Arabidopsis VARIEGATED 3 encodes a chloroplast-targeted, zinc-finger protein required for chloroplast and palisade cell development

Henrik Næstad1, Agnethe Holm1, Tom Jenkins1, H. Bjørn Nielsen2, Cassandra A. Harris3, Michael H. Beale3, Mathias Andersen4, Alexandra Mant5, Henrik Scheller6, Bilal Camara6, Ole Mattsson1 and John Mundy1,*

1Institute of Molecular Biology, Copenhagen University, Øster Farimagsgade 2A, 1353 Copenhagen K, Denmark
2BioCentrum, Technical University of Denmark, Building 208, 2800 Lyngby, Denmark
3Long Ashton Research Station, University of Bristol, Weston Road, Bristol, BS41 9AF, UK
4Research Center Foulum, PO Box 50, 8839 Tjele, Denmark
5Institute of Plant Biology, Royal Veterinary and Agricultural University, Thorvaldsensvej 40, 1871 Frederiksberg, Denmark
6CNRS, Institut de Biologie Moleculaire, Rue de General Zimmer 12, 67084 Strasbourg, France

*Author for correspondence (e-mail: mundy@biobase.dk)

Summary
The stable, recessive Arabidopsis variegated 3 (var3) mutant exhibits a variegated phenotype due to somatic areas lacking or containing developmentally retarded chloroplasts and greatly reduced numbers of palisade cells. The VAR3 gene, isolated by transposon tagging, encodes the 85.9 kDa VAR3 protein containing novel repeats and zinc fingers described as protein interaction domains. VAR3 interacts specifically in yeast and in vitro with NCED4, a putative polyene chain or carotenoid dioxygenase, and both VAR3 and NCED4 accumulate in the chloroplast stroma. Metabolic profiling demonstrates that pigment profiles are qualitatively similar in wild type and var3, although var3 accumulates lower levels of chlorophylls and carotenoids. These results indicate that VAR3 is a part of a protein complex required for normal chloroplast and palisade cell development.

Key words: VAR3, NCED4, Transposon tagging, Chloroplast development, Stromal subcellular localization

Introduction
Chloroplast functions are dependent upon proteins encoded both within the plastid and nuclear genomes. Nuclear genes whose mutations affect overall chloroplast function encode proteins involved in photosynthesis, plastid metabolism, chloroplast and host cell biogenesis, or in nuclear-chloroplast traffic and signaling. Signaling between the nucleus and chloroplasts regulates both plastid and nuclear gene transcription (reviewed by Mullet, 1993; Barkan et al., 1995). An example of chloroplast-to-nucleus signaling is the decreased transcription of nuclear Lhcb (CAB) and RbcS genes when plants are treated with the carotenoid biosynthetic inhibitor norflurazon that leads to photo-oxidized, nonfunctional plastids. However, gun (genome uncoupled) mutants constitutively express Lhcb genes under such conditions. Analysis of gun mutants indicates that the tetrapyrrole intermediate Mg-protoporphyrin (Mg-ProtoIX) acts as a chloroplast-to-nucleus signal that regulates expression of nuclear genes encoding chloroplast-targeted proteins (Strand et al., 2003).

Several classes of mutation result in variegated plants that have leaves consisting of normal green and also white or yellow sectors in which chloroplast development is retarded or disrupted. For example, variegation is seen in plants with unstable mutations that affect chloroplast function, such as those due to nuclear transposition events (Chatterjee et al., 1996; Keddie et al., 1996). In addition, variegation may be caused by the induction of defective plastids in some cells by stable, loss-of-function mutations in nuclear genes (reviewed by Rodermel, 2001; Sakamoto, 2003). Such mutant alleles that have been characterized in Arabidopsis include chloroplast mutator [chm (Abdelnoor et al., 2003)], yellow variegated 1 [var1: Sakamoto et al., 2002] and var2 (Takechi et al., 2000) and immutans (im). VAR1 and VAR2 encode thylakoid proteins related to the bacterial chaperone metalloprotease FtsH. FtsH represents a family of 12 different nuclear genes, nine of which encode FtsH proteins located in the chloroplast. However, only mutations of VAR1 or VAR2 have been shown to cause typical leaf variegation and sensitivity to photoinhibition. This and other data indicate that VAR1 and VAR2 play important roles in the repair cycle of photosystem II thylakoid membranes (Sakamoto et al., 2003).

The formation of albino sectors in the im mutant is enhanced by increased light intensity. IM is a chloroplast-targeted, alternative oxidase involved in electron transfer associated with the desaturation of phytoene, a carotenoid intermediate, via phytoene desaturase (Carol et al., 1999; Wu et al., 1999). Carotenoids function both as accessory pigments in the photosynthetic apparatus, and in photo-protection by quenching reactive oxygen species. Thus, loss of IM function renders plastids susceptible to photo-oxidation. Interestingly, the promoter of nuclear-encoded phytoene desaturase is induced by
inhibitors of chlorophyll and carotenoid biosynthesis (Corona et al., 1996). This suggests that carotenoids serve as plastid signals regulating nuclear gene expression.

Genetic lesions affecting chloroplast development do not always alter leaf morphology. However, Arabidopsis im and several non-variegated recessive mutations [pac1 (Reiter et al., 1994); clal (Crowell et al., 2003); cuel (Streetfield et al., 1999)] exhibit perturbed chloroplast and leaf development, including failure in palisade cell expansion. As these genes encode chloroplast proteins, mutations in them may affect cell development because of incomplete chloroplast biogenesis. While most such mutations lead to decreased Lhcb and RbcS mRNA levels, pac1 accumulates normal levels of Lhcb mRNA. This indicates that pathways mediating plastid developmental signals are separable from pathways that regulate expression of nuclear-encoded photosynthesis genes.

We describe a recessive, stably tagged Arabidopsis mutant, designated variegated 3 (var3), that exhibits variegation and markedly reduced palisade cell numbers, but normal levels of Lhcb and RbcS mRNAs. The VAR3 gene and corresponding full-length cDNA were isolated and shown to encode the novel VAR3 zinc-finger protein. Genetic, molecular and biochemical analyses indicate that VAR3 interacts with, and may thereby affect the activity of, chloroplast localized enzymes.

Materials and Methods
Isolation and characterization of var3
F3 progeny of transposants generated in ecotype Ler (Sundaresan et al., 1995) were examined for visible mutant phenotypes by growing 12 plants per line in soil. Scanning electron microscopy (SEM) of leaves was performed using standard protocols. For transmission electron microscopy (TEM), leaves were fixed overnight at 5°C in 0.1 M phosphate-buffered saline (pH 7.0) 2.5% glutaraldehyde and 2% paraformaldehyde, postfixed in 1% OsO4, dehydrated in acetone, followed by infiltration with epoxy resin (SPURR). Ultra thin sections were put on Formvar-covered whole grids and examined in a Jeol 100CX microscope.

Gene expression in var3
Total RNA from 2-week-old wild-type and var3 plantlets exhibiting extensive variegation under constant illumination (435 μmol m⁻² s⁻¹) was prepared for mRNA blot hybridizations and reverse transcription (RT) using standard protocols (RNA-gents total RNA; Promega). RT-PCR was performed with a kit (Perkin Elmer, N808-0069) using an RT primer (5’CATTTTCGTGTCGTTCAGAGCA) and forward primer (5’ATGAAACCTCACCAGACTCTCCTCC). EF-1α control primers were 5’GTTCATACCAATTTGTTGTCATGG and 5’GAGTACTTGGGGGTAGTGCCGTACC.

For photosynthetic gene expression analysis under photo-oxidative conditions, total RNA was prepared from Arabidopsis Ler and var3 seedlings according to the method of Logemann et al. (Logemann et al., 1987) and grown for 10 days in continuous white light (435 μmol m⁻² s⁻¹) on 1x MS medium, 2% sucrose plates with or without 5 μM norflurazon. For RT-PCR, 1 μg of template total RNA was DNase treated according to manufacturer’s protocols ( Gibco-BRL). Lhcb1 primers for RT-PCR were 5’GGTTACAGTGCAGTGAAGAGAC and 5’ACACTACAGACAGACATG. RT-PCR was performed with 25 cycles to ensure that products would remain within the linear range. EF-1α control primers were again used for the control, and RT-PCR was performed with 27 cycles of amplification.

