Identification, cloning and characterization of an ultrapetala transcription factor CsULT1 from Crocus: a novel regulator of apocarotenoid biosynthesis

Nasheeman Ashraf1*, Deepti Jain2 and Ram A Vishwakarma3

Abstract

Background: Crocus sativus is a triploid sterile plant with long red stigmas which form commercial saffron. Saffron is the site for synthesis and accumulation of apocarotenoids like crocin, picrocrin and safranal which are responsible for its color, flavour and aroma making it world’s most expensive spice. These compounds are formed by oxidative cleavage of zeaxanthin by carotenoid cleavage dioxygenases. Although the biosynthetic pathway of apocarotenoids is known to a considerable extent, the mechanism that regulates its tissue and developmental stage specific expression is not known.

Results: In the present work, we identified, cloned and characterized ultrapetala transcription factor called CsULT1 from Crocus. The gene contains an 80 amino acid long conserved SAND domain. The CsULT1 transcript was more abundant in stigma and showed increase in expression from pre anthesis stage till anthesis and decreased in post anthesis stage which corroborated with the accumulation pattern of crocin indicating its possible role in regulation of apocarotenoid biosynthesis. CsULT1 was found to be transcriptionally active and localized in nucleus. Its expression is induced in response to phytohormones like auxin, methyljasmonate and salicylic acid. Overexpression of CsULT1 in Crocus calli resulted in enhanced expression of key pathway genes like phytoene synthase (PSY), phytoene desaturase (PDS), beta carotene hydroxylase (BCH) and carotenoid cleavage dioxygenases (CCDs) indicating its role in regulation of apocarotenoid biosynthesis.

Conclusion: This work presents first report on isolation and characterization of ultrapetala gene from Crocus. Our results suggest that CsULT1 is a novel regulator of Crocus apocarotenoid biosynthesis. We show for the first time involvement of plant SAND domain proteins in regulating secondary metabolic pathways.

Keywords: Ultrapetala, Crocus, Stigma, Saffron, Carotenoids, Apocarotenoids, SAND domain

Background

Crocus sativus L. (Iridaceae) is a sterile triploid plant propagated vegetatively through corms [1]. The desicced stigma of C. sativus forms saffron and is source of various carotenoids and unique compounds called apocarotenoids which are produced by oxidative tailoring of carotenoids [2]. Apocarotenoids are synthesized in a number of plants including maize, tomato, Arabidopsis, Cricus etc. but Crocus finds a special place because it is the only plant which produces crocin, picrocrocin and safranal in significant quantities [3]. The saffron apocarotenoids are formed by zeaxanthin cleavage [4] followed by specific glycosylation steps [5]. Because of the presence of these unique apocarotenoids Crocus stands apart from other related crops and is considered as one of the world’s costliest spices [6]. Besides, saffron apocarotenoids also have tremendous pharmacological properties and have been used for the treatment of a wide range of cancers [7,8].
Carotenoids and their cleavage products are synthesized by plastid localized methylerythritol phosphate (MEP) pathway. Biosynthesis of these compounds is regulated throughout the life cycle of a plant and dynamic changes occur in their composition to match the prevailing developmental requirements and response to external environmental stimuli [9]. Although the carotenoid biosynthetic pathway has been studied to a considerable extent in many plants including *Crocus* but the fundamental knowledge regarding the regulation of carotenogenesis in plant cells is still in its infancy [10]. In *Crocus*, apocarotenoids are synthesized only in stigma part of the flower and that too in developmental stage specific manner, but nothing is known about the mechanism that regulates its synthesis. Therefore, it will be quite interesting to take a step towards unravelling the regulatory pathway of carotenoid/apocarotenoid biosynthesis in *Crocus*.

