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Chloroplast cold-resistance is mediated by the acidic domain of the RNA

binding protein CP31A 2 Ayako Okuzaki^{#\$1}, Marie-Kristin Lehniger^{1\$}, Jose M Muino², Benjamin Lenzen¹, Thilo Rühe³, Dario 3 Leister³, Uwe Ohler², Christian Schmitz-Linneweber^{1*} 4 5 ¹ Molecular Genetics, Humboldt-University Berlin, Philippstr.13, 10115 Berlin, Germany ² Computational Regulatory Genomics, Humboldt-University Berlin / Max Delbrück Centre for Molecular 6 Medicine, Philippstr.13, 10115 Berlin, Germany 7 ³ Plant Molecular Biology, Department of Biology, Ludwig Maximilian University of Munich, Munich, 8 9 Germany. 10 11 # current address: College of Agriculture, Tamagawa University, Tamagawagakuen 6-1-1, Machida-shi, 12 Tokyo, Japan 13 \$ equally contributed 14 *corresponding author: 15 Christian Schmitz-Linneweber phone: +49 30 2093-49700 16 17 fax: +49 30 2093-49701 18 Email: smitzlic@rz.hu-berlin.de 19 20 Running title: An acidic domain for chloroplast cold resistance 21 Key words: chloroplast, RNA processing, RNA binding, acidic domain, RNA editing, RNA stability, 22 organelle, Arabidopsis thaliana Author contributions 23 C.S.-L. conceived the original research plan; D.L. supervised the experiments; A.O. generated the 24 transgenic plant lines and performed their analysis. M.-K. L. performed the RIP-Seq and microarray 25 experiments; J.M.M. analyzed the RIP-Seq data. B.L. performed the SSMART analysis; T.R. performed 26 27 mutant analyses. C.S.-L., D.L. and U.O. supervised the project, designed the experiments and analyzed the data; C.S.-L. wrote the article with contributions of all the authors; C.S.-L. agrees to serve as the author 28 responsible for contact and ensures communication. 29 30 One sentence summary:

Cold exposure induces increased RNA association of the RRM protein CP31A, which mediates cold-

resistance of *Arabidopsis thaliana* via its acidic domain

Abstract

Chloroplast RNA metabolism is characterized by long-lived mRNAs that undergo a multitude of post-transcriptional processing events. Chloroplast RNA accumulation responds to environmental cues, foremost light and temperature. A large number of nuclear-encoded RNA-binding proteins (RBPs) are required for chloroplast RNA metabolism, but we do not yet know how chloroplast RBPs convert abiotic signals into gene expression changes. Previous studies showed that the chloroplast ribonucleoprotein 31A (CP31A) is required for the stabilization of multiple chloroplast mRNAs in the cold, and that the phosphorylation of CP31A at various residues within its N-terminal acidic domain (AD) can alter its affinity for RNA *in vitro*. Loss of CP31A leads to cold sensitive plants that exhibit bleached tissue at the center of the vegetative rosette. Here, by applying RIP-Seq, we demonstrated that CP31A shows increased affinity for a large number of chloroplast RNAs *in vivo* in the cold. Among the main targets of CP31A were RNAs encoding subunits of the NDH complex and loss of CP31A lead to reduced accumulation of *ndh* transcripts. Deletion analyses revealed that cold-dependent RNA binding and cold resistance of chloroplast development both depend on the AD of CP31A. Together, our analysis established the AD of CP31A as a key mediator of cold acclimation of the chloroplast transcriptome.

Introduction

Chloroplasts contain genetic information that is essential for photosynthesis. The expression of this information is realized by a unique mixture of ancestral bacterial and derived eukaryotic features (Barkan, 2011). Chloroplast gene expression adapts to various environmental changes, including light and temperature (e.g. Klein, 1991; Mentzen and Wurtele, 2008; Cho et al., 2009; Castandet et al., 2016). Contributions to such acclimation processes have been described on the transcriptional level (Pfannschmidt, 2003; Tsunoyama et al., 2004), but post-transcriptional processes are likely to dominate (Deng and Gruissem, 1987; Eberhard et al., 2002; Udy et al., 2012). One key change in posttranscriptional processes between chloroplast and their bacterial ancestors are the vastly increased RNA half lives in the organelle. In bacteria, transcription and translation are usually directly coupled and mRNAs have short half-lives (in the range of minutes; Selinger et al., 2003). In chloroplasts, the half-lives of mRNAs are long (in the range of hours) and untranslated RNAs accumulate in large amounts (Klaff and Gruissem, 1991; Germain et al., 2012). The turnover rates of chloroplast RNAs change in response to developmental and environmental cues, which is suggesting that RNA stability is regulated in chloroplasts (Deng et al., 1989; Klaff and Gruissem, 1991; Biehl et al., 2005; Bollenbach et al., 2007; Germain et al., 2013; Manayski et al., 2018), but the underlying regulatory factors are largely unknown. Possible candidates for regulators of RNA stability are pentatricopeptide repeat (PPR) proteins and chloroplast ribonucleoproteins (cpRNPs). PPR proteins specifically associate with one or few RNAs (Barkan and Small, 2014), while cpRNPs are

