of-function mutation in a conservative cytochrome P450 domain. It was shown that the divergence between F3’H and F3’5’H genes occurred 129 million years ago (MYA) before the emergence of monocot and dicot plant species. The F3’H copy approximately occurred 80 MYA; the appearance of F3’5’H copies occurred 8, 36 and 91 MYA. qRT-PCR analysis revealed the tissue-specific activity for some copies of the studied genes. The F3’H-1 gene was transcribed in aleurone layer, lemma and pericarp (with an increased level in the coloured pericarp), whereas the F3’H-2 gene was expressed in stems only. The F3’5’H-1 gene was expressed only in the aleurone layer, and in a coloured aleurone its expression was 30-fold higher. The transcriptional activity of F3’5’H-2 was detected in different tissues with significantly higher level in uncoloured genotype in contrast to coloured ones. The F3’5’H-3 gene expressed neither in stems nor in aleurone layer, lemma and pericarp. The F3’5’H-4 gene copy was weakly expressed in all tissues analysed.

Conclusion. F3’H and F3’5’H-coding genes involved in anthocyanin synthesis in H. vulgare were identified and characterised, from which the copies designated F3’H-1, F3’H-2, F3’5’H-1 and F3’5’H-2 demonstrated tissue-specific expression patterns. Information on these modulators of the anthocyanin biosynthesis pathway can be used in future for manipulation with synthesis of diverse anthocyanin compounds in different parts of barley plant. Finding both the copies with tissue-specific expression and

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a copy undergoing pseudogenization demonstrated rapid evolutionary events tightly related with functional specialization of the duplicated members of the cytochrome P450-dependent monooxygenases gene families.

**Subjects** Genetics, Molecular Biology, Plant Science

**Keywords** Flavonoid pigments, Anthocyanin biosynthesis, Gene duplication, *Hordeum*, Gene evolution, Near-isogenic lines, P450, CYP75

**INTRODUCTION**

Plant phenolic compounds flavonoids and their coloured derivatives anthocyanins are secondary metabolites providing important functions (*Grotewold, 2006a; Grotewold, 2006b*). Flavonoids are ubiquitously present in plant cells. They are involved in the regulation of developmental processes, in the protection against biotic and abiotic stress and in the attraction of seed dispersers and pollinators (*Khlestkina, 2013; Pourcel et al., 2007; Landi, Tattini & Gould, 2015*). Due to their antioxidant activity, these compounds are also useful for the health of plant foods consumers—humans and animals (*Khoo et al., 2017; Chaves-Silva et al., 2018*).

Cytochrome P450 (also called CYP) proteins, named for the absorption band at 450 nm, are one of the largest proteins superfamilies (*Werck-Reichhart & Feyereisen, 2000*). These proteins are found in all organisms from protists to mammals, but their number has exploded in plants. Flavonoid 3′-hydroxylation (*F3′H, CYP75B, EC 1.14.13.21*) and flavonoid 3′, 5′-hydroxylation (*F3′5′H, CYP75A, EC 1.14.13.88*) are cytochrome P450-dependent monooxygenases that require NADPH as a co-factor (*Tanaka & Brugliera, 2013*). These enzymes are involved in the biosynthesis of anthocyanin compounds—glycosylated forms of anthocyanidins producing by the flavonoid biosynthesis pathway (Fig. 1). *F3′H* and *F3′5′H* compete for substrate recruitment and hydroxylate 3′ or 3′5′ position of dihydroflavonols for the parallel synthesis of delphinidin and cyanidin, the precursors of blue and reddish-purple pigments (*Tanaka, Brugliera & Chandler, 2009; Tanaka & Brugliera, 2013*). Barley (*Hordeum vulgare* L.) is an important agricultural crop. In addition to the photosynthetic pigments giving a green colour, barley produces pigments that form diverse colouration patterns of different parts of plant. Purple and blue anthocyanins are accumulated in barley grains in the pericarp and aleurone layer, respectively (*Adzhieva et al., 2015; Shoeva, Strygina & Khlestkina, 2018*). Despite the fact that the genes coding the enzymes involved in anthocyanin biosynthetic pathway is well understood at the genetic and molecular level, the least studied genes in this branch are *F3′H* and *F3′5′H*. Because of useful properties of anthocyanin compounds, the study of genes involved in the anthocyanin biosynthesis is important. Previously, the presence of one *F3′H* gene copy (*F3′H-1*) expressing in genotype with purple pericarp was shown, as well as the presence of one *F3′5′H* copy (*F3′5′H-1*) with aleurone specific expression (*Shoeva et al., 2016; Strygina, Börner & Khlestkina, 2017*). Since the fact of tissue-specific activity of these genes and the fact that these anthocyanin compounds can be accumulated in other parts of the plant, it was concluded that there should be other copies of the *F3′H*
**MATERIALS & METHODS**