The major goal of the present study was to identify transcription factors that regulate apocarotenoid biosynthesis in saffron. It is a well established fact that stigma part of the *Crocus* flower is the actual site for synthesis of many important apocarotenoids [11,12], however, we still attempted to study pattern of crocin accumulation (crocin being an important metabolite) in different parts of the flower and at different stages of stigma development. We also selected five transcription factors belonging to five different gene families from saffron gene database [1] and investigated their temporal and spatial expression profile. The results demonstrated that *ULTRAPETALUM* (*ULT*) gene shows higher expression in stigma tissue and the expression increased till the day of anthesis and subsequently decreased. This expression profile matched with the accumulation pattern of crocin in saffron thereby indicating a possible role of this gene in regulating biosynthesis of apocarotenoids. The *ULT* encodes a small cysteine rich protein containing a B-box like motif and a SAND domain, a DNA binding motif previously reported only in animal transcription factors [13]. This transcription factor has been proposed to act as regulator of developmental gene expression. In *Arabidopsis*, it functions in floral stem cell termination pathway [14]. *ULT* has been shown to act as trithorax group chromatin remodelling factor which regulates function of Agamous locus during stem cell termination [15]. Recently it has been demonstrated that *ULT1* acts as an antirepressor that promotes transcriptional activation by antagonizing PcG-induced histone methylation and, via physical interaction with *ATX1* that deposits H3K4me3 activating marks, promotes an open chromatin conformation to recruit proteins involved in transcriptional initiation and elongation [16]. More recently *ULT* was found to be involved in gynoecium formation [17]. *ULT1* function thus represents a novel chromatin-mediated mechanism that activates genes controlling stem cell fate in plants. This observation expands the repertoire of plant epigenetic regulators involved in developmental pathways and suggests involvement of chromatin mediated pathways in controlling dynamics of transcription during such pathways.

In this report we describe identification, isolation and characterization of *ULT* gene, *CsULT1*, from *Crocus sativus*. *CsULT1* is preferentially expressed in stigma and induced by phytohormones such as MJ, SA, 2,4-D. Further, *CsULT1* is localized in nucleus and is transcriptionally active. *Crocus* transformation has not yet been established. Here we studied transient overexpression of *CsULT1* in *Crocus* calli and observed that its overexpression upregulates some key carotenoid/apocarotenoid pathway genes. This work represents, to our knowledge, the first functional characterization of a *C. sativus* *ULT* gene and also first report on a transcriptional regulator of apocarotenoid biosynthetic pathway.

**Methods**

**Plant material**  
*Crocus sativus* was grown in the experimental farm at Indian Institute of Integrative medicine (IIIM), Srinagar, India (longitude: 34°5′24″ N; latitude: 74°47′24″ and altitude 1585 m above sea level). It was used as source plant material for the present study. The voucher specimen was deposited at Janaki Ammal Herbarium (RRLH), IIIM, Jammu. The details of the specimen are: (Accession number: 22893; Accession date: 12/01/2015; name of collector: Nasheeman Ashraf; Place of collection: IIIM, Srinagar Farm; Date of collection: 01/01/2015). For tissue specific expression profiling, on the day of flower opening, tepals, anthers and stigma were collected from flowers separately, frozen in liquid nitrogen and stored in ~80°C till further use. For developmental stage specific expression, stigma was collected at three different stages viz three days before anthesis, on the day of anthesis and two day after anthesis. For hormone treatments, flowers were grown in pots and were mist sprayed with 100 μM methyljasmonate, 1 mM salicylic acid, 50 μM 2, 4-D and 100 μM ABA. Tissue samples were collected after 12 and 24 h of hormone treatment. For overexpression studies, calli overexpressing *CsULT1* and vector control calli were taken for RNA isolation. For each experiment, tissue from three biological replicates was pooled in.

**Sample preparation and HPLC analysis**  
Crocin analysis was done as described by Moraga et al. [12]. For extract preparation, 0.5 mg tissue from tepals, anthers and stigma (collected at three different stages) was crushed with a micropestle in 700 μl Tris–HCl (50 mM, pH 7.5 containing 1 M NaCl), and incubated for 10 minutes on ice. This was followed by addition of
700 μl of chloroform. The extract was then incubated on ice for an additional 10 min. Centrifugation at 3000 g for 5 min at 4°C was done to separate the phases. The lower chloroform phase was evaporated and the dried residues were stored together with the upper aqueous phases at –80°C until high-performance liquid chromatography (HPLC) analysis. The LCMS apparatus of Nexera UHPLC (130 MPa) equipped with MS-8030 (Shimadzu) was used for the Study and data was generated using lab solutions software. Enable RP-C18 column (250 mm × 4.6 mm, 5 μm) was used. The injection volume was 5 μl and flow rate 0.3 ml/min. Mobile Phase A (Water and Acetonitrile ratio 1:1) and mobile phase B (0.1% Acetic acid in water) were used in a linear gradient flow and column temperature was set at 75°C initially.

