PROGRESS REPORT

Stefan Hörtensteiner

The loss of green color during chlorophyll degradation a prerequisite to prevent cell death?

Received: 15 December 2003 / Accepted: 10 January 2004 / Published online: 8 April 2004 © Springer-Verlag 2004

Keywords Cell death \cdot Chlorophyll degradation \cdot Non-fluorescent chlorophyll catabolite \cdot Pheophorbide *a* oxygenase \cdot Red chlorophyll catabolite reductase \cdot Senescence

During plant senescence, chlorophyll (Chl) is degraded to non-fluorescent Chl catabolites (NCCs; Fig. 1a). These linear tetrapyrroles accumulate in the vacuoles of senescing cells and, in many plant species, represent the final products of Chl catabolism (Matile et al. 1988; Kräutler 2003). Despite the billions of tons of Chl disappearing this way every year and the fascinating autumnal color change of deciduous trees resulting from it, most reactions underlying conversion of Chl to NCC have only recently been elucidated (for recent reviews, see Hörtensteiner 1999; Takamiya et al. 2000; Kräutler 2003). Chl breakdown is a multi-step pathway (Fig. 1a) aiming to increase pigment solubility and to abolish the photodynamic properties of Chl by complete disruption of the conjugated π -electron system. Taking this into account, Chl breakdown can be apostrophized as Chl detoxification (Hörtensteiner 1999).

As inferred from the structures of NCCs (Kräutler et al. 1991), the most remarkable structural change is the oxygenolytic opening of the porphyrin macrocycle of Chl. This reaction is catalyzed by the joint action of two enzymes, pheophorbide *a* oxygenase (PaO) and red Chl catabolite reductase (RCCR) converting pheophorbide (pheide) *a* to a primary fluorescent Chl catabolite (pFCC; Fig. 1a; Rodoni et al. 1997a). Several lines of evidence suggest that the two enzymes interact during catalysis, thereby channeling the first porphyrin cleavage

S. Hörtensteiner

Institute of Plant Sciences, University of Bern, Altenbergrain 21, 3013 Bern, Switzerland E-mail: shorten@ips.unibe.ch Tel.: +41-31-6313156 Fax: +41-31-6314942 product, red Chl catabolite (RCC). Thus, in vitro, RCC does not accumulate in the absence of PaO. In addition, RCCR is sensitive to oxygen, although PaO requires O_2 for incorporation into pheide a (Rodoni et al. 1997a; Wüthrich et al. 2000). PaO has been demonstrated to be a non-heme iron-containing monooxygenase, that specifically introduces one oxygen atom of O_2 at the α methine bridge of pheide a (Hörtensteiner et al. 1995, 1998). In addition, PaO is specific for pheide a with pheide b inhibiting in a competitive manner (Hörtensteiner et al. 1995). PaO is located at the inner envelope membrane of senescing chloroplasts (Matile and Schellenberg 1996). In contrast, RCCR is a soluble chloroplast protein (Wüthrich et al. 2000), suggesting that the site of conversion of pheide *a* to pFCC is at the stromal periphery of the envelope. RCCR stereospecifically reduces the C1/C20 double bond of RCC, thereby forming two possible C1 stereoisomers of pFCC. The source of RCCR determines which one is formed; thus, for example, RCCR of Arabidopsis thaliana produces a different isomer than RCCR isolated from tomato (Hörtensteiner et al. 2000b).