**Identification and structural analysis of duplicated genes**

The homologous nucleotide sequences of $F3'H$ and $F3'5'H$ genes in the barley genome with Bowman’s near-isogenic lines (NILs) contrasting in anthocyanin pigmentation: ‘BW’ (Bowman), ‘PLP’ (purple lemma and pericarp) and ‘BA’ (intense blue aleurone) were studied. The aim of this study was the identification and characterization of the $F3'H$ and $F3'5'H$ genes copies in the barley genome with Bowman’s near-isogenic lines (NILs) contrasting in anthocyanin pigmentation: ‘BW’ (Bowman), ‘PLP’ (purple lemma and pericarp) and ‘BA’ (intense blue aleurone).

The homologous nucleotide sequences of $F3'H-1$ (GenBank: AK362052) and $F3'5'H-1$ (GenBank: MF679159) were found in databases IPK Barley BLAST Server (https://webblast.ipk-gatersleben.de/barley_ibsc/), BARLEX (http://apex.ipk-gatersleben.de/apex/?p=284:10) and EnsemblPlants (http://plants.ensembl.org/index.html) using BLASTN search ($p$-value = 0.001). The multiple sequence alignment was made using Multalin (http://multalin.toulouse.inra.fr/multalin/). The gene structure was predicted using FGENESH+ software (http://www.softberry.com/berry.phtml?topic=fgenes_plus&group=programs&subgroup=gfs) using predicted polypeptide sequences of $F3'H-1$ and $F3'5'H-1$. The available promoter sequences were analysed with New PLACE database (https://sogo.dna.affrc.go.jp/cgi-bin/sogo.cgi?lang=en&pj=640&action=page&page=newplace). The annotation of the functional domains was carried out using InterPro: protein sequence analysis & classification (https://www.ebi.ac.uk/interpro/).

Modelling of the tertiary structure of the predicted amino acid sequences was performed using SWISS-MODEL (https://swissmodel.expasy.org/). Using MEGA v6.06 software (http://www.megasoftware.net) with 1,000 bootstrap replicates to assess the branch support the construction of the UPGMA tree, the calculation of $Ka/Ks$ ratio and calculation of divergence time was performed. The calibration of timeclock was based on divergence time.
Table 1  Gene-specific primers used for qPCR analysis of structural $F3'H$ and $F3'5'H$ genes of barley.

| Gene   | Forward primer (5′→3′) | Reverse primer (5′→3′) | PCR product length (bp) |
|--------|------------------------|------------------------|-------------------------|
| $F3'H$-1 | GCCAGGGAGTTCAAGGACA | CTCGCTGATGAATCCGTCCA | 168                     |
| $F3'H$-2 | AGGATAATCGCCCAGAGAAGG | GCCATGCGCCACTCCAC | 203                     |
| $F3'5'H$-1 | ATCGCATGTCGTGGCTATG | GCCGAGTTCACCATCATTTC | 143                     |
| $F3'5'H$-2 | CACAGACCTCAACATCAAAGC | TCCATCTCGCCTGTGCT | 138                     |
| $F3'5'H$-3 | GAACGCGGTTCAGACAT | TCCATCTCGCCTGTGCT | 148                     |
| $F3'5'H$-4 | GAACGACGACGCGAGAC | CGCCATTGCCACTCCAC | 109                     |

between barley and maize (50–60 MYA) (Salse et al., 2009; Cheng et al., 2012; Subburaj et al., 2016) and potato and petunia (30 MYA) (Kamenetzky et al., 2010).

