

Identification of alleles of carotenoid pathway genes important for zeaxanthin accumulation in potato tubers

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Abstract We have investigated the genetics and molecular biology of orange flesh colour in potato (*Solanum tuberosum* L.). To this end the natural diversity in three genes of the carotenoid pathway was assessed by SNP analyses. Association analysis was performed between SNP haplotypes and flesh colour phenotypes in diploid and tetraploid potato genotypes. We observed that among eleven beta-carotene hydroxylase 2 (*Chy2*) alleles only one dominant allele has a major effect, changing white into yellow flesh colour. In contrast, none of the lycopene epsilon cyclase (*Lcye*) alleles seemed to have a large effect on flesh colour. Analysis of zeaxanthin epoxidase (*Zep*) alleles showed that all (diploid) genotypes with orange tuber flesh were homozygous for one specific *Zep* allele. This *Zep* allele showed a reduced level of expression. The complete genomic sequence of the recessive *Zep* allele, including the promoter, was determined, and compared with the sequence of other *Zep* alleles. The most striking difference was the presence of a non-LTR retrotransposon sequence in intron 1 of the recessive *Zep* allele, which was absent in all other *Zep* alleles investigated. We hypothesise that the presence of this large sequence in intron 1 caused

the lower expression level, resulting in reduced *Zep* activity and accumulation of zeaxanthin. Only genotypes combining presence of the dominant *Chy2* allele with homozygosity for the recessive *Zep* allele produced orange-fleshed tubers that accumulated large amounts of zeaxanthin.

Keywords *Solanum tuberosum* · SNP analysis · Allele mining · Orange flesh · Carotenoids · Zeaxanthin epoxidase

Abbreviations

Chy2	Beta-carotene hydroxylase 2
Lcye	lycopene epsilon cyclase
Zep	Zeaxanthin epoxidase
AMD	Age-related macular degeneration
eQTL	Expression quantitative trait locus
POCI	Potato oligo chip initiative
SNP	Single nucleotide polymorphism
CAPS	Cleaved amplified polymorphic sequence
BAC	Bacterial artificial chromosome
PGSC	Potato genome sequencing consortium
SIFT	Sorting intolerant from tolerant
PolyPhen	Polymorphism phenotyping
qRT-PCR	Quantitative real-time polymerase chain reaction
LTR	Long terminal repeat

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Introduction

Flesh colour in most tetraploid potato cultivars ranges from white via cream and yellow to dark yellow. This yellow

colour is caused by the presence of specific carotenoids. A small number of cultivars have red or blue/purple flesh, caused by the presence of anthocyanins.

The main carotenoids present in cultivated potato are lutein, violaxanthin, zeaxanthin and antheraxanthin (Breithaupt and Bamedi 2002; Brown et al. 1993; Iwanzik et al. 1983; Nesterenko and Sink 2003). Beta-carotene, the precursor of vitamin A, is almost absent in *S. tuberosum* genotypes or closely related *Solanum* species (Breithaupt and Bamedi 2002). Carotenoids are recognised as important health promoting ingredients of the human diet. Some have antioxidant properties, and are supposedly beneficial in preventing cancer, cardiac disease, and eye diseases (Krinsky et al. 2004). Lutein and zeaxanthin are thought to be important in the human diet to prevent age-related macular degeneration (AMD; Moeller et al. 2006; Seddon et al. 1994; Snodderly 1995). Lutein and zeaxanthin are components of the *macula lutea* in the human eye (Handelman et al. 1988), protecting the retina against damaging irradiation, but they have to be replenished constantly. As humans can not produce lutein and zeaxanthin themselves they have to be consumed by eating carotenoid-rich plant products. Lutein is present in high amounts in dark green leafy vegetables such as spinach and kale. Zeaxanthin, however, is less abundant in most vegetables (Sommerburg et al. 1998).

In tetraploid potato lutein is present in relatively large amounts, whereas zeaxanthin is present in lower amounts (Breithaupt and Bamedi 2002; Nesterenko and Sink 2003). However, some *Solanum* species closely related to *S. tuberosum* have high zeaxanthin content (Andre et al. 2007). These are known as ‘Papa Amarilla’ because of their deep yellow or orange-fleshed tubers. These landraces grown by indigenous farmers in the Andean region belong to the diploid species *S. stenotomum*, *S. goniocalyx* and *S. phureja* (Brown et al. 2007; Burgos et al. 2009). Brown et al. (2007) and Brown (2008) observed a relationship between ploidy level and total carotenoid content in 38 native South American cultivars. Significantly higher mean levels of total carotenoids were observed in diploid cultivars compared with tetraploid cultivars. Morris et al. (2004) describe a diploid high carotenoid-accumulating *S. phureja* accession (DB375\1, or ‘Inca Dawn’) that predominantly contains zeaxanthin, but has a lower yield than tetraploid *S. tuberosum* cultivars (Bradshaw and Ramsay 2005). Kobayashi et al. (2008) bred a diploid potato variety with orange flesh and very high zeaxanthin content. This variety was derived from *S. phureja*. It would be interesting to obtain high-yielding orange-fleshed tetraploid potato cultivars to aid in the recommended daily uptake of zeaxanthin, as potatoes and potato products constitute a considerable part of the human diet in the Western world.

Yellow flesh colour in potato is mainly dependent on the presence of a dominant allele (Fruwirth 1912) at the *Y* (Yellow) locus. The *Y* locus has been mapped on chromosome 3 of potato by Bonierbale et al. (1988). The most likely candidate for the gene involved in yellow flesh colour is beta-carotene hydroxylase (abbreviated to *Bch* or *Chy2*; Brown et al. 2006). This gene has been mapped at the same position as the *Y* locus (Thorup et al. 2000).

Until now, the gene(s) responsible for the orange tuber flesh colour in diploid *Solanum* species are unknown. Brown et al. (1993) observed progeny with orange flesh colour and high levels of zeaxanthin in a hybrid population of *S. phureja*-*S. stenotomum*. They suggested that the orange phenotype was caused by a dominant *Or* allele at or close to the *Y* locus on chromosome 3 of potato. However, this was not corroborated by later research, as Brown (2008) reported. Lack of transmissibility outside the immediate ‘Papa Amarilla’ gene pool negated the hypothesis that the expression of *Or* was consistent with a strong dominant monogenic inheritance. In cauliflower an *Or* gene was cloned, responsible for orange-coloured curds (Lopez et al. 2008). This gene was found not to be involved in the carotenoid biosynthesis pathway, but to control chromoplast differentiation, resulting in the sequestering of large amounts of carotenoids.

This paper describes DNA polymorphisms among haplotypes of three candidate genes involved in the carotenoid pathway in monoploid, diploid and tetraploid potato genotypes, and explains the inheritance of yellow and orange potato tuber flesh colour.

