Genetic architecture controlling variation in grain carotenoid composition and concentrations in two maize populations

Catherine B. Kandianis · Robyn Stevens · Weiping Liu · Natalia Palacios · Kevin Montgomery · Kevin Pixley · Wendy S. White · Torbert Rocheford

Received: 20 April 2013 / Accepted: 12 August 2013 / Published online: 17 September 2013 © The Author(s) 2013. This article is published with open access at Springerlink.com

Abstract
Key message Genetic control of maize grain carotenoid profiles is coordinated through several loci distributed throughout three secondary metabolic pathways, most of which exhibit additive, and more importantly, pleiotropic effects.

Abstract The genetic basis for the variation in maize grain carotenoid concentrations was investigated in two F2:3 populations, DEexp × CI7 and A619 × SC55, derived from high total carotenoid and high β-carotene inbred lines. A comparison of grain carotenoid concentrations from population DEexp × CI7 grown in different environments revealed significantly higher concentrations and greater trait variation in samples harvested from a subtropical environment relative to those from a temperate environment.

Communicated by M. Xu.

Electronic supplementary material The online version of this article (doi:10.1007/s00122-013-2179-5) contains supplementary material, which is available to authorized users.

C. B. Kandianis · R. Stevens · K. Montgomery Department of Crop Sciences, University of Illinois, Urbana, IL 61801, USA e-mail: ceb19@cornell.edu

C. B. Kandianis Department of Pediatrics, USDA-ARS Children’s Nutrition Research Center, Baylor College of Medicine, Houston, TX 77030, USA

K. Montgomery Montgomery Consulting, Maroa, IL 61756, USA

R. Stevens U.S. Agency for International Development, Washington, DC 20523, USA

T. Rocheford Department of Agronomy, Purdue University, West Lafayette, IN 47907, USA e-mail: trochefo@purdue.edu

W. Liu · W. S. White Department of Food Science and Human Nutrition, Iowa State University, Ames, IA 50011, USA

Genotype by environment interactions was significant for most carotenoid traits. Using phenotypic data in additive, environment-specific genetic models, quantitative trait loci (QTL) were identified for absolute and derived carotenoid traits in each population, including those specific to the isomerization of β-carotene. A multivariate approach for these correlated traits was taken, using carotenoid trait principal components (PCs) that jointly accounted for 97% or more of trait variation. Component loadings for carotenoid PCs were interpreted in the context of known substrate-product relationships within the carotenoid pathway. Importantly, QTL for univariate and multivariate traits were found to cluster in close proximity to map locations of loci involved in methyl-erythritol, isoprenoid and carotenoid metabolism. Several of these genes, including lycopene epsilon cyclase, carotenoid cleavage dioxygenase1 and beta-carotene hydroxylase, were mapped in the segregating populations. These loci exhibited pleiotropic effects on α-branch carotenoids, total carotenoid profile and β-branch carotenoids, respectively. Our results confirm that several
QTL are involved in the modification of carotenoid profiles, and suggest genetic targets that could be used for the improvement of total carotenoid and β-carotene in future breeding populations.

Introduction

Selection of yellow grain varieties is noted as a hallmark of maize domestication (Doebley et al. 2006). Yellow pigmentation is attributed to an accumulation of carotenoids in the endosperm resulting from a gain of function mutation in the first, rate-limiting enzyme in the carotenoid pathway, phytoene synthase (y1, psy1) (Palaisa et al. 2004) (Fig. 1). In the 1930s, the discovery of increased nutrition in yellow maize grain (Mangelsdorf and Fraps 1931) led to selection of pigmented grain as a desirable quality trait for both human food and animal feed (Bauernfeind et al. 1981; Weber 1987). It is now widely known that plant-based carotenoids, the source of this yellow pigment in maize grain, can provide dietary vitamin precursors and antioxidants that are essential to human health.

Provitamin A (proVA) carotenoids form the retinyl structural base for vitamin A molecules through cleavage reactions specific to animal metabolism (Fierce et al. 2008; Leuenberger et al. 2001). ProVA carotenoids can be found in many plant-based foods and include β-carotene (2 retinyl groups), β-cryptoxanthin (1 retinyl group) and α-carotene (1 retinyl group). Lutein and zeaxanthin, dihydroxy-xanthophyll conversion products of these proVA carotenoids in plants, are also prevalent in vegetative and seed tissues and are an abundant source of antioxidants. ProVA carotenoids provide vitamin A activity which is involved in immune function, protection of vision systems, and cellular differentiation (Stephensen 2001; Whitcher et al. 2001), and dihydroxy-xanthophylls are believed to aid in the prevention of macular degeneration and visual acuity (Chucair et al. 2007; Seddon et al. 1994), prompting the implementation of various nutritional interventions involving increased consumption of plant-based carotenoids. Consumption of several of these plant metabolites is associated with research outcomes indicative of health improvement (Bouis et al. 2011; Li et al. 2010; Tang et al. 2009). Therefore, from a health and nutritional standpoint both the quantity and quality of carotenoid profiles are important when plant-based foods are being used to satisfy dietary requirements.

Maize grain carotenoid concentrations are among the highest produced in cereals (Howitt and Pogson 2006), and exhibit considerable diversity in the composition of grain carotenoid profiles with respect to the predominant carotenoids (lutein and zeaxanthin), proVA carotenoids (α-carotene, β-carotene and β-cryptoxanthin) and other non-proVA carotenoids (zeinoxanthin) (Harjes et al. 2008).

Genetic analyses conducted with populations segregating in seed color (Chandler et al. 2013; Islam 2004) and carotenoid profiles (Chander et al. 2008; Stevens 2007; Wong et al. 2004) have demonstrated that compositional and concentration differences in seed carotenoids are quantitatively inherited. Biochemical characterization of various maize endosperm color mutants originally classified by Robertson (1975) has helped to identify various maize-specific homologs of carotenoid pathway genes reported in bacteria or model crop species (Fig. 1). Interestingly, concomitant decreases in lutein and zeaxanthin are observed when selection for increased ProVA is imposed with crtRB1 (Yan et al. 2010; data not shown), indicating that the accumulation of carotenoid metabolites can be competitive due to a related biochemical origin or to an altered flux through the pathway.

![Image](Image)

Fig. 1 Biochemical pathways contributing to carotenoid biosynthesis. Carotenoids are derived from products of glycolysis (dark gray fill) and isoprenoid biosynthesis (light gray fill). Substrate is committed to the carotenoid biosynthesis pathway (white fill) by phytoene synthase (PSY). Pathway enzymes are shown in bold, carotenoid intermediates measured in this study in italics, and all other intermediates in normal font. Enzymes defined as: DXS 1-deoxy-d-xylulose-5-phosphate synthase, DXR DXP reductoisomerase, HDS 4-hydroxy-3-methylbut-2-eno1-y1 diphosphate synthase, HDR 4-hydroxy-3-methylbut-2-eno1-y1 diphosphate reductase, IPP isopentyl pyrophosphate isomerase, GGPPS geranyl geranyl pyrophosphate synthase, PSY phytoene synthase, PDS phytoene desaturase, Z-ISO 15-cis zeta carotene isomerase, ZDS zeta carotene desaturase, CRTISO carotenoid isomerase, LCYβ lycopene epsilon cyclase, LCYβ lycopene beta cyclase, CRTRB1 beta-carotene hydroxylase, ZEP zeaxanthin epoxidase, VDE1 violaxanthin de-epoxidase, CCD1 carotenoid cleavage dioxygenase 1. Abbreviated intermediates are: MEP dimethylerythritol 4-phosphate, DMAPP dimethylally1 diphosphate, IPP isopentyl diphosphate, GGPP geranyl geranyl pyrophosphate. References and genetic location for all pathway genes are provided in Supplemental Table 7.
Genetic mapping studies have also indicated that a significant number of the QTL involved with carotenoid accumulation in grain exhibit pleiotropic effects on multiple carotenoid traits (Harjes et al. 2008; Yan et al. 2010). This suggests that these QTL could control the substrate-product conversion of multiple metabolites in the pathway, therein revealing potential pathway control points. Reactions involved in glycolysis and isoprenoid biosynthesis appear to have a large impact on the magnitude of carotenoid accumulation in a manner similar to PSY1 (Estevez et al. 2001; Lois et al. 2000; Sandmann et al. 2006). Carotenoid production occurs within plastid organelles, in close proximity to the site of glycolysis. As depicted in Fig. 1, three-carbon molecules from glycolysis are committed to the methyl-erythritol phosphate (MEP) pathway through deoxyxylulose synthase (DXS) and deoxyxylulose reductase (DXR) (Lichtenthaler et al. 1997). The MEP pathway synthesizes five-carbon isoprenoids and subsequently produces isoprene isomers to form a common substrate pool shared by secondary metabolite pathways synthesizing plant sterols, terpenes, gibberellins, and carotenoids (Estevez et al. 2001).

Naturally occurring allelic variation for some of these pathway steps has been shown to affect variation in carotenoids in a wide variety of maize germplasm. Considering recent advances in our understanding of metabolic pathway reactions potentially involved with variation in carotenoid concentration and composition, it is likely that genes involved upstream of carotenoid synthesis will be prime candidates for the regulation of multiple carotenoid traits, while genes involved within carotenoid synthesis will be candidates for compositional differences. In this study, we investigate the genetic architecture underlying carotenoid concentration and compositional differences in F$_{2:3}$ maize populations A619 × SC55 and DEexp × CI7, two genetic populations derived from high total carotenoid and high β-carotene parent inbreds. We evaluate carotenoid trait profiles and identify QTL across temperate and subtropical environments, map genes of the carotenoid biosynthesis pathway relative to these QTL, and employ a multivariate approach to QTL mapping of carotenoid traits as a possible method to identify major regulatory control points for carotenoid accumulation.