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generalists that bind to a large number of mRNAs (Kupsch et al., 2012; Teubner et al., 2017). Two cpRNPs named CP29A and CP31A were previously shown to be required for the accumulation of chloroplast RNAs in the cold, which makes them interesting candidates to be mediators of global RNA stability during the acclimation to changing temperature conditions (Kupsch et al., 2012). The cpRNP protein family consists of ten members in Arabidopsis thaliana (Ruwe et al., 2011). All cpRNPs are targeted to the chloroplast post-translationally, and dedicated import receptors appear to be responsible for their transport across the chloroplast envelope (Li and Sugiura, 1990; Grimmer et al., 2014). cpRNPs are highly regulated proteins that react to various external and internal signals, particularly light, which controls both their expression and their protein modification state (summarized in Ruwe et al., 2011). Several cpRNPs have been identified as phosphoproteins (Reiland et al., 2009) and an N-terminally acetylated isoform of CP29A was shown to respond rapidly to changes in light and developmental stages (Wang et al., 2006). The phosphorylation of cpRNPs can alter their RNA-binding characteristics in vitro (Lisitsky and Schuster, 1995; Loza-Tavera et al., 2006), but evidence for condition-dependent RNA binding of cpRNPs or any other chloroplast RBPs in vivo are lacking. Several molecular functions have been suggested for cpRNPs. In vitro, a tobacco homolog of Arabidopsis CP31A has been shown to support RNA editing of multiple sites (Hirose and Sugiura, 2001). Other cpRNPs have been reported to be required for the 3'-end processing of several mRNAs (Schuster and Gruissem, 1991; Hayes et al., 1996; Schuster et al., 1999). They also support the ribozymatic maturation of a viroid RNA genome (Daros and Flores, 2002). The most notable and general function of cpRNPs is however their role in stabilizing mRNAs. In vitro, cpRNPs protect the mRNA encoding the D1 subunit of photosystem II against degradation (Nakamura et al., 2001). In vivo, they are required for the accumulation of a multitude of mRNAs (Kupsch et al., 2012; Teubner et al., 2017). Co-immunoprecipitation analyses have demonstrated that cpRNPs are associated with multiple chloroplast RNAs (Kupsch et al., 2012; Teubner et al., 2017), and that they prefer unprocessed (unspliced) RNAs over mature forms (Nakamura et al., 1999). Together, the data obtained from these functional analyses show that cpRNPs have a broad target range and contribute to various RNA-processing events and to RNA stabilization. All cpRNPs share a similar design with two RNA recognition motifs (RRMs) that are preceded by a domain rich in acidic residues (acidic domain; AD). While the RRMs are well characterized RNA binding domains (Li and Sugiura, 1991; Ye and Sugiura, 1992; Lisitsky et al., 1995), the role of the AD remains to be determined. The cpRNP CP31A stands out from the family in having a particularly long acidic domain, which contains two phosphorylated serine residues within a short repeat element of five amino acids (Reiland et al., 2009). CP31A's AD was suggested to be important for RNA editing based on *in vitro* assays (Hirose and Sugiura, 2001). The AD of the spinach homolog of CP31A is supportive of RNA binding in

vitro, but does not bind RNA itself (Lisitsky and Schuster, 1995). Recently, plastid casein kinase II (pCKII) was demonstrated to phosphorylate the acidic domain of CP31A in vitro (Schonberg et al., 2014). Genetic analyses have demonstrated that *Arabidopsis* CP31A supports RNA editing at multiple sites and modulates the stability of multiple mRNAs (Tillich et al., 2009; Kupsch et al., 2012). An RNA strongly affected in cp31a mutants was the ndhF mRNA, which encodes a subunit of the NDH complex. CP31A binds to the 3'-UTR of ndhF and is required for the generation of the ndhF 3'-terminus (Kupsch et al., 2012). Mutants of CP31A display cold sensitivity: the germination rate of null mutants is reduced and their newly emerging leaf tissue bleaches at 8°C (Kupsch et al., 2012). All analyzed proteins of the photosynthetic apparatus are reduced in this defective tissue, which is at least in part due to multiple defects in RNA processing (e.g., RNA splicing, RNA editing, and intercistronic processing), but is likely primarily caused by the strong reduction of multiple chloroplast mRNAs (Kupsch et al., 2012). However, it remains unclear how CP31A mechanistically confers cold resistance and whether it directly perceives cold as a signal. We hypothesized that the AD could act as a regulatory domain for RNA binding and cold-responsiveness. We therefore analyzed the ability of CP31A to associate with RNA in response to cold and assessed the role of the AD for CP31A's ability to bind and process chloroplast RNA. Our results demonstrate that the AD of CP31A is essential for plant cold acclimation, at least in part because of its supportive role for cold-dependent RNA binding.

Results

The RRM domains of CP31A are sufficient for RNA editing of CP31A-dependent sites

The binding of RBPs to their RNA targets can be altered by protein modifications. For several members of the cpRNP family, phosphorylation has been demonstrated to alter RNA binding *in vitro* (Lisitsky and Schuster, 1995; Loza-Tavera et al., 2006; Reiland et al., 2009). As phosphoproteomic analyses identified several phosphorylation sites within the acidic N-terminal domain of CP31A (Reiland et al., 2009; Schonberg et al., 2014), we decided to test the function of this acidic phosphodomain by deletion mutagenesis. We designed three T-DNA based constructs to express CP31A variants in *Arabidopsis* (Fig. 1A). The first two expressed variants lack the AD. One was driven by the 35S promoter, while the other was driven by the endogenous genomic region upstream of the transcriptional start site, presumably encompassing the unknown *CP31A* promoter (12769646 to 12767953 of the *Arabidopsis thaliana* chromosome 4 sequence; nucleotides –1 to –1694 relative to the *CP31A* start codon). These constructs were introduced into a *cp31a*-deficient background (homozygous *cp31a*-1 null allele) by *Agrobacterium*-mediated gene transfer. In addition, the full-length protein driven by its native promoter was used as a positive control. All plant lines retrieved showed normal development and were indistinguishable from wt plants under normal growth conditions (see below).

We assessed CP31A protein accumulation in our transgenic lines by using an antibody that reacts to two peptides: one situated in the linker between the two RRM domains and another at the C-terminus of the protein (Kupsch et al. 2012). The antibody thus detects both the full-length as well as the AD-deletion proteins. We observed accumulation of proteins that were of the appropriate size for all transgenic lines (Fig. 1B). These data demonstrate that the fulllength and AD-deficient proteins were expressed and specifically detected by the antibody. We quantified signals from three independent transgenic lines for each construct (Fig. 1B, C). As expected, there is variability in expression between individual lines, which can be attributed to position effects of the transgene insertion site. In sum, these analyses demonstrate that all three constructs were successfully expressed CP31A-deficient in a background.

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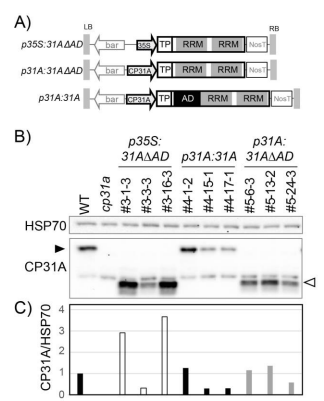


Figure 1: Construction of AD deletion mutants.

A) Schematic overview of constructs used to complement cp31a null mutants. bar = selectable marker; black arrows indicate promoters. TP = transit peptide. NosT = the terminator of the nos gene. LB/RB = borders of the Agrobacterium T-DNA.