Standard protocols for microarray analysis by the Arabidopsis Functional Genomics Consortium were followed (afgc.stanford.edu/afgc_html/AFGCProtocols-fev2001.pdf), and the var3 hybridization data is available under proposal #944-DCA-505-GEJ-7. A data set (http://genome-www4.stanford.edu/MicroArray/SMD/) of intensity signals from 14 wild-type arrays (CH1D_MEAN values of ExpID 11595, 10658, 11601, 10651, 11592, 9755, 10653, and CH2D_MEAN values of ExpID 11598, 10659, 11602, 10652, 11594, 8031, 10654) was used to estimate gene-specific variation in wild-type hybridizations, and two arrays for comparing var3 mutant and wild type (ExpID 10653 and 10654, CH1D_MEAN and CH2D_MEAN).

All data are from array print batch CIW-2_05-00. This data set was normalized by fitting a smoothed spline to the correlation between the cumulative density distribution of a virtual average array (one channel) and a given array channel. The spline function was subsequently applied to the array data points (Workmann et al., 2002). The 14 wild-type arrays exhibited consistent intensity measurements and allowed estimates of gene-specific standard deviation (std14Xwt). The geometric mean of the z-like score (int4Xwt/int2Xwt/std14Xwt) from the two arrays was then used to rank the genes in Fig. 2C.

Pigment profiling
Wild-type and var3 seeds were germinated on MS medium containing 3% sucrose, and incubated at 22°C under constant light (435 μmol m⁻² s⁻¹). Approximately 140 seedlings were harvested at the 2-4 leaf stage, freeze-dried, and five replicate methanol extractions prepared for each type of seedling using the method of Fraser et al. (Fraser et al., 2001). Extracts were analyzed for carotenoids and chlorophylls by HPLC with a photo-diode array detector according to Bramley (Bramley, 1992). Peak assignments were made by co-injection with authentic standards. Unknown carotenoids and chlorophylls were suggested by the UV spectra of the peaks.

VAR3 gene, cDNA and var3 complementation
DNA for inverse PCR and Southern blotting was isolated with the DENasy kit (Qiagen). Genomic sequence flanking the Ds insertion was amplified by inverse PCR. 50 ng of var3 DNA was digested with EcoRI (20 μl volume) and self-ligated with E. coli T4 ligase (New England Biolabs). Two μl of the ligation was used as template with transposon/GUS primers (5’CCAGAATCTGATCCCAACGCCGT-CG and 5’CCGGCGATTGTGAGCAGAGAGG). The VAR3 gene was annotated from sequences surrounding the Ds insertion site using BLAST (www.ncbi.nlm.nih.gov/BLAST), NetPlantGene (cbs.dtu.dk/services/NetPGen), Psort (psort.nibb.ac.jp), ChloroP (cbs.dtu.dk/services/ChloroP), PFSCAN (irec.isb.ch/software/PFSCAN), PFAM (sanger.ac.uk/Software/Pfam/search.shtml), and ClustalW with Bclustal (clustalw.genome.ad.jp). Primers including the predicted VAR3 start and stop codons (5’TATTAAGATCTAGATGAAACTCCACCAGACTCTGACTCCTCC and 5’TATTAAGATCTAGATGAAACTCCACCAGACTCTGACTCCTCC) were used to amplify a var3 cDNA by PCR using λ-phage DNA isolated from a cDNA library (CD4-6, Arabidopsis Biological Resource Center, Ohio, USA) as template with annealing at 55°C for 5 minutes and elongation at 68°C for 5 minutes. A 2277 bp fragment was amplified and cloned into a pCR-blunt vector (Invitrogen, k2700-20) and sequenced using an ABI 3100 sequencer. Complementation of var3 was performed with a genomic fragment of 4,941 bp, including 1,541 bp 5’ to the VAR3 start codon and 539 bp 3’ to the stop codon (nt. 56,462 to 51,521 of clone MVA3, AB006706) isolated by long range PCR using ecotype Landsberg DNA template and primers 5’GACCCAGAAAACAAAGGGTGTTGCGG and 5’CTGTCCTCTCTAAACGAGCGCTTGTTTCCG. The product was cloned in the Smal site of pCAMBIA 3303 carrying a BASTA resistance marker (thaliana.botany.wisc.edu/cambia), and the resultant construct transferred via Agrobacterium into var3 plants by vacuum infiltration (Bechtold et al., 1993). T1 BASTA resistant, wild-type plants were selected and shown to carry the VAR3 transgene, and to be homozygous for the var3 insertion allele by PCR. Segregation analysis of T2 BASTA resistant T1 plants
were homozygous for kanamyacin resistance carried on \textit{Ds}, and that the \textit{var3} phenotype segregated 1:3 wild type.

\textbf{VAR3 localization}

An \textit{NcoI}-linked \textit{VAR3} cDNA fragment containing the ORF without the stop codon was generated by PCR with \textit{Pfu} (Promega) and \textit{VAR3} linker-primers (5'TATTAACCATTGGAATTCTGGCAGTATGAGATG-3', 5'TATTTAACCATAGCTTCCTCTTCCTCTTCCTC; 5'TATTTAACCATTGGAATTCTGGCAGTATGAGATG-3', and the \textit{VAR3} cDNA template. This fragment was cloned in \textit{NcoI} of binary vector \textit{pCAMBIA} 1303 carrying a hygromycin resistance marker, generating a single ORF encoding \textit{VAR3}, \textit{GUS} and GFP under control of the 35S promoter. This construct was transformed via \textit{Agrobacterium tumefaciens} strain GV3101 into wild type and \textit{var3}. T1 plants were selected on hygromycin and analyzed for phenotype, \textit{GUS} activity and GFP fluorescence. Two wild-type lines were also shown by PCR to contain \textit{GUS} carried on \textit{pCAMBIA} 1303 and to be homozygous for \textit{Ds} in \textit{var3}. These lines were analyzed for GFP fluorescence using a Zeiss LSM510 microscope.

\textbf{VAR3 interaction analyses}

A cDNA library was used according to the manufacturer’s instructions (Clontech, FL4000AB). The full-length \textit{VAR3} cDNA bait was cloned as a \textit{BamHI}-linked fragment with primers 5'TATTAAGGACCAGCTTTGAGACGGTTGCAATG-3', 5'TATTAAGGACCAGCTTTGAGACGGTTGCAATG-3', and \textit{VAR3} cDNA template. This fragment was cloned in \textit{BamHI} of binary vector \textit{pGBD-C1} that was introduced into strain PJ69A-4A (James et al., 1996). This strain was transformed with library cDNA and screened for prototrophic growth on medium lacking tryptophan, leucine, histidine and adenine for 4 days at 30°C. A total of 15\textsuperscript{6} transposants generated with a \textit{Ds} element. \textit{VAR3} displayed normal skotomorphogenesis, and the cotyledons developed normally in the light, but the first true leaves exhibited variegation seen as yellow areas (compare Fig. 1A and B). This variegation became indistinct in older leaves so that the only phenotypic difference compared with wild type (\textit{Ler}) at flowering was pale flower bud sepals in \textit{var3} (Fig. 1D and E). Variegation in \textit{var3} differed from that in \textit{im} (Fig. 1B, E compared with C, F) which consisted of pronounced, light-dependent sectoring. Light microscopy of \textit{var3} leaves showed that yellow areas contained few if any palisade mesophyll cells, which resulted in a rough leaf surface (compare Fig. 1G and H). Scanning electron microscopy confirmed that these areas collapsed during critical point drying (Fig. 1J).

To study formation of the yellow areas, \textit{Ler} and \textit{var3} plants were grown under high or low light intensities (16 hours light, 435 \textmu mol photons m\textsuperscript{-2}s\textsuperscript{-1} and 125 \textmu mol m\textsuperscript{-2}s\textsuperscript{-1}) and \textit{im} plants were grown under low light as a reference. Pictures were taken at 2- or 3-day intervals and leaf size and the relative size of the yellow areas were estimated by image analysis. In some of the emergent first to fourth leaves, up to 80% of the leaf area could be yellow, but their relative area decreased at a rate comparable to that of the leaf growth rate. Subsequent leaves exhibited markedly less patchiness. A decrease in the relative area of white sectors was also seen in \textit{im}, but both the change in their percentage area and rate of decrease were lower (not shown).