Gene expression analysis using quantitative real time PCR

Total RNA was extracted from pooled tissue using TRIzol reagent and used for cDNA synthesis by Reverse Transcription kit (Fermentas) following manufacturer's instructions. qRT-PCR was performed in triplicates in ABI StepOne Real time (Applied biosystems). The reaction was carried out in a total volume of 20 μl, consisting of 10 μl of 2X SYBR Green Master Mix, 0.2 μM (each) gene specific primers for all the genes studied and 100 ng of template cDNA. The cycling parameters were 95°C for 20 s, followed by 40 cycles of 95°C for 15 s and 60°C for 1 min. The sequence of all the primers used in this study is given in Additional file 1. The specificity of each primer pair was validated by a dissociation curve (a single peak is given in Additional file 1. The specificity of each primer pair was validated by a dissociation curve (a single peak was observed for each primer pair) (Additional file 2). The relative quantification method (ΔΔCT) was used to evaluate quantitative variation between the replicates examined. The amplification of actin cDNA was used as an endogenous control to normalize all data.

Cloning of full length CsULT1 gene

The partial clone of CsULT1 was obtained using cDNA synthesized from Crocus flower RNA and primers (ULT-F and ULT-R) designed from EST sequence (cr:saCl000502:1) present in NCBI (www.ncbi.nlm.nih.gov/nucest). Sequence analysis of the partial clone revealed that it has the 3′end and only 5′end needs to be amplified in order to obtain the full length clone. Thus the full length cDNA clone of CsULT1 was obtained by performing 5′RACE using gene specific primer (ULT-5′) and UAP primer provided with the 5′RACE kit (Clontech) following manufacturer's instructions. The amplified product was run on 1% agarose gel and purified with gel extraction kit (Qiagen). The purified product was then cloned in the pGEM-T Easy vector and sequenced. For the amplification of full length clone, gene specific primers were designed from the full length nucleotide sequence as obtained from alignment of partial clone and the 5′RACE product. The full length cDNA clone was amplified by PCR using cDNA as template and the gene specific primer pair (CsULT-F and CsULT-R). The PCR product was run on 1% agarose gel, purified by gel extraction kit (Qiagen) and subsequently cloned into the pGEM-T Easy vector. The cycling conditions used were 3 min at 94°C, 30 cycles (30s at 94°C, 30 s at 60°C and 1 min at 72°C) and final extension for 10 min at 72°C. The nucleotide sequence of CsULT1 was submitted to GenBank and the accession number is KM670459.

Sequence analyses

The full length nucleotide sequence of CsULT1 was translated using Translate tool (http://web.expasy.org/translate/) and the properties of deduced amino acid sequence were estimated using ProtParam (http://web.expasy.org/protparam/). Multiple sequence alignment and phylogenetic analysis was performed using the ClustalW with the default parameters through the service of the European Bioinformatics Institute (http://www.ebi.ac.uk/Tools/msa/clustalw2).

Subcellular localization

The subcellular localization of CsULT1 was studied by performing transient expression assay in onion epidermal cells. For this, CsULT1 with restriction sites for NcoI and SpeI was amplified using ULTCam-F and ULTCam-R1 primer pair and fused in frame with 5′ terminus of GFP reporter gene in pCAMBIA-1302. The cycling parameters were same as described above. The fusion construct of CsULT1-GFP was bombarded on to the onion peels using biolistic gene delivery device PDS-1000/He (Bio-Rad, USA). The onion peels were then incubated for 24 hours before visualizing in confocal microscope.

Transactivation assay

Full length protein coding sequence of CsULT1 was cloned in yeast (Saccharomyces cerevisiae) expression vector pGBK7T (Clontech) at Nd6-EcoRI site to express CsULT1 protein fused to GAL4 DNA-binding domain (GAL4-BD). The primers used were ULTGBKT-F and ULTGBKT-R and cycling parameters are same as described above. The resulting construct was transformed into Y187 yeast strain. The positive transformants were selected onto synthetic medium lacking tryptophan and leucine. Cells from two independent transformants were collected and assayed for β-galactosidase activity by using ortho-nitrophenyl-β-D-galactoside (ONPG) as substrate as described in clontech manual (PT3024-1).