Materials and methods

Plant materials

For sequence analyses DNA was used from five monoploid potato genotypes: 7322 (H7322 or AM79.7322, originally from G. Wenzel, Institut für Genetik, Grünbach, Germany, see: De Vries et al. 1987; Hovenkamp-Hermelink et al. 1988), M5 and M38 (851-5 and 851-38, Uijtewaal 1987), M47 and M133 (1022 M-47 and 1022 M-133, Hoogkamp et al. 2000). DNA from 20 monoploid *S. phureja* and *S. chacoense* clones was obtained from Richard Veilleux (Blacksburg, Virginia, USA, see Lightbourn and Veilleux 2007). DNA was isolated from eleven diploid genotypes: C (USW5337.3, Hanneman and Peloquin 1967), E (77.2102.37, Jacobsen 1980), RH88-025-50 and RH90-038-21 (Park et al. 2005), RH89-039-16 and SH83-92-488 (Roupe van der Voort et al. 1997; Van Os et al. 2006), 87.1024/2 and 87.1029/31 (Jacobsen et al. 1989), G254 (Uijtewaal et al. 1987), R5 (EJ92-6486-19, from cross 87.1024/2 × EJ91-6104-19) and 413 (transformant of

interdihaploid H²260; Binding et al. 1978; De Vries-Uijtewaal et al. 1989). Three diploid orange-fleshed genotypes were analysed: cultivars ‘Papa Pura’ and ‘Andean Sunrise’ (provided by Agrico Research BV), and *S. phureja* ‘Yema de Huevo’ (obtained from Enrique Ritter, Vitoria, Spain, see Ritter et al. 2008). Two diploid populations were analysed in which orange-fleshed progeny segregated: the C×E population (Jacobs et al. 1995) and the IvP92-030 population (cross G254 × SUH2293, from Ronald Hutten, Lab. of Plant Breeding, Wageningen University). Furthermore, orange-fleshed diploid genotype IvP01-84-19 from the cross 96-4622-20 × IvP92-027-9 (Ronald Hutten) was included. Additionally, a set of 225 tetraploid cultivars was used (D’hoop et al. 2008; D’hoop 2009). In this set no genotypes with orange-fleshed tubers were present. Flesh colour of the tetraploids was determined in a field experiment in 2006 (D’hoop 2009). Flesh colour values were on an ordinal scale ranging from 4 (=white) to 9 (=orange) according to the Dutch Catalogue of Potato Varieties (www.nivap.nl).

DNA isolation

Genomic DNA from the monoplloid and diploid genotypes was isolated from leaf tissue according to the CTAB method from Rogers and Bendich (1988). DNA from the tetraploid cultivars was isolated according to Van der Beek et al. (1992).

PCR amplification and sequencing

Amplicons for sequencing were generated from 50 ng genomic DNA template. PCR amplifications were performed in 50 or 25 µl reactions using 1 u of Taq polymerase, 1× reaction buffer, 200 nM dNTP and 250 nM of each primer. Standard cycling conditions were: 4 min initial denaturation at 94°C, followed by 35 cycles of 30 s denaturation at 94°C, 30 s annealing at 55°C and 30 s to 1 min extension at 72°C. Reactions were finished by 7 min incubation at 72°C. For CAPS marker analysis 30 cycles were used. Most PCRs were performed with SuperTaq Polymerase buffer and enzyme (Applied Biosystems). PCR products were examined for quality on ethidium bromide-stained agarose gels. PCR products were directly sequenced on ABI377 or ABI3700 sequencers at Greenomics (Wageningen University and Research Centre) using the dideoxy chain-termination method and ABI PRISM Reaction Kit. One or both of the amplification primers were used as sequencing primers. For SNP analysis of the *Chy2* gene primers CHY2ex4F (5'-CCATAGACCAAGAGAAGGAC C-3') and Beta-R822 (5'-GAAAGTAAGGCACGTTGGCA AT-3') were used. For SNP analysis of the *Lcye* gene primers AWLCYe1 (5'-AAAAGATGCAATGCCATTCGAT-3')

and AWLCYe2 (5'-GAAATACTCGGGGTACTTGAAC-3') were used. For SNP analysis of the *Zep* gene primers AWZEP9 (5'-GTGGTCTTGAGAATGGACAAC-3') and AWZEP10 (5'-CACCAGCTGGTTCATTGTAAAA-3') were used. As CAPS marker for *Chy2* the 308-bp CHY2ex4F + Beta-R822 PCR product was cleaved with *AluI*. The 163-bp fragment was indicative for the presence of *Chy2* allele 3. As CAPS marker for *Lcye* the AWLCYe1 + AWLCYe2 PCR product was digested with *SsiI*, which distinguished allele 2 from alleles 1 and 3. By digesting the AWLCYe1 + AWLCYe2 PCR product with *HpyCH4IV* allele 1 could be distinguished from alleles 2 and 3.

SNP analysis and bioinformatics

PCR reactions included a mixture of templates reflecting the different alleles in DNA samples from heterozygous diploid and tetraploid genotypes. Trace files from directly sequenced PCR products were analysed for secondary peaks, indicative for SNPs, with the Vector NTI software package from Invitrogen. Homology searches were performed at the NCBI webpage (<http://www.ncbi.nlm.nih.gov/>), the TGI webpage (<http://compbio.dfci.harvard.edu/tgi/cgi-bin/tgi/Blast/index.cgi>), the SGN webpage (<http://sgn.cornell.edu/>), and the PGSC webpage (restricted access; <http://bacregistry.potatogenome.net/pgscreg/main.py>). To study possible effects of amino acid changes on functionality of the protein the programs SIFT (Sorting Intolerant From Tolerant, see <http://sift.jcvi.org/> [Ng and Henikoff 2006]) and PolyPhen (Polymorphism Phenotyping, see <http://coot.embl.de/PolyPhen/>) were used. Classification of the non-LTR retrotransposon sequence in *Zep* allele 1 was performed at <http://www.girinst.org/RTphylogeny/RTclass1>.

Genetic mapping and QTL analysis in the diploid C×E population

A C×E genetic map using 94 C×E progeny was made based on an earlier version of the map (Celis-Gamboa 2002) with additional SNP markers (Anithakumari et al. 2010) using mapping software Joinmap 4.0[®] (Van Ooijen 2006). QTL analysis of quantitative data was performed using the software package MapQTL[®] Version 5.0 (Van Ooijen 2004).