Exploiting the biochemical characteristics of PaO, 21 candidate genes for PaO in Arabidopsis were identified recently by using functional genomics (Pružinská et al. 2003). One of them, Accelerated cell death 1 (Acd1), was subsequently shown to exhibit PaO activity in vitro after expression in Escherichia coli. The properties of the heterologously expressed protein were identical to native PaO. In contrast, a homologue of ACD1, At4g25650, did not exhibit PaO activity (Pružinská et al. 2003). Thus, Acd1 encodes Arabidopsis PaO (AtPaO). AtPaO is a Rieske-type iron-sulfur cluster-containing oxygenase. In Arabidopsis, five Rieske-type oxygenases are present which have rather diverse functions (Gray et al. 2002). Besides PaO and At4g25650, the function of which is unknown, chlorophyll a oxygenase (CAO; Tanaka et al. 1998) and choline monooxygenase (CMO; Rathinasabapathi et al. 1997) contain Rieske centers. In addition, Tic55, a component of the protein import machinery at the inner envelope (TIC) also belongs to this group of



Fig. 1 a Pathway of chlorophyll degradation during senescence. Depicted are the structures of the Chl catabolites pheophorbide *a* (*pheide a*), red Chl catabolite (*RCC*), primary fluorescent Chl catabolite (*pFCC*), and non-fluorescent Chl catabolite (*NCC*). The key reaction is catalyzed by the joint action of pheide *a* oxygenase (*PaO*) and RCC reductase (*RCCR*) without release of the intermediate, RCC. Relevant carbon atoms and the α -methine bridge that is cleaved by PaO are labeled. *R*₁, *R*₂ and *R*₃ in the NCCs indicate species-specific differences (Kräutler 2003). **b** Leaves from wild-type maize at a senescent stage (*left*) and from *lls1* showing a lesion mimic phenotype (*right*)

oxygenases (Calibe et al. 1997), although an enzymatic (oxygenase) activity has not been demonstrated in this case. Rieske-type oxygenases are widely distributed in pro- and eukaryotes. In all cases, electrons necessary to drive the redox cycle of the Rieske center irons are provided from reduced ferredoxin (Fd; Schmidt and Shaw 2001). Accordingly, PaO, CAO and CMO are Fddependent enzymes (Schellenberg et al. 1993; Rathinasabapathi et al. 1997; Tanaka et al. 1998). Fd is kept in the reduced state by the activity of a reductase. In higher plants, the nature of this reductase is unknown so far, but possibly Fd-NADPH oxidoreductase (FNR) is involved. It has been shown that another protein of the protein import machinery at the inner envelope, Tic62, is able to bind FNR (Küchler et al. 2002).

The RCCR gene has been identified using classical protein purification and PCR-based cloning strategies (Rodoni et al. 1997b; Wüthrich et al. 2000). RCCR is a novel protein that does not have high homology to other reductases, but is distantly related to a family of Fddependent bilin reductases, necessary for the biosynthesis of phycobilins and the phytochrome chromophore (Frankenberg et al. 2001). In contrast to other Fddependent enzymes, RCCR and the bilin reductases appear to lack a metal or flavin cofactor. Thus, electron transfer is believed to occur directly from reduced Fd to the respective substrates (Frankenberg and Lagarias 2003). In this respect, RCCR would rather be active as a "chaperone", mediating the interaction of Fd and RCC (at this stage still bound to PaO), and controlling the regio- and stereoselective reduction (Kräutler 2003). When comparing the amino acid sequences of different RCCRs, no obvious domains can be identified that could be responsible for the stereospecificity. Therefore, chimeric proteins that were composed of portions of RCCR from tomato and Arabidopsis were produced in E. coli and their stereospecificity analyzed. It turned out that a Phe-to-Val exchange at position 218 was sufficient to change the stereospecificity of the Arabidopsis RCCR (I. Anders and S. Hörtensteiner, unpublished). Interestingly, Phe²¹⁸ lies adjacent to a stretch of four amino acids that is absent in the bilin reductases (Frankenberg et al. 2001).