Materials and methods

Genetic materials

Two mapping populations were developed from maize inbred lines selected for superior and complementary carotenoid composition and total carotenoid profiles. This information is based on the multi-year surveys of the Goodman-Buckler Diversity Panel (Harjes et al. 2008). The first population consists of 103 F$_{2:3}$ progeny derived from the cross of DEexp × CI7. DEexp was developed by Dr. Jim Hawk at the University of Delaware by selfing from a southeastern U.S. region Pioneer Hi-bred F$_1$ hybrid. CI7, developed by the USDA-ARS, is derived from the backcross L317/33-16/L317. Both lines yield high seed β-carotene concentrations, and also have high total carotenoid concentrations relative to lines screened in the Goodman-Buckler Diversity Panel (Supplemental Figure 1). Specific combining effects for β-carotene have been observed in the DEexp × CI7 hybrid, including high-parent heterosis for β-carotene in some, but not all, years (T. Rocheford, data not shown).

The second population consists of 227 F$_{2:3}$ progeny derived from the cross of A619 × SC55. A619 is an early maturing line developed in Minnesota, derived from the backcross A171/OH43/OH43, and exhibits a high total carotenoid profile relative to most lines in the Goodman Diversity Panels in 2003 and 2005 (top 25 and 20 % respectively, Supplemental Figure 1). SC55 was developed in South Carolina and is characterized by proportionally high β-carotene in grain. Favorable specific combining ability for high β-carotene concentration was observed in hybrid seed of the A619 × SC55 cross (Stevens 2007) and other materials (Egesel et al. 2003).

The above F$_1$ crosses were self-pollinated to create the F$_2$ generation. The F$_2$ seed from a single ear for each cross was planted, and plants were hand self-pollinated to create ears of F$_{2:3}$ derived seed.

Field evaluation

In 2005, the 103 F$_{2:3}$ lines of DEexp × CI7 and 227 F$_{2:3}$ lines of A619 × SC55 were grown in two replicates in each of two environments, the University of Illinois Urbana-Champaign and the International Maize and Wheat Improvement Center (CIMMYT), El Batan, Mexico (hereafter noted as Illinois environment and Mexico environment, respectively), using alpha (0, 1) incomplete block designs. The families were planted in single row plots of 5 m length, with 76 cm spacing between rows. Each plot was thinned to a density of approximately 15 plants per 5 m, or 43,000 plants ha$^{-1}$. Seven to nine plants were sib-pollinated within each row. Growing seasons varied by environment as follows: Urbana, IL, May–October; El Batan, Mexico, May–November. Fluctuation in average monthly temperature and precipitation recorded at both locations in 2005 is shown in Supplemental Figure 2.

After storage at room temperature for approximately 4 months, seed was bulked for each plot at shelling and an aliquot of approximately 10 g of seed from each plot was...
stored at −80 °C until carotenoid extraction could be performed on a plot mean basis. Due to field technical complications, the Illinois replicates of population A619 × SC55 were not included this study.

DNA extraction and genotyping

DNA extraction was performed as described by Mikkilineni and Rocheford (2003). DNA samples prepared from F2,3 family seed bulks were used for genotyping. A parental survey using 748 publicly available microsatellite (SSR) markers from MaizeGDB was performed. Pioneer Hi-Bred performed in-kind genotyping on both DEexp × CI7 and A619 × SC55 populations using polymorphic markers identified in the parental survey in addition to several proprietary markers, designated by the prefix “pio”. Additional SSR markers genotyped at the University of Illinois were added to the genetic map. Allele-specific markers lcy -MZA, crrRB1-InDel4, and ccdl-5p were used to map three genes implicated in carotenoid synthesis or degradation: lycopene epsilon cyclase (lcy; NM_001153368.1), beta-carotene hydroxylase (crrRB1; NM_001112437), and carotenoid cleavage dioxygenase 1 (ccdl; DQ100346), respectively. Information for functional marker assays designed for this study (lcy and ccdl) and those derived from Yan et al. (2010) (crrRB1) is listed in Supplemental Table 1.

Phenotypic data collection

Carotenoid extraction from finely ground maize kernels was performed at Iowa State University as previously described (Li et al. 2007b). Briefly, the method involves mild heat treatment (50 °C × 15 min) in methanol followed by extraction with tetrahydrofuran and a 5-min saponification with methanolic potassium hydroxide at room temperature. After washing with water, the carotenoids are partitioned into hexane/methylene chloride (5:1 v/v), which is then evaporated to dryness under vacuum. The dried extract is reconstituted in methyl-tert-butyl ether followed by methanol and an aliquot is injected into the HPLC system. An internal standard, β-apo-8’-carotenal, was added during the extraction. Carotenoid separation was performed on a C-30 YMC Carotenoid Column, with gradient separation (methanol/methyl-tert-butyl ether) and photodiode array detection. Carotenoids were quantified using internal standard calibration curves. Measured carotenoid traits included lutein, zeaxanthin, zeinoxanthin, β-cryptoxanthin, α-carotene, trans and cis isomers of β-carotene, phytoene and trans-phytofluene. The summation of all carotenoids with absorption spectra in the visible range (lutein, zeaxanthin, zeinoxanthin, β-cryptoxanthin, α-carotene, total β-carotene) is termed total colored carotenoid in subsequent analyses. Isomers of β-carotene measured in this study include all-trans, 9-cis, 13-cis, and 15-cis β-carotene, and are represented by concentration (μg g−1 DW) and proportion (ratio of isomer to total β-carotene). The ratio of colorless to colored carotenoids is defined by the proportion of phytoene plus phytofluene to total colored carotenoid. Proportion of alpha branch carotenoids (lutein, zeinoxanthin, α-carotene) to beta branch carotenoids (zeaxanthin, β-cryptoxanthin, β-carotene) is defined by α:β branch. Provitamin A was calculated according to the number of retinyl groups associated with all measured carotenoids, resulting in the expression ProVA = β-carotene + 0.5*β-cryptoxanthin + 0.5*α-carotene. All other ratios are comprised of absolute concentrations.

Genetic map construction

Genetic maps for both populations were constructed using JoinMap® Version 3 (Ooijen and Voorrips 2001). Genotype data were screened for segregation distortion; markers with genotype segregation significantly different from the predicted 1:2:1 Mendelian ratio were removed. Maps were created from individual linkage groups using a LOD threshold of P < 0.001 and recombination frequency of 0.49 using Haldane’s map function. The final linkage map for DEexp × CI7 consisted of 109 markers, spaced with an average interval length of 15 cM over a total of 1,486 cM. The final linkage map for A619 × SC55 consisted of 117 markers, spaced with an average interval length of 16.2 cM over a total of 1,728.6 cM. Genetic maps for both populations are shown in Supplemental Figure 3.

Phenotypic data analyses

All analyses were carried out using statistical procedures in SAS version 9.2 (SAS Institute 2008). Trait means and ranges by environment and for combined environments were obtained through Proc MEANS. Product-moment correlations of raw trait data were performed using Proc CORR. Analysis of variance for trait data was modeled according to $y_{ijkl} = \mu + a_l + b_j + \delta_{ijkl} + \chi_{i} + (\alpha \gamma)_{il} + e_{ijkl}$ where $y$ is the carotenoid phenotype (μg g−1) of an individual in the $l$th environment, $j$th replicate, $k$th block and $l$th genotype, $\mu$ is the population mean, $a$ is the effect of the $l$th environment, $b$ is the effect of the $j$th replication in the $l$th environment, $\delta$ is the effect of the $k$th block in $j$th replication of the $l$th environment, $\chi$ is the effect of the $l$th genotype (or family), $(\alpha \gamma)$ is the interaction effect of the $l$th environment and the $l$th genotype, and $e$ is the experimental error. The analyses were performed with Proc MIXED and all model effects were considered random. Variance component estimates for genetic variance ($\sigma^2_{ge}$), genotype by environment interaction variance ($\sigma^2_{Tre}$), and error variance
additive by additive interactions or dominance effects were tested using the marker set selected by stepwise regression. Model fit criteria including Akaike’s Information Criterion (AIC) and adjusted $R^2$ were used to select models accounting for greatest phenotypic variation with fewest estimated parameters. For most traits in both populations, an additive model produced an adequate fit to the data.

Adjusted $R^2$ values from the final multiple regression model express the proportion of phenotypic variation adjusted by the number of estimated model parameters. Additive effects are noted for each QTL, and reflect the partial regression coefficient of the marker genotype on the phenotype. These effects are expressed as a deviation from the second inbred parents, which in this study are CI7 and SC55. Effects associated specifically with the $crtRB1$ locus have been previously noted in population DEexp × C17 by Yan et al. (2010).

**Results**

Comparison of descriptive statistics for traits across locations

Eight carotenoid traits (lutein, zeaxanthin, zeinoxanthin, $\beta$-cryptoxanthin, $\alpha$-carotene, $\beta$-carotene) plus colorless carotenoids phytoene and phytofluene were measured in par-

ent inbreds and F$_{2,3}$ progeny for population DEexp × C17 across two environments and for population A619 × SC55 in one environment (Mexico). Six derived carotenoid traits including total carotenoid concentration, total colored carotenoid concentration, colored:colorless carotenoid ratio, $\beta$-carotene conversion ($\beta$-car: $\beta$-cry ratio), $\alpha$:$\beta$ branch conversion ($\alpha$:$\beta$ ratio) and provitamin A (ProVA) were calculated from measured carotenoid traits (Supplemental Figure 4). DEexp and CI7 parent inbreds differed significantly ($\alpha = 0.05$) in lutein, $\beta$-carotene and phytoene concentrations in both environments (Fig. 2). High total carotenoid concentrations were observed in A619, DEexp and CI7, while SC55 had high $\beta$-carotene concentrations relative to other experimental germplasm planted in the same environment year (Supplemental Figure 1), confirming the selection criteria used for these parents.