Moreover, all \textit{im} rosette leaves formed could be equally affected, and demarcations between their green and white sectors remained distinct.

\textbf{Results}

\textbf{Characterization of the \textit{var3} mutant}

\textit{VAR3} phenotype was identified in a population of stable transg sporting generated with a \textit{Ds} gene trap (Sundaresan et al., 1995). \textit{VAR3} was inherited as a recessive allele that segregated with kanamyacin resistance carried on the \textit{Ds} element. \textit{VAR3} displayed normal skotomorphogenesis, and the cotyledons developed normally in the light, but the first true leaves exhibited variegation seen as yellow areas (compare Fig. 1A and B). This variegation became indistinct in older leaves so that the only phenotypic difference compared with wild type (\textit{Ler}) at flowering was pale flower bud sepals in \textit{var3} (Fig. 1D and E). Variegation in \textit{var3} differed from that in \textit{im} (Fig. 1B, E compared with C, F) which consisted of pronounced, light-dependent sectoring. Light microscopy of \textit{var3} leaves showed that yellow areas contained few if any palisade mesophyll cells, which resulted in a rough leaf surface (compare Fig. 1G and H). Scanning electron microscopy confirmed that these areas collapsed during critical point drying (Fig. 1J).

\textbf{Chloroplast import assays}

\textit{VAR3} cDNA encoding the full-length precursor of \textit{VAR3} (pre\textit{VAR3}) was amplified by PCR and cloned in pGEM-T (Promega). The plasmid pGADT7:N Ced4 encoding the full-length precursor of NCED4 (preNCED4) was partially digested with \textit{NcoI}, to remove the vector HA epitope tag, and then religated. \textit{VAR3} and NCED4 were transcribed with T7 RNA polymerase and translated in vitro using a coupled transcription-translation rabbit reticulocyte lysate kit, (Promega) in the presence of L-[\textsuperscript{35}S]methionine (Amersham Biosciences). Plasmid constructs encoding wheat pre33K [33 kDa subunit of the oxygen-evolving complex (Kirwin et al., 1989)] and pea \textit{RbcS} precursor were transcribed in vitro using SP6 RNA polymerase, and translated using a wheat germ lysate system (Promega), in the presence of L-[\textsuperscript{35}S]methionine.

\textit{Pea} (\textit{var}, Kelvedon Wonder) chloroplasts were isolated and import assays performed according to the method of Robinson and Mant (Robinson and Mant, 2002). Briefly, intact chloroplasts equivalent to 50 \textmu g chlorophyll were mixed with 8 mM MgATP, 5 mM unlabeled L-methionine, 5 mM unlabeled L-cysteine, 5 mM MgCl\textsubscript{2} and 12.5 \mu l translation mixture, all in HS buffer (330 mM sorbitol, 50 mM Heps-KOH pH 8.0), final volume 150 \mu l. Assays were incubated for 60 minutes at 26°C in an illuminated water bath (150 \textmu mol photon m\textsuperscript{-2}s\textsuperscript{-1}). Incubations were halted by the addition of 1 ml ice-cold HS, followed by centrifugation at 3000 \textgreek{g} for 5 minutes at 4°C. Chloroplast pellets were resuspended in 120 \mu l HS and 30 \mu l removed, representing total chloroplasts. The remaining 90 \mu l chloroplasts were incubated for 40 minutes at 4°C with 0.2 mg ml\textsuperscript{-1} thermolysin (Protease type X; Sigma), 2.5 mM CaCl\textsubscript{2} in a final volume of 250 \mu l HS. The protease digestion was halted with 10 mM EDTA and one third of the sample removed (protease-treated chloroplasts sample). The remaining chloroplasts were centrifuged at 3000 \textgreek{g} for 5 minutes at 4°C, and resuspended in lysis buffer (10 mM Heps-KOH, pH 8.0, 5 mM MgCl\textsubscript{2}, 10 mM EDTA). After 5 minutes incubation on ice, the lystate was centrifuged at 20,000 \textgreek{g} for 5 minutes at 4°C. The supernatant was the stromal extract, and the pellet, thylakoid and envelope membranes. Membranes were resuspended in lysis buffer without EDTA and divided into equal portions. One portion was analyzed directly, the other after incubation with 0.2 mg ml\textsuperscript{-1} thermolysin, 2.5 mM CaCl\textsubscript{2} for 40 minutes on ice. Thermolysin incubation was stopped by addition of 10 mM EDTA and centrifugation at 20,000 \textgreek{g} for 5 minutes at 4°C. All chloroplast fractions were mixed with protein sample buffer and heated to 95°C for 4 minutes before analysis by SDS-PAGE and fluorography.

\textbf{Arabidopsis \textit{VARIEGATED 3}}

\textbf{4809}
Confocal microscopy revealed that var3 variegation started at early stages of chloroplast development shortly before the leaves emerged at the shoot apex (Fig. 1K). At this stage, areas of cells with normally developed chloroplasts were mingled with areas with partially developed and undeveloped plastids, as judged by the red fluorescence of chlorophyll and precursors and by plastid size (Fig. 1L-N). Since normal cells were found abutting cells in the yellow areas, the effect of VAR3 appeared to be cell-autonomous. This pattern of yellow areas was retained in expanded leaves (Fig. 1O,P), and rudimentary lamellae could be seen in partially developed chloroplasts and ‘vacuoles’ of swollen proplastids within their cells. The presence of such small chloroplasts with reduced numbers of thylakoids was confirmed in these cells by transmission electron microscopy (not shown). In contrast, cells containing normal chloroplasts together with clearly undeveloped plastids were not observed. This indicates that var3 cells are not heteroplastidic.

Gene expression in var3
The levels of expression of Lhcb1 and other genes involved in photosynthetic reactions may be regulated by plastid-derived signals including chlorophyll precursors and carotenoids (reviewed by Rodermel, 2001). This initially suggested that the expression levels of nuclear genes like Lhcb1 or RbcS might be reduced in var3 because of its defective chloroplasts. However, northern blotting with total var3 RNA revealed only a slightly less accumulation of Lhcb1 and RbcS mRNAs than in wild type (Fig. 2A). In order to determine whether var3 represented a gun mutant, expression of Lhcb1 was investigated under photo-oxidative conditions in the presence of 5 μM norflurazon. Both Ler and var3 seedlings accumulated Lhcb1 under nonphotobleaching conditions. However, Lhcb1 mRNA was not detected under photo-oxidative conditions in the presence norflurazon (Fig. 2B). In contrast, control EF-1α (elongation factor 1 alpha) expression was detected in both the presence and absence of norflurazon. This indicates that VAR3 is not involved in the Mg-ProtoIX plastid-to-nucleus signaling pathway.

To gain insight into the effect of the var3 mutation on global gene expression, expression profiling of wild-type versus var3 was performed with microarrays of some 8,000 cDNAs. The dataset was analyzed using the standard deviation of signal...
transcription and translation, as well as rRNAs from both
mRNAs of genes encoding proteins involved in chloroplast
monitoring by the microarray was not significantly different

slide 2 (z-score=(intvar3-intwt)/std14wt). Dots in first and third
control. (B) RT-PCR detection of
most likely to be differentially expressed in
wild-type arrays in the database (Fig. 2C). This identified genes
intensities of specific clones estimated from the values for 14
arrays from which the
quadrants represent consistent measurements of clones from both
conditions (5
photo-oxidative conditions (green, wtG and
var3
A
wt
var3
B
wt
wt
var3
var3
EF-1α
Lhcb1
RbcS
EF-1α
Lhcb1
EF-1α

D

Acc#   Description Mean  Z  Fold change
T21734  C  rpoB   7.8  4.8
Z30079  NC  Lipase/hydrolase-like  3.3  3.0
N95969  M  28S rDNA  3.9  2.0
T04571  NC  Protein Kinase APK1A  2.9  2.2
T43023  NC  HSP70-9a  2.5  1.6
N95975  M  18S rRNA  2.5  1.7
583560  C  60S Ribosomal Protein L16  2.2  1.6
T41263  C  60S rib1  2.2  2.4
N95957  NC  RNA Binding Protein a  2.2  1.7
T22251  NC  Elongation Factor  2.1  1.5
AA002288  C  ABQDC