Cloning of *Zep* promoter sequence

The promoter sequence of *Zep* allele 1 was obtained by Genome Walking using the Universal GenomeWalker kit (Clontech) and the BD Advantage 2 PCR Enzyme System (BD Biosciences). DNA from diploid genotype R5 (homozygous for *Zep* allele 1) was used as template. Four libraries were made using *DraI*, *EcoRV*, *StuI* and *ScaI* enzymes. The

first PCR was performed with primer AP1 and gene-specific primer AWZEPGW1 (5'-TTCTGTGGAACCTTCAAATC ACCGTTA-3'). The nested PCR was performed with primer AP2 and gene-specific primer AWZEPGW2 (5'-GCCCATTTCCAAGCTCCTACAAGGTA-3'). The *DraI*- and *StuI*-libraries yielded a 1.1 and 1.8-kb PCR fragment, respectively. The 1.8-kb PCR fragment of the *StuI* library was sequenced, and proved to contain a *DraI* restriction site at the expected position.

Cloning intron 1 of *Zep* allele 1

To obtain the intron 1 sequence of *Zep* allele 1 primers AWZEP25 (5'-CTGGCTGCATCACTGGTCAAAG-3') and AWZEP20 (5'-TCATTTCATAATTGTATCCTCCC-3') were used. The Expand High Fidelity PCR System (Roche Applied Science) was used to obtain the 4.7-kb PCR fragment. This fragment was cloned into pGEM-Teasy (Promega). Plasmid DNA was isolated using the Promega Wizard Plus minipreps DNA Purification system. DNA from three independent colonies was first sequenced using the T7, AWZEP25 and AWZEP20 primers, and subsequently with primers designed on the obtained sequences.

Measurement of carotenoids

From 94 progeny of the diploid C×E population carotenoids were extracted and analysed by HPLC with photodiode array (PDA) detection, according to the protocol described by Bino et al. (2005). In short, 0.5 g FW of ground and frozen tuber material was extracted with methanol/chloroform/1 M NaCl in 50 mM Tris (pH 7.4) in a ratio of 2.5:2:2.5 (v:v:v) containing 0.1% butylated hydroxytoluene (BHT). After centrifugation, the samples were re-extracted with 1 ml chloroform (+BHT). The chloroform fractions were combined, dried under a flow of N₂ gas and taken up in ethyl acetate containing 0.1% BHT. Carotenoids present in the extracts were separated by HPLC using an YMC-Pack reverse-phase C30 column and analysed by PDA detection with wavelength range set from 240 to 700 nm. Eluting compounds were identified based on their absorbance spectra and co-elution with commercially available authentic standards (neoxanthin, violaxanthin, antheraxanthin, lutein, zeaxanthin, β -cryptoxanthin, ϵ -carotene, α -carotene, β -carotene, ζ -carotene, δ -carotene, polycopene and all-trans lycopene. Limit of detection was about 5 μ g per 100 g FW and technical variation (6 independent extractions and analyses of the same tuber powder) was less than 8%.

In addition to the measurement of individual carotenoids yellowness of the tuber flesh of the 94 C×E progeny was determined by spectrophotometry. The same carotenoid extraction used for HPLC analysis was measured with a

Perkin Elmer UV/MS Spectrometer Lambda 10. The peak area from 380 to 515 nm was determined with the UV WinLab software from Perkin Elmer using the 525–580 nm measurement as baseline. The peak area in the yellow spectrum is referred to as absorbance between 380 and 515 nm.

Quantitative RT-PCR

Total RNA of 23 selected genotypes of the C×E population was isolated from mature tubers as described by Bachem et al. (1998). mRNA was purified using the RNeasy mini kit (Qiagen) and reverse transcribed using the iScript cDNA synthesis kit from Bio-Rad. Relative expression level of the *Zep* locus was determined by real-time quantitative reverse transcriptase PCR (qRT-PCR) on an iQ detection system (Bio-Rad) according to the Bio-rad iQ SYBR Green Supermix protocol. The primer sequences used for the analysis were StZEP_RT_F (5'-AAGTGCCGAGTCAGGAAGCC-3') from exon 7 and StZEP_RT_R (5'-CAAGTCCGACGCCAAGATAAGC-3') from exon 8. Potato elongation factor 1- α (*EF1 α*) primers were used for relative quantification (Nicot et al. 2005). Relative quantification of the target RNA expression level was performed using Bio-rad iQ5 analysis program.

Accession numbers

Sequence data from this article can be found in the EMBL/GenBank data libraries under accession numbers HM013963 (potato *Chy2* genomic sequence), HM013964 (potato *Zep* allele 1 genomic sequence) and HM013965 (potato *Zep* allele 2 genomic sequence). The potato *Lcye* genomic sequence is available in the Third Party Annotation Section of the DDBJ/EMBL/GenBank databases under the accession number TPA: BK007065. The sequence of the PCR fragment obtained with primers AWLCYe1 and AWLCYe2 for allele 4 is available under accession number HM011105. Accession numbers for the BAC sequences are AC216345 (tomato BAC LE_HBa-11D12), AC238165 (potato BAC RH091F11), AC238104 (RH071N14), AC238398 (RH196E12), AC238240 (RH132J19) and AC215383 (tomato BAC C02HBa0104A12).

Results

Allelic variation for the *beta-carotene hydroxylase 2* gene (*Chy2*)

We started our investigation of the genetic requirements for orange potato tuber flesh by analysing whether

orange-fleshed diploid potato genotypes contain different *Chy2* alleles than white- or yellow-fleshed tetraploid cultivars. *Chy2* was observed to be an important gene involved in tuber flesh colour in two separate studies performed on diploid populations (Brown et al. 2006; Kloosterman et al. 2010). RNA expression analysis using the 44 K POCI array (Kloosterman et al. 2008) resulted in the identification of an eQTL for yellow tuber flesh colour on potato chromosome 3 at a similar position as the *Chy2* gene. Further analysis indicated that in the diploid C×E population the parent C-specific allele of the *Chy2* gene was correlated with yellow flesh. This allele shows higher expression than the other two alleles segregating in the C×E population (Kloosterman et al. 2010). Goo et al. (2009) also observed a higher expression level of *Chy2* associated with yellow flesh colour in a small number of potato cultivars.

In order to study allelic variation for the *Chy2* gene in tetraploids we first determined the complete genomic sequence of the *Chy2* allele in monohaploid ($2n = x = 12$) potato genotype 7322, as only mRNA and EST sequences were known for the potato *Chy2* gene. The obtained 2,255-bp genomic sequence (Fig. 1) contains seven exons. Next, we performed direct sequencing analyses of a PCR fragment obtained with primers CHY2ex4F and Beta-R822, spanning exon 4, intron 4 and exon 5 of the *Chy2* gene (Fig. 1). We used DNA of four additional potato monohaploids, 20 *S. phureja* and *S. chacoense* monohaploids, and 11 diploids. From these sequences in total eight different haplotypes could be determined (Supplemental Table S1; alleles 1–7 and 11).