Plant material, RNA extraction, cDNA synthesis

Plant material exploited for gene expression analysis included the cultivar Bowman of barley *H. vulgare* and two Bowman’s near-isogenic lines (NILs): ‘BW’ (Bowman, NGB22812), ‘PLP’ (purple lemma and pericarp, NGB22213) and ‘BA’ (intense blue aleurone, NGB20651). The set of the lines was provided by the Nordic Gene Bank (NGB, http://www.nordgen.org). PLP NIL have reddish-purple pericarp and stems due to the presence of *PLP* loci (chromosomes 2AL and 7HS); BA NIL have blue aleurone layer to the presence of *BA* loci (chromosomes 4HL and 7HL). The plants for RNA extraction from aleurone layers, pericarps, lemmas and stems were grown in ICG Greenhouse Core Facilities (Novosibirsk, Russia) under a 12 h photoperiod at 20–25 °C. The experiments were conducted in three replicates for each genotype. Aleurones and pericarps were cut out with a scalpel from grains at early dough stage maturity. RNAs from aleurone layers, pericarps, lemmas and stems (collected at the end of flowering) were extracted using a RNeasy Mini Kit (QIAGEN, Hilden, Germany) followed by DNase treatment with RNase-free DNase set (QIAGEN, Hilden, Germany). To obtain single-stranded cDNA samples total RNA was converted in a 20-µL reaction mixture from a template consisting of 0.4 µg of total RNA using a RevertAid First Strand cDNA Synthesis Kit (Thermo Fisher Scientific Inc., Waltham, MA, USA).

Primer design and qRT-PCR

Gene-specific primer pairs were constructed using Oligo Primer Analysis Software v.7 (https://www.oligo.net/) based on sequences found in IPK Barley BLAST Server (Table 1). The qRT-PCR was based on a SYNTOL SYBR Green I kit (Syntol, Moscow, Russia). The amplifications were performed in an ABI Prism 7,000 Sequence Detection System (Applied Biosystems, http://www.lifetechnologies.com). PCR was performed in a 15-µL reaction mixture under the following conditions: 1 cycle −15 min at 95 °C; 40 cycles −15 s at 94 °C, 30 s at 60 °C, 30 s at 72 °C. The construction of PCR product melting curves under the conditions: 15 s at 95 °C; 15 s at 60 °C; 15 s at 95 °C. The reference sequence was *Ubiquitin* gene; primers were suggested in (Himi & Noda, 2005). The raw data is in File S1. Each sample was run in three technical replications. The differences among the lines were tested by Mann–Whitney U-test ($p \leq 0.05$).


**RESULTS**

**Identification of F3’Hs and F3’5’Hs in barley genome**

Amid all identified highly homologous protein-coding sequences to F3’H-1 and F3’5’H-1 genes with 85.5–97.1% identity in functionally significant domains in the barley genome, only six encode proteins belonging to the CYP75 protein class (File S2). Among the revealed genes, one CYP75B-like copy of the F3’H-1 gene sequence (1HL; GenBank: AK362052) located on chromosome 6HS was found (File S2). The gene was designated F3’H-2. Its predicted full coding nucleotide sequence shares 69.6% identity with F3’H-1. Three CYP75A gene sequences were identified using F3’5’H-1 gene sequence (4HL; GenBank: MF679160): two highly homologous gene copies designated F3’5’H-2 (6HL) and F3’5’H-3 (6HS) with a level of identity 82.6% and 83.0%, respectively, and one copy designated F3’5’H-4 with 63.0% identity (7HS).

**Study of the structural organisation of the F3’H and F3’5’H genes**

All F3’H and F3’5’H genes identified in the current study in H. vulgare genome consist of two exons with the exception of F3’5’H-1 having three exons. Analysis of the promoter elements for the annotated genes (~600 bp upstream to the ATG start site) revealed many motives responsible for light-dependent activation (especially in F3’H-1 and F3’5’H-1), as well as Myb-dependent and Myc-dependent elements required for genes involved in the biosynthesis of flavonoid compounds (Fig. 2A, File S3). Unlike other copies, F3’5’H-2 and F3’5’H-3 have only one light-induced promoter element (GATA-box).

Amino acid sequences alignment with framing functional domains are shown in Fig. 2B. All the identified genes have a Cytochrome P450 domain (E-class, group I; IPR002401), however, F3’5’H-3 gene copy carries a frameshift indel mutation, which results in the truncation of the functional Cytochrome P450 domain in the middle and affects the tertiary protein structure (Fig. 2B, File S4). These sequences also possessed the conserved domains of flavonoid hydroxylase, including proline-rich region, heme binding domain, oxygen binding motif, hydroxylation activity site (CR1), EXXR motif and substrate recognition sites (SRS) (Fig. 3). Six functional SRSs, that are important for the determination of substrate specificity in CYP75 proteins, were determined in the predicted amino acid sequences of barley F3’Hs and F3’5’Hs. In F3’5’H-3 only three SRS, proline-rich and CR1 motifs are present (Fig. 3). All other barley CYP75s have not lost their functional domains.