The rather complex PaO/RCCR reaction represents a key step of the entire Chl catabolic pathway. Thus, PaO activity is restricted to senescence (Schellenberg et al. 1993; Hörtensteiner et al. 1995), whereas activities of other enzymes of the pathway, such as chlorophyllase (catalyzing the initial removal of phytol from Chl) or RCCR are constitutive (Trebitsh et al. 1993; Rodoni et al. 1997a; Jakob-Wilk et al. 1999). Surprisingly, both PaO mRNA and protein are present in non-senescent leaf tissue (Grav et al. 2002: Pružinská et al. 2003). Although PaO expression is up-regulated to some extent upon senescence induction, at the same time the increase in activity is a magnitude higher (Pružinská et al. 2003). From this it is concluded that PaO is regulated on the posttranscriptional level as well. So far, the nature of this proposed regulation has not been elucidated.

The joint reaction of PaO and RCCR is responsible for the loss of green pigment color. In this respect, it is most important for Chl detoxification during senescence. The importance of Chl catabolism for plant survival can be inferred from the analysis of Chl catabolic mutants. Different mutants have been identified that are defective in either PaO or RCCR. These include *Ara*- bidopsis accelerated cell death 1 (acd1; Greenberg and Ausubel 1993) and maize lethal leaf spot 1 (lls1; Gray et al. 1997), in which the maize homologue of AtPaO is affected (Fig. 1b), and Arabidopsis acd2, which is defective in RCCR (Greenberg et al. 1994; Mach et al. 2001). All of these mutants develop cell death lesions on their leaves in an age-dependent fashion. The phenotype is similar to the induction of defense reactions in pathogen resistance; thus, respective mutants are termed lesion mimic mutants (Mach et al. 2001). The affected genes were believed to be involved in a cell death suppression mechanism either directly (affecting a signal cascade?) or through, for example, the removal of toxic molecules (Greenberg and Ausubel 1993; Gray et al. 1997). In favor of the latter was the finding that in *lls1* a cell death-inducing signal was derived from plastids and lesion formation was light dependent (Gray et al. 2002). Indeed, both PaO and RCCR mutants accumulate Chl catabolites (pheide a in *lls1* and *acd1*, and RCC in *acd2*) upon dark-induced senescence. In addition, the content of these catabolites positively correlates with cell death progression of the respective mutants (Pružinská et al. 2003; A. Pružinská and S. Hörtensteiner, unpublished results). Thus, in these mutants, the accumulation of photoreactive Chl catabolites can be suggested to cause the production of reactive oxygen species that in turn induce cell death (Mach et al. 2001). Surprisingly, other mutants have been described that do not develop an apparent cell death phenotype although they also accumulate pheide a due to reduced PaO activities (Vicentini et al. 1995; Thomas et al. 1996). Besides pheide a, these mutants also accumulate chlorophyllides, indicating that they have a genetic defect that is different from *acd1* or *lls1*. On the other hand, several lesion mimic mutants have been identified that are affected in genes of Chl biosynthesis (Hu et al. 1998; Meskauskiene et al. 2001; Ishikawa et al. 2001).

Altogether, it can be concluded that functional Chl metabolism, i.e. biosynthesis, turnover and degradation, is important to prevent the accumulation of photodynamic intermediates. Furthermore, it becomes obvious that Chl degradation via the PaO/RCCR pathway is a vitally important process during plant senescence. Quite likely it is also involved in cellular responses to a variety of stresses that are linked to Chl breakdown, such as the hypersensitive reaction. The photodynamic properties of Chl enable the conversion of light energy to chemical energy during photosynthesis, but during senescence, photodynamism may turn into a threat. Thus, parallel to the "invention" of Chl and the evolution of oxygenic photosynthesis, plants evolved a mechanism for the detoxification of Chl. In unicellular photosynthesizers, such as Chlorella protothecoides, RCCR is absent and, consequently, RCC-like compounds are excreted into the medium (Engel et al. 1991; Hörtensteiner et al. 2000a). However, the development of multicellular plants required, in addition to PaO, the appearance of RCCR to enable the safe disposal of Chl catabolites inside the vacuole.