Plant expression vector and transformation of Crocus calli

The CsULT1 gene with NcoI and SpeI restriction sites was PCR amplified using ULTCam-F and ULTCam-R2 primer pair and cloned into pCAMBIA1302 vector, containing
CaMV 35S promoter. For the transient expression, the *Crocus* calli were arranged at the center of petri dish and the biolistic gene delivery device PDS-1000/He (Bio-Rad, USA) was used for transgene delivery via microprojectile bombardment. Plasmid DNA (at the concentration of 1 μg/μL) was coated on the surface of gold particles and bombarded on to the calli. The particle delivery system was adjusted to 1100 psi of helium pressure and 27 mmHg of vacuum pressure inside the chamber. After bombardment, the calli were transferred to fresh media and after five days they were again transferred to media containing hygromycin for selection of transgenic structures. After 10 days, the calli which did not harbour the *CsULT1*-pCAMBIA construct turned blackish whereas the ones with the gene construct looked fresh and were used for further experimental studies. The transgenic calli were screened by genomic PCR. For this, genomic DNA was isolated from independent *CsULT1* overexpression and empty control calli using the DNeasy Plant mini kit (Qiagen). The presence of *CsULT1* was confirmed by genomic PCR using gene specific primer and reverse primer corresponding to GFP. Transgenic and control calli were used for measuring the transcript levels of few carotenoid pathway genes including *PSY* (GenBank accession: AJ888514), *PDS* (GenBank accession: AY183118), *BCH* (GenBank accession: AJ937791), *CCD4b* (GenBank accession: EU523663.1) and *CCD2* (GenBank accession: KJ541749) using quantitative real time PCR as described above.

**Results**

**Analysis of Crocin in different tissues and developmental stages**

Since crocin is the most important metabolite in saffron and responsible for its coloring property, we measured its quantity in different parts of *Crocus* flower and at various stages of stigma development (pre anthesis, anthesis and post anthesis) using HPLC. Results indicated that crocin was present only in stigma part of flower. We were not able to detect crocin in other parts of the flower like tepals and anther. Further, its content showed increasing trend from pre anthesis to anthesis stage and later again decreased after anthesis (Figure 1). This was in confirmation with earlier reports [12] where they have shown that the major apocarotenoids like crocin and picrocrocin are detected in orange stage and increased rapidly during the following stages of stigma development till they reached maximum in scarlet stage at anthesis. This confirms that stigma is the site for synthesis and accumulation of major *Crocus* apocarotenoids and their synthesis is congruent with development of stigma reaching highest at anthesis stage.

**Isolation and expression profiling of *CsULT1***

We aimed at identification of transcription factors which regulate biosynthesis of *Crocus* apocarotenoids. Towards this, five transcription factors belonging to different gene families were selected from saffron ESTs (www.ncbi.nlm.nih.gov/nucest). The selected genes were *Myb* (cr.saCl000348:1), MADS box (cr.saCl001329:1), *WRKY* (cr.saCl000652:1), Zinc finger (cr.saCl000359) and *ULT* (cr.saCl000502:1). The expression pattern for all these genes was investigated in various tissue types and at different developmental stages using quantitative real time PCR. Our results demonstrated that a *ULT* transcription factor showed higher induction in stigma part of the flower and its expression increased till the day of anthesis and then subsequently decreased (Figure 2a and b). This expression pattern corroborated with the accumulation pattern of apocarotenoids suggesting involvement of this gene in regulating biosynthesis of these compounds. Among other genes studied, only *Myb* showed higher expression in stigma as compared to other flower parts, however, its expression at different developmental stages of stigma did not match with the pattern of apocarotenoid accumulation (Additional file 3).