**Evolutionary analysis of CYP75 genes**

The number of non-synonymous substitutions per non-synonymous sites (Ka), the number of synonymous substitutions per synonymous sites (Ks) and the Ka/Ks ratio for CYP75 genes of barley were calculated. Synonymous and non-synonymous substitution rates ranged between 0.541–0.685 and 0.269–0.461 for identified paralogs, respectively (File S5). Using the formula Ka/Ks, it was predicted that F3’H and F3’5’H paralogs may be under stabilising selection (Ka/Ks is close to 0.5) with the exception of F3’5’H-3. This copy may experience neutral selection since the Ka/KsF3’5’H-3 is close to one (File S5).

The phylogeny of F3’H and F3’5’H genes was analysed using complete coding sequences of identified genes from genome of barley and other angiosperm species. The phylogenetic
Figure 2 (A) Diverse promoter structure of F3'H and F3'5'H genes in barley; (B) structure analysis of identified genes. Promoter analysis was performed using New PLACE database. Orange, light-dependent motifs, blue, bHLH-type transcription factors binding elements, green, MYB-type transcription factors binding elements. Analysed protein motif is Cytochrome P450, E-class, group I (IPR002401).

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The tree indicated that F3'H and F3'5'H families form two separate clusters (Fig. 4, blue and purple clusters, respectively); within each one clearly divided into two groups—monocot and dicot plant species. It was assumed that F3'H and F3'5'H genes are the results of duplication and neofunctionalization of the single CYP75 gene in a genome of the common ancestor of monocot and dicot plant species. The analysis of genetic similarity and the divergence time calculation revealed that this event occurred about 129 million years ago (MYA) (Fig. 4) shortly before the monocots and dicots divergence (estimated time is 110–116 MYA).

In addition, we calculated the time of segmented duplications in H. vulgare genome with the formation of paralogous gene copies (Fig. 4). The F3'H copy apparently occurred...
### Proline-rich region

| SRS1 | SRS2 | SRS3 |
|------|------|------|
| 150  | 350  | 551  |

### Heme binding domain

| SRS4 | SRS5 |
|------|------|
| 450  | 551  |

#### Figure 3

The conserved domains of the F3′H and F3′5′H protein sequences in barley. Critical motifs are indicated with blue rectangles. Blue letters, oxygen binding motif. CR1, hydroxylation activity site. SRS, substrate recognition site.

before the divergence of Triticeae tribe from rice and maize about 80 MYA. In contrast, the F3′H in barley genome was duplicated at least three times: 91, 36 and 8 MYA. Thus, the last formation of the F3′H copy occurred after the separation of Hordeum genera from the common Triticeae ancestor (approximately 9–11 MYA Cheng et al., 2012; Subburaj et al., 2016).

#### Analysis of the F3′H and F3′5′H genes expression

Comparative analysis of relative gene expression levels was performed using RNAs isolated from the aleurone layer, pericarp, lemma and stems of the Bowman’s near-isogenic lines (NILs) contrasting in anthocyanin pigmentation: ‘BW’ (Bowman, NGB22812), ‘PLP’ (purple lemma and pericarp, NGB22213) and ‘BA’ (intense blue aleurone, NGB20651) (File S6). It was found, that the F3′H-1 gene was expressed in aleurone layer, pericarp and lemma with an increased expression level in a pigmented pericarp of ‘PLP’ (3.6 times higher than in uncoloured one) (Fig. 5). A tissue-specific expression was detected for the F3′5′H-1 gene only in aleurone layer (Fig. 5). Expression of the F3′5′H-1 gene only in aleurone layer was confirmed (Fig. 5). It was shown that in pigmented aleurone of ‘BA’ this gene was expressed 30 times actively than in uncoloured aleurone of ‘BW’. F3′5′H-2 was strongly expressed in pericarp and aleurone layer of ‘BW’ in comparison to coloured ones (9.3 and 12.7 times higher, respectively) (Fig. 5). Expression of the F3′5′H-3 gene was not detected in analysed tissues. The gene F3′5′H-4 was weakly expressed in all studied tissues with slight expression increasing in the pigmented stems (Fig. 5).