We observed a correlation between presence of a single haplotype—allele 3—and yellow flesh colour in a number of diploids. Heterozygosity for allele 3 is sufficient for yellow flesh colour, indicating this is a dominant allele. This observation corroborates the results of genetic analysis of the *Chy2* (or *Bch*) gene by Brown et al. (2006). Furthermore, we observed that SNP 142C distinguished *Chy2* allele 3 from all other haplotypes. As this SNP is unique to one haplotype it is a so-called ‘haplotype tag SNP’ (Johnson et al. 2001), hereafter referred to as ‘tag

SNP’. Allele 3 is most probably identical to the dominant allele B described by Brown et al. (2006), because sequencing of allele 3 showed the presence of allele B-specific primer sequence YellowF1. This allele is considered to be the dominant Y allele at the *Yellow* (*Y*) locus first postulated by Fruwirth (1912).

Next, we performed a SNP analysis on the DNA of a set of 225 tetraploid potato cultivars (D’hoop 2009). This set aims to represent the most important potato cultivars of the last 150 years in terms of acreage and/or value as progenitor, mainly from Europe, but also from the USA, Canada, and some other continents. The same PCR fragment that was analysed for the monohaploids and diploids was amplified in the tetraploids. Direct sequencing was performed, which resulted in the discovery of three additional alleles (Supplemental Table S1; alleles 8, 9, 10). The dosage of SNP 142C (i.e. allele 3) was determined from the sequence trace files. Dosage of allele 3 could also be estimated by using a CAPS marker assay (Supplemental Fig. S1), in which of all *Chy2* alleles only allele 3 yielded a 163-bp fragment. The allele 3 dosage was related to the flesh colour value (Table 1). A flesh colour value of 5.5 or lower represents white flesh, whereas a flesh colour value higher than 5.5 indicates yellow flesh. As is shown in Table 1 presence or absence of *Chy2* allele 3 is correlated with flesh colour: the group of cultivars lacking allele 3 have a mean value of 5.1 (white flesh), whereas the cultivars simplex, duplex, triplex or quadruplex for allele 3 have a mean value higher than 6 (yellow flesh). The data suggest a dosage effect of allele 3 as the quadruplex genotypes have the highest mean value. However, this group consists of only two genotypes. Although there is a clear correlation between presence of *Chy2* allele 3 and yellow flesh, the intensity of the yellow flesh colour shows considerable variation within the different dosage groups, as shown in Fig. 2. This was also observed by Brown et al. (2006).

To evaluate if *Chy2* alleles other than allele 3 have a (small) influence on flesh colour the *Chy2* allele composition was determined for 199 of the 225 tetraploid potato

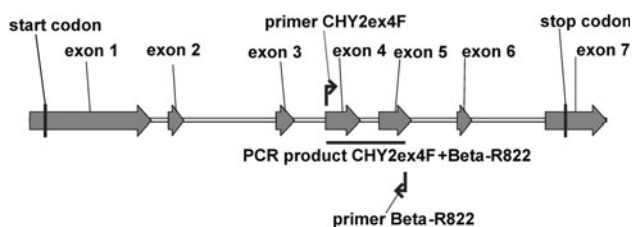


Fig. 1 Schematic representation of the genomic sequence of the potato *beta-carotene hydroxylase 2* (*Chy2*) gene. The 2,255-bp of allele 1 of monohaploid 7322 is shown. The PCR product analysed for the presence of SNPs is indicated

Table 1 Relation between the number of *Chy2* alleles and tuber flesh colour

<i>Chy2</i> allele 3 dosage	Mean tuber flesh colour value	Flesh colour	Number of genotypes
0× (nulliplex)	5.0	White	69
1× (simplex)	6.3	Yellow	71
2× (duplex)	6.5	Yellow	46
3× (triplex)	6.6	Yellow	11
4× (quadruplex)	7.4	Yellow	2

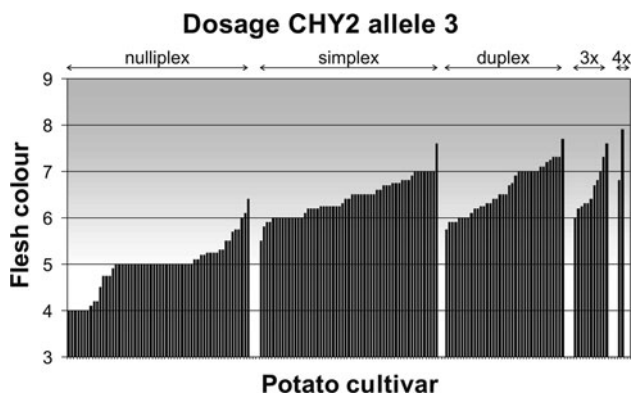


Fig. 2 Variation in flesh colour value in classes of tetraploid potato genotypes with 0 \times , 1 \times , 2 \times , 3 \times , and 4 \times *Chy2* allele 3. Flesh colour value equal to or below 5.5 is considered to correspond to white flesh, values above 5.5 are indicative of yellow flesh

genotypes by analyzing tag SNPs. Large differences in allele frequency were observed. Four major alleles were observed: alleles 5 (35%), 3 (26%), 1 (20%) and 2 (13%). Four minor alleles were observed: alleles 6 (4%), 10 (2%), 8 (0.8%) and 9 (0.3%). Alleles 4, 7 and 11 were not observed in the tetraploid *S. tuberosum* cultivars and seem to be restricted to diploid germplasm. Minor allele 6 has been present in the *Solanum tuberosum* gene pool for a long time, as it is observed in nine cultivars released before 1900. Minor alleles 8, 9 and 10 seem to be novelties in the *S. tuberosum* gene pool, because these alleles are only present in cultivars released to the market after 1960, directly descending from backcross introgression material with late blight or cyst nematode resistance. For example, allele 10, containing an indel (1-nt deletion) in the analysed PCR fragment, is present in a number of cultivars derived from VTN 62-33-3 (1962), suggesting that this might be an *S. vernei* allele (see Kort et al. 1972).

None of the alleles 1, 2, 5, 6, 8, 9 or 10 were related to yellow flesh colour. Furthermore, none of these alleles influenced flesh colour value within the white/creamy flesh colour class, nor within the yellow flesh colour class. Thus, the variation in intensity of yellow flesh colour can not be explained by the composition of the other *Chy2* alleles in the simplex, duplex and triplex allele 3 groups.