Full length *ULT* was cloned by RT-PCR and 5’ RACE and was named as *CsULT1* [GenBank accession number: KM670459]. The gene contains 708 bp open reading frame encoding for 235 amino acids long protein (Additional file 4) with a predicted molecular mass of 26.5kD and pl 8.32. Domain search revealed presence of conserved SAND domain in *CsULT1* which normally consists of evolutionarily conserved 80 to 100 amino acid long DNA binding motif [18]. The sequence alignment of ULTs from various organisms has revealed two conserved cores in SAND domains viz TPxxFE and KDWK. While TPxxFE is perfectly conserved among all ULT proteins in plants, KDWK shows variability at primary level however, the secondary structure is conserved [19]. Alignment of *CsULT1* with other plant orthologs (Figure 3) showed high sequence homology with ULT from *Phoenix* (79.57%), *Vitis vinifera* (78%), *Populus* (76%) and *Medicago* (75%). Sequence alignment showed that ULT proteins show significant homology along the entire length of the protein except at the extreme N terminus. Moreover, TPxxFE motif was present in *CsULT1* and was conserved among all the proteins used for alignment. Phylogenetic analysis of selected sequences placed *CsULT1* close to its homolog from *Phoenix* (Figure 4).

**Subcellular localization of *CsULT1***

In order to have a preliminary understanding about the mechanism underlying the regulatory activity of *CsULT1*, its subcellular localization was investigated. Programs like Prosite and PSORT revealed absence of any sorting signal and predicted *CsULT1* to be localized in cytosol. For confirming the localization experimentally, *CsULT1* was cloned in frame with GFP reporter gene. The expression of the fusion gene construct *CsULT1*-GFP was driven by the 35S promoter of cauliflower mosaic virus (*CaMV*—
The fusion gene was introduced into onion (*Allium cepa*) epidermal cells by particle bombardment. While the control GFP accumulated throughout the cell, *CsULT1*-GFP was localized in the nucleus (Figure 5). This might be because of the fact that ULT proteins are small enough and can diffuse passively into the nucleus through the nuclear pores [15].

**Transactivation assay**

To investigate the ability of *CsULT1* to activate transcription, a transient expression assay was performed using a GAL4-responsive reporter system in yeast cells. For this, the full-length coding region of *CsULT1* was fused to the GAL4 DNA-binding domain (BD) to generate pGBK7-*CsULT1-BD* construct which was then transformed into yeast strain Y187. The transformants were assayed for their ability to activate transcription from the GAL4 upstream activation sequence. The transformed yeast cells harboring pGBK7-*CsULT1-BD* construct grew well in SD medium lacking tryptophan and leucine, and showed β-galactosidase activity, whereas cells containing pGBK7 (negative control) showed no β-galactosidase activity (Figure 6). This data confirmed transcriptional activity of *CsULT1*.

**Induction of *CsULT1* by phytohormones**

To investigate the effect of various phytohormones on expression of *CsULT1*, Crocus flowers were treated with...
Figure 2 qRT-PCR analysis of CsULT1 expression. (A) in different tissues of C. sativus (B) at different developmental stages. Transcript levels were normalized by actin. Data are means and SD from three biological replicates.

Figure 3 Multiple sequence alignment of CsULT1. The deduced amino acid sequence of CsULT1 is aligned with homologs from other plants.
salicylic acid (SA), methyljasmonate (JA), 2,4-D and abscisic acid (ABA) in a time course study. The expression of CsULT1 was measured by qPCR using RNA isolated from treated tissue samples. Compared with the uninduced control, CsULT1 expression increased in response to all these hormones used (Figure 7). In response to SA treatment, expression of CsULT1 increased up to 122 fold (log 7 fold) at 24 hr post treatment while in response to JA, the expression enhanced approximately up to 150 fold (log 7 fold) at 12 h post treatment. CsULT1 showed maximum change in expression in response to 2,4-D where it showed 175 fold (log 7.4) induction at 12 hr of treatment. However, there was not much significant change in expression in response to ABA.

Transient over-expression of CsULT1 in C. sativus calli increases MEP pathway gene expression

For gaining an understanding on the role of CsULT1 in Crocus apocarotenoid biosynthesis, the gene was transiently expressed in Crocus calli under the control of CaMV-35S promoter. The presence of transgene in transiently transformed calli was confirmed by genomic PCR. In the CsULT1-overexpressing calli, the CsULT1 gene was expressed 2.5 fold higher than the empty vector control. Further, we checked expression of few of the MEP pathway genes and observed that PSY, PDS, BCH, CsCCD4b and CsCCD2 genes showed upregulation in CsULT1 overexpressing calli (Figure 8a). PSY and PDS catalyze the initial rate limiting steps in carotenoid biosynthetic pathway. Further, BCH is involved in the formation of zeaxanthin from beta carotene [11] and this zeaxanthin acts as the substrate for the formation of Crocus apocarotenoids by CsCCD2 enzyme [20]. CsCCD4b (another member of CCD gene family) is also involved in the formation of apocarotenoids from carotenoid substrates. Therefore, enhanced expression of the

![Image](image-url)
above mentioned genes may result in increased zeaxanthin pool which may subsequently be tailored to form apocarotenoids.