Heritabilities on an entry mean basis ($h^2$) for carotenoid concentrations were high in population DEexp × C17: phytoene, 0.68; phytofluene, 0.71; $\alpha$-carotene, 0.71; total $\beta$-carotene, 0.59; zeinoxanthin, 0.82; $\beta$-cryptoxanthin, 0.79; lutein, 0.74; zeaxanthin, 0.75. Both concentrations ($h^2 = 0.56–0.64$) and relative proportions ($h^2 = 0.54–0.74$) of the $\beta$-carotene isomers were found to be highly heritable. The effect of genotype was significant ($\alpha = 0.01$) for all carotenoids in population DEexp × C17, and

Results

Comparison of descriptive statistics for traits across locations

Eight carotenoid traits (lutein, zeaxanthin, zeinoxanthin, $\beta$-cryptoxanthin, $\alpha$-carotene, $\beta$-carotene, phytoene and phytofluene) and four $\beta$-carotene isomers (9-9is, 13-9is, 15-9is and all-trans $\beta$-carotene) were measured in parent inbreds and F$_{2,3}$ progeny for population DEexp × C17 across two environments and for population A619 × SC55 in one environment (Mexico). Six derived carotenoid traits including total carotenoid concentration, total colored carotenoid concentration, colored:colorless carotenoid ratio, $\beta$-carotene conversion ($\beta$-car: $\beta$-cry ratio), $\alpha$:$\beta$ branch conversion ($\alpha$:$\beta$ ratio) and provitamin A (ProVA) were calculated from measured carotenoid traits (Supplemental Figure 4). DEexp and CI7 parent inbreds differed significantly ($\alpha = 0.05$) in lutein, $\beta$-carotene and phytoene concentrations in both environments (Fig. 2). High total carotenoid concentrations were observed in A619, DEexp and CI7, while SC55 had high $\beta$-carotene concentrations relative to other experimental germplasm planted in the same environment year (Supplemental Figure 1), confirming the selection criteria used for these parents.

Heritabilities on an entry mean basis ($h^2$) for carotenoid concentrations were high in population DEexp × C17: phytoene, 0.68; phytofluene, 0.71; $\alpha$-carotene, 0.71; total $\beta$-carotene, 0.59; zeinoxanthin, 0.82; $\beta$-cryptoxanthin, 0.79; lutein, 0.74; zeaxanthin, 0.75. Both concentrations ($h^2 = 0.56–0.64$) and relative proportions ($h^2 = 0.54–0.74$) of the $\beta$-carotene isomers were found to be highly heritable. The effect of genotype was significant ($\alpha = 0.01$) for all carotenoids in population DEexp × C17, and

---

**Principal component analysis**

Principal component analysis (PCA) was conducted for each population using Proc PRINCOMP in SASv9.2 using a matrix of trait BLUPs for colored carotenoids (lutein, zeaxanthin, zeinoxanthin, $\beta$-cryptoxanthin, $\alpha$-carotene, $\beta$-carotene) plus colorless carotenoids phytoene and phytofluene.

**Linkage analysis by composite interval mapping**

Quantitative trait locus analysis was performed using composite interval mapping (CIM) with PLABQTL software (Utz and Melchinger 1993). This mapping methodology is based on the Haley-Knott regression method for marker intervals (Haley and Knott 1992), and is supplemented by cofactor selection (Jansen and Stam 1994; Zeng 1994). Univariate (single trait BLUP values) and multivariate (PC score) traits were mapped according to the following model: $y_{ijkl} = \mu + a_{ij} + \beta_{k} + \beta_{k}X_{ij} + \sum_{k \neq l} g_{k} z_{kj} + \epsilon_{ijkl}$ where $y$ is the trait value of family $j$, $a_{ij}$ is the intercept, $B_{i}$ is the genetic effect of the QTL located within the interval between marker $i$ and marker $i + 1$, $x_{ij}$ is a coded variable indicating the genotype of the marker interval (where maximally the value of parent 1 is −1, and parent 2 is 1), $\sum_{k \neq l} g_{k} z_{kj}$ is the summation of selected cofactor effects where $g_{k}$ is the partial regression coefficient of the trait value on marker cofactor $k$ and $z_{kj}$ is a coded variable for the genotype at cofactor $k$.

A LOD threshold equivalent to an experiment-wise Type I error rate of $\alpha = 0.05$ was used as the initial model selection criterion (Lander and Botstein 1989) for both populations; to obtain the threshold, 1,000 permutations of the trait data were run using a model of specified gene action, cofactor selection (Doerge and Churchill 1996), and a 2 cM scan. This corresponded to a threshold of LOD 3.5 for DEexp × C17 analyses and LOD 2.9 for A619 × SC55 analyses. Regression models accounting for digenic additive by additive interactions or dominance effects were tested using the marker set selected by stepwise regression. Model fit criteria including Akaike’s Information Criterion (AIC) and adjusted $R^2$ were used to select models accounting for greatest phenotypic variation with fewest estimated parameters. For most traits in both populations, an additive model produced an adequate fit to the data.

Adjusted $R^2$ values from the final multiple regression model express the proportion of phenotypic variation adjusted by the number of estimated model parameters. Additive effects are noted for each QTL, and reflect the partial regression coefficient of the marker genotype on the phenotype. These effects are expressed as a deviation from the second inbred parents, which in this study are CI7 and SC55. Effects associated specifically with the $crtRB1$ locus have been previously noted in population DEexp × C17 by Yan et al. (2010).
The genotype × environment (G×E) interaction effects were significant for lutein, β-carotene, phytoene and phytofluene. The environment effect was significant (P < 0.05) for all traits in the DEexp × CI7/IL population except α-carotene. Carotenoid concentrations were larger in the Mexico than in the Illinois environment, with the exception of colorless carotenoids (phytoene and phytofluene) which were smaller in Mexico (Fig. 2, Supplemental Fig 4, Supplemental Fig 5). Trait concentrations and ratios were normally distributed in the DEexp × CI7 F₂₋₃ population and transgressive segregation for all traits was observed in both directions for both environments. This contrasted sharply

Fig. 2 Distribution of measured carotenoid traits by population and environment. Carotenoid concentrations measured for individuals in each population-environment combination are represented by frequency histograms proportional to the size of each population. Data are shown by population-environment for A619 × SC55/MX (blue solid line), DEexp × CI7/IL (red dashed line), and DEexp × CI7/MX (green dashed line), respectively. Parent profiles are listed within respective bar graphs by population-environment and significantly different means (α = 0.05) are noted by (asterisk) (color figure online)
with trait distributions from the A619 × SC55 F2:3 population in the Mexico environment, in which distributions for nearly all carotenoids were skewed toward the direction of the SC55 parent profile, and transgressive segregation was observed for all carotenoids at the higher end of trait distributions. Across both populations, cis isomers of β-carotene constituted a much smaller proportion of total β-carotene relative to the all-trans isomer (50–80 % of total), with relative proportions of: 9-cis, 10-25 %; 13-cis, 12–20 %; 15-cis, 2-6 % (Supplemental Fig 5).

Correlation and principal component analyses for carotenoid concentrations

Pairwise correlations among carotenoid traits were positive and significant (α = 0.05) among all colored carotenoids for the DEexp × CI7 F2:3 families (Table 1), except for β-carotene and lutein (r = −0.32). Small but significant negative correlations were observed between colorless carotenoids (phytoene and phytofluene) and several colored carotenoids. Correlations among carotenoid concentrations were positive among nearly all colored and colorless carotenoids, and no traits were negatively correlated in the A619 × SC55 population (Table 1).

For DEexp × CI7 F2:3 progeny, three PCs accounted for 97 % of the total variance in the trait set from the Mexico environment (Table 2). PC1 accounted for the majority of the variance (90 %, λ = 8.59) almost entirely through variation in lutein (0.99). PC2 explained 5 % of the trait variance (λ = 0.51) through substantial positive loadings in colorless carotenoids (phytoene, 0.91; phytofluene, 0.33) and β-carotene (0.22), whereas PC3 explained a similar amount of variation (4 %, λ = 0.36) through positive loadings for lutein (0.23) and β-carotene (0.95), and a negative loading for phytoene (−0.18).

Four PCs explained the variation in the carotenoid trait matrix for A619 × SC55, with PC1 and PC2 accounting for 49 % (λ = 5.92) and 39 % (λ = 4.73) of the trait variance in the Mexico environment, respectively, through substantial loadings in lutein, zeaxanthin and phytoene. PC3 and PC4 accounted for 7 and 3 % of the variation, respectively, and had substantial loadings for lutein, zeaxanthin and β-carotene. Eigenvectors for the identified A619 × SC55 PCs demonstrated both synergistic and antagonistic effects through positive and negative loadings.

Loadings for PC traits were interpreted in the context of the biochemical pathway. In DEexp × CI7, most of the pathway variations were attributed to lutein, whereas smaller trait correlations appeared to be involved in apportioning carotenoid branch precursors to β-carotene (PC2) or could be involved in competition between phytoene precursors and downstream carotenoids (PC3). A619 × SC55 PCs were interpreted as a description of synergistic fluxes (PC1) or competitive allocation (PC2) of phytoene substrate to the largest carotenoid pools (lutein and zeaxanthin). PC3 described a competition between lutein and β-carotene versus zeaxanthin, whereas PC4 described a competition between α and β branch carotenoids.

Quantitative trait locus mapping in DEexp × CI7 population

For the DEexp × CI7 population, marker-genotype association with trait BLUPs from Mexico identified from 2 to 8 QTL explaining significant variation for the various carotenoid concentrations and ratios (Table 3, Supplemental Tables 2 and 3). Multiple regression models accounted for 20.3–55.3 % (R2adj) of the phenotypic variation in individual

### Table 1 Correlations between kernel carotenoid concentrations within A619 × SC55, and DEexp × CI7 F2:3 populations

| Population: A619 × SC55 | lut | zea | zein | βcry | αcar | βcar | total | phyene | phyflu |
|-------------------------|-----|-----|------|------|------|------|-------|--------|--------|
| lut                     |     |     |      |      |      |      | 0.67  | 0.36   | 0.45   |
| zea                     |     |     |      |      |      |      | 0.31  | 0.70   | 0.14   |
| zein                    |     |     |      |      |      |      | 0.79  | 0.41   | 0.33   |
| βcry                    |     |     |      |      |      |      | 0.15  | 0.33   | 0.43   |
| αcar                    |     |     |      |      |      |      | 0.19  | 0.19   | 0.48   |
| βcar                    | -0.32 | ns |      |      |      |      | 0.48  | 0.19   | 0.19   |
| total                   | 0.95 | 0.35 | 0.33 |      |      |      | 0.93  | 0.30   | 0.30   |
| phyene                  |     |     |      |      |      |      | 0.05  | 0.19   | 0.24   |
| phyflu                  |     |     |      |      |      |      | -0.24 | -0.24  | 0.97   |

Significant positive correlations shown in white; significant negative correlations shown in light gray; non-significant correlations are indicated by ns. Significance tested at a = 0.05. Traits are coded for lutein (lut), zeaxanthin (zea), zeinoxanthin (zein), β-cryptoxanthin (bcry), α-carotene (acar), β-carotene (bcar), total colored carotenoid (total), phytoene (phyene) and phytofluene (phyflu)
carotenoid concentrations, and 31.8–58.6 % ($R^2_{adj}$) of variation in ratios.