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DISCUSSION

Gene duplication is an important evolutionary mechanism providing a source of genetic material for the specialization or the new gene function appearance through the mutations and selection (Proulx, 2011; Magadum et al., 2013). Evolution by gene duplication has arisen as a general principle of biological evolution, which is apparent from the prevalence of duplicated genes in all genomes of sequenced organisms (Ohno, 1970). Gene copies have occurred as a result of segmental duplications (duplication of individual genomic regions) or polyploidization (whole genome duplications) (Ohno, 1970; Lynch et al., 2001; Eichler & Sankoff, 2003).

Gene duplicates can expect one of the possible fates: pseudogenization (PG), subfunctionalization (SF) or neofunctionalization (NF) (Ohno, 1970). In the PG process, one of the gene copies loses its function after degenerate mutation acquiring, for example, in the promoter region. The NF process proposes that one gene copy retains the ancestral function while the other gets a novel function. The SF is a major process of divergence with differential division of ancestral gene functions (Ohno, 1970).

In plants, the pattern of the SF leading to tissue-specific expression is frequent. For instance, regulatory genes coding bHLH/Myc-type transcription factors controlling the
The relative level of gene expression

The relative level of gene expression

Figure 5  Expression of F3'H and F3'5'H genes in barley cv Bowman and Bowman NILs contrasting for anthocyanin pigmentation. (A) F3'H-1. (B) F3'H-2. (C) F3'5'H-1. (D) F3'5'H-2. (E) F3'5'H-4. Barley tissues: aleurone layer, pericarp, lemma and stems. The data is presented as mean value ± standard error. An asterisk (*) indicates a statistically significant difference between Bowman and its NIL, with p ≤ 0.05 (Mann–Whitney U-test).

Full-size [DOI: 10.7717/peerj.6266/fig-5]

Anthocyanins biosynthesis in barley grain divide their functions: HvMyc1 gene (located in 2HL chromosome) regulates accumulation of anthocyanin pigments in pericarp while HvMyc2 gene (located in 4HL chromosome) provides the biosynthesis of anthocyanin pigments in aleurone layer like in PLP and BA NILs, respectively (Jende-Strid, 1993; Cockram et al., 2010; Strygina, Börner & Khlestkina, 2017). As an example of tissue-specification of structural anthocyanin biosynthesis genes flavanone 3-hydroxylase (F3'H) genes in Triticum aestivum genome could be considered: the copy designated TaF3H-B2 is transcribed specifically in roots of bread wheat while the TaF3H-B1 gene copy is not expressed in roots but it is expressed in other different parts of the plant (Khlestkina et al., 2013).

In the flavonoid biosynthesis pathway, F3'H and F3'5'H are important enzymes controlling the hydroxylation at the 3' and 5' of reddish-purple and blue pigments, respectively (Tanaka, Brugliera & Chandler, 2009; Tanaka & Brugliera, 2013). In most plants, F3'H and F3'5'H genes are present in low-copy number. In the current work, we have identified duplicated copies of F3'H and F3'5'H genes in H. vulgare genome. We have shown that the divergence between F3'H and F3'5'H genes from the common ancestor’s CYP75 gene occurred 129 MYA, which occurred based on our calculations 13–19 MYA years earlier than the appearance of monocot and dicot plant species (110–116 MYA according to our calculations; 90–165 MYA according to different estimates Chaw et al., 2004; Herron et al., 2009; Cheng et al., 2012). The duplication of F3'H and F3'5'H in barley genome took place several times: the F3'H copy arose approximately 80 MYA while the appearance of F3'5'H copies occurred 8, 36 and 91 MYA (Fig. 4). Thus, the first acts of duplication of both genes occurred before the origin of the family Poaceae (Gramineae) (Kellogg, 2001).
The ratio of non-synonymous ($K_a$) to synonymous ($K_s$) substitutions is used to determine the direction of natural selection after duplication: $K_a/K_s > 1$ implies positive selection, $K_a/K_s < 1$ means stabilising selection, $K_a/K_s = 1$ indicates neutral selection (Kondrashov et al., 2002). Analysis of duplicated $F3'H$ and $F3'5'H$ genes indicated that most of the identified gene copies are under stabilising selection. The exception is $F3'5'H-3$ gene copy, which is supposed to be a pseudogene due to the mutation in the coding part of the gene, which breaks the reading frame and changes the protein structure. In addition, we did not detect its transcriptional activity in analysed tissues.