Chy2 allele composition was determined in diploid orange-fleshed genotypes ‘Papa Pura’, ‘Andean Sunrise’, ‘Yema de Huevo’, IvP92-030-11 and IvP01-84-19. Although they all contained one *Chy2* allele 3, they differed in the other allele, and no novel allele absent from the tetraploid gene pool was discovered. Therefore, we concluded that other genes—possibly involved in the carotenoid biosynthetic pathway—influenced the intensity of the yellow flesh colour.

Allelic variation for the *lycopene epsilon cyclase* gene (*Lcye*)

The lycopene epsilon cyclase gene product is required for the synthesis of α -carotene, the precursor of lutein (see Tanaka et al. 2008). Silencing of the *Lcye* gene resulted in a significant increase in the beta-carotenoids (Diretto et al. 2006). An increase in the level of beta-carotene and zeaxanthin is expected to result in a darker yellow flesh.

To be able to analyse allelic variation for the *Lcye* gene in potato we needed information on the genomic sequence of the *Lcye* gene. Tomato BAC LE_HBa-11D12 (from chromosome 12) contains the tomato homologue of *Lcye*. Using this sequence four potato BAC clones from diploid genotype RH89-039-16 containing the potato *Lcye* homologue were retrieved from the Potato Genome Sequencing Consortium (PGSC) database: RH091F11, RH071N14, RH196E12 and RH132J19. The 7,000-bp genomic sequence of the *Lcye* gene contains 11 exons (Fig. 3). Primers AWLCYe1 and AWLCYe2 were designed, which amplified a fragment spanning exon 7 to exon 9 of the *Lcye* genomic sequence (Fig. 3). PCR fragments from the monoplids and diploids were directly sequenced, and 5 different alleles could be observed (Supplemental Table S2). Potato BACs RH091F11 and RH071N14 contained allele 4, whereas two BACs from the homologous chromosome (RH196E12-4 and RH132J19-7) contained allele 1 of diploid genotype RH89-039-16.

Two alleles (allele 2 and allele 5) contain a T at SNP position 545. This nucleotide causes a change of amino acid 401 of the *Lcye* protein from S (serine) to F (phenylalanine). To analyse the effect of this amino acid change we consulted two software programs: SIFT and PolyPhen. According to the SIFT program an F at position 401 is not tolerated, and according to the PolyPhen program it is possibly damaging. Diploid genotype C contains *Lcye* alleles 2 and 3, while genotype E contains *Lcye* alleles 1 and 2. Thus, C and E have *Lcye* allele 2 in common. Therefore, 25% of the progeny of a cross between these

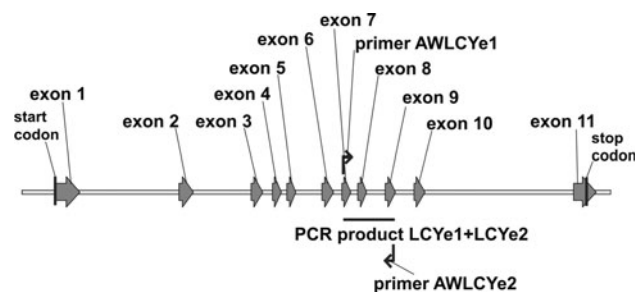


Fig. 3 Schematic representation of the genomic sequence of the potato *lycopene epsilon cyclase* (*Lcye*) gene. A 7,000-bp sequence from BAC RH091F11 from diploid genotype RH89-039-16 is shown. The PCR product analysed for the presence of SNPs is indicated

genotypes is expected to be homozygous for allele 2. Some of the progeny of the C×E cross have a much higher flesh colour value than the yellow parent C, i.e. some have orange flesh. To investigate whether homozygosity of *Lcye* allele 2 leads to a difference in flesh colour 94 C×E progeny plants were analysed for their *Lcye* allele composition by CAPS marker assays. Data were used to localize the *Lcye* gene on the C×E linkage map. As expected, *Lcye* mapped to a position on chromosome 12 close to the STM2028 microsatellite marker and the SUS4 gene at the Southern distal end of the chromosome.

Dosage of *Lcye* allele 2 and dosage of *Chy2* allele 3 were plotted against flesh colour value (Fig. 4). This figure shows a strong correlation between presence of *Chy2* allele 3 and a high value for flesh colour, whereas dosage of *Lcye* allele 2 does not seem to have an influence on flesh colour. This suggests that a different gene than *Lcye* must have an influence on intensity of yellow flesh colour.

Allelic variation for the zeaxanthin epoxidase gene (*Zep*)

Another candidate gene for orange tuber flesh in potato is the *Zep* gene. *Zep* is involved in the conversion of zeaxanthin into antheraxanthin, and in the conversion of antheraxanthin into violaxanthin (Tanaka et al. 2008). Silencing of the *Zep* gene in potato resulted in transformants with higher zeaxanthin levels, and increased total carotenoid contents (Römer et al. 2002). Similarly, Morris et al. (2004) observed an inversed trend between the level of *Zep* transcript and tuber carotenoid content in a range of potato germplasm.

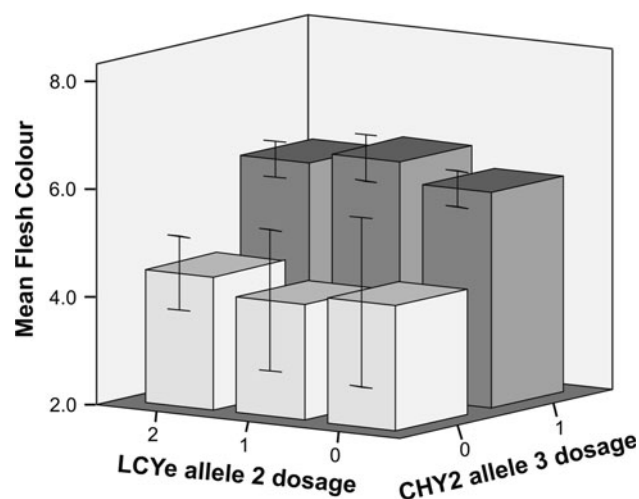


Fig. 4 Relation between *Chy2* allele 3 dosage, *Lcye* allele 2 dosage, and phenotypic value of tuber flesh colour in the diploid C×E mapping population. Error bars ± 2 SE

A genomic sequence of the tomato *Zep* gene was found to be present in BAC C02HBa0104A12.1, anchored to tomato chromosome 2 (Supplemental Fig. S2), which is in agreement with the map position of the pepper (*Capsicum*) *Zep* gene on chromosome 2 (Thorup et al. 2000). Using the tomato BAC sequence and potato *Zep* cDNA sequence DQ206629 primers were designed allowing amplification and sequencing of the complete potato *Zep* gene (Fig. 5). Using primer combination AWZEP9 + AWZEP10 five different alleles could be distinguished in the monoploid and diploid *S. tuberosum* genotypes (Supplemental Table S3, alleles 1–5). The AWZEP9 + AWZEP10 PCR product, spanning exon 3 to exon 5, showed the presence of a relatively large indel in intron 4. In *Zep* allele 1 a sequence of 49 bp is absent, which is present in alleles 2, 3, 4 and 5. Therefore, *Zep* allele 1 could be distinguished from the other alleles by gel electrophoresis. The AWZEP9 + AWZEP10 PCR product of allele 1 is 535 bp long, whereas this PCR product is 584 bp for the other alleles (Supplemental Fig. S3b).