Although DEexp and C17 carotenoid profiles derived from the Mexico environment were very similar (Fig. 2), significant QTL were detected for several traits in the corresponding $F_{2:3}$ progeny (Table 3, Supplemental Table 2). Variation in concentrations and trait ratios for colorless carotenoids was attributed to 4–8 main effect QTL which predominantly were located on chromosomes 3 and 4. A larger proportion of the favorable additive effects at these QTL were contributed by DEexp relative to C17. Eight QTL were detected for β-carotene, several of which individually accounted for more than 10 % of the variation for this trait; most of the positive additive effects for these QTL were also derived from DEexp. Variation in lutein, the carotenoid differing most between inbreds DEexp and C17, was explained by only two large QTL detected at chromosome 3 (pos. 70, positive CI7 effect of 1.19 μg g⁻¹) and chromosome 8 (pos. 66, positive DEexp effect of 2.81 μg g⁻¹). Epistatic effects were significant in only two derived traits, α:β branch ratio and colorless:colored carotenoid ratio. While it would be expected that metabolic QTL originating from the same pathway would yield more epistatic interactions, it is likely that the sample size of this population prevented detection of such effects.

Fewer carotenoid QTL were detected for the DEexp × C17 in Illinois than described above for the Mexico environment. This was attributed to larger trait averages and variances in Mexico than Illinois (Fig. 2). A single, large effect QTL on chromosome 8 (pos. 52–54, $R^2_{part} = 40.3$) explained variation in lutein, and although this QTL did not map to the exact pos. as those accounting for variation in lutein in the Mexico environment (Mexico, pos. 66; Illinois pos. 52–54), the magnitude of additive effect and source of parental contribution of the lutein QTL were nearly identical across the two environments, suggesting that the same locus could be operating in both environments. Similarly, QTL for zeaxanthin, β-cryptoxanthin, total colored, colorless:colored ratio, βcar:βcry ratio, α:β branch ratio accounting for trait variation in the DEexp × C17 population grown in Illinois were also detected in the Mexico environment, and had consistent parent of origin and genetic effects (Supplemental Table 3, Table 3). No loci were found to account for significant variation for α-carotene, β-carotene, phytoene or phytofluene (Supplemental Table 3) for DEexp × C17 in the Illinois environment.

QTL included in the final regression models were found to overlap among carotenoid traits known to be related by way of sequence in metabolic pathways (i.e. colorless or β-cyclized carotenoids) or through extent of hydroxylation (i.e. carotenes or xanthophylls). Colorless carotenoids phytoene and phytofluene shared common QTL at chromosomes 3 (pos. 74; pos. 142), 4 (pos. 52), and 10 (pos. 54). β-cyclized carotenoids (β-carotene, β-cryptoxanthin and zeaxanthin) shared common QTL at chromosomes 2 (pos. 66), 8 (pos. 80), and 10 (pos. 56–58). A β-branch relationship was exposed through a chromosome 10 QTL, which increased the β-carotene:β-cryptoxanthin ratio through the DEexp allele (ratio effect = 0.96, $R^2_{part} = 26.3$ %) yet increased zeaxanthin through the C17 allele. α-and β-carotenes were both affected by a QTL on chromosome 2 (pos. 64–66) and another on chromosome 5 (pos. 82–84) with both positive effects derived from the DEexp alleles. The most pervasive pleiotropic effects on the entire pathway were detected at chromosome 2 (pos. 32–40) and chromosome 3 (pos. 68–72), which both increased total colored carotenoids in addition to many individual carotenoids.

QTL mapping for PC1, which accounted for 90 % of the variance among the carotenoid trait matrix, revealed two major QTL at chromosomes 3 (pos. 70) and 8 (pos. 66) to account for 37.8 % of the total phenotypic variation at the Mexico environment (Table 5). These effects were similar to those found for lutein in the same environment, and

---

**Table 2** Principal component parameters for DEexp × C17 and A619 × SC55 $F_{2:3}$ populations describing carotenoid trait variation

| Population          | DEexp × C17 | A619 × SC55 |
|---------------------|-------------|-------------|
|                     | PC1  | PC2  | PC3  | PC1  | PC2  | PC3  | PC4  |
| **Eigenvalues ($\lambda$)** | 8.59 | 0.51 | 0.36 | 5.92 | 4.73 | 0.80 | 0.42 |
| **Total variance (%)**   | 0.90 | 0.05 | 0.04 | 0.49 | 0.39 | 0.07 | 0.03 |
| **Eigenvectors**          |       |       |      |      |      |      |      |
| Lutein                | 0.99 | 0.08 | 0.05 | 0.57 | 0.71 | −0.33| −0.25|
| Zeaxanthin            | 0.02 | −0.12| 0.23 | 0.26 | 0.31 | 0.75 | 0.49 |
| Zeinoxanthin          | 0.04 | −0.01| 0.05 | 0.03 | 0.06 | 0.02 | 0.02 |
| β-cryptoxanthin       | 0.00 | −0.01| 0.03 | 0.05 | 0.04 | 0.15 | 0.08 |
| α-carotene            | 0.01 | 0.00 | 0.05 | 0.01 | 0.00 | −0.04| 0.03 |
| β-carotene            | −0.07| 0.22 | 0.95 | 0.05 | 0.00 | −0.56| 0.83 |
| Phytoene              | −0.06| 0.91 | −0.18| 0.76 | −0.63| 0.00 | −0.03|
| Phytofluene           | −0.02| 0.33 | −0.07| 0.15 | −0.09| 0.04 | −0.05|
Table 3  QTL detected by composite interval mapping for carotenoid composition traits in DEexp × CI7 population, Mexico environment, 2005

| Chr. Bin | Contig | Interval | Pos. | LOD | Add | $R^2$ (%) | LOD | Add | $R^2$ (%) | LOD | Add | $R^2$ (%) |
|----------|--------|----------|------|-----|-----|-----------|-----|-----|-----------|-----|-----|-----------|
| 3.04/05  | 120/125| umc1683–umc1102| 74 | 11.66 | −0.40 | 20.9 | 7.68 | −0.12 | 12.6 |
| 3.08     | 146    | umc1273–pio6 | 142 | 5.06  | 0.20 | 9.6  | 4.62 | 0.06 | 3.7  |
| 4.00     | pio7–pio8| 0 | 4.18 | −0.22 | 11.6 | |
| 4.03/04  | 158/164| adh2–umc2061| 52 | 5.91  | −0.23 | 9.8  | 4.93 | −0.14 | 17.7 |
| 6.00/02  | 260/271| umc1018–umc1083| 46 | 9.11  | −0.30 | 16.7 | |
| 9.07/08  | 391    | bnlg1375–umc1505 | 112 | 5.18  | 0.23 | 9.3  | |
| 10.06    | 415    | CrRB1–bnlg1028 | 54 | 10.75 | −0.41 | 23.0 | 9.17 | −0.13 | 13.6 |

Genetic environment, significance and effect of selected QTL for each trait are listed. Indicated are LOD (Logarithm of Odds), Add (Additive effect) and $R^2$ (%) (coefficient of partial determination). Significant digenic interactions detected between main effect QTL are indicated at the bottom of the list of main effects; contributing main effects are marked by asterisk(s).

coincided with PC1 QTL detected in the Illinois environment (chromosome 3, pos. 72, $R^2_{adj} = 14.9$; chromosome 8, pos. 66, $R^2_{adj} = 43.8$; Supplemental Table 3).

Variation in β-carotene isomer concentrations and proportions mapped to 3–8 loci for all-trans, 9-cis and 13-cis β-carotene, accounting for 25–60.5 % of the phenotypic variation (Supplemental Table 4). No QTL were significantly associated with 15-cis β-carotene in this population. Several QTL accounted for variation in more than one isomer, including loci on chromosomes 5 (pos. 92–98; 13-cis, all-trans), 7 (pos. 70–76; 9-cis, 13-cis, all-trans), and 10 (pos. 18–52; 9-cis, 13-cis, all-trans). Similarly, overlapping effects for more than one proportion trait were observed for QTL on chromosomes 2 (pos. 77–90; 9-cis, all-trans), 5 (pos. 94–96; 9-cis, all-trans), and 8 (pos. 70–80; 9-cis, 13-cis, all-trans).
Quantitative trait locus mapping in A619 × SC55 population

Trait BLUPs from the Mexico environment were used to identify significant marker genotype-phenotype associations for the A619 × SC55 population. A range of 3–6 QTL was found to explain 21.6–56.2 % of the variation for individual carotenoid concentrations, and 37.2–47.4 % for carotenoid ratios (Table 4, Supplemental Table 5). Fewer additive by additive interactions were detected in the A619 × SC55 population than in DEexp × CI7. A model for total colored carotenoids explained 53.9 % of the phenotypic variation through four main effects and one epistatic interaction. Variation in two traits, β-carotene and...