B) Western analysis of transgenic lines. Blots were probed with a HSP70 antibody to control for loading and with an antiserum recognizing CP31A. The position of the full-length protein and the truncated, AD-less version are indicated by filled and open arrowheads, respectively.

C) Quantification of CP31A signals normalized to HSP70 signals (from (B)).

Since CP31A has been shown to support

RNA editing at 13 specific sites (Tillich et al., 2009), we investigated these and additional sites in our complementation lines by performing next-generation sequencing of cloned amplified cDNA (Bentolila et al., 2013), testing a total of 16 sites on 10 amplicons. This included sites that were previously shown not to be affected in *cp31a* mutants. Two wt and two *cp31a-1* null mutant plants served as controls. The lowest read coverage observed for any individual site was above 500 reads per site and was found for site *ndhD* 116281 (number refers to the position in the *Arabidopsis* chloroplast genome), while the average coverage for all sites and all genotypes analyzed was 3132 reads per site.

Of the 13 sites previously described to be dependent on CP31A, we confirmed that RNA editing of 11 sites was reduced in *cp31a*-1 mutants (Fig. 2A). Additional defects were found for two previously unreported

sites in *ndhB* (94999, 96579), whereas no defect was observed for the previously reported sites, *petL* (65716) and *ndhB* (95225). We speculate that these differences reflect differences in growth conditions: our plants were grown under long-day conditions (16-h light/8-h dark), whereas the previous study used short-day conditions (8-h light/16-h dark; Tillich et al., 2009). Importantly, all our complementation lines, irrespective of the utilized promoter or the presence of the AD showed wt-like editing states at most analyzed sites (Fig. 2A).

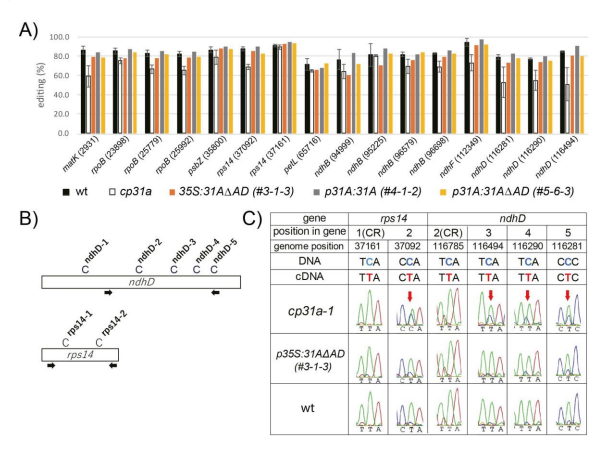


Figure 2: Analysis of RNA editing sites in *cp31a* complementation lines.

A) Summary of results from amplicon sequencing of RNA editing sites in complementation lines. Editing sites were amplified by PCR and sequenced on an Illumina platform using barcoded adapters that allowed to eliminate duplicate amplifications. Each individual cDNA sequence is scored for its editing status and the frequency of edited versus non-edited cDNAs is shown. Error bars represent the standard deviation calculated from two replicate experiments with WT and *cp31a* mutants, respectively. Numbers refer to the position of the editing site in the *Arabidopsis thaliana* chloroplast genome.

- B) Schematic overview of two genes with their editing sites that were assessed by Sanger sequencing of amplified cDNAs. Arrows indicate positions of primers used for cDNA amplification.
- C) Excerpts from electropherograms showing fluorescence signals for base triplets with the editing site always at the center. Bases marked by red arrows are targets of CP31A. Green traces refer to T signals, blue to C signals and red to A signals. The expected edited (on cDNA) and unedited (on DNA) triplets are shown above. CR = control sites not affected by CP31A according to Tillich et al (2009).

We confirmed this finding by independent amplification and Sanger sequencing of selected RNA-editing sites of the *ndhD* and *rps14* mRNAs. The amplicons we sequenced harbored both, CP31A-dependent and

CP31A-independent editing sites (Fig. 2B). Our sequencing confirmed that in *cp31a* mutants, sites *ndhD*-3, 4 and 5, *rps14*-2 show reduced editing efficiency when compared to wt. This defect is complemented in the AD-less CP31A protein (Fig. 2C). The successful complementation demonstrates that the RRM domains are sufficient for the RNA-editing function of CP31A.

The AD of CP31A is not essential for, but is supportive of the stability of the *ndhF* mRNA

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We next analyzed the RNA-stabilizing function of CP31A by focusing on ndhF, which is its major target under normal growth conditions (Tillich et al., 2009). Using RNA gel-blot hybridization, we quantified the amount of ndhF mRNA in the transgenic lines. The accumulation of this transcript is at least partially restored in all complementing lines relative to the *cp31a* mutant (Fig. 3A, B). Even in a line with very little CP31A accumulation and lacking the AD, a signal for *ndhF* is visible (#3-3-3. Fig. 1, Fig. 3A,B). This indicates that the RRM domains were sufficient to stabilize the ndhF mRNA. However, we observed marked differences in the efficiency of *ndhF* stabilization between the full-length and AD-deficient proteins. Lines accumulating large amounts of the AD-less CP31A protein had only little more *ndhF* mRNA than lines expressing much less full-length CP31A protein (compare for example line #3-1-3 with #4-17-1 in Fig. 1B and Fig. 3A). This was visualized by calculating the ratio of ndhF mRNA levels to

CP31A protein levels (i.e., the stabilization

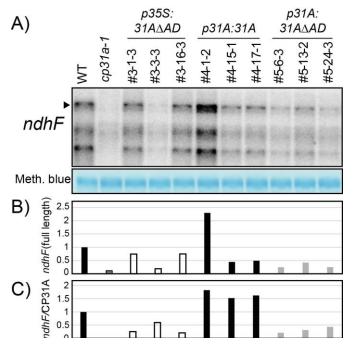


Figure 3: Analysis of the accumulation of the *ndhF* mRNA in *cp31a* complementation lines.

A) RNA gel blot analysis of the *ndhF* mRNA in various *cp31a* complementation lines. 4 μg total cellular RNA were separated by denaturing agarose gel electrophoresis, transferred to a nylon membrane and probed for *ndhF*. The transcript labelled with an arrowhead represents the full-length *ndhF* mRNA, whereas smaller signals correspond to degradation products (Kupsch et al. 2012). The nylon membrane was labeled with methylene blue to control for equal loading (excerpt with 23S rRNA shown below blot).