Fig. 2. Gene expression in wild type and var3. (A) RNA blot of 20 μg total RNA from wild type (wt) and var3 probed with labeled cDNAs of CAB (X56062), RbcS (X13611; Lhcb1) and EF-1α (16260) control. (B) RT-PCR detection of Lhcb1 in wt and var3 under non-photo-oxidative conditions (green, wtG and var3G) and photo-oxidative conditions (5 μM norflurazon, wtG and var3G) compared with EF-1α control. (C) Scatter plot of z-score of slide 1 versus z-score of slide 2 (z-score=(intvar3-intwt)/std14wt). Dots in first and third quadrants represent consistent measurements of clones from both arrays from which the var3 sample was measured higher or lower relative to wt, respectively. Dots in second and fourth quadrants represent inconsistent measurements. The line represents a geometric mean of 4 std., equal to the most extreme, inconsistent measurement (geometric mean of absolute values). (D) The table shows the top genes ranked after the geometric mean of z-scores. C, chloroplast encoded; M, mitochondrion encoded; N, nuclear encoded; NC, nuclear encoded chloroplast targeted. The protein was formerly designated HSP70-7 but is now known as Hsc70-2 (Sung et al., 2001).

intensities of specific clones estimated from the values for 14
wild-type arrays in the database (Fig. 2C). This identified genes
most likely to be differentially expressed in var3 based on the
geometric mean of the z-score from the two arrays (Fig. 2D).
This confirmed that expression of Lhcb1 and RbcS mRNAs
monitored by the microarray was not significantly different in
var3 and Ler. In addition, it indicated that the levels of
mRNAs of genes encoding proteins involved in chloroplast
transcription and translation, as well as tRNAs from both
chloroplasts and mitochondria, were elevated in var3 compared
to wild type. This indicates that coordinated gene expression in
the plastids, mitochondria and nucleus was perturbed in var3
(Hedtke et al., 1999). For example, the most significantly
elevated mRNA was rpoB or chloroplast-encoded RNA
polymerase subunit 2. rpoB expression is dependent upon DAG,
a nuclear gene whose unstable loss of function produces a
phenotype similar to that of var3 (Chatterjee et al., 1996). In
addition, mRNA levels of a chloroplastic heat shock protein and
a chaperone protein were elevated, and both of these types of
proteins have been shown to associate with carotenoid
metabolizing enzymes (Al-Babili et al., 1996; Bonk et al., 1997).
This is of interest as VAR3 was shown to interact with homologs
of 9-cis-exoperoxycarotenoid dioxygenase (NCED, below).

Isolation of VAR3
An EcoRI site approximately 200 bp from the Ds right border
was used to isolate chromosomal sequence flanking Ds by
inverse PCR. Product sequencing identified the site of Ds
integration at nucleotide 54,785 of P1 clone MAV3 (AB006706) on chromosome V (Fig. 3A). The insertion site
was confirmed by PCR using var3 DNA as template with primers specific to each transposon end and to flanking
genomic sequences. Sequencing revealed an 8 bp target site
duplication typical of Ds insertions (Fig. 3A).
To identify the VAR3 gene, 6 kb of clone MAV3 sequence
(ecotype Columbia, Col) flanking the Ds insertion site was
analyzed with NetPlantGene. This predicted four intron splice
sites in five flanking regions with the codon capacity of exons.
Exon assembly yielded a 2,274 bp open reading frame (ORF)
encoding the VAR3 protein (Fig. 4A). The first predicted exon
contained the
Ds
insertion after Ala46, indicating that
VAR3
starts with a
stop codon. This sequence
is identical to the sequence of a subsequently deposited EST
(AV539640) isolated from a cDNA library from root mRNAs.

Fig. 3. VAR3 locus and mRNA. (A) VAR3 wild type, sequence of
exon one (Var3), sequence of var3 allele with Ds left border (in gray)
(var3 left) and var3 with Ds right border (gray) and the 8-bp repeat
underlined (var3 right). Numbers give nucleotide position from the
same sequence of wild type Col-O (complement of P1 clone MAV3,
AB006706). (B) RT-PCR detection of VAR3 mRNA in wild type (wt)
and var3 (lanes 2 and 4) compared to control EF-1α mRNA (lanes 1
and 3). Molecular markers are shown on both sides of the gel.

Isolation of VAR3
An EcoRI site approximately 200 bp from the Ds right border
was used to isolate chromosomal sequence flanking Ds by
inverse PCR. Product sequencing identified the site of Ds
integration at nucleotide 54,785 of P1 clone MAV3 (AB006706) on chromosome V (Fig. 3A). The insertion site
was confirmed by PCR using var3 DNA as template with primers specific to each transposon end and to flanking
genomic sequences. Sequencing revealed an 8 bp target site
duplication typical of Ds insertions (Fig. 3A).
To identify the VAR3 gene, 6 kb of clone MAV3 sequence
(ecotype Columbia, Col) flanking the Ds insertion site was
analyzed with NetPlantGene. This predicted four intron splice
sites in five flanking regions with the codon capacity of exons.
Exon assembly yielded a 2,274 bp open reading frame (ORF)
encoding the VAR3 protein (Fig. 4A). The first predicted exon
contained the
Ds
insertion after Ala46, indicating that
VAR3
starts with a
stop codon. This sequence
is identical to the sequence of a subsequently deposited EST
(AV539640) isolated from a cDNA library from root mRNAs.
Completion of the *Arabidopsis* genome sequence produced an identical annotation of the *VAR3* gene (At5g17790).

The *var3* allele and *var3* complementation

Northern hybridization with the *VAR3* cDNA as probe failed to detect *VAR3* mRNAs in wild type and *var3*. In contrast, *VAR3* mRNA was detectable by reverse transcription PCR in wild type but not in *var3* (Fig. 3B). RT-PCR also detected low levels of *VAR3* mRNA in wild-type leaves and roots (not shown) from which the *VAR3* EST was isolated. These results indicate that *VAR3* mRNA is rare in wild type, and that the *Ds* insertion in *var3* produces a truncated mRNA that does not accumulate to detectable levels.
To confirm that the Ds insertion in VAR3 was responsible for the var3 phenotype, a wild-type VAR3 genomic fragment was introduced into the var3 mutant to examine complementation. The VAR3 genomic fragment, including the transcribed region and 1500 bp of 5′ upstream and 500 bp of 3′ downstream sequence, was cloned into pCAMBIA3300 carrying BASTA resistance, and transformed via A. tumefasciens into homozygous kanamycin-resistant var3 plants. Phenotypically wild-type, BASTA-resistant T1 progeny were shown by PCR to be homozygous for the var3 Ds allele, and to carry the complementing VAR3 transgene. As expected, T2 progeny of these selfed T1 transformants were kanamycin resistant and segregated 3:1 for BASTA resistant, wild-type growth versus var3 variegation. We conclude that loss of VAR3 function produces the var3 mutant phenotype.

Interestingly, one other T1 transformant showed somatic complementation such that about half of the plant appeared wild type, while the other half exhibited variegation as var3. PCR confirmed that the wild-type half tissue carried the VAR3 transgene, while the variegated regions did not (data not shown). This confirmed that the effect of VAR3 on plastid and palisade development is cell-autonomous.

VAR3 protein and homologs

The VAR3 protein contains 758 amino acids with a molecular mass of 85.9 kDa (Fig. 4A). Database homology searches revealed that VAR3 contains two tandem C2X10C2 zinc-finger domains. These domains comprise a conserved 30 amino acid residue consensus (X2GWDIX2CXnNFARRX2CXRCX2-PRPEX2; pFAM00641) characterized in RAN binding protein 2 (RanBP2) and other nucleoporins (Fig. 4B). These proteins are involved in nuclear trafficking, and the zinc fingers of RanBP2 bind Ran-GDP (Nakielny et al., 1999). This suggests that the similar domains of VAR3 may be involved in protein-protein interactions. VAR3 also contains three longer 50 amino acid repeated sequences in the C-terminal region (Fig. 4B). Between these repeats are two shorter, proline-rich repeats that contain the minimal consensus signature of proteins interacting with WW (-P-P-X-Y-) or SH3 (-P-X-P-X-) domains (Pawson and Scott, 1997).