Various phytohormones were shown to induce expression of 
CsULT1 which in turn induced expression of key pathway genes of carotenoid metabolism. Therefore we were keen to investigate change in expression of pathway genes in response to phytohormone application. It was observed that SA, JA and 2,4-D induced expression of carotenoid pathway genes (Figure 8b) therefore indicating their possible role in mediating the function of CsULT1 in regulating carotenoid/apocarotenoid biosynthesis. Taken together, these results suggest that CsULT1 has a role in regulating metabolic flux towards the biosynthesis of apocarotenoids in *Crocus*.

**Discussion**

*Crocus* is known to accumulate large amounts of the apocarotenoids like crocetin (and its glycosylated forms, crocins), picrocrocins and saffranal in stigma part of the flowers. The proposed biosynthetic pathway is initiated through the symmetric cleavage of zeaxanthin at the 7,8/7,8 positions by a CCD2 enzyme [20]. The two cleavage products formed are 3-OH-β-cyclocitril and crocetin.

![Figure 6 Transactivation analysis of CsULT1 by β-galactosidase assay.](image)

Vec represents empty vector control and ULT-1 and ULT-2 represent two independent colonies used for the assay. Values are taken as average of three independent experiments of the transformants and presented as fold increase in activity.

![Figure 7 qRT PCR showing relative transcript level of CsULT1 in response to various hormones.](image)

Transcript levels were normalized by actin transcript level. Error bars indicate SD of three replicates.
dialdehyde which are further dehydrogenated and glycosylated to yield picrocrocin and crocins respectively. Aim of our study was to identify transcription factors regulating synthesis of these *Crocus* apocarotenoids. For this, we used combined approach of transcript and metabolite profiling. Since crocetin (which is subsequently converted into crocin) and picrocrocin are products of same cleavage step and crocin is more stable, we investigated crocin levels in different parts of *Crocus* flower and in stigma collected at three different developmental stages (pre anthesis, anthesis and post anthesis). We could detect crocin only in stigma while in other parts it was below detection levels. Further, crocin content increased from pre anthesis stage to anthesis and later decreased post anthesis (Figure 1). In earlier reports also same trend has been described for crocin accumulation [12].

Domain analysis showed that *CsULT1* contains a SAND domain which represents conserved 80-residue amino acid sequence and is found in a number of nuclear proteins, many of which function in chromatin-dependent transcriptional control [13]. These include proteins linked to various human diseases, such as the Sp100 (Speckled protein 100 kDa), NUDR (Nuclear DEAF-1 related), GMEB (Glucocorticoid Modulatory Element Binding) proteins and AIRE-1 (Autoimmune regulator 1) proteins [18]. Many of these proteins have been shown to bind DNA, but no clear sequence or structural relationship to known DNA binding motifs has been established. Based on the conservation of positively charged residues, including a characteristic

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**Figure 8** Relative expression levels of selected carotenoid pathway genes (A) in *CsULT1* overexpressing calli (B) in response to various phytohormones. The actin gene was used as an internal control. Each relative gene expression represents the average of three replicates with error bars representing SD.
KDWK sequence motif, the SAND domain has been suggested to mediate the DNA binding of these proteins.

In animals SAND domain-containing proteins are found in nucleus or cytoplasm, or have dual localization being present in both nucleus and cytoplasm [21-23]. In confirmation with this data pertaining to animal proteins, plant ultratapetal proteins with SAND domains are also demonstrated to localize to both the nucleus and the cytosol [15]. However, our study showed that CsULT1 is localized in nucleus (Figure 5). SAND domain containing proteins have been found to be transcriptionally active and are involved in regulation of gene expression [17]. We also investigated transcriptional activity of CsULT1 and our results demonstrated that it activated beta galactosidase enzyme proving that it is transcriptionally active (Figure 6).