- B) Quantification of the full-length *ndhF* transcript labeled with an arrowhead in (A).
- C) Ratio of *ndhF* mRNA levels over CP31A protein levels (from Fig. 1C).

efficiencies; Fig. 3C), which clearly showed that the full-length proteins outperformed the AD-less proteins in stabilizing the *ndhF* mRNA. Together, these results show that the AD supports but is not essential for stabilizing the *ndhF* mRNA and that the RRM domains are capable of performing this task by themselves.

A)

The AD is required for cold-stress tolerance

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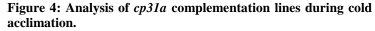
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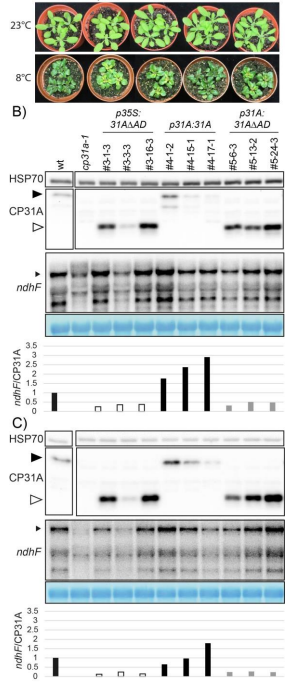
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To test the impact of the AD on CP31A-mediated cold tolerance, we challenged 2-week-old plants from the transgenic lines with low temperature (8°C) for 5 weeks. Under this treatment, the *cp31a*-null lines are known to display bleaching of the freshly emerging tissue at the center of the leaf rosette (Fig. 4A). Similar losses of pigment were also found in the lines expressing the AD-less CP31A, but not in wt plants or in mutant plants complemented with the full-length protein (Fig. 4A). The pigment deficiency has been shown to be paralleled by a global reduction of the chloroplast-encoded proteins required for photosynthesis in cp31a mutants (Kupsch et al., 2012). This reduction of photosynthetic proteins was also observed in plants expressing the AD-less CP31A while plants expressing full-length CP31A proteins, accumulated protein amounts similar to those seen in wt plants (Suppl. Fig. 1). Next, we tested the accumulation of CP31A and *ndhF* in these lines under cold exposure. We separately harvested full leaves, as well as the bleached center part of rosettes in case molecular defects are restricted to the phenotypically conspicuous tissue. However, both tissues showed the same tendency: In line with the findings at normal growth temperatures, the *ndhF*/CP31A ratio was consistently higher for plants expressing full-length CP31A versus plants expressing the AD-less protein (Fig. 4B/C and Suppl. Fig. 2).



A) Phenotypes of complementation lines grown at 23° C or challenged with 8° C for five weeks. Note the bleached tissue in the center of the rosette in *cp31a* deficient plants and in plants lacking the AD.



p35S:31A∆AD

#3-1-3

ср31а

p31A:31A p31A:31AΔAD

#4-1-2

B) Analysis of CP31A and HSP70 protein accumulation and *ndhF* mRNA accumulation in *cp31a* complementation lines. Here, the entire rosette leaves were used for protein and RNA extraction. Panels from top to bottom: immunoblot analysis of HSP70 proteins as loading control; immunoblot analysis of CP31A proteins; RNA gel blot analysis of *ndhF*; Methylene blue stain of cytosolic rRNA as loading control; Ratio of quantified *ndhF* and CP31A signals. Filled and open arrowheads indicate full-length and AD-less CP31A, respectively, while the small arrowhead denotes the full-length *ndhF* transcript.

C) Same analysis as in (B), but with tissue only from the center of the rosette (bleached area in cp31a null mutants).

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These experiments show that the AD of CP31A supports the accumulation of chloroplast RNA and is essential for conferring cold resistance to emerging leaf tissue in *Arabidopsis*.

CP31A associates with multiple mRNAs with a preference for transcripts encoding subunits of the NDH complex, and is required for *ndh* mRNA accumulation at standard growth temperatures

RBPs of various origins and organisms show differential expression under temperature changes, and genetic studies indicate that they are important for temperature-dependent RNA processing (Lin et al., 2010; Malay et al., 2011; Gotic et al., 2016). That said, few studies have examined whether the binding of RBPs to their RNA targets is modified in response to external and internal cues in vivo. Since CP31A is required for cold resistance, we tested whether its ability to bind RNA is modified at low temperature and also assayed the importance of the AD for this response. Previous studies using low-resolution techniques could only identify whole transcripts as targets of CP31A (Kupsch et al., 2012). We here used a RIP-Seq approach to obtain better resolution (Suppl. Fig. 3A). For this, wt Arabidopsis seedlings and plants expressing the AD-less CP31A protein in a cp31a background were grown for 14 days under normal growth conditions and then subjected to formaldehyde cross-linking. For cold tests, seedlings were grown for 13 days at standard growth temperature (21°C), transferred to 4°C for 24 hours, and were then subjected to cross-linking and immunoprecipitation (IP; Kupsch et al., 2012). We used such a short incubation time in the cold to avoid pleiotropic effects that can be expected in the bleached tissue resulting from longer cold challenges. The efficiency of precipitation was comparable between samples grown at normal and low temperatures (Suppl. Fig. 3B). RIP-Seq libraries were prepared in duplicate from input and pellet samples (Suppl. Fig. 3C). We tested the reproducibility of the RIP-Seq experiments by calculating pairwise correlation coefficients across all samples (Suppl. Fig. 3D). We found strong correlations among the input samples (average Pearson coefficient (R): 0.95), and the pellet samples from the IP (average Pearson coefficient (R): 0.97). As expected, input and pellet samples formed separate clusters in our correlation analysis (Suppl. Fig. 3D). This analysis demonstrates high reproducibility between biological replicates of our RIP-Seq assay.