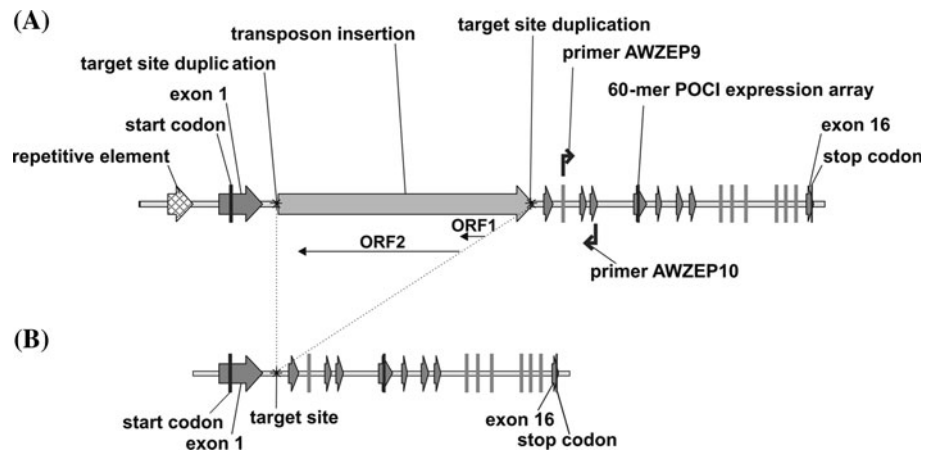
Recessive inheritance of orange tuber flesh

Zep allele composition was determined in diploid orange-fleshed genotypes ‘Papa Pura’, ‘Andean Sunrise’, ‘Yema de Huevo’, IvP92-030-11 and IvP01-84-19. All five genotypes proved to be homozygous for *Zep* allele 1. As these five genotypes are not closely related to each other this suggests the involvement of *Zep* allele 1 in the orange flesh phenotype.

Genetic evidence for the involvement of *Zep* allele 1 in orange flesh colour was obtained from cosegregation in the diploid IvP92-030 population (progeny of the cross between diploids G254 and SUH2293). This population was analysed for both *Chy2* and *Zep* allele compositions (Supplemental Fig. S3). Parent G254 contained *CHY2* alleles 2 and 6, while parent SUH2293 contained *Chy2* alleles 3 and 5. Progeny with allele combinations 2 + 3, 2 + 5, 3 + 6 and 5 + 6 were observed (Supplemental Fig. S3a). Both parents G254 and SUH2293 contained *Zep* alleles 1 and 2. Progeny with allele combinations 1 + 1, 1 + 2 and 2 + 2 were obtained in numbers compatible with the expected 1:2:1 ratio (Supplemental Fig. S3b). Only progeny plants IvP92-030-9 and IvP92-030-11, containing *Chy2* allele 3 and homozygous for *Zep* allele 1, showed the orange-fleshed phenotype. This suggests a model in which presence of dominant *Chy2* allele 3 and homozygosity for recessive *Zep* allele 1 are required to obtain an orange-fleshed potato.

To investigate this further progeny of the C×E cross was analysed for *Zep* allele composition. Both C and E parents contain *Zep* alleles 1 and 2. A CAPS marker was

Fig. 5 Schematic representation of the genomic sequence of the potato *zeaxanthin epoxidase* (*Zep*) gene. **a** 11,000-bp sequence of allele 1, including promoter and coding region; **b** 6,008-bp sequence of allele 2, including promoter and coding region



developed to easily distinguish both *Zep* alleles. For this, PCR product AWZEP9 + AWZEP10 was digested with the enzyme *Hin6I*. The PCR product of *Zep* allele 1 remained undigested, whereas the PCR product of allele 2 was digested into fragments of 439 and 145 bp. Ninety-four C×E progeny were analysed with this CAPS marker. The *Zep* gene was mapped in the C×E population on chromosome 2 in a similar position as the one on tomato chromosome 2 (Supplemental Fig. S2). Dosage of *Zep* allele 1 and dosage of *Chy2* allele 3 were related to flesh colour value (Fig. 6). This figure shows that homozygosity of *Zep* allele 1 in combination with presence of *Chy2* allele 3 results in a significantly higher mean flesh colour value. A QTL analysis for absorbance in the yellow spectrum in the C×E population resulted in a highly significant QTL on chromosome 2, on the same position as the *Zep* gene (B. Kloosterman, pers.comm.).

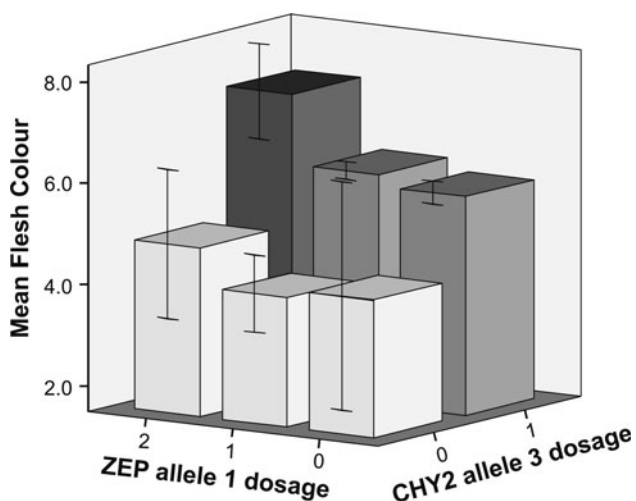


Fig. 6 Relation between *Chy2* allele 3 dosage, *Zep* allele 1 dosage, and flesh colour value in the diploid C×E population. Error bars ± 2 SE

Role of *Zep* allele 1 on zeaxanthin levels

For 88 C×E progeny the amounts of individual carotenoids were determined. A small number of progeny (10 genotypes) proved to contain relatively high levels of zeaxanthin (>250 μg per 100 g fresh weight). These progeny invariably contained *Chy2* allele 3 and were homozygous for *Zep* allele 1. In Fig. 7 the relation between zeaxanthin content and absorbance in the yellow spectrum is displayed for four classes of genotypes. Y or y represent the dominant or recessive *Chy2* allele, respectively, and Z or z represent the dominant or recessive *Zep* allele. Y is *Chy2* allele 3, and z is *Zep* allele 1. This figure shows that zeaxanthin only accumulates in considerable amounts in genotypes homozygous for *Zep* allele 1 (zz, filled symbols). When dominant *Chy2* allele 3 is present (Yzz) the zeaxanthin level is higher than when this allele is absent (yzz). These results