Proteins with overall sequence similarity to VAR3 are apparently only present in plant genomes including the monocot, rice (Gl:13161434; BAB33004). Arabidopsis has three VAR3 homologs with similarities in their N-terminal regions, the zinc fingers, and some of the repeats in the C-terminal region (Fig. 4A,B). Two homologs (AAF79711 and AAG52346) are divergent only in the C-terminal half, while another (AAG51125) shares overall similarity to VAR3. VAR3 and these homologs may therefore have a modular structure composed of different protein-protein interaction domains.

The ChloroP algorithm predicted that VAR3 contains a 79 residue, N-terminal chloroplast targeting signal. ChloroP also predicted chloroplast target peptides for two of the Arabidopsis homologs (AAG51125 and AAF79711; Fig. 4B), but not for the third (AAG52346). Interestingly, the homolog (AAG51125) most similar to VAR3 also contains two putative, bipartite nuclear localization signals. It is therefore possible that VAR3 protein family members are differentially compartmentalized within cells.

VAR3 sub-cellular localization

To investigate VAR3 subcellular localization, var3 was transformed with a construct carrying a hygromycin resistance marker and a translational fusion between VAR3, E. coli β-glucuronidase (GUS) and the Aequorea victoria green fluorescent protein (GFP) with 6x HIS tag (Var3-GUS-mGFP5-6xHIS) under control of the constitutive CaMV 35S promoter. Twelve wild-type looking, hygromycin-resistant T1 plants were selected that produced significant GUS reporter activity measured histochemically. T2 progeny of these lines proved to be kanamycin resistant and to segregate 3:1 hygromycin resistant wild type to hygromycin sensitive var3. This was confirmed for two lines by PCR for homozygosity of the var3 insertion allele and for the VAR3 reporter transgene. Western blotting with an anti-GFP antibody detected the expected 182 kDa Var3-GUS-mGFP5-6xHIS fusion protein in Ni-NTA affinity purified proteins from total extracts of these plants (data not shown). These results indicate that expression of the triple fusion protein complemented var3.

Since VAR3 constitutes the N-terminal region of this functional triple fusion, this reporter is probably localized within cells similarly to wild-type VAR3 protein. Confocal microscopy was therefore used to examine the subcellular localization of GFP fluorescence in lines expressing GUS;GFP from the pCAMBIA 1303 control construct, and those expressing the complementing triple reporter. This showed that GFP fluorescence from the GUS;GFP control was localized throughout cells and not limited to the chloroplasts identified by their red autofluorescence (Fig. 5, panels 1-4 compared with 5-8). In contrast, GFP fluorescence from the triple VAR3:GUS;GFP reporter was clearly localized to chloroplasts (Fig. 5. panels 9-12). Closer inspection of this pattern in numerous cells revealed that although GFP fluorescence was visible throughout chloroplasts, it was concentrated in foci. These results indicate that the complementing VAR3 fusion accumulated in chloroplasts. The presence of GFP foci suggest it may accumulate in a suborganellar complex or compartment, although these foci may result from inappropriate aggregation of the fusion or GFP moieties because of their overexpression from the 35S promoter or to instability of the protein.

VAR3 interacts in yeast and in vitro with chloroplast-targeted proteins

VAR3 contains three different repeated domains, one of which is similar to the zinc-finger of RanBP2 that binds Ran-GDP (Nakielny et al., 1999). In the absence of biochemical data on VAR3 function, we used directed yeast two-hybrid assays to test whether VAR3 interacts with Arabidopsis Ran1, its GTP-binding form Ran-GTP, GDP-binding Ran-GDP, Ran2, as well as RanBP1a, RanBP1b and RanGAP1 controls (Haizel et al., 1997). None of these exhibited interaction with VAR3, although control transformations with RanBP1a interacted with Ran1 and Ran2, measured by adenine and histidine prototrophy and β-galactosidase assay (data not shown). This indicates that VAR3 zinc-fingers do not bind Arabidopsis Ran.

This prompted us to screen a yeast two-hybrid cDNA library for proteins that interact with VAR3. This screen of a library from mRNA of rosette plants identified 51 clones out of 1.5×107 transformants screened with full-length VAR3 as bait. These preys were sorted into three groups of plasmids by...
hybridization and restriction mapping. Sequencing revealed that the two smallest groups of clones were out of frame fusions to unclassified proteins (At5g07890 and AAG52339), and were not examined further.

The largest group of clones encoded \textit{At}NCED4 (At4g19170; AL021687), a chloroplast-targeted member of a family of enzymes similar to nine-

_cis_-epoxycarotenoid dioxygenase (NCED) (Iuchi et al., 2001). Fig. 6A shows that the prey plasmid expressing 258 amino acids of the NCED4 C terminus fused to the Gal4 activation domain permitted growth of the yeast strain carrying the bait plasmid expressing the \textit{VAR3} Gal4 DNA-binding domain fusion, and that this was not due to either auto-activation of the nutritional markers by the NCED4-Gal4AD fusion, or to the \textit{VAR3}-Gal4BD fusion. These results indicate that \textit{VAR3} interacts with NCED4 in yeast. In contrast, the strain carrying \textit{VAR3} and the IM alternative oxidase was incapable of prototrophic growth, indicating that \textit{VAR3} does not interact with IM.

To confirm these results, epitope-tagged, full-length \textit{VAR3} and the 259 amino acid N-terminal part of NCED4 from the cDNA library screen were synthesized by coupled in vitro transcription/translation and examined for interaction in vitro. This showed that \textit{VAR3}, NCED4 (residues 336-595) and human lamin C control (LAM) were each of the expected size, and were immunoprecipitated with antibodies against the \textit{VAR3} N-terminal cMYC tag, or by the NCED4 or LAM N-terminal HA epitope tags (Fig. 6B, lanes 1-3). In addition, NCED4 was co-immunoprecipitated with c-Myc tagged \textit{VAR3} by anti-c-Myc antibody, while LAM was not (Fig. 6B, lanes 4 and 5). This indicates that \textit{VAR3} is not a ‘sticky’ protein, and that \textit{VAR3} and NCED4 interact in vitro. The smaller NCED4 products, which probably result from premature termination as they are immunoprecipitated via the N-terminal HA epitope tag (lane 2), may bind to \textit{VAR3} or to each other (lane 4). The same results were obtained with the full-length NCED4 protein (not shown).

In vitro chloroplast import assays were used to confirm that \textit{VAR3} and NCED4 are targeted to chloroplasts (Fig. 6C). In vitro-translated, radiolabelled \textit{VAR3} and NCED4 were incubated with isolated, intact chloroplasts, and the chloroplasts fractionated post-import to determine the sub-plastid location of these proteins. For both \textit{VAR3} and NCED4, import was accompanied by processing to a smaller size, confirming that \textit{VAR3} and NCED4 encode precursor proteins. The smaller proteins were not a result of thermolysin processing of unimported \textit{VAR3} and NCED4, as demonstrated by control digestion of the in vitro translation mixture (lanes \textit{Tr}+). Fractionation of the chloroplasts showed that \textit{VAR3} and NCED4 are located in the stroma, as is the case for the control stromal protein RbcS (SSU). A very small proportion of \textit{VAR3} and NCED4 co-localized with thylakoid membranes, but thermolysin digestion of the membranes demonstrated that this was surface associated, in contrast to the thylakoid lumen control 33 kDa subunit of the oxygen-evolving complex (33 k, Fig. 6C). These results confirm the results of the in vivo \textit{VAR3}:GUS:GFP localization (Fig. 5). They also confirm that \textit{Ar}NCED4, like the maize NCED VP14 (Tan et al., 2001), is chloroplast localized.