For the analysis of RIP-Seq reads, we followed established procedures originally developed for ChIP-Seq (Bardet et al., 2011; Muino et al., 2011). We started by identifying transcript regions with an enrichment of RNA in pellet samples over input samples. In total, 75 different binding regions were identified as significant in at least one of the four experiment, which are henceforth called binding sites (BSs). The 75 BSs of CP31A are located in 44 transcripts of diverse functionality. Of these 44 transcripts, 31 were also identified by previous RIP-Chip experiments (Suppl. Tab. 1, Suppl. Fig. 4). The difference observed is likely due to methodological dissimilarities. Purified chloroplast stroma was used for the previous RIP-Chip experiments without a cross-link, while here, we used cross-linked frozen total leaf tissue in the RIP-seq. Given these technical differences, it was gratifying to see a similar set of transcripts enriched in the two types of RIP experiments (Suppl. Tab. 1). We next focused on those BSs located within a gene body and

thus can be clearly assigned to a specific gene. By contrast, binding sites within intergenic regions cannot be easily assigned to individual genes due to the polycistronic nature of chloroplast transcripts. Assigned BSs are found in genes for the photosynthetic complexes as well as for the gene expression apparatus (Tab. 1). We compared the actual distribution of BSs with the calculated numbers of BSs expected for each functional category (if random binding in the transcriptome is assumed; Tab. 1). This demonstrated that the NDH complex is the only functional category overrepresented among RIP-Seq targets (12 found versus 3 expected; Tab. 1). We also found an unexpected large number of binding sites antisense to coding regions of known genes (Tab. 1). Although antisense RNAs are known to exist in chloroplasts, their levels do not (with a few exceptions) reach those of sense RNAs and their functions (if any) remain unclear (Nakamura et al., 2003; Hotto et al., 2010; Hotto et al., 2011; Sharwood et al., 2011). Consequently, the functional significance of CP31A's association with antisense RNAs remains unclear as well.

Table 1: Summary of target sites of CP31A across all samples analyzed in RIP-Seq experiments

	Unique CP31A binding sites ¹		Unique CP31A target genes ²		Length of potential target genes (bp) ³		Expected unique binding sites ⁴	
	sense	antisense	sense	antisense	sense	antisense	sense	antisense
NDH complex	12	0	6	0	13002	13002	3	3
Photosynthesis (except <i>ndh</i> genes)	6	3	5	3	23644	23644	6	6
Other metabolism	4	0	2	0	4580	4580	1	1
Ribosome	0	5	0	5	22778	22778	6	6
RNA polymerase	3	1	3	1	11170	11170	3	3
tRNA	2	1	2	1	10196	10196	2	2
Other	2	9	2	4	24177	24177	6	6
All BS	29	19	20	14			27	27
Total BSs in intergenic regions	27		NA		44931		21	
Total BSs	75		34				7	75

¹ this includes binding sites identified in all samples; rRNAs were excluded from calling binding sites

depends only on the combined length of all genes within a class)

It is however interesting that at least for the main target category, the *ndh* genes, a preference for sense sites is evident (Tab. 1), suggesting that CP31A is relevant for *ndh* mRNA expression. We therefore analyzed

² gene annotations according to Araport11

³ sum of the length of all genes within a category

⁴ number of target sites expected in a gene class if the 75 binding sites are randomly distributed in the entire chloroplast genome and if the abundance of transcripts is considered equal between all genes (i.e.

1%^{2%}3%_{-2%}

wt/cp31a > 1.5

119 probes

Cytochrome b/f complex

ср31а

NDH complex

ATP synthase

75%

PSII

■ PSI

ndhF

ndhD

2-

ndhK

ndhB

B)

10%

10%

4%

32%

8%

16%

All probes

2274 probes

■ ycf1

■ vcf2

sia4

crr2

others

gene expression

wt(1/2) wt(1/4)

A)

chloroplast RNA levels in cp31a versus wt samples at normal temperatures using an oligonucleotide tiling array that represents the entire chloroplast genome of Arabidopsis thaliana. We scored only probes whose RNA levels were at least one third lower in the mutant compared to wt. Of all exon probes in the array, 16% represent *ndh* sequences. Importantly, among exon probes whose signals were decreased by at least 0.6-fold in the mutants, 75% contained *ndh* sequences (Fig. 5A, B). No other functional category showed enrichment in this analysis. To confirm these results, we performed RNA gel-blot hybridization experiments for four *ndh* genes that represent the four *ndh* operons in the chloroplast genome (Fig. 5C). We analyzed cp31a mutant RNAs alongside RNAs from seedlings with impairment in the expression of SIGMA FACTOR 4 (SIG4), which is specifically required for the transcription of the ndhF mRNA 2005), and **CHLORORESPIRATORY** (Favory REDUCTION 2 (CRR2), which is required for the accumulation of monocistronic *ndhB* transcripts (Hashimoto et al., 2003). As expected, the control mutants showed specific defects for their known target RNAs (Fig. 5C). By contrast, the *ndhK* transcripts accumulated to normal levels in these two control lines, and the ndhD mRNA was only slightly decreased (Fig. 5C). In contrast and consistent with our microarray results, the cp31a mutants displayed reductions in all four *ndh* transcripts analyzed. The decrease was most pronounced for ndhF, strong for ndhK and ndhB, and somewhat weaker for ndhD (Fig. 5C). Collectively, these findings indicate that CP31A stabilizes mRNAs from all four *ndh* operons, suggesting that CP31A regulates *ndh* mRNAs as a group.

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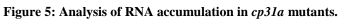
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A) Summary of microarray analyses of 14 days old wt and cp31a

mutant seedlings. Relative abundance of exon probes showing at least 1.5 fold stronger signals in wt versus *cp31a* mutants. Three replicate microarray hybridizations were analyzed and assigned to different gene categories.

B) Relative distribution of all exon probes on the microarray to different gene categories.

C) RNA gel blot analysis of 14 days old Arabidopsis seedlings. 4 µg RNA from from wt, *cp31a* mutants, and two control mutants (*crr2* and *sig4*) together with dilutions of wt samples (1/2 and 1/4) were probed with radiolabeled RNA probes against four different *ndh* genes. The resulting autoradiographs are always shown with the corresponding methylene blue stains of the membranes (below). The 2 kb marker band is shown as a reference.

Arabidopsis CP31A prefers U-rich sequences in vivo

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Prior *in vitro* experiments showed that a CP31A homolog in tobacco prefers homopolymers of G and U (Li and Sugiura, 1991). We asked whether a sequence preference can be identified in our dataset, i.e. *in vivo*, as well. We used the SSMART motif finder (Munteanu et al., 2018) to query BSs in our RIP-Seq data for common sequences. This uncovered consensus motifs with a clear preference for Us (Fig. 6A; Suppl. Fig. 5). We next compared the frequencies of all possible trinucleotides within significant CP31A BSs against those in all chloroplast coding sequences. Indeed, we found that UUU was the most strongly enriched trinucleotide in this comparison, followed by a number of other U-rich trinucleotides (Fig. 6B). We therefore conclude that CP31A preferentially binds U-rich sequences. As U-rich trinucleotides are the most common trinucleotides in the AT-rich chloroplast genome, our findings suggest that CP31A has evolved to allow a broad RNA target range in the chloroplast transcriptome.