References

- Calibe A, Grimm R, Kaiser G, Lübeck J, Soll J, Heins L (1997) The chloroplastic protein import machinery contains a Riesketype iron–sulfur cluster and a mononuclear iron-binding protein. EMBO J 16:7342–7350
- Engel N, Jenny TA, Mooser V, Gossauer A (1991) Chlorophyll catabolism in *Chlorella protothecoides*. Isolation and structure elucidation of a red bilin derivative. FEBS Lett 293:131–133
- Frankenberg N, Lagarias JC (2003) Phycocyanobilin:ferredoxin oxidoreductase of *Anabaena* sp. PCC 7120. J Biol Chem 278:9219–9226
- Frankenberg N, Mukougawa K, Kohchi T, Lagarias JC (2001) Functional genomic analysis of the HY2 family of ferredoxindependent bilin reductases from oxygenic photosynthetic organisms. Plant Cell 13:965–978
- Gray J, Close PS, Briggs SP, Johal GS (1997) A novel suppressor of cell death in plants encoded by the Lls1 gene of maize. Cell 89:25–31
- Gray J, Janick-Bruckner D, Bruckner B, Close PS, Johal GS (2002) Light-dependent death of maize *lls1* cells is mediated by mature chloroplasts. Plant Physiol 130:1894–1907
- Greenberg JT, Ausubel FM (1993) *Arabidopsis* mutants compromised for the control of cellular damage during pathogenesis and aging. Plant J 4:327–341
- Greenberg JT, Guo A, Klessig DF, Ausubel FM (1994) Programmed cell death in plants: a pathogen-triggered response activated coordinately with multiple defense functions. Cell 77:551–563
- Hörtensteiner S (1999) Chlorophyll breakdown in higher plants and algae. Cell Mol Life Sci 56:330–347
- Hörtensteiner S, Vicentini F, Matile P (1995) Chlorophyll breakdown in senescent cotyledons of rape, *Brassica napus* L.: enzymatic cleavage of phaeophorbide a in vitro. New Phytol 129:237–246
- Hörtensteiner S, Wüthrich KL, Matile P, Ongania K-H, Kräutler B (1998) The key step in chlorophyll breakdown in higher plants. Cleavage of pheophorbide *a* macrocycle by a monooxygenase. J Biol Chem 273:15335–15339
- Hörtensteiner S, Chinner J, Matile P, Thomas H, Donnison IS (2000a) Chlorophyll breakdown in *Chlorella protothecoides*: characterization of degreening and cloning of degreening-related genes. Plant Mol Biol 42:439–450
- Hörtensteiner S, Rodoni S, Schellenberg M, Vicentini F, Nandi OI, Qiu Y-L, Matile P (2000b) Evolution of chlorophyll degradation: the significance of RCC reductase. Plant Biol 2:63–67
- Hu G, Yalpani N, Briggs SP, Johal GS (1998) A porphyrin pathway impairment is responsible for the phenotype of a dominant disease lesion mimic mutant of maize. Plant Cell 10:1095–1105
- Ishikawa A, Okamoto H, Iwasaki Y, Asahi T (2001) A deficiency of coproporphyrinogen III oxidase causes lesion formation in *Arabidopsis*. Plant J 27:89–99
- Jakob-Wilk D, Holland D, Goldschmidt EE, Riov J, Eyal Y (1999) Chlorophyll breakdown by chlorophyllase: isolation and functional expression of the *Chlase1* gene from ethylene-treated *Citrus* fruit and its regulation during development. Plant J 20:653–661
- Kräutler B (2003) Chlorophyll breakdown and chlorophyll catabolites. In: Kadish KM, Smith KM, Guilard R (eds) The porphyrin handbook, vol 13. Elsevier, San Diego, pp 183–209
- Kräutler B, Jaun B, Bortlik K-H, Schellenberg M, Matile P (1991) On the enigma of chlorophyll degradation: the constitution of a secoporphinoid catabolite. Angew Chem Int Ed Engl 30:1315– 1318
- Küchler M, Decker S, Hörmann F, Soll J, Heins L (2002) Protein import into chloroplasts involves redox-regulated proteins. EMBO J 21:6136–6145
- Mach JM, Castillo AR, Hoogstraten R, Greenberg JT (2001) The Arabidopsis-accelerated cell death gene ACD2 encodes red chlorophyll catabolite reductase and suppresses the spread of disease symptoms. Proc Natl Acad Sci USA 98:771–776