The genes encoding $F3'H$ showed a precise tissue-specific activity likewise $TaF3H$ genes of bread wheat: $F3'H-1$ is expressed in aleurone layer, pericarp and lemma, while $F3'H-2$ is transcriptionally active in stems only (Fig. 5). Besides, increasing of the expression level were observed in tissues with reddish-purple pigmentation (pericarp and stems) apparently provided by cyanidin derivatives (these identifications are putative due to the absence of a biochemical study of the gene products). The increase of relative expression level of $F3'H-1$ in the aleurone layer or lemma was not detected in BA and PLP NILs (Fig. 5). In these tissues, there are almost or completely no cyanidin derivatives, which is evident from the phenotype of these lines (File S6). An increase in the level of gene expression in anthocyanin-pigmented plant tissues is a common feature of genes in anthocyanins biosynthesis pathway in cereals (Shoeva et al., 2015; Shoeva et al., 2016; Shoeva & Khlestkina, 2015). For example, in the pericarp of purple-grained PLP line the expression level of flavonoid biosynthesis structural genes ($CHS$, $CHI$, $F3H$, $F3'H$, $DFR$, $ANS$) was significantly higher than in the uncoloured Bowman, that led to total anthocyanin content increase in PLP line identified by ultra-performance liquid chromatography (HPLC) (Shoeva et al., 2016).

Among the $F3'5'H$ copies, only two have a tissue-specific activity: $F3'5'H-1$ and $F3'5'H-2$. The $F3'5'H-1$ copy was expressed only in the aleurone layer, and the level of its activity was much higher in the blue aleurone compared to the uncoloured one (Fig. 5). Aleurone-specific expression of this gene was noted earlier, and it was shown that $F3'5'H-1$ is one of the key regulators of the aleurone layer pigmentation (Strygina, Börner & Khlestkina, 2017). The copy designated $F3'5'H-2$ was expressed only in the barley grain. Moreover, the expression of this gene is much higher in the aleurone layer and pericarp in the green BW line in comparison to coloured ones (Fig. 5). The $F3'5'H-4$ gene copy was expressed in all tissues analysed. Since there are almost no light-dependent elements in the promoters of $F3'5'H-2$ and $F3'5'H-4$, it can be assumed that these gene copies encode for different isoenzymes specialised in the synthesis of such flavonoid compounds as catechin available in the barley at the high level (McMurrough, Loughrey & Hennigan, 1983; Madigan, McMurrough & Smyth, 1994). Alike specialization was demonstrated earlier for such organisms as tea plant and its relatives (Punyasiri et al., 2004; Jin et al., 2017). These results suggest the SF and diversification of $F3'H$s and $F3'5'H$s in the barley genome.

**CONCLUSIONS**

$F3'H$ and $F3'5'H$-coding genes involved in anthocyanin synthesis in *Hordeum vulgare* L. were identified and characterised. One $F3'H$ ($F3'H-2$) and three $F3'5'H$s ($F3'5'H-2$, $F3'5'H-3$, and $F3'5'H-4$) were identified.
were described for the first time. The subfunctionalization expressing in tissue-specific activity of \( F3'S'H-1, F'H-2, F3'S'H-1 \) and \( F3'S'H-2 \) genes was shown. It was also found that the \( F3'S'H-3 \) gene, carrying frame-shift indel mutation, is the pseudogenic duplicate. Finding both the copies with tissue-specific expression and the \( F3'S'H-3 \) copy undergoing pseudogenization demonstrated rapid evolutionary events tightly related with functional specialization of the duplicated members of the cytochrome P450-dependent monooxygenases gene families. The results obtained are important for understanding of the features of flavonoid biosynthesis regulation in barley.

### ADDITIONAL INFORMATION AND DECLARATIONS

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**Competing Interests**
The authors declare there are no competing interests.

**Author Contributions**
- Alexander V. Vikhorev performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Ksenia V. Strygina conceived and designed the experiments, performed the experiments, analyzed the data, contributed reagents/materials/analysis tools, prepared figures and/or tables, authored or reviewed drafts of the paper, approved the final draft.
- Elena K. Khlestkina conceived and designed the experiments, contributed reagents/materials/analysis tools, authored or reviewed drafts of the paper, approved the final draft.

**Data Availability**
The following information was supplied regarding data availability:
Raw data is available in the Supplemental Materials.

**Supplemental Information**
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.6266#supplemental-information.
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