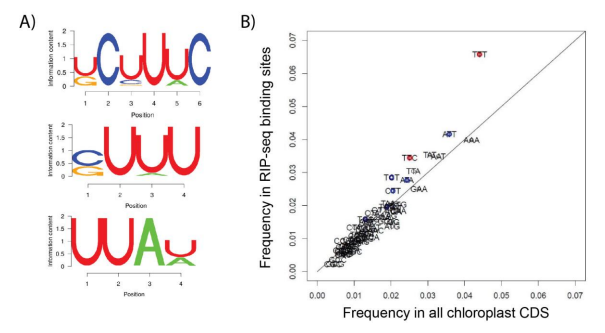


Figure 6: Analysis of sequence elements in CP31A BSs

A) Top scoring consensus motifs identified in CP31A BSs. The SSMART RNA motif finder was run on all significant BSs detected in wt plants grown in cold conditions, as under these conditions, the number of BSs was the highest. The contribution of each BS to the consensus motifs was calculated according to its adjusted p-value - the more significant the BS, the stronger its impact on the consensus.

B) Overrepresentation of the UUU(TTT) trinucleotide in CP31A BSs. The scatterplot shows the frequency of all possible trinucleotide sequences in the CP31A BSs regions (y-axis) versus the trinucleotide frequency in all chloroplast coding sequences. Trinucleotides with significant enrichment in RIP-Seq BS regions are marked in red (p<0.01) or blue (p<0.05).

CP31A shows an increased association with RNA after cold exposure

We hypothesized that CP31A might impact cold resistance in *Arabidopsis* plants through temperaturedependent changes in its affinity for some or all of its target RNAs, which could affect their stability, processing, and/or translation. We therefore compared BSs according to how reliable their enrichment in a given dataset is, based on its FDR value. Using the FDR values, we first focuses on the effect of cold on RNA binding of full-length CP31A and then analyzed the effect of the loss of the AD on binding (next chapter). A majority of BSs were found with high significance in CP31A precipitates in the cold, but were not significantly enriched at higher temperature. This was already visible for some sites in the coverage graphs, as shown in Fig. 7A for binding sites within the rpl2 mRNAs, or for a site in the intergenic region between the rps14 and psbZ genes. Many BSs show at least some read coverage at both temperatures, but the FDR analysis helped to separate significant from non-significant BSs. The FDR level is displayed for all 75 binding sites as a heat map in Suppl. Fig. 6 and is summarized in Fig. 7B. Most importantly, 30 BSs show significant enrichment with full-length CP31A at 4°C, but not at 21°C (Fig. 7B). Only 9 sites show the opposite behavior, that is, significant enrichment only at 21°C, but not at 4°C. Thus many more sites showed increased versus decreased association with CP31A after cold exposure. Three of the BSs gained in the cold were located in the *ndhF* mRNA, which is known to be a prime target of CP31A (Kupsch et al., 2012). Four more CP31A BSs in *ndh* transcripts were cold-dependent as well (Suppl. Fig. 6). In general, ndh mRNAs appeared to be important targets of CP31A under both temperature conditions (Suppl. Fig. 6). Taken together, the results of our temperature-dependent RIP-Seq analysis demonstrate that RNA association of full-length CP31A is higher under cold exposure than at normal temperature, and that ndh mRNAs are prime targets under both normal and low temperatures.

The AD of CP31A supports cold-dependent RNA binding

The RRM domain is a highly versatile protein domain that functions in the context of adjacent protein regions. Often, linkers between RRMs or protein domains outside the RRM domains support RNA interactions in addition to the canonical RRM contacts (Lunde et al., 2007). Given this, we used RIP-Seq to test whether the RNA-binding behavior of the AD-less CP31A protein differed from that of the full-length protein. Using the same strategy described for our binding-site analysis with the full-length protein, we scored significant BSs in plants expressing the AD-less variant grown under both normal and cold conditions. We found that the AD-less version of CP31A bound far fewer RNAs than its full-length counterpart (Fig. 7B, Suppl. Fig. 6). Under normal growth temperatures, we identified 41 significant BSs (FDR < 0.05) for the full-length protein and only 10 for the AD-less version (Fig. 7B). Interestingly, the AD-less protein is still responsive to cold: The AD-less protein associated with 18 sites in the cold that are not found to co-precipitate significantly with the protein at normal temperatures (Fig. 7B). On the other hand, AD-less CP31A bound 35 fewer sites than the full-length protein after cold exposure (Fig. 7B). This demonstrates that the AD supports RNA binding under both conditions. Together, our results show (i) that the two RRM domains still react to lowered temperatures in their ability to bind RNA, and (ii) that the AD substantially supports but is not essential for RNA binding.

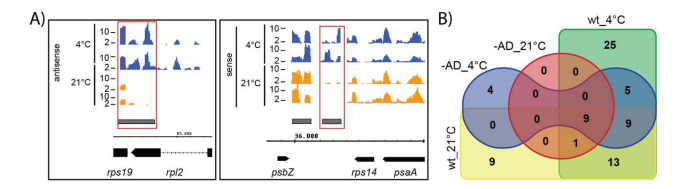


Figure 7: RIP-Seq uncovers cold-induced RNA binding of CP31A.

A) Visualization of three examples of temperature-dependent CP31A binding sites. Top track shows the ratio of precipitated versus input reads of RIP-Seq experiments of wt plants grown at 4°C (blue) or 21°C (orange, for two biological replicates at each growth temperature). The track below represents in grey boxes the identified binding sites. The bottom track shows the gene models in black and an intron in *rpl2* as a hatched line. The numbers on the thin black line refer to positions on the *Arabidopsis* chloroplast genome. The red rectangles indicate temperature-dependent CP31A BSs.

B) Venn diagram summarizing the occurrence of binding sites in the two genotypes and the two temperature regimes used for growing the plants. For each genotype-temperature combination, only BSs with an FDR < 0.05 were counted.