Table 4 QTL detected by composite interval mapping for carotenoid composition traits in A619 × SC55 population, Mexico environment, 2005

| Chr. Bin | Contig | Interval | Pos. | LOD | Add R² (%) | LOD | Add R² (%) | LOD | Add R² (%) |
|----------|--------|----------|------|-----|------------|-----|------------|-----|------------|
| 1.11     | 64     | umc2242–umc1979 | 178  | 3.30 | 0.50 | 3.2 | 3.13 | 0.16 | 7.9 |
| 3.07     | 142    | pio_5–umc1489  | 134  | 5.91 | 0.16 | 8.3 |
| 5.06     | 251    | pio_9–umc2013   | 144  | 2.91 | 0.06 | 1.4 |
| 7.02     | 297    | umc1068–bnlg1094 | 46   | 13.10 | 1.45 | 23.8 | 22.07 | 0.34 | 32.5 |
| 7.04     | 323/325| umc1944–umc1125 | 120  | 4.30 | 0.80 | 9.1 |
| 9.08     | 391    | zct128–umc1505  | 166  | 6.59 | 1.33 | 19.6 | 8.07 | 0.25 | 19.6 |

Interaction

| Chr. Bin | Contig | Interval | Pos. | LOD | Add R² (%) | LOD | Add R² (%) | LOD | Add R² (%) |
|----------|--------|----------|------|-----|------------|-----|------------|-----|------------|
| 2.04     | 77     | umc1541–pio_4 | 128  | 5.63 | −0.12 | 10.1 |
| 2.05     | 90     | pio_4–umc1459 | 130  | 6.74 | −1.00 | 33.5 |
| 2.07     | 91/105 | bnlg1396–dupsr25 | 172  | 5.97 | 0.34 | 10.2 |
| 4.02     | 156    | pio_08–phi295450 | 32   | 6.26 | 0.39 | 13.5 |
| 4.03     | 158/176| adh2–umc1142   | 68   | 4.34 | −0.27 | 6.6 |
| 5.03     | 212/217| umc2035–umc2295 | 72   | 10.07 | 0.38 | 17.5 |
| 6.05     | 285/287| umc1805–umc1859 | 90   | 5.03 | 0.30 | 10.9 |
| 7.00     | 293/296| umc1241–umc1068 | 46   | 3.24 | −0.08 | 6.7 |
| 9.07     | 391    | umc1675–umc2099 | 136  | 4.32 | −0.49 | 11.6 |
| 9.08     | 391    | zct128–umc1505 | 164  | 4.07 | −0.11 | 12.4 | 3.25 | −0.41 | 7.4 |
| 10.03    | 400    | umc2017–pio_14  | 62   | 3.33 | −0.06 | 2.5 |
| 10.05    | 414    | umc1506–CrR-B1  | 88   | 10.20 | −0.21 | 17.6 | 4.20 | −0.46 | 10.4 |
| 10.06    | 417    | CrR-B1–umc1993  | 94   | 6.43 | 0.37 | 14.0 | 0.79 | 19.4 |

Interaction

| Chr. Bin | Contig | Interval | Pos. | LOD | Add R² (%) | LOD | Add R² (%) | LOD | Add R² (%) |
|----------|--------|----------|------|-----|------------|-----|------------|-----|------------|
| 1.03     | –      | phi339017–pio_1 | 72   | 5.75 | 0.20 | 14.6 |
| –        | –      | pio_7–pio_8   | 0    | 6.84 | 0.03 | 11.7 |
| 5.02     | 208/212| umc1587–umc2035 | 64   | 6.06 | 0.03 | 8.3 | 9.29 | 0.20 | 16.0 |
| 6.05     | 283    | pio_10–umc1114 | 76   | 3.85 | −0.10 | 4.5 |
| 8.04     | –      | umc1343–LCYe  | 88   | 5.93 | 0.03 | 9.1 |
| 8.05     | 354    | LCYe–umc1340   | 92   | 5.21 | 0.64 | 10.0 |
| 9.07     | 391    | umc1675–umc2099 | 136  | 3.77 | −0.46 | 1.4 |
| 9.07     | 391    | CCD1–zct128   | 162  | 3.57 | −0.08 | 3.3 | 9.78 | −1.74 | 18.5 |

Interaction

Genetic environment, significance and effect of selected QTL for each trait are listed. Indicated are LOD (Logarithm of Odds), Add (Additive effect) and R² (%) (coefficient of partial determination). Significant digenic interactions detected between main effect QTL are indicated at the bottom of the list of main effects; contributing main effects are marked by asterisk(s).
Table 5 QTL detected by composite interval mapping for carotenoid principal components in DEexp × CI7 and A619 × SC55 population, Mexico environment, 2005

| Pop. | Trait | Model $R^2_{adj}$ | Chr. Bin | Contig | Interval | Pos. | LOD | Add | $R^2$ (%) |
|------|-------|-------------------|----------|--------|----------|------|-----|-----|-----------|
| D × C | PC1 | 37.8 | 3.04 | 116/120 | bnlg1019–umc1683 | 70 | 6.19 | 1.20 | 10.2 |
|      |     |       | 8.04/07 | 353/363 | umc1343–bnlg1828 | 66 | 4.05 | -2.83 | 31.2 |
|      | PC2 | 52.1 | 1.03/04 | 12/14 | phi109275–umc2217 | 100 | 3.52 | -0.22 | 8.3 |
|      |     |       | 2.00/02 | 68/69 | umc2246–bnlg1017 | 0 | 3.74 | -0.23 | 9.6 |
|      |     |       | 3.01 | 111 | umc1892–umc1814 | 28 | 7.47 | 0.31 | 13.5 |
|      |     |       | 3.04/3.05 | 120/125 | umc1683–umc1102 | 74 | 3.99 | -0.64 | 24.7 |
|      |     |       | 3.05 | 131 | umc2265–pio5 | 90 | 3.97 | 0.40 | 11.3 |
|      |     |       | 3.08 | 146 | umc1273–pio6 | 142 | 10.07 | 0.25 | 14.5 |
|      |     |       | 4.03/04 | 158/164 | adh2–umc2061 | 50 | 4.92 | -0.29 | 16.0 |
|      |     |       | 6.00/02 | 260/271 | umc1018–umc1083 | 40 | 9.24 | -0.28 | 18.1 |
|      |     |       | 7.02 | 298 | pio9–umc2327 | 66 | 4.53 | -0.13 | 3.3 |
|      |     |       | 9.07/08 | 391 | bnlg1375–umc1505 | 112 | 4.34 | 0.28 | 12.8 |
|      |     |       | 10.06 | 415 | CrtRB1–bnlg1028 | 54 | 15.62 | -0.53 | 34.2 |
|      | PC3 | 68.0 | 2.00/02 | 68/69 | umc2246–bnlg1017 | 0 | 3.78 | -0.18 | 11.2 |
|      |     |       | 2.04 | 77 | umc1465–pio4 | 64 | 9.14 | -0.30 | 29.5 |
|      |     |       | 4.06 | 172/181 | umc1895–umc2027 | 68 | 4.33 | 0.22 | 19.7 |
|      |     |       | 5.03 | 210/219 | umc2060–umc1692 | 82 | 12.54 | -0.42 | 38.8 |
|      |     |       | 5.06/09 | 251/254 | umc2198–umc2209 | 178 | 4.53 | -0.25 | 22.0 |
|      |     |       | 7.02 | 298/301 | umc2327–phi034 | 74 | 8.60 | -0.33 | 31.7 |
|      |     |       | 8.07/09 | 363/366 | bnlg1828–umc1663 | 94 | 3.56 | 0.25 | 15.1 |
|      |     |       | 9.04/9.05 | 385 | umc1107–umc1094 | 52 | 3.76 | 0.18 | 10.6 |
|      |     |       | 10.02/03 | 392/397 | umc1576–umc1367 | 22 | 4.10 | -0.25 | 16.3 |
| A × S | PC1 | 60.4 | 1.11 | 64 | umc2242–umc1979 | 182 | 5.18 | -0.55 | 4.8 |
|      |     |       | 7.00 | 293–296 | umc1241–umc1068 | 44 | 14.02 | -1.30 | 24.3 |
|      |     |       | 7.04 | 323/325 | umc1944–umc1125 | 122 | 2.93 | 0.49 | 4.8 |
|      |     |       | 9.07 | 391 | umc1675–umc2099 | 138 | 3.81 | -0.95 | 11.9 |
|      |     |       | 9.08 | 391 | zct128–umc1505 | 166 | 11.04 | -1.63 | 23.7 |
|      | PC2 | 31.1 | – | – | pio_7–pio_8 | 6 | 5.79 | 0.63 | 5.1 |
|      |     |       | 5.03 | 212/217 | umc2295–bnlg1892 | 76 | 4.15 | 0.85 | 9.4 |
|      |     |       | 7.04 | 323 | pio_11–umc1944 | 118 | 4.06 | -0.63 | 5.8 |
|      |     |       | 8.02 | 329/345 | umc1034–phi115 | 64 | 3.36 | 0.70 | 6.6 |
|      |     |       | 9.07 | 391 | pio_13–umc1675 | 116 | 3.95 | -1.43 | 20.4 |
|      | PC3 | 40.9 | 2.02 | 70 | umc1934–zca381 | 88 | 3.29 | -0.29 | 6.6 |
|      |     |       | 2.04 | 77 | umc1541–pio_4* | 128 | 8.94 | -0.68 | 23.3 |
|      |     |       | 4.08 | 188/192 | umc2187–umc1559* | 140 | 3.07 | 0.24 | 4.9 |
|      |     |       | 10.06 | 417 | Crtr-B1–umc1993 | 94 | 5.32 | -0.37 | 10.3 |
|      |     |       | Interaction* | – | – | – | -0.31 | 3.1 |
|      | PC4 | 36.1 | 2.08 | 108 | dupssr25–umc1736* | 184 | 4.51 | 0.2 | 4.2 |
|      |     |       | 4.02 | 156 | pio_08–phi295450** | 28 | 4.78 | 0.2 | 6.8 |
|      |     |       | 5.03 | 212/217 | umc2035–umc2295* | 72 | 8.02 | 0.3 | 16.2 |
|      |     |       | 6.02 | 271/276 | umc1178–umc1918** | 34 | 4.23 | 0.2 | 4.6 |
|      |     |       | 6.05 | 285/287 | umc1805–umc1859 | 90 | 6.01 | 0.3 | 11 |

Genetic environment, significance and effect of selected QTL for each trait are listed. Indicated are LOD (Logarithm of Odds), Add (Additive effect) and $R^2$ (%) (coefficient of partial determination).
β-carotene:β-cryptoxanthin ratio, was more fully explained by models including dominance effects than only additive effects (data not shown).

Since parent inbred profiles between A619 and SC55 differed for almost all carotenoids, significant QTL were examined for pleiotropic effects. A major effect located on chromosome 9 (pos. 162–166) significantly affected phytoene (1.33 μg g⁻¹, $R^2_{part} = 19.6\%$), phytofluene (0.25 μg g⁻¹, $R^2_{part} = 19.6\%$), zeaxanthin (0.08 μg g⁻¹, $R^2_{part} = 3.3\%$), lutein (1.74 μg g⁻¹, $R^2_{part} = 18.5\%$), β-cryptoxanthin (0.21 μg g⁻¹, $R^2_{part} = 17.6\%$) and zeaxanthin (0.46 μg g⁻¹, $R^2_{part} = 10.4\%$). Reduction in each of these carotenoid traits was associated with the SC55 allele, which agreed with the low carotenoid profile of inbred SC55.