Plant developmental, metabolic and stress pathways have been shown to be influenced and controlled by various phytohormones. In order to gain an insight about effect of different phytohormones on CsULT1 expression, we investigated effect of SA, JA, 2,4D and ABA on expression of CsULT1. Our results indicated that CsULT1 is induced in response to all the hormones studied. However, effect of SA, JA and 2,4-D was much more profound than ABA (Figure 7). Jasmonates are known elicitors of plant secondary metabolism and trigger extensive transcriptional reprogramming which ultimately leads to activation of whole metabolic pathway [24]. Induction of CsULT1 in response to JA might be part of this transcriptional activation scenario which as a final outcome leads to activation of carotenoid metabolic pathway. SA has been reported to induce expression of many carotenogenesis related genes [25]. Since CsULT1 is a probable regulator of carotenogenesis, its induction in response to SA is thus in confirmation with earlier reports. Auxin has been demonstrated to have a profound effect on stigma development [26]. Therefore enhanced expression of CsULT1 in response to 2,4-D treatment might suggest a parallel role of auxin in regulating stigma development via a vis apocarotenoid biosynthesis in Crocus. Although ABA treatment also enhanced expression of CsULT1, it was much less as compared to other hormones.

Several attempts have been made to establish Crocus transformation but no success has been achieved so far. Lack of transformation protocol is a limitation for functional characterization of genes in Crocus. The site of apocarotenoid biosynthesis is Crocus stigma, however, many of the pathway genes are expressed in callus also. Considering the limitation of transformation system, we transiently overexpressed CsULT1 in Crocus calli by particle bombardment in order to confirm its role in regulating carotenoid/apocarotenoid pathway. The expression of CsULT1 in transgenic calli was 2.5 fold as compared to vector control. This value is not good enough but since transformation in Crocus has not been established and callus is not the actual site of gene expression, the reported increase in expression can be considered as significant. Further, the expression analysis of a few carotenoid pathway genes was carried out and the results showed increase in expression level of PSY and PDS genes which catalyze initial rate limit steps of this pathway. This suggests role of CsULT1 in regulating carotenoid biosynthesis in Crocus. Till so far there is only one report on regulation of PSY gene expression by phytochrome biosynthesis (PIF1) which binds directly to PSY promoter and thereby regulates carotenoid accumulation during daily cycles of light and dark in mature plants [27]. Another member of AP2 gene family (RAP2.2) binds to of PSY promoter and is shown to modestly regulate the transcript levels of PSY and PDS in Arabidopsis [28]. Also BCH which is involved in conversion of beta carotene into zeaxanthin showed enhanced expression in transgenic calli (Figure 8a). In earlier reports CsCCD4b was considered responsible for cleaving zeaxanthin to produce apocarotenoids. However, later it was shown to cleave beta carotene at the 9,10 and/or the 9,10 positions, yielding beta-ionone. Recently a new isoform of CCDs (CsCCD2) was identified and isolated from Crocus and was shown to cleave zeaxanthin sequentially at 7,8 and 7,8 double bonds suggesting that CsCCD2 catalyzes the first dedicated step in crocin biosynthesis [20]. We investigated change in expression of CsCCD4b as well as CsCCD2 in transgenic calli and observed that their expression was enhanced around 4 and 5 fold respectively (Figure 8a). This suggests that apart from regulating synthesis of crocin and picrocrocin from zeaxanthin, CsULT1 also plays role in regulating biosynthesis of other apocarotenoids including beta ionone. Thus CsULT1 might regulate expression of more than one members of CCD gene family. Except for PIF1 and RAP2.2 no other transcription factors have been identified till so far which regulate expression of genes involved in carotenogenesis in plants. Therefore the present work will form a platform for enhancing our knowledge on regulation of this important pathway.