Discussion

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CP31A co-regulates ndh genes

During the evolution of chloroplasts from cyanobacterial ancestors, operon structures were disrupted and operons were shuffled. Many chloroplast operons therefore include genes with different functions. The ndh genes, for example, are separated into four transcriptional units in Arabidopsis and are mixed with genes from other functional categories. This lack of conservation of operon structures suggests that transcriptional units are less important than other processes for *ndh* gene regulation in the chloroplast context, giving way to post-transcriptional processes. Translation plays an important role for regulation (Zoschke and Bock, 2018), but it remains unclear, how post-transcriptional co-regulation is achieved prior to translation. We herein demonstrate that CP31A associates with many *ndh* mRNAs under both normal and low temperatures. All analyzed ndh mRNAs were reduced in cp31a mutants under normal growth conditions, indicating that the CP31A-ndh interaction is functionally important. Since the loss of CP31A does not impede transcription (Kupsch et al., 2012), we conclude that the protein is required for the stability of *ndh* mRNAs. An alternative explanation is that the loss of one RNA of the NDH complex has an indirect, hierarchical effect on the other ndh RNAs that yields the observed reduction in all ndh mRNAs. This phenomenon has been well described for hierarchical protein synthesis cascades in chloroplasts (Choquet et al., 2001), but it has not been previously shown to impact mRNA synthesis or stability. Moreover, the likelihood of this scenario is weakened by our observation that the losses of ndhF or ndhB in the sig4 or crr2 mutants, respectively, were not followed by the loss of all ndh mRNAs. Taken together, our data indicate that CP31A combines ndh mRNAs from different genomic loci into a post-transcriptional operon in the chloroplast. Similar post-transcriptional operons, or "RNA regulons", have been well-described in fruit flies, budding yeast, and mammalian cells (Keene, 2007). In most cases, these RNA operons function in the combined translation and/or stabilization of the participating RNAs (Gerber et al., 2004; Lykke-Andersen and Wagner, 2005; Townley-Tilson et al., 2006). Given the RNA-stabilizing roles of cpRNPs in general and of CP31A in particular, we propose that CP31A adjusts the stability of the *ndh* transcripts as a group.

The AD contributes to the ability of CP31A to recognize RNA

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CP31A has two canonical RRM motifs, which contain the expected aromatic amino acid residues in the RRM's RNP1 and RNP2 motifs known to be crucial for RNA binding (Ruwe et al., 2011). Here, we show that these RRMs are sufficient to edit chloroplast RNAs and stabilize the *ndhF* mRNA. Still, compared to the wt protein, more AD-less protein is needed to stabilize a similar level of *ndhF* mRNA. Moreover, loss of the AD decreases the ability of CP31A to bind RNA. Thus, the AD domain supports but is not essential for the RNA-binding and RNA-stabilizing functions of CP31A. We do not yet know how the AD achieves its effects at the molecular level. Two non-exclusive possible modes of action are (i) a direct involvement in protein-RNA interactions, and (ii) a role in protein-protein interactions for the recruitment of additional RNA processing factors. Both modes of action were already described in non-plant systems for acidic domains (Lunde et al., 2007). Such functional diversity of an AD has been described for two structural relatives of CP31A, the non-plant RNA binding proteins hnRNP Q and hnRNP R, that also have an Nterminal AD and C-terminal RRM motifs (Geuens et al., 2016). The AD of hnRNP R was shown to support RNA binding and is essential for the role of hnRNP R in the re-initiation of transcription (Fukuda et al., 2013). Similarly, the AD of hnRNP Q forms part of the RNA interaction surface (Hobor et al., 2018) and in addition interacts with the editing factor Apobec (Quaresma et al., 2006). These examples show that it is conceivable that the AD of CP31A contributes to protein-protein interactions (e.g., with other RNA-binding proteins) and also contributes directly to RNA binding. This could serve to increase the affinity for RNA and mediate RNA stabilization and RNA-processing events. A future structural analysis of CP31A bound to RNA would greatly help to understand the mechanistic function of CP31A's AD.

Cold-dependent RNA association of CP31A

Many RBPs show differential association with RNA depending on physico-chemical conditions *in vitro*, but relatively little is known about how external and internal cues modulate the target spectrum and target affinity of an RBP *in vivo*. Changed binding to RNA due to external signals has been shown for mammalian RNA binding proteins (Rousseau et al., 2002; Cloutier et al., 2018), but even here, rarely on a genome-wide scale (Benegiamo et al., 2018), and comparable data are missing for plants. Our present results on CP31A show that plant RNA-binding proteins can change their RNA target profiles in response to an external signal,

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here low temperatures, but it remains unclear how this differential RNA binding is induced. Cold is expected to change RNA structures and CP31A can bind dsDNA *in vitro*, albeit with less affinity than ssDNA and ssRNA (Li and Sugiura, 1991). It is also notable that spinach cpRNPs have been implicated in the establishment of the structured 3'-ends of several mRNAs *in vitro* (Schuster and Gruissem, 1991; Lisitsky et al., 1995; Hayes et al., 1996). These RNA termini require the formation of stem-loop RNA structures, and cpRNP-like proteins have been found to associate with such stem-loops in UV cross-linking experiments (Stern et al., 1989; Chen and Stern, 1991; Danon and Mayfield, 1991; Nickelsen and Link, 1993). Thus, the binding of CP31A to RNAs under cold exposure could reflect the emergence of RNA structures that become more stable at lower temperatures. An *in vivo* analysis of the structure of CP31A target sites under different temperatures could help address the question of why CP31A associates more extensively with RNA in the cold.