Pleiotropic effects were observed within pathway branches. A major effect QTL on chromosome 7 (pos. 46) was found to explain 23.8 % of the variation in phytoene and 32.5 % of the variation in phytofluene, with the A619 allele associated with higher levels. A QTL at chromosome 5 (pos. 70–76) augmented total colored carotenoids (1.00 μg g⁻¹, $R^2_{part} = 8.1\%$) and β-carotene (0.38 μg g⁻¹, $R^2_{part} = 17.5\%$) through the SC55 allele, and also decreased colorless:colored carotenoids through the same allele (0.12 μg g⁻¹, $R^2_{part} = 7.3\%$). Competition within the β-branch appeared to be mediated by a QTL on chromosome 10 (pos. 88–94), associated with increased β-carotene (0.37 μg g⁻¹, $R^2_{part} = 14.0\%$) and β-carotene:β-cryptoxanthin (ratio effect = 4.57, $R^2_{part} = 40\%$), and decreased β-cryptoxanthin (0.21 μg g⁻¹, $R^2_{part} = 17.6\%$) and zeaxanthin (0.46 μg g⁻¹, $R^2_{part} = 10.4\%$) through the SC55 allele. α-branch carotenoids were significantly affected by a QTL located on chromosome 8 (pos. 88–92), which accounted for significant variation in α-carotene, lutein and α/β branch ratio.

Use of allele-specific markers for specific carotenoid biosynthesis genes permitted the comparison of gene map locations with detected QTL effects in this population. Lycopene epsilon cyclase (lcy) was mapped to chromosome 8, 92 cM, coinciding with trait effects for several α-branch traits including α-carotene (pos. 88), lutein (pos. 88) and α/β branch ratio (pos. 92). Beta-carotene hydroxylase (crtRB1) mapped to chromosome 10, 93 cM, close to QTL effects for β-cryptoxanthin (pos. 88), zeaxanthin (pos. 88), β-carotene:β-cryptoxanthin (pos. 88) and β-caroten (pos. 94). Carotenoid cleavage dioxygenase 1 (cdd1) mapped to chromosome 9, 145 cM, between two clusters of QTL at pos. 136 (affecting lutein and zeaxanthin) and pos. 162–166 (affecting phytoene, phytofluene, β-cryptoxanthin, zeaxanthin, zeinoxanthin, lutein and total carotenoids).

Genetic effects accounting for variation in four multivariate traits (PCs) were mapped in A619 × SC55. QTL associated with variation in all PC traits consisted of

![Diagram](image)

**Fig. 3** Network of metabolic QTL influencing grain carotenoid concentrations in populations DEexp × C17 and A619 × SC55. QTL are shown for univariate (single carotenoid concentration) and multivariate (PC) traits mapping to locations of genes known to be in the carotenoid or precursor pathways (overlap delimited by black squares). Correlated trait effects are shown for all PC traits, where colored squares indicate the carotenoids which are best described by a given PC; the presence of only blue squares shows that these carotenoid variation described by a single PC covaries proportionally, whereas presence of blue and red squares indicates an inverse covariation between the carotenoids described by that PC (color figure online).
models with 4–5 main effect QTL and few epistatic interactions (Table 5). Models for multivariate traits were found to account for more phenotypic variation (range 31.1–60.4 %) than the univariate carotenoid traits (21.6–56.2 %). Variation in PC1, consisting of significant and positive loadings for phytoene, lutein and zeaxanthin, was explained by five QTL with positive effects predominantly originating from the A619 genome. The QTL at chromosome 9 (pos. 166) with large, pleiotropic effects were observed to affect this PC trait. The PC2 trait, describing antagonism between colorless and colored carotenoids was explained by five QTL, including one observed to have pleiotropic effects on β-branch carotenoids in univariate analysis at chromosome 5 (pos. 76). The PC3 trait, which describes antagonistic variation between zeaxanthin with lutein and β-carotene, was affected by four QTL, one which mapped in close proximity to the crtRB1 locus on chromosome 10, and another which affected β-branch traits on chromosome 2 (pos. 128). Competition between α- and β-branches, described by PC4, was affected by five main effect QTL and two digenic interactions. Positive effects at all of these QTL were small, and were contributed by SC55 (Table 5). Univariate and multivariate traits overlapped extensively, as shown in Supplemental Figures 6 and 7.

A range of 3–7 loci accounted for 8.2–36.5 % of the variation in β-carotene isomer concentration and proportion traits in population A619 × SC55 (Supplemental Table 6). As observed in population DEexp × CI7, several loci cosegregated with variation in more than one isomer concentration including: chromosome 5 (pos 72, 13-cis and all-trans), 10 (pos 92, CrtRB1 marker; 13-cis and all-trans). Loci with pleiotropic effects on β-carotene isomer proportion were identified on chromosomes 1 (pos 12; 9-cis, all-trans), 4 (pos 186; 9-cis, all-trans), 5 (pos 72; 9-cis, 15-cis, all-trans) and 6 (pos 34-38; 9-cis, 15-cis, all-trans).

Comparison of detected QTL to known biosynthesis genes

A total of 63 main effect QTL for carotenoid univariate traits were found to span 29 marker intervals for the DEexp × CI7 F_{2:3} population in the Mexico environment, in contrast to 22 unique main effects for the multivariate principal component traits. For the A619 × SC55 population, a total of 50 main effect QTL across 29 marker intervals were found to explain phenotypic variation in univariate traits, as compared to 19 main effect QTL for multivariate traits across 18 marker intervals. The map locations for univariate and PC traits often overlapped, but QTL that were unique to PC traits were detected in both populations (Supplemental Figure 6 and 7). Many of the detected QTL clusters were found within proximity of mapped carotenoid biosynthesis genes (Andorf et al. 2010; Supplemental Table 6). Although genetic maps for both populations did not provide sufficient resolution to definitively conclude that the known genes and detected QTL mapped within proximity to the same environment (as gauged by chromosome bin assignment), inferences for these gene-phenotype relationships were strengthened by comparing the affected traits with the known metabolic functions of previously identified genes, overlapping detection between univariate and multivariate traits, and detection in previous mapping studies (Fig. 3).

Overall substrate increases to the carotenoid pathway in population DEexp × CI7 were described by PC2, which mapped to chromosome bins also containing genes involved in carbon substrate allocation to the MEP pathway (DXR, bin 3.04) and isoprenoid substrate allocation to the carotenoid pathway (PSY1, bin 6.01). PC3 described the balance between colorless carotenoids and colored pathway intermediates, and mapped near genes involved with carotenoid conversion within the main branch (ZDS, bin 7.02) in addition to conversion of main branch carotenoids to α and β branch carotenoids (LCYβ, bin 5.04). PSY2, an isoform of phytoene synthase in bin 8.07, was also found to map near PC3 QTL. The largest trait variation was observed for lutein, and was represented by the PC1 trait. This PC trait mapped to chromosomes bins containing DXR (bin 3.04) and LCYε (bin 8.05), both of which are involved in carotenoid synthesis by committing carbon substrate to the carotenoid pathway, and to the α-branch, respectively.

Multivariate traits for A619 × SC55 also described overall substrate increases to the pathway through PC1 and allocation between colorless substrate and colored intermediates through PC2. These traits mapped close to genes that are involved in the allocation of substrate precursors to the carotenoid pathway (HDS, bin 5.03; GGPPS1, bin 7.04) as well as one that removes carotenoids from the pathway through degradation (CCD1/wcl1, bin 9.07). Balance between the accumulation of β-carotene and zeaxanthin, explained by PC3, had associated QTL mapping proximal to genes directly involved in zeaxanthin metabolism including crtRB1(bin 10.06) and ZEP1(bin 2.04) as well as carotenoid conversion by CrtISO1 (bin 4.08). PC4 described competition between the α- and β-pathway branches; although QTL for this trait were found to map within the same chromosome bins containing genes allocating substrate to the entire pathway (GGPPS1, bin 2.08; DXX1, bin 6.05).

Discussion

Inferences to metabolic functions for detected QTL

The genetic architecture for high β-carotene and total carotenoids as well as various individual and compositional traits was investigated in two F_{2:3} populations,
DEexp × CI7 and A619 × SC55. Through map location comparisons of the detected QTL with those reported for genes of the MEP, isoprenoid, and carotenoid pathways, we concluded that a large proportion of the effects detected in this study are likely derived from metabolic QTL. Confidence in the association of known biosynthesis genes with detected QTL was highest when the trait affected by the QTL was known to be a product or substrate of the associated biosynthesis gene (Fig. 3). Major effect QTL previously identified in other genetic mapping studies also support the association between variation in specific carotenoid traits with locations of putative metabolic QTL (Fig. 3; Islam 2004; Wong et al. 2004; Chander et al. 2008; Harjes et al. 2008). Metabolic function has been confirmed by association analyses for several of these loci including lcy (Harjes et al. 2008), crtRB1 (Yan et al. 2010) and CCD1 (Kandianis and Rocheford, data not shown). We emphasize, however, that inferences should be confirmed through additional genetic analyses utilizing functional markers that are in complete and near complete linkage disequilibrium with the alleles associated with trait variation (Andersen and Rubsterstedt 2003), and/or through the use of genetic mapping with high density, high throughput markers.