The \textit{Arabidopsis} genome contains seven NCED-like genes, [\textit{NCED1-6} and 9 (Iuchi et al., 2001)], and two more distantly related genes (\textit{NCED7} and 8). To examine whether \textit{VAR3} binding was specific to NCED4, directed two-hybrid assays were performed with the \textit{VAR3} bait and NCED4 (BAB01336) and NCED5 preys (At1g30100) This revealed that \textit{VAR3} interacted with NCED5, but not with NCED3 (data not shown). These results indicate that \textit{VAR3} specifically interacts with
certain NCED family members, including NCED4 and 5. NCED4 and 5 are not transcriptionally upregulated upon drought stress, and neither perform the cleavage of 9-cis-epoxycarotenoids (Iuchi et al., 2001). In contrast, NCED3 performs this cleavage reaction and is therefore involved in the biosynthesis of the plant hormone abscisic acid.

Fig. 6. VAR3 protein interaction and targeting. (A) Yeast two-hybrid interactions. Left panel shows growth of yeast strains carrying different plasmids on minimal medium lacking Trp, Leu, His and adenine. Right panel identifies fusion proteins expressed from the DNA-binding (BD bait) and activation domain (AD prey) plasmids in the yeast strains shown on the left. LAM is human lamin C, TD1 and VA3 (B) Co-immunoprecipitation (Co-IP) of VAR3 and NCED4. (Left to right) Lane 1, IP of cMyc-tagged VAR3; lane 2, IP of HA-tagged NCED4 (partial length open reading frame isolated from the cDNA library screen); lane 3, IP of HA-tagged LAM; lane 4, Co-IP of NCED4 with cMyc-tagged VAR3; lane 5, lack of Co-IP of control LAM with cMyc-tagged VAR3. Proteins and antibodies are noted above. (C) In vitro chloroplast import assay. Isolated, intact chloroplasts were incubated with in vitro translated, radiolabeled precursor proteins, and chloroplasts were then fractionated and samples analyzed by SDS-PAGE and fluorography. Gel lanes are: Tr, in vitro translation mixture; Tr+, translation mixture digested with thermolysin; C, total chloroplasts; C+, thermolysin-treated chloroplasts; S, stroma; T, thylakoid membranes; T+, thermolysin-treated thylakoids. Pre- and m- denote precursor and mature protein, respectively. SSU is a RbcS stromal marker and 33 K is the 33 kDa subunit of the oxygen-evolving complex, thylakoid lumen marker. The sizes (kDa) of precursor and mature forms of VAR3 and NCED4 are indicated.

drought stress, and neither perform the cleavage of 9-cis-epoxycarotenoids (Iuchi et al., 2001). In contrast, NCED3 performs this cleavage reaction and is therefore involved in the biosynthesis of the plant hormone abscisic acid.

var3 accumulates reduced levels of pigments

The variegated im mutant suffers from light dependent photo-oxidative damage because of a defect in carotenoid biosynthesis that leads to the accumulation of phytoene (Carol et al., 1999; Wu et al., 1999). Since VAR3 interacts with NCED homologs, var3 variegation might also be due to an alteration in carotenoid biosynthesis or metabolism. To examine this, we performed comparative profiling, by HPLC-diode array, of carotenoids and chlorophylls in var3 and wild-type plants. There was no qualitative difference in the pigment profiles of var3 and wild-type when grown in 24-hour or 4-hours/day light regimes. However, under both light regimes there were significant reductions in the levels of all detected carotenoids, as well as a less severe reduction in chlorophylls a and b (Table 1). In both wild type and var3 under the 4 hours/day light regime, one tetrapyrrole peak, tentatively assigned as the chlorophyll precursor protochlorophyllide, increased significantly compared to the levels in plants grown in continuous light. These results indicate that variegation in var3 is unlike that in im, and is not due to loss of specific photoprotective pigments.

Discussion

The biogenesis of chloroplasts and the development of leaf cells containing them are regulated by intracellular signals between the plastid and the nucleus. Since most chloroplast protein functions are nuclear encoded, many nuclear mutations result in defective chloroplast biogenesis and lethal albinism (Barkan et al., 1995). In contrast, viable variegated mutants produced by stable nuclear mutations are rare. We identified the variegated3 (var3) mutant in a screen of transposant lines generated with a modified maize Ds system (Sundaresan et al., 1995). Molecular analysis revealed that the element excised from the DsG1 donor site on chromosome II and re-inserted with 8 bp flanking repeats into the VAR3 gene on chromosome V. Ds insertion produced a truncated VAR3 mRNA that was undetectable in var3 by RT-PCR. Complementation of var3 by transformation with a wild-type VAR3 genomic fragment demonstrated that loss of VAR3 expression is the cause of mutant variegation.

Chloroplasts in green areas of var3 leaves were similar to those in wild type, while the yellow areas contained no, or only partially developed, chloroplasts. These characteristics are similar to those of other stable Arabidopsis mutants chloroplast mutator, im, var1 and var2, as well as in unstable mutants such as Arabidopsis albino 3 (Sundberg et al., 1997), Antirrhinum olive and maize HCF 106 (Barkan et al., 1995). Since these mutants have partially developed but defective chloroplasts, the corresponding genes may be involved in the synthesis or assembly of the photosynthetic machinery, or in essential plastid metabolic functions.

Measurements of photosynthetic pigments revealed no qualitative differences in pigments in var3 compared to wild-type. However, carotenoid levels in var3 were 35-57% of that
in wild type, while chlorophyll a levels in var3 were approximately 70% of wild type under high light regimes. This indicates that loss of VAR3 function has a greater effect upon carotenoid levels than upon chlorophyll levels. Under low light regimes, total carotenoids and chlorophylls were significantly reduced in both var3 and wild type grown in continuous light. An exception to this, in both lines, was levels of a tetrapyrrole, tentatively assigned as protochlorphyllide. This precursor accumulates in low light regimes when chlorophyll biosynthesis is down-regulated via attenuation of protochlorophyllide oxidoreductase. In this respect, var3 retains the ability to modulate chlorophyll biosynthesis as does the wild type. Under low light regimes, the majority of carotenoid and chlorophyll (a+b) levels in var3 were reduced in concert with those in wild type. Although this may indicate a role for VAR3 in modulating lutein oxidation, no confirmatory accumulation of lutein, or decrease in neoxanthin, was observed. In summary, the profiling indicated that VAR3 does not affect production of individual pigments.

Reductions in var3 pigment levels in whole leaves are not accompanied by relative decreases in mRNAs of major nuclear-encoded photosynthetic genes such as Lhcb (CAB) and RbcS. Therefore, mRNA accumulation patterns in var3 appear similar to those of pac1, and of the gun mutants in which the expression of chloroplast-targeted, nuclear gene products is uncoupled from chloroplast development (Mochizuki et al., 2001; Strand et al., 2003; Möller et al., 2001). An explanation for this class of mutants is that they affect levels of Mg-ProtoIX that represses Lhcb and RbcS expression in the absence of functional chloroplasts. Since the var3 mutation primarily affects carotenoid levels, carotenogenic compounds may be involved in such signaling (Rodermel, 2001). However, the lack of Lhcb transcript accumulation in var3 under photo-oxidative conditions indicates that var3 is not a gun mutant. Thus, loss of VAR3 function does not appear to affect the levels of tetrapyrrole or carotenogenic intermediates that regulate the expression of nuclear genes encoding chloroplast-targeted proteins.

Transcriptome analysis indicated that the levels of mRNAs encoding plastid and mitochondrial transcription and translation components are elevated in var3. The increase in rpoB mRNA levels is interesting because the Antirrhinum nuclear gene DAG is required for rpoB expression, such that unstable dag mutants exhibit variegation because of the resulting block in chloroplast differentiation (Chatterjee et al., 1996). A simple explanation for the increases in these mRNAs in var3 is that they reflect a mechanism by which plastids attempt to increase overall protein synthesis to compensate for the failure of photosynthetic functions (Mullet, 1993). Alternatively, VAR3 might affect the production or transduction of signals regulating chloroplast gene expression, perhaps via its interaction with chloroplast localized enzymes (below). In any event, the interdependency of chloroplast and mitochondrial metabolic pathways may also explain why the expression of mitochondrial genes encoding translational components appear to be affected in var3. Such increases in mitochondrial gene expression have been described in albino barley leaves carrying the recessive nuclear abo gene mutation (Hedtke et al., 1999). Such effects are also opposite but analogous to that produced by mutation of the nuclear gene CHM causing mitochondrial genome rearrangements that affect chloroplast function (Abdelnoor et al., 2003). Taken together, these results suggest that signals involved in coordinating mitochondrial and plastid activities are perturbed in var3. The micro array results also revealed that the levels of mRNAs encoding a chloroplastic heat shock protein, cpHsc70-2, and a chaperone subunit, Cpn-60α, were elevated in var3. These proteins are likely orthologs of Narcissus pseudonarcissus chloroplastic HSP70 and Cpn60 chaperonin that are components of oligomeric assemblies with phytoene desaturase, and at least two other carotenoid biosynthetic enzymes (Bonk et al., 1997; Al-Babili et al., 1996). While circumstantial, these results suggest an additional link between VAR3 function and carotenoid metabolism.