What is the functional consequence of the increased RNA association of CP31A under cold exposure? Given that chloroplast RNAs are lost in the bleached tissue of cp31a mutants, we propose that an increased interaction with target transcripts could impact the resilience of these transcripts against RNases. An open question however is, which of the many CP31A – transcript interactions in the cold is key to avoid bleaching. Although it is likely that many of these interactions have an incremental effect on the phenotype, we can make some guesses about major contributors. Loss of *ndh* mRNAs, although a prime target of CP31A, is unlikely to be responsible for the bleaching phenotype, since ndh genes are downregulated in the cold in Arabidopsis (Ivanov et al., 2012), and mutants devoid of the NDH complex do not show bleaching in the cold (Li et al., 2004; Yamori et al., 2011). By contrast, there is a number of other CP31A target RNAs that are known to be essential for chloroplast development and cold-dependent loss of their processing and stabilization could well explain bleaching. This includes mRNAs for subunits of the ATP synthase (atpA, atpB), of the cytochrome $b_0 f$ complex (e.g. petB, petD), of the photosystems (e.g. psaA, psbB), as well as mRNAs for subunits of the plastid-encoded RNA polymerase (rpoB, rpoC2). Loss of the expression of any of these genes leads to defective chloroplasts including pale phenotypes (Hajdukiewicz et al., 1997; Rogalski et al., 2006; Stoppel et al., 2011; Manavski et al., 2015; Chen et al., 2016). Furthermore, many mutants affected in chloroplast translation are more sensitive to cold than wt (Rogalski et al., 2008; Fleischmann et al., 2011; Gu et al., 2014; Wang et al., 2016; Zhang et al., 2016; Paieri et al., 2018; Pulido et al., 2018). This includes mutants of chloroplast RNA binding proteins that affect translation and that show cold-dependent bleaching of the center of the Arabidopsis rosette (Wang et al., 2016), which is strikingly similar to the phenotype of cp31a mutants in the cold. CP31A binds to several mRNAs coding for ribosomal proteins (e.g. rps18), thus being potentially important for the production of the translational machinery in the cold. Taken together, we hypothesize that CP31A becomes important for the expression of a combination of its target mRNAs in the cold; their compromised expression in cold-treated cp31a mutants leads to loss of chloroplast development and thus bleaching. Complementation analyses and time-resolved analysis of the accumulation of mRNAs in *cp31a* mutants during cold treatment could reveal, which mRNAs are key targets for CP31A during cold acclimation.

Methods

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Plant growth

Arabidopsis thaliana Columbia-0, *cp31a-1* T-DNA insertion mutants (Tillich et al., 2009) and AD-deletion mutants were grown on soil with a 16-h-light/8-h-dark cycle at 23°C in a CLF growth cabinet at 120 μmol·m⁻²·s⁻¹. For cold stress application, temperatures were lowered to 8°C when plants reached an age of two weeks. Cold exposure was then continued for another five weeks to allow the emergence of sufficient new tissue prior to harvest. For RIP-seq experiments plants were grown on a soil/vermiculite 4:1 mixture at 21°C for 14 days (normal conditions). Cold stress was applied after 13 days at 21°C with a 24 h exposure to 4°C.

Vector construction and production of transgenic plants

A full length Arabidopsis CP31A cDNA clone was prepared from total Arabidopsis RNA. The genomic region encoding the acidic domain (+241 to +447 bp from the start codon of the open reading frame) was deleted as follows: The 5' region of CP31A (1-240 bp) and the 3' region of CP31A (448-990 bp) were amplified with the primers 31A_gateway_for and A31-minusADrev and A31-minusAD_for and 31A gateway rev, respectively. These 5' and 3' regions were joined by restriction-based ligation into a pBluescript vector. The p35SP:31A△AD binary vector for constitutive expression of CP31A was constructed as follows: The 31A△AD fragment was amplified with the primer set (SmaI-31AF01; 31A-SmaIR01) and ligated into pJET1.2 cloning vector (Thermo Scientific), and the SmaI digested product was integrated into a SmaI site between the 35S promoter and Nos terminator sequences of the binary vector pGL1 (provided by Dr. Boris Hedtke, HU Berlin), which also contains the bar expression cassette. For producing the p31AP:31A vector, the CP31A 5' UTR and promoter region (1,694 bp of the 5' region of CP31A; 12769646 to 12767953 of the Arabidopsis thaliana chromosome 4 sequence; Acc. No. CP002687.1) was cloned with the specific primer set (CP31A5'up-1694; CP31A-CDS5'_rev). Then, the CP31A cDNA was combined with the promotor by Gibson cloning using the primers (XhoI-31Apro F; Cp31A-CDS5' rev; CP31A-ADtest for; 31A-SmaIR01) and cloned into pJET1.2, yielding pXhoI-31AP:31A-SmaI. The 35S promoter was removed and replaced with multi cloning site including XhoI and SmaI sites, yielding vector pGL1-MCS. XhoI/SmaI digestion products from pXhoI-31AP:31A-SmaI were inserted into XhoI/SmaI sites of pGL1-MCS, yielding p31AP:31A vector. p31AP:31A△AD was constructed in an analogous way. These three vectors were introduced into the Agrobacterium GV3103 strain independently, and integrated into the knock-out mutant *cp31a-1* (SALK_109613; Tillich et al., 2009) by the floral-dip method. The T₀ generation

530 was grown on 15 cm diameter plastic plates filled with soil (mixture of horticulture soil and vermiculite). 1/2000 diluted BASTA was sprayed on the plants several times in order to select for BASTA resistant plants. 531 BASTA resistant plants were transferred to plastic pots filled with soil and cultivated at 23°C, 8-h-dark/16-532 h-light at a light intensity of 120-130 µmol·m⁻²·s⁻¹. 533 534 **Immunoblot analysis** 535 Total protein (15 µg) were extracted from fully developed leaves grown at normal conditions and 536 immunological analysis were carried out and antibodies were used as previously reported (Kupsch et al., 537 2012). Immunoblot analysis for the RIP-seq experiments was performed in the same way with protein 538 samples taken from the input, supernatant and pellet fraction of the co-immunoprecipitations. 539 **RIP-Seq analysis** 540 For RIP-seq experiments Arabidopsis thaliana Columbia-0 and AD-deletion mutants under the native promoter (#5-7-1, #5-13-2) were harvested in duplicates after 14 days and flash-frozen in liquid nitrogen. 541 542 The flash-frozen plant material was grinded under liquid nitrogen to a homogeneous powder. Between 250 543 to 350 mg plant material was suspended in 3 ml RIP-seq lysis buffer containing formaldehyde (50 mM 544 HEPES-KOH pH 8.0, 200 mM KCl, 5 mM MgCl₂, 5 mM CaCl₂, 0.5% Nonidet P-40, 0.5% Sodium deoxycholate, 1x cOmpleteTM, EDTA-free Protease Inhibitor Cocktail (Roche), 100 U RiboLock RNase 545 546 Inhibitor (ThermoFisher Scientific), 1% formaldehyde) per 1 g plant material. Crosslinking was performed 547 for 10 min under rotation at room temperature and stopped by the addition of 125 mM glycine and a 5 min 548 incubation. The plant extract was centrifuged for 10 min at 20,000xg and 4°C to remove insoluble plant 549 material and