Use of trait correlations in QTL analyses has been reported to result in increased power of QTL resolution, increased precision of parameter estimates, and the simplification of simultaneous hypothesis testing for multiple traits (Gilbert and Le Roy 2004). Multitrait mapping approaches using multivariate techniques (Korol et al. 1995) or transformation of correlated traits to canonical variables which are then used univariately (Mangin et al. 1998) have also been successfully employed. As information describing the size and system of effects from pleiotropic QTL could be lost through the estimation of single trait effects (Jiang and Zeng 1995), there was reason to attempt QTL discovery in this system of correlated traits by pairing univariate and multivariate analyses. Principal component (PC) traits identified using the variance–covariance matrix for eight carotenoids were mapped by composite interval analysis, leading to the detection of 2–8 QTL for each principal component. Using this strategy, we were able to identify loci simultaneously affecting multiple univariate carotenoid traits, which was suggestive of a pleiotropic QTL. As such, functional associations for the detected suite of QTL suggest that carotenoid concentrations are heavily influenced by (1) the allocation of carbon substrate to the carotenoid pathway, and (2) removal of carotenoids from the pathway through CCD1-facilitated degradation or ZEP1-mediated conversion. In contrast, β-carotene concentrations are largely attributed to (1) the allocation of carotenoid substrate from the main branch supplied by phytoene through the bifurcation point at lycopene, (2) competition between α- and β-branches and (3) conversion within the β-branch.
been driven by the potential to form provitamin A, and particularly β-carotene (Harjes et al. 2008; Welsch et al. 2010; Yan et al. 2010; Ye and Beyer 2000; Babu et al. 2013). As not all proV A carotenoids are equally absorbed and converted to vitamin A (Tanumihardjo et al. 2010; Davis et al. 2008), proV A bioavailability has emerged as a nutritionally valued trait of similar importance to total proV A concentration.

This study describes natural variation in the β-carotene isomers of maize grain, proV A carotenoids which have been shown to differ in bioavailability. β-carotene in foods is often reported as the compilation of one or more β-carotene isomers, including the most common all-trans form, and the less prevalent but more oil-soluble 9-, 13- and 15-cis isomers (Schieber and Carle 2005). cis- Isomerization of β-carotene is noted to be an artifact of food processing (Achir et al. 2011; Mordi et al. 1993), but also naturally occurs to varying degrees in foods (Khoo et al. 2011). The isomeric state of dietary β-carotene is nutritionally important as its conformation considerably influences its bioavailability. trans isomers of β-carotene are preferentially absorbed (Deming et al. 2002; Stahl et al. 1995). Once absorbed, the 9-cis isomer of β-carotene is primarily converted to all-trans forms of vitamin A (Maeda et al. 2011). Regarding provitamin A activities, however, 13-cis and 9-cis β-carotene only maintain 53 and 38 % of the activity conferred by trans β-carotene, respectively (Schieber and Carle 2005).

Results from this study reveal a genetic basis for β-carotene isomerization in maize grain, specifically for traits of concentration and proportion of 9-cis, 13-cis, 15-cis and all-trans β-carotene. As of yet, there is no evidence for enzymatic conversion of trans β-carotene or other carotenoid precursors to cis β-carotene. Considering that carotenoids are antioxidants, it is possible that isomerization of trans β-carotene under light, heat or oxidative stress could give rise to sizable amounts of cis β-carotene. It is also possible that β-carotene isomerization could indeed be facilitated by an isomerase, similar to those which facilitate isomerization of pro-lycopene carotenoids (Chen et al. 2010; Isaacson et al. 2004). This implies that genetic factors influencing cis β-carotene accumulation could include: (1) metabolic QTL affecting synthesis or degradation of trans β-carotene; (2) as of yet unknown metabolic QTL specific to the formation of cis β-carotene and; (3) factors conferring molecular stability of trans β-carotene which could influence the degree to which non-enzymatic trans–cis isomerization occurs, perhaps with changing cellular conditions. Accordingly, we found several QTL with pleiotropic effects for more than one isomer trait to colocalize with known metabolic candidates including ZEP1 (Chr. 2), LCYβ (Chr. 5), LCYε (Chr. 8), and CrtrB1 (Chr. 10).

Recommendations for future carotenoid breeding studies

The QTL results presented here are consistent with the concept that modification of grain carotenoid profiles primarily originates from variation in the carotenoid biosynthesis pathway, or metabolism that directly creates and allocates substrate to the carotenoid pathway. Association mapping of sequence variation in candidate genes from the carotenoid biosynthesis pathway has revealed that carotenoid composition is significantly altered by genes regulating α- and β-branch allocation through LyCε and β-branch conversion through CrtrB1. From the associations of mapped QTL effects with map locations of known biosynthesis genes, there is reason to expect that increased total carotenoids can be achieved through altered metabolism, and that biochemical reactions upstream of the carotenoid pathway must be targeted to attain this increase. Characterizing allelic variation representative of MEP and isoprenoid pathways through loci such as DXR should be highly informative and useful in association analysis tests and marker-assisted selection studies for total carotenoid grain profiles. Some of the genes in these pathways were detected by PCs representing significant and positive loadings of the major colored carotenoids and phytene; however, in the case where phytene and colored carotenoids negatively covary, it may be necessary to include measurement of colorless carotenoids (phytene and phytofluene) for further genetic analyses.

Factors contributing to efflux from the carotenoid pathway also appear to affect total carotenoid accumulation, through select carotenoids on a specific pathway branch (ZEP) or through non-specific reduction of nearly all carotenoids (CCD1). Compositional differences have been achieved through modification of intra-pathway steps including cyclization reactions (LCYε, LCYβ) and hydroxylation reactions (CRtrB1). Given that most of the phenotypic variation for grain carotenoids in maize exists in lutein, zeaxanthin, zeinoxanthin, β-cryptoxanthin and β-carotene, the steps identified here primarily affect these intermediates and will likely account for most of the phenotypic variation upon selection of the desired alleles. Marker-assisted selection strategies for β-carotene and provitamin A concentrations are already employing variation at LyCε and CrtrB1 to attain desired carotenoid profiles. Given the results of this study, there is merit to further investigate the ZEP locus and LyCε for compositional modification, and to evaluate the influence of environment on the metabolic activity of these pathway enzymes.

Acknowledgments We thank DuPont Pioneer for in-kind genotyping, Robert Meeley of DuPont Pioneer for additional A619 and SC55 sequence data, and Don McCarty for sequence information relative to CCD1/wc1. We also thank Ines Carvalho and Patricia Bermudez for their assistance in data assembly and quality control, Jerry Chandler.
and Don Roberts for their excellent field support, and Michael Grusak and G. R. Johnson for critical review of this manuscript. Funding for this project was provided by HarvestPlus, National Science Foundation (NSF) DBI-0922493 and Patterson Chair Endowment to T.R.

Conflict of interest The authors declare that they have no conflict of interest.

Ethical standards Experiments comply with the current laws of the United States of America and Mexico.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References

Achir N, Pénicaud C, Avallone S, Bohuon P (2011) Insight into β-carotene thermal degradation in oils with multiresponse modeling. JAOCSc 88(12):2035–2045

Andersen JR, Lubberstedt T (2003) Functional markers in plants. Trends Plant Sci 8:554–560

Andorf CM, Lawrence CJ, Harper LC, Schaeffer ML, Campbell DA, Sen TZ (2010) The Locus Lookup tool at MaizeGDB: Identification of genomic regions in maize by integrating sequence information with physical and genetic maps. Bioinformatics 26(3):434–436. Art No bp556

Auldridge ME, Block A, Vogel JT, Dahney-Smith C, Mila L, Bouzayen M, Magallanes-Lundback M, DellaPenna D, McCarty DR, Klee HJ (2006) Characterization of three members of the Arabidopsis carotenoid cleavage dioxygenase family demonstrates the divergent roles of this multifunctional enzyme family. Plant J 45:982–993

Babu R, Rojas NP, Gao S, Yan J, Pixley K (2013) Validation of the effects of molecular marker polymorphisms in LcyE and CrtrB1 on provitamin A concentrations for 26 tropical maize populations. Theor Appl Genet 126:389–399

Bauermeier JD, Adams CR, Marusich WL (1981) Carotenoids as colorants and vitamin A precursors. Academic Press, New York

Bouis HE, Hotz C, McClafferty B, Meenakshi JV, Pfeiffer WH (2011) Biofortification: a new tool to reduce micronutrient malnutrition. Food Nutr Bull 32(1 Suppl):S31–S40

Buckner B, Kelso TL, Robertson DS (1990) Cloning of the Y1 locus of maize, a gene involved in the biosynthesis of β-carotene. Plant Cell 2:867–876

Buckner B, Miguel PS, JanickBuckner D, Bennezet JL (1996) The Y1 locus gene of maize codes for phytoene synthase. Genetics 143:479–488

Chander S, Guo YQ, Yang XH, Zhang J, Lu XQ, Yan JB, Song TM, Rocheford TR, Li JS (2008) Using molecular markers to identify two major loci controlling carotenoid contents in maize grain. Theor Appl Genet 116:223–233

Chandler K, Lipka AE, Owens BF, Li H, Buckler ES, Rocheford T, Gore MA (2013) Genetic analysis of visually scored orange kernel color in maize. Crop Sci 53(1):189–200

Chen Y, Li F, Wurtzel ET (2010) Isolation and characterization of the Z1-ISO gene encoding a missing component of carotenoid biosynthesis in plants. Plant Physiol 153(1):66–79

Chuaquir AJ, Rotstein NP, SanGiovanni JP, During A, Chew EY, Politi LE (2007) Lutein and zeaxanthin protect photoreceptors from apoptosis induced by oxidative stress: relation with docosahexaenoic acid. Invest Ophthalmol Vis Sci 48:5168–5177

Davis C, Jing H, Howe JA, Rocheford T, Tanumihardjo SA (2008) beta-Cryptoxanthin from supplements or carotenoid-enhanced maize maintains liver vitamin A in Mongolian gerbils (Meriones unguiculatus) better than or equal to beta-carotene supplements. Br J Nutr 100:786–793

Deming DM, Teixeira SR, Erdman JW (2002) All-trans β-carotene appears to be more bioavailable than 9-cis or 13-cis β-carotene in gerbils given single oral doses of each isomer. J Nutr 132(9):2709–2712

Doebley JF, Gaut BS, Smith BD (2006) The molecular genetics of crop domestication. Cell 127:1309–1321

Doege RW, Churchill GA (1996) Permutation tests for multiple loci affecting a quantitative trait. Genetics 142:249–259

Egesel CO, Wong JG, Lambert RJ, Rocheford TR (2003) Combining ability of maize inbreds for carotenoids and tocopherols. Crop Sci 43:818–823