Carotenoids are involved in many biological functions including plastid biogenesis, photosynthesis, photomorphogenesis etc. Carotenoid metabolic pathway is also linked with many other pathways like ABA and GA biosynthesis. Therefore, carotenoid metabolism might be regulated at multifaceted levels in plants. Further, because of this close coordination of many pathways, content and composition of carotenoids is important. Thus biosynthesis of carotenoids and their turn-over to produce apocarotenoids needs to be tightly regulated in order to maintain their steady levels in plants. Hormones are known to play key roles in regulating various metabolic pathways. They also help in coordinating
interplay between various pathways. Our results also demonstrated effect of phytohormones on CsULT1 expression. In order to further our understanding on mechanism of regulation of carotenoid biosynthesis by CsULT1, we also investigated effect of various phytohormones on carotenoid pathway genes particularly those which showed enhanced expression in CsULT1 over-expressing calli. Our results indicated that expression of all the genes studied was enhanced in response to SA, JA and 2, 4-D. however, there was no significant increase in expression of these genes in response to ABA (Figure 8b). Earlier also, SA and JA have been shown to affect expression of many carotenogenic pathway genes [29]. If considered individually, PSY was induced more in response to SA and JA while PDS expression increased more under the influence of SA and 2,4-D. BCH showed much higher response to 2,4-D than other hormones while CCD4b and CCD2 showed more or less similar trend in expression in response to all the hormones except ABA. ABA did not cause any significant effect on expression of carotenoid pathway genes. Earlier, exogenous application of ABA was shown to repress transcript levels of many chloroplast genes [30]. It was also found to reduce chlorophyll levels and also endogenous ABA levels [31]. This might be the reason that we did not observe any significant increase in expression levels of carotenoid pathway genes in response to ABA. Thus in general, change in the expression of these genes followed the same trend as that of CsULT1. This suggests that these phytohormones might play a role in mediating effect of CsULT1 on regulation of carotenoid metabolism. Further, different genes showed varied induction levels in response to different hormones which might be because of the fact that control of plant metabolic pathways is a complex phenomenon and network of cross-communicating hormone signaling pathways are involved so as to maintain overall metabolite homeostasis within the plant system.

Taken together, these results suggest that CsULT1 plays role in diverting metabolic flux towards enhanced production of apocarotenoids. Recently ULT in Arabidopsis has been shown to be involved in gynoecium development and patterning [17]. In Crocus, apocarotenoid biosynthesis is congruent with stigma development. This indicates that CsULT1 might have a parallel role in regulating Crocus gynoecium (stigma) development via a vis apocarotenoid biosynthesis. However, this needs to be further verified by experimentation in Crocus itself or in other alternative plant system.

Conclusions
In this study an ultrapetala transcription factor from Crocus, CsULT1, was identified, cloned and characterized. The CsULT1 transcript was expressed more in stigma till flower anthesis. Application of phytohormones like 2,4D, JA and SA enhanced CsULT1 expression. The gene is nuclear localized and is transcriptionally active. Moreover, ectopic expression of this gene altered expression of some important genes of carotenoid/apocarotenoid pathway confirming that CsULT1 plays a regulatory role in Crocus apocarotenoid biosynthesis. Furthermore, the hormones which affected expression of CsULT1 were also shown to enhance expression of pathway genes indicating their role in mediating regulatory role of CsULT1.

Additional files

Additional file 1: Table S1. List of primer sequences.
Additional file 2: Figure S1. Melt curve depicting single peak in qRT-PCR.
Additional file 3: Figure S2. qRT-PCR analysis of different Crocus transcription factors.
Additional file 4: Figure S3. Nucleotide and deduced amino acid sequence of CsULT1.

Abbreviations
ULT: Ultrapetala; PSY: Phytoene synthase; PDS: Phytoene desaturase; CCD: Carotenoid cleavage dioxygenase; BCH: Beta cyclohydroxylase; MEP: Methylerythritol phosphate.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
NA: designed and performed experiments and wrote the manuscript; DJ: performed experiments; RAV: conceptualized experiments and did critical revision of manuscript. All authors read and approved the final manuscript.

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Author details
1Plant Biotechnology Division, CSIR- Indian Institute of Integrative Medicine, Jammu J&K 180001, India. 2National Institute of Plant Genome Research, Aruna Asaf Ali Marg, New Delhi 110067, India. 3National Institute of Plant Gene Research, Aruna Asaf Ali Marg, New Delhi 110067, India. 4Medicinal Chemistry Division, CSIR- Indian Institute of Integrative Medicine, Canal Road, Jammu J&K 180001, India.

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