- Matile P, Schellenberg M (1996) The cleavage of pheophorbide *a* is located in the envelope of barley gerontoplasts. Plant Physiol Biochem 34:55–59
- Matile P, Ginsburg S, Schellenberg M, Thomas H (1988) Catabolites of chlorophyll in senescing barley leaves are localized in the vacuoles of mesophyll cells. Proc Natl Acad Sci USA 85:9529–9532
- Meskauskiene R, Nater M, Goslings D, Kessler F, op den Camp R, Apel K (2001) FLU: a negative regulator of chlorophyll biosynthesis in *Arabidopsis thaliana*. Proc Natl Acad Sci USA 98:12826–12831
- Pružinská A, Anders I, Tanner G, Roca M, Hörtensteiner S (2003) Chlorophyll breakdown: pheophorbide a oxygenase is a Rieske-type iron–sulfur protein, encoded by the accelerated cell death 1 gene. Proc Natl Acad Sci USA 100:15259–15264
- Rathinasabapathi B, Burnet M, Russell BL, Gage DA, Liao PC, Nye GJ, Scott P, Golbeck JH, Hanson AD (1997) Choline monooxygenase, an unusual iron–sulfur enzyme catalyzing the first step of glycine betaine synthesis in plants: prosthetic group characterization and cDNA cloning. Proc Natl Acad Sci USA 94:3454–8
- Rodoni S, Mühlecker W, Anderl M, Kräutler B, Moser D, Thomas H, Matile P, Hörtensteiner S (1997a) Chlorophyll breakdown in senescent chloroplasts. Cleavage of pheophorbide a in two enzymic steps. Plant Physiol 115:669–676
- Rodoni S, Vicentini F, Schellenberg M, Matile P, Hörtensteiner S (1997b) Partial purification and characterization of red chlorophyll catabolite reductase, a stroma protein involved in chlorophyll breakdown. Plant Physiol 115:677–682

- Schellenberg M, Matile P, Thomas H (1993) Production of a presumptive chlorophyll catabolite in vitro: requirement for reduced ferredoxin. Planta 191:417–420
- Schmidt CL, Shaw L (2001) A comprehensive phylogenetic analysis of Rieske and Rieske-type iron sulfur proteins. J Bioenerg Biomembr 33:9–26
- Takamiya K, Tsuchiya T, Ohta H (2000) Degradation pathway(s) of chlorophyll: what has gene cloning revealed? Trends Plant Sci 5:426–431
- Tanaka A, Ito H, Tanaka R, Tanaka NK, Yoshida K, Okada K (1998) Chlorophyll a oxygenase (CAO) is involved in chlorophyll b formation from chlorophyll a. Proc Natl Acad Sci USA 95:12719–12723
- Thomas H, Schellenberg M, Vicentini F, Matile P (1996) Gregor Mendel's green and yellow pea seeds. Bot Acta 109:3–4
- Trebitsh T, Goldschmidt EE, Riov J (1993) Ethylene induces de novo synthesis of chlorophyllase, a chlorophyll degrading enzyme, in *Citrus* fruit peel. Proc Natl Acad Sci USA 90:9441– 9445
- Vicentini F, Hörtensteiner S, Schellenberg M, Thomas H, Matile P (1995) Chlorophyll breakdown in senescent leaves: identification of the biochemical lesion in a *stay-green* genotype of *Festuca pratensis* Huds. New Phytol 129:247–252
- Wüthrich KL, Bovet L, Hunziker PE, Donnison IS, Hörtensteiner S (2000) Molecular cloning, functional expression and characterisation of RCC reductase involved in chlorophyll catabolism. Plant J 21:189–198