Estevez JM, Cantero A, Reindl A, Reichler S, Leon P (2001) 1-deoxy-D-xylulose-5-phosphate synthase, a limiting enzyme for plastidic isoprenoid biosynthesis in plants. J Biol Chem 276:22901–22909

Fierce Y, Vieira MdM, Piantedosi R, Wyyss A, Blaner WS, Paik J (2008) In vitro and in vivo characterization of retinoid synthesis from beta-carotene. Arch Biochem Biophys 472:126–138

Galpaz N, Wang Q, Menda N, Zamir D, Hirschberg J (2008) Abscisic acid deficiency in the tomato mutant high-pigment 3 leading to increased plastid number and higher fruit lycopene content. Plant J 53:717–730

Gilbert H, Le Roy P (2004) Power of three multitrait methods for QTL detection in crossbred populations. Genet Sel Evol 36:347–361

Hable WE, Oishi KK, Schumaker KS (1998) Viviparous-5 encodes phytoene desaturase, an enzyme essential for abscisic acid (ABA) accumulation and seed development in maize. Mol Gen Genet 257:167–176

Haley CS, Knott SA (1992) A simple regression method for mapping quantitative trait loci in line crosses using flanking markers. Heredity 69:315–324

Hallauer AR, Miranda JB (1988) Quantitative genetics in maize breeding. Iowa State University Press, Ames

Harjes CE, Rocheford TR, Bai L, Brutnell TP, Kandianis CB, Sowinski SG, Stapleton AE, Vallabhaneni R, Williams M, Wurtzel ET, Yan J, Buckler ES (2008) Natural genetic variation in lycopene epsilon cyclase tapped for maize biofortification. Science 319:330–333

Howitt CA, Pogson BJ (2006) Carotenoid accumulation and function in seeds and non-green tissues. Plant Cell Environ 29:435–445

Isaacson T, Ohad I, Beyer P, Hirschberg J (2004) Analysis in vitro of phytoene desaturase, an enzyme essential for abscisic acid (ABA) biosynthesis pathway in plants. Plant Physiol 136(4):4246–4255

Islam SN (2004) Survey of carotenoid variation and quantitative trait loci mapping for carotenoid and tocopherol variation in maize. Dissertation or Thesis, University of Illinois at Urbana-Champaign

Jansen RC, Stam P (1994) High-resolution of quantitative traits using RFLP linkage maps. Genetics 136:1447–1455

Jiang CJ, Zeng ZB (1995) Multiple-trait analysis of quantitative trait loci employing correlated trait complexes. Genetics 140:1111–1127

Khoo H-, Prasad KN, Kong K-, Jiang Y, Ismail A (2011) Carotenoids in seeds and non-green tissues. Plant Cell Environ 29:435–445

Kim JS, Nam Y, Blaner WS, Paik J (2007) In vitro and in vivo characterization of retinoid synthesis from beta-carotene. Arch Biochem Biophys 472:126–138

Lander ES, Botstein D (1989) Mapping mendelian factors under quantitative trait loci employing RFLP linkage maps. Genetics 136:1447–1455

Menges JL, Henry CA, Blaner WS, Paik J (2007) In vitro and in vivo characterization of retinoid synthesis from beta-carotene. Arch Biochem Biophys 472:126–138

Sen TZ (2010) The Locus Lookup tool at MaizeGDB: Identifi- cation of genomic regions in maize by integrating sequence information with physical and genetic maps. Bioinformatics 26(3):434–436. Art No bp556

Theor Appl Genet (2013) 126:2879–2895

© Springer
Leuenberger MG, Engeloch-Jarret C, Woggon WD (2001) The reaction mechanism of the enzyme-catalyzed central cleavage of beta-carotene to retinal. Angew Chem Int Ed 40:2614–2617

Li F, Murillo C, Wurtzel ET (2007a) maize Y9 encodes a product essential for 15-cis-zeta-carotene isomerization. Plant Physiol 144:1181–1189

Li S, Tayie FAK, Young MF, Rocheford T, White WS (2007b) Retention of provitamin A carotenoids in high beta-carotene maize (Zea mays) during traditional African household processing. J Agric Food Chem 55(26):10744–10750

Li S, Nugroho A, Rocheford T, White WS (2010) Vitamin A equivalence of the beta-carotene in beta-carotene-biofortified maize porridge consumed by women. Am J Clin Nutr 92:1105–1112

Lichtenhalter HK, Rohmer M, Schwender J (1997) Two independent biochemical pathways for isopentenyl dipiphosphate and isoprenoid biosynthesis in higher plants. Physiol Plant 101:643–652

Lois LM, Rodriguez-Concepcion M, Gallego F, Campos N, Boronat A (2000) Carotenoid biosynthesis during tomato fruit development: regulatory role of 1-deoxy-D-xylulose 5-phosphate synthase. Plant J 22:503–513

Lopez AB, Van Eck J, Conlin BJ, Paolillo DJ, O'Neill J, Li L (2008) Effect of the cauliflower Or transgene on carotenoid accumulation and chromoplast formation in transgenic potato tubers. J Exp Bot 59:213–223

Maeda T, Perusek L, Palczewski K, Von Linig J (2011) Dietary 9-cis-β, β-carotene fails to rescue vision in mouse models of leber congenital amaurosis. Mol Pharmacol 80(5):941–952

Mangelsdorf PC, Fraps GS (1931) A direct quantitative relationship between vitamin A and the number of genes for yellow pigmentation. Science 73:241–242

Mangin B, Thoquet P, Grimsley N (1998) Pleiotropic QTL analysis. Plant J 106(7):1326–1332

Matthews PD, Luo RB, Wurtzel ET (2003) Maize phytoene desaturase and zeta-carotene desaturase catalyse a poly-Z desaturation pathway: implications for genetic engineering of carotenoid content among cereal crops. J Exp Bot 54:2215–2230

Mikkilineni V, Rocheford TR (2003) Sequence variation and genomic organization of fatty acid desaturase-2 (fad2) and fatty acid desaturase-6 (fad6) DNAs in maize. Theor Appl Genet 106(7):1326–1332

Mordi RC, Walton JC, Burton GW, Hughes L, Ingold KU, Lindsay DA, Moffatt DJ (1993) Oxidative degradation of β-carotene and β-Apo-8'-carotenal. Tetrahedron 49(4):911–928

Ooijen JWV, Voorrips RE (2001) JOINMAP version 3.0, software for the calculation of genetic linkage maps. Plant Research International, Wageningen

Palaisa K, Morgante M, Tingey S, Rafalski A (2004) Long-range selection reveals the 5-cis geometrical isomer. J Nat Genet 125(8):2128–2133

Phillips RJ (2001) Carotenoids A, infection, and immune function. Annu Rev Nutr 21:167–192

Pombo BR, Gallacher AE (1993) PLABQTL: a computer program to map QTL 1.2

Schieber A, Carle R (2005) Occurrence of carotenoid cis-isomers in food: technological, analytical, and nutritional implications. Trends Food Sci Technol 16(9):416–422

Steddon JM, Ajani UA, Sperduto RD, Hiller R, Blair N, Burton TC, Farber MD, Gragoudas ES, Haller J, Miller DT, Yannuzzi LA, Willett W (1994) Dietary carotenoids, vitamin-A, vitamin-C, and vitamin-E, and advanced age-related macular degeneration. JAMA 272:1413–1420

Singh M, Lewis PE, Hardeman K, Bai L, Rose JKC, Mazourek M, Chomet P, Brunnell TP (2003) Activator mutagenesis of the pink scutellum/1/viviparous7 locus of maize. Plant Cell 15:874–884

Stahl W, Schwarz W, Von Laar J, Sies H (1995) All-trans β-carotene preferentially accumulates in human chylomicrons and very low density lipoproteins compared with the 9-cis geometrical isomer. J Nutr 125(8):2128–2133

Stephens CB (2001) Vitamin A, infection, and immune function. Annu Rev Nutr 21:167–192

Stevens RL (2007) Genetic and QTL analysis of carotenoid variation in two mapping populations of maize. Dissertation or Thesis, University of Illinois at Urbana-Champaign

Tang G, Qin J, Dolnikowski GG, Russell RM, Grusak MA (2009) Golden rice is an effective source of vitamin A. Am J Clin Nutr 89:1776–1783

Tanumihardjo SA, Palacios N, Piixley KV (2010) Provitamin a carotenoid bioavailability: what really matters? Int J Vitam Nutr Res 80(4–5):336–350

Utz HF, Melchinger AE (1993) PLABQTL: a computer program to map QTL 1.2

Vallabhaneni R, Wurtzel ET (2009) Timing and biosynthetic potential for carotenoid accumulation in genetically diverse germplasm of maize. Plant Physiol 150:562–572

Vallabhaneni R, Gallagher CE, Lucciardello N, Cuttriss AJ, Quinlan RF, Wurtzel ET (2009) Metabolite sorting of a germplasm collection reveals the Hydroxylase 3 locus as a new target for maize provitamin A biofortification. Plant Physiol 151(3):1635–1645

Weber EJ (1987) Carotenoids and tocots of corn grain determined by HPLC. J Am Oil Chem Soc 64:1129–1134

Welsch R, Arango J, Bär C, Salazar B, Al-Babili S, Beltrán J, Chavarriaga P, Ceballos H, Tohme J, Beyera P (2010) Provitamin a accumulation in cassava (Manihot esculenta) roots driven by a single nucleotide polymorphism in a phytoene synthase gene. Plant Cell 22(10):3348–3356

Whitner JP, Srinivasan M, Upadhayay MP (2001) Cornal blindness: a global perspective. Bull World Health Organ 79:214–221

Wong JC, Lambert RJ, Wurtzel ET, Rocheford TR (2004) QTL and candidate genes phytoene synthase and zeta-carotene desaturase associated with the accumulation of carotenoids in maize. Theor Appl Genet 108:349–359

Yan J, Kandianis CB, Harjes CE, Bai L, Kim E, Yang X, Skinner DJ, RF, Wurtzel ET (2009) Corneal blindness: a global perspective. Bull World Health Organ 79:214–221

Ye X, Beyer P (2000) Engineering the provitamin A (β-carotene) biosynthetic pathway into (carotenoid-free) rice endosperm. Science 287(5451):303–305

Zeng ZB (1994) Precision mapping of quantitative trait loci. Genetics 136:1457–1468