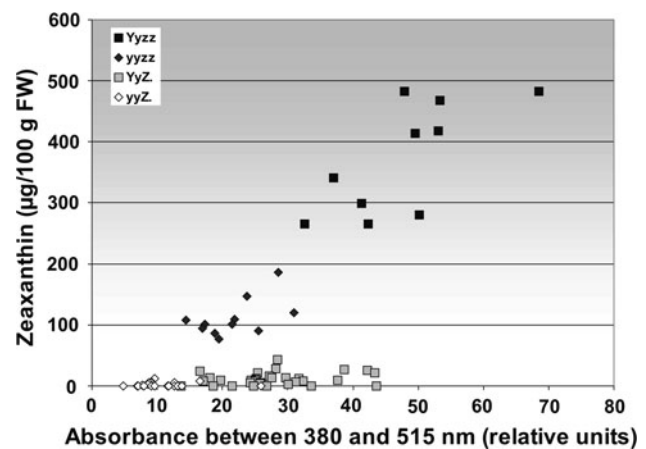


Fig. 7 Relation between zeaxanthin content and absorbance in the yellow spectrum in the diploid C×E population. Zeaxanthin content in $\mu\text{g}/100$ g fresh weight. Four genotypic classes are indicated, with Y/y representing *Chy2* alleles, and Z/z representing *Zep* alleles. Y is dominant *Chy2* allele 3; z is recessive *Zep* allele 1

suggest that orange flesh colour indicates the presence of a zeaxanthin level of more than 250 $\mu\text{g}/100$ g fresh weight tuber.

ZEP allele 1 is expressed at a low level

Expression analysis using the 44 k POCI array indicated that tuber RNA from parents C and E showed a similar level of hybridization to the *Zep*-derived 60-mer oligo (B. Kloosterman, pers. comm.). Both parents are heterozygous for *Zep*, containing alleles 1 and 2. Tuber RNA from C \times E progeny homozygous for *Zep* allele 2 showed a higher level of hybridization with the *Zep* oligo than both parents, whereas tuber RNA from C \times E progeny homozygous for *Zep* allele 1 showed a lower level of hybridization than both parents. This may reflect a difference in homology of the *Zep* alleles with the oligo. It was found that the 60-mer oligo on the POCI array (Kloosterman et al. 2008) is identical to a sequence in exon 6 of *Zep* allele 1, while there is one mismatch with the sequence in *Zep* allele 2. If the mismatch would result in a lower level of hybridization it would be expected that RNA from progeny homozygous for *Zep* allele 2 would show a lower level of hybridization. However, the opposite was observed. Therefore, the array results suggest that *Zep* allele 2 is expressed at a higher level than *Zep* allele 1. This was confirmed by quantitative RT-PCR (Fig. 8): diploid C \times E progeny homozygous for ZEP allele 1 showed a significantly lower level of expression than C \times E progeny homozygous for ZEP allele 2. Heterozygous progeny displayed an intermediate level of expression. A similar observation was made by Morris et al. (2004) who found that high carotenoid-accumulating diploid *S. phureja* genotype

DB375\1 (later renamed cultivar ‘Inca Dawn’) showed low expression of the *Zep* gene. They observed an inverse relationship between zeaxanthin transcript level and total carotenoid content in a range of potato germplasm.

To investigate the reason for the lower expression level of *Zep* allele 1 a 1.8-kb fragment containing the promoter of this allele was obtained by Genome Walking and was sequenced. A BLASTN analysis revealed that the 5' part of this sequence contained a repetitive element (present on several chromosomes of *S. tuberosum* and *S. lycopersicum*). An analysis of *cis*-regulatory elements showed that the ZEP promoter contains light-regulated, phytochrome-regulated, and water stress-regulated boxes, as well as hypo-osmolarity-responsive and sugar-repression elements. Subsequently, this sequence was compared with the promoter sequence of *Zep* allele 2 (as present in monoploid M133), see Supplemental Fig. S4. Although a number of SNPs was observed, no obvious differences were found that could explain the different expression levels.

Next, the complete genomic sequence, including all exons and introns, was determined for both *Zep* alleles 1 and 2 (Fig. 5 and Supplemental Fig. S4). The exon sequences were translated into protein sequences and aligned to each other (Supplemental Fig. S5). Although a number of amino acid changes were observed, especially in the first exon, no obvious amino acid change was found predicting a non-functioning enzyme according to SIFT. Alignment of the deduced amino acid sequences of *Zep* alleles 1 and 2 with *Zep* protein sequences of other Solanaceous species (Supplemental Fig. S6) indicated that the differences in amino acids between *Zep* alleles 1 and 2 mostly occurred in the least conserved regions.

However, we observed a large difference in size in the first intron. Intron 1 in *Zep* allele 2 is 389 bp in size, comparable with the 438-bp intron 1 of the tomato *Zep* genomic sequence. In contrast, the size of intron 1 in *Zep* allele 1 is 4,509 bp. By comparing the sequences of the first intron of *Zep* alleles 1 and 2 we observed that a 4,102-bp non-LTR retrotransposon-like sequence had integrated in intron 1 of allele 1, causing a target site duplication of 18 bp. Analysis of this sequence showed the presence of two open reading frames (ORFs) in the DNA strand antisense relative to the promoter (Fig. 5a). The first ORF contains an endonuclease/ exonuclease domain. The second ORF contains a non-LTR retrotransposon reverse transcriptase domain. Alignment of the reverse transcriptase domain of the retrotransposon in intron 1 in *Zep* allele 1 (*ZEPphur*) with the domains of LINE-1 like retrotransposons from other species is shown in Supplemental Fig. S7. Analysis of the protein sequence of ORF2 at the non-LTR retrotransposon classification webpage (Kapitonov et al. 2009) revealed that the retrotransposon in *Zep* allele 1 belongs to the RTE clade, and is closely related