VAR3 encodes a novel protein containing two zinc-finger domains conserved in nucleoporins and RNA binding proteins (pFAM00641). Such fingers in nucleoporins interact with RAN (Nakielny et al., 1999), although yeast two-hybrid assays did not reveal interactions between VAR3 and Arabidopsis nucleoporins or RAN. This suggests that this class of zinc fingers in different proteins mediate interactions to various proteins. VAR3 also contains long C-terminal repeats with proline rich regions reminiscent of motifs interacting with SH3 and WW domains, suggesting the existence of similarly folded protein sub-domains. Database searches indicate that VAR3 homologs are restricted to plants, suggesting they perform functions specific to plants. Their modular structure is evidenced in the two VAR3 homologs containing only some of the conserved domains. A third homolog is similar to VAR3 throughout its length, and may therefore perform VAR3 redundant functions.

In addition to the var3 phenotype, two lines of evidence...
indicate that VAR3 is required for normal chloroplast function. First, VAR3 has a chloroplast target peptide that enabled import to chloroplast stroma, and a VAR3 reporter fusion that complemented the var3 phenotype was localized to chloroplasts. Second, yeast two-hybrid screening revealed that VAR3 specifically interacted with chloroplast localized NCED4 and 5, two of seven Arabidopsis proteins homologous to 9-cis-epoxycarotenoid dioxygenases (Schwartz et al., 1997; Iuchi et al., 2001; Tan et al., 2001). The interaction between VAR3 and NCED4 was confirmed in vitro, and both were shown to accumulate in stromal fractions by chloroplast import assays. These interactions are apparently specific, because VAR3 does not interact with the chloroplast-localized IM alternative oxidase involved in carotenoid biosynthesis, or with NCED3. NCED3 mRNA, and mRNAs of the maize VP14, avocado PaNCED1 and bean PvNCED1 orthologs, are induced by water stress (Schwartz et al., 1997; Qin and Zeevaart, 1999; Iuchi et al., 2001). These enzymes catalyze cleavage of 9-cis-epoxycarotenoids to generate 2-cis,4-trans-xanthoxin, the major precursor of abscisic acid (ABA). Neither Arabidopsis VAR4 or 5 are directly involved in ABA biosynthesis since their mRNAs do not accumulate during desiccation, and recombinant NCED4 and 5 proteins do not cleave 9-cis-epoxycarotenoids in vitro (Iuchi et al., 2001). Interestingly, phylogenetic sequence analysis indicated that NCED4 is more closely related to Cucurbita maxima CsZCD than to other Arabidopsis members of the NCED family. CsZCD catalyzes the 7,8(7’,8’) cleavage of zeaxanthin leading to the formation of chloroplast-derived apocarotenoids including crocin and safranal (Bouvier et al., 2003). Nonetheless, cleavage products of zeaxanthin, trans-lutein or trans-violaxanthin were not detected in in vitro or cellular assays with recombinant NCED expressed in E. coli. The enzymatic activity and physiological substrates or products of NCED 4 therefore remained to be determined.

In summary, our data are consistent with a role for VAR3 as a component of a protein complex that functions in a metabolic pathway required for chloroplast development. By analogy to var1 and var2 (Sakamoto, 2003; Sakamoto et al., 2003), the mechanism of VAR3 variegation, including effects specific to cell-type and developmental stages, may also involve a redundant activity provided by the VAR3 homologs identified here. The striking effect of the var3 mutation on the development of palisade cells in true leaves, but not in cotyledons, is similar to that seen in tomato dcl1 (Keddie et al., 1996) and Arabidopsis var2 and cuel (Chen et al., 2000; Stretfiel et al., 1999). VAR3 may therefore be required for both chloroplast development and palisade cell morphogenesis, although its stromal localization suggests that defects in palisade cell elongation are a secondary effect of incomplete chloroplast development. Nonetheless, these secondary effects may include the disruption of specific signals, including carotenogenic products, required for palisade morphogenesis (Rodermel, 2001). Such products may signal diverse developmental processes, as posited for those of the Arabidopsis MAX4 dioxygenase-like gene (Sorefan et al., 2003) that may control synthesis of a metabolite(s) that acts downstream of auxin to inhibit shoot branching.

We thank Pierre Carol (Grenoble), Ferenc Nagy (Szeged), Colin Robinson (Warwick), and Mitsuru Akita (East Lansing) for cDNAs encoding IM, RanBP and Ran and RanGAP, wheat pre33k, and pea pre-SSU, respectively. Peter Bramley (London) is thanked for carotenoid standards. The work was supported by grants to H.B.N. (Danish Research Fund and Novozymes), I.M. [EU QLGG-CT-1999-00551 and Danish Research Councils (9001412, 53002008, 5300250)], M.A. (Danish Food Directorate KUL-97-DIF-6) and M.B. (BBSRC, IGF-GARNet project).

References

Abdelnoor, R. V., Yule, R., Elo, A., Christensen, A. C., Meyer-Gaunz, G., Mackenzie, S. A. (2003). Substoichiometric shifting in the plant mitochondrial genome is influenced by a gene homologous to MutS. Proc. Natl. Acad. Sci. USA 100, 5968-5973.

Al-Babili, S., von Lintig, J., Haubruck, H. and Beyer, P. (1996). A novel, soluble form of phytoene desaturase from Narcissus pseudonarcissus chromoplast is Hsp70-complexed and competent for flavinylation, membrane association and enzymatic activation. Plant J. 9, 601-612.

Barkan, A., Voelker, R., Mendel-Hartvig, J., Johnson, D. and Walker, M. (1995). Genetic analysis of chloroplast biogenesis in higher plants. Physiol. Plant. 93, 163-170.

Bechtold, N., Ellis, J. and Pelletier, G. (1993). In planta agrobacterium mediated gene transfer by infiltration of adult Arabidopsis plants. C.R. Acad. Sci., Paris, Life Sci. 316, 1191-1199.

Bonk, M., Hoffmann, B., von Lintig, J., Schledz, M., Al-Babili, S., Hobeika, E., Kleinig, H. and Beyer, P. (1997). Chloroplast import of four carotenoid biosynthetic enzymes in vitro reveals differential fates prior to membrane binding and oligomeric assembly. Eur. J. Biochem. 247, 942-950.

Bouvier, F., Suire, C., Mutterer, J. and Camara, B. (2003). Oxidative remodeling of chloroplast carotenoids: identification of the carotenoid dioxygenase CsCCD and CsZCD genes involved in Cocus secondary metabolite biogenesis. Plant Cell 15, 47-62.

Bramley, P. M. (1992). Analysis of carotenoids by high performance liquid chromatography and diode-array detection. Phytochem. Anal. 3, 97-104.

Carl, P., Stevenson, D., Bisanz, C., Breitenbach, J., Sandmann, G., Mache, R., Coupland, G. and Kuntz, M. (1999). Mutations in the Arabidopsis gene IMMUTANS cause a variegated phenotype by inactivating a chloroplast terminal oxidase associated with phytoene desaturation. Plant Cell 11, 57-68.

Chatterjee, M., Sparvoli, S., Edmunds, C., Garosi, P., Findlay, K. and Martin, C. (1996). DAG, a gene required for chloroplast differentiation and palisade development in Antirrhinum majus. EMBO J. 15, 4194-4207.

Chen, M., Choi, Y., Voytas, D. F. and Rodermel, S. (2000). Mutations in the Arabidopsis VAR2 locus cause leaf variegation due to the loss of a chloroplast Fritz protease. Plant J. 22, 303-313.