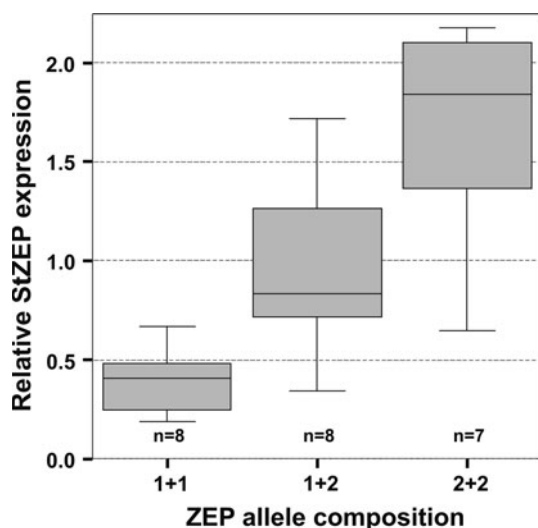


Fig. 8 Quantitative RT-PCR of *Zep* alleles. Relative expression level of the *Zep* gene for C \times E progeny homozygous for allele 1, heterozygous (allele 1 + allele 2), or homozygous for allele 2. *Zep* allele 1 results in a lower expression level than *Zep* allele 2

to the RTE1_ZM retrotransposon from *Zea mays* (Obukhanych and Jurka 2007). A BLASTN search of this retrotransposon sequence using the available potato genomic sequences at the PGSC webpage revealed homologous sequences on all 12 potato chromosomes, except chromosome 7. A similar search at the NCBI webpage using the high throughput genomic sequences (HTGS) database showed the presence of homologous retrotransposon sequences in several Solanaceous species, e.g. *Solanum*, *Nicotiana*, *Petunia* and *Capsicum* species. A BLASTN search using EST databases revealed that transcription of sequences homologous to the *ZEPphur* retrotransposon occurs.

Occurrence of ZEP allele 1 in the tetraploid potato gene pool

To determine the frequency of *Zep* allele 1 in the tetraploid potato gene pool PCR with primers AWZEP9 + AWZEP10 was performed using DNA from a set of 221 tetraploid potato cultivars (D'hoop et al. 2008) and some additional cultivars. From 230 genotypes that yielded a PCR product only 5 contained *Zep* allele 1, all in simplex. These were genotypes Black 1256, Prevalent (descendent from Black 1256), Producent (descendent from Prevalent), Lady Claire and Pallas (both descendents from *S. phureja* PHUR 71-464-7). This indicates that *Zep* allele 1 is a rare allele in the tetraploid potato gene pool (frequency 0.5%).

Zep allele composition was determined in 111 tetraploid potato genotypes not containing *Zep* allele 1, by sequencing the AWZEP9 + AWZEP10 PCR product. Four additional alleles were observed besides the alleles present in the monoploid and diploid genotypes (Supplemental Table S3). Alleles 2 (36%), 3 (25%), 4 (14%) and 5 (20%) were major alleles, whereas alleles 6 (0.9%), 7 (0.5%), 8 (1.6%) and 9 (1.6 %) were minor alleles. The minor alleles 6–9 are all present in tetraploid potato cultivars released after 1960. Alleles 3–9 had an intron 1 of similar size as intron 1 in allele 2. Therefore none of these alleles contained the transposon insertion present in allele 1.

Discussion

We conclude that homozygosity for *Zep* allele 1 in the presence of dominant *Chy2* allele 3 is causing the orange flesh colour phenotype, due to high levels of zeaxanthin. Furthermore, we conclude that *Zep* allele 1 is a recessive allele. The accumulation of zeaxanthin does not seem to be caused by impaired function of the *Zep* protein resulting from amino acid changes, as we observed no obvious amino acid changes in allele 1 compared with allele 2. Rather, the accumulation of zeaxanthin seems to result

from a lower steady state mRNA level of *Zep*, as determined by qRT-PCR.

Morris et al. (2004) observed an inverse relationship between the *Zep* transcript level and the total tuber carotenoid content. They investigated transcript level by quantitative RT-PCR using primers designed on the basis of tomato *Zep* cDNA sequence Z83835. However, DNA polymorphisms between potato and tomato can easily distort such analyses. Their forward primer contains an A at position 18 instead of G, as present in potato *Zep* cDNA DQ206629 and potato EST CK278242. We observed only G at this position in our *Zep* alleles, including allele 1, indicating a SNP in *Zep* sequences between tomato and potato close to the 3' end of the forward primer. This mismatch may have a considerable influence on overall level of amplification in the RT-PCR experiment. We performed qRT-PCR on a diploid population segregating for *Zep* alleles 1 and 2, using primers without mismatches, and observed a clear difference in expression levels between genotypes homozygous for allele 1, heterozygous (allele 1 + 2) and homozygous for allele 2.

A small number of SNPs was observed in the promoter sequence of allele 1 compared with the sequence of allele 2, which may explain the difference in expression level between the two alleles. However, we think it is more plausible that the difference in expression level is caused by the large retrotransposon insertion in the first intron of allele 1. This large insertion may cause inefficient splicing of the pre-mRNA into mature mRNA, or alternative splicing caused by cryptic splice sites. Hanson (1989) reported that efficient intron splicing in plants may be constrained by intron length. Ohmori et al. (2008) reported that integration of a transposon in intron 4 of rice gene *DL* resulted in reduced expression of the gene. Similarly, Gazzani et al. (2003) and Michaels et al. (2003) observed that weak alleles of the *Arabidopsis thaliana* *FLC* gene showing reduced expression contained a transposon in the first functional intron. A lower expression level of the zeaxanthin epoxidase gene results in the accumulation of zeaxanthin, at the expense of antheraxanthin, violaxanthin and neoxanthin (Tanaka et al. 2008). As zeaxanthin is relatively orange coloured (depending on concentration and milieu), while antheraxanthin, violaxanthin and neoxanthin are (light) yellow, this explains the orange flesh phenotype of the potato genotypes homozygous for *Zep* allele 1 (and containing *Chy2* allele 3).

Potato genotypes homozygous for *Zep* allele 1 do not show the wilted phenotype as observed in *Arabidopsis*, *Nicotiana plumbaginifolia* and tomato *Zep* mutants (Duckham et al. 1991; Galpaz et al. 2008; Marin et al. 1996), in which synthesis of plant hormone abscisic acid (ABA), a downstream metabolite of the carotenoid pathway, is compromised. Therefore, we conclude that the

reduced expression of *Zep* allele 1 does not result in complete absence of ABA.

Römer et al. (2002) achieved increased zeaxanthin levels in potato tubers by genetic engineering. However, consumer acceptance of GMO cultivars is very low. Use of the natural variant *Zep* allele 1 allows classical breeding for orange-fleshed potato. *Zep* allele 1 probably is an *S. phureja* *Zep* allele, as it is almost absent in the *S. tuberosum* gene pool, and only present in a few tetraploid potato genotypes with *S. phureja* in their ancestry. This means that breeding of a tetraploid potato cultivar with orange tuber flesh (with high zeaxanthin content) is a challenging task.

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