Corona, V., Aracri, B., Kosturkova, G., Bartley, G. E., Pitto, L., Gionetti, L., Scolnik, P. A. and Giuliano, G. (1996). Regulation of a carotenoid biosynthesis gene promoter during plant development. Plant J. 9, 505-512.

Crowell, N., Packard, C., Pierson, C., Giner, J.-L., Downes, B. and Chary, N. (2003). Identification of an allele of CLA1 associated with variegation in Arabidopsis thaliana. Physiol. Plant 118, 29-37.

Fraser, P. D., Pinto, M. E. S., Holloway, D. E. and Bramley, P. M. (2001). Application of high performance liquid chromatography with photo-diode array detection to the metabolic profiling of plant isoprenoids. Plant J. 24, 513-528.

Haizel, T., Merkle, T., Aniko, P., Erzsebet, F. and Nagy, F. (1997). Characterization of proteins that interact with the GTP-bound form of the regulatory GTPase Ran in Arabidopsis. Plant J. 11, 93-103.

Hedtkle, B., Wagner, I. B. W., Wagner, I. B. T. and Hess, W. R. (1999). Intergenellar crosstalk in higher plants: impaired chloroplast development affects mitochondrial gene and transcripts levels. Plant J. 19, 635-643.

Logemann, J., Schell, J. and Willmitzer, L. (1987). Improved method for the isolation of RNA from plant tissues. Anal. Biochem. 163, 16-20.

Iuchi, S., Kobayashi, M., Taji, T., Naramoto, M., Seki, M., Kato, T., Tabata, S., Kakubari, Y., Yamaguchi-Shinozaki, K. and Shinozaki, K. (2001). Regulation of drought tolerance by gene manipulation of 9-cis-epoxycarotenoid dioxygenase, a key enzyme in abscisic acid biosynthesis in Arabidopsis. Plant J. 27, 323-333.

James, P., Halliday, J. and Craig, E. A. (1996). Genomic libraries and a host strain designed for highly efficient two-hybrid selection in yeast. Genetics 144, 1425-1436.

Keddie, J. S., Carroll, B., Jones, J. D. G. and Gruissem, W. (1999). Analysis of chloroplast development and palisade cell morphogenesis in leaves. EMBO J. 15, 4208-4217.
Kirwin, P. M., Meadows, J. W., Shackleton, J. B., Musgrove, J. E., Elderfield, P. D., Mould, R., Hay, N. A. and Robinson, C. (1989). ATP-dependent import of a lumenal protein by isolated thylakoid vesicles. EMBO J. 8, 2251-2255.

Mochizuki, N., Brusslan, J. A., Larkin, R., Nagatani, A. and Chory, J. (2001). Arabidopsis genomes uncoupled 5 (GUN5) mutant reveals the involvement of Mg-chelatase H subunit in plastid-to-nucleus signal transduction. Proc. Natl. Acad. Sci. USA 98, 2053-2058.

Mulder, J. E. (1993). Dynamic regulation of chloroplast transcription. Plant Physiol. 103, 309-313.

Möller, S. G., Kunkel, T. and Chua, N.-H. (2001). A plastidic ABC protein involved in intercompartmental communication in light signaling. Genes Dev. 15, 90-103.

Nakielny, S., Shaikh, S., Burke, B. and Dreyfuss, G. (1999). Nup153 is an M9-containing mobile nucleoporin with a novel Ran-binding domain. EMBO J. 18, 1982-1995.

Pawson, T. and Scott, J. D. (1997). Signaling trough scaffold, anchoring, and adaptor proteins. Science 278, 2075-2080.

Qin, X. and Zeekavart, J. A. D. (1999). The 9-cis-epoxy-carotenoid cleavage reaction is the key regulatory step in abscisic acid biosynthesis in water-stressed bean. Proc. Natl. Acad. Sci. USA 96, 15354-15361.

Reiter, R. S., Coomber, S. A., Bourett, T. M., Bartley, G. E. and Scolnik, P. A. (1994). Control of leaf and chloroplast development by the Arabidopsis gene pale cress. Plant Cell 6, 1253-1264.

Robinson, C. and Mant, A. (2002). Import of proteins into isolated chloroplasts and thylakoid membranes. In Molecular Plant Biology Volume Two: A Practical Approach (ed. P. Gilmartin and C. Bowler), pp. 123-146. Oxford, UK: Oxford University Press.

Rödermel, S. (2001). Pathways of plastid-to-nucleus signaling. Trends Plant Sci. 6, 471-478.

Sakamoto, W. (2003). Leaf-variegated mutations and their responsible genes in Arabidopsis. Genes Genet. Syst. 78, 1-9.

Sakamoto, W., Tamura, T., Hanba-Tomita, Y., Sodmergen, and Murata, M. (2002). The VAR1 locus of Arabidopsis encodes a chloroplastic FtsH and responsible for leaf variegation in the mutant alleles. Gene Cells 7, 769-780.

Sakamoto, W., Zaltsman, A., Adam, Z. and Takahashi, Y. (2003). Coordinated regulation of complex formation of YELLOW VARIEGATED 1 and YELLOW VARIEGATED 2, chloroplastic FtsH metalloproteases involved in the repair cycle of photosystem II in Arabidopsis thylakoid membranes. Plant Cell 15, 2843-2855.

Schwartz, S. H., Tan, B. C., Gage, D. A., Zeevaart, J. A. and McCarty, D. R (1997). Specific oxidative cleavage of carotenoids by VP14 of maize. Science 276, 1872-1874.

Sorefan, K., Booker, J., Haurogne, K., Goussot, M., Bainbridge, K., Foo, E., Chatfield, S., Ward, S., Beveridge, C., Rameau, C. et al. (2003). MAX4 and RMS1 are orthologous dioxygenase-like genes that regulate shoot branching in Arabidopsis and pea. Genes Dev. 17, 1469-1474.

Strand, A., Asami, T., Alonso, J., Ecker, J. R. and Chory, J. (2003). Chloroplast to nucleus communication triggered by accumulation of Mg-chelatase IX. Nature 421, 79-83.

Streathfield, S. J., Weber, A., Kinsman, E. A., Hausler, R. E., Li, J., Beilinmiller, D. P., Kaiser, W. M., Pyke, K. A., Flugge, U. I. and Chory, J. (1999). The phosphoenolpyruvate/phosphate translocater is required for phenolic metabolism, palisade cell development and plastid-dependent nuclear gene expression. Plant Cell 11, 1609-1621.

Sundaresan, V., Springer, P., Volpe, T., Haward, S., Jones, J. D. G., Dean, C., Ma, H. and Martienssen, R. (1995). Patterns of gene action in plant development revealed by enhancer trap and genetrap transposable elements. Genes Dev. 9, 1797-1810.

Sundberg, E., Slagter, J. G., Fridborg, L., Cleary, S. P., Robinson, C. and Coupland, G. (1997). ALBINO3 and Arabidopsis nuclear gene essential for chloroplast differentiation, encodes a chloroplast protein that shows homology to proteins present in bacterial membranes and yeast mitochondria. Plant Cell 9, 717-730.

Sung, D. Y., Vierling, E. and Guy, C. L. (2001). Comprehensive expression profile analysis of he Arabidopsis Hsp70 gene family. Plant Physiol. 126, 789-800.

Tan, B. C., Cline, K. and McCarty, D. R. (2001). Localization and targeting of the VP14 epoxy-carotenoid dioxygenase to chloroplast membranes. Plant J. 27, 373-382.

Takechi, K., Sodmergen, Murata, M., Motoyoshi, F. and Sakamoto, W. (2000). The YELLOW VARIEGATED (VAR2) locus encodes a homologue of FtsH, an ATP-dependent protease in Arabidopsis. Plant Cell Physiol. 41, 1334-1346.

Workman, C., Jensen, L. J., Jarmer, H., Berka, R., Gautier, L., Nielsen, H. B., Saxild, H., Nielsen, C., Brunak, S. and Knudsen, S. (2002). A new non-linear normalization method to reduce variability in DNA microarray experiments. Genome Biol. 3:research0048.

Wu, D., Wright, D. A., Wetzel, C., Voytas, D. and Rödermel, S. (1999). The IMMUTANS variegation locus of Arabidopsis defines a mitochondrial alternative oxidase homolog that functions during early chloroplast biogenesis. Plant Cell 11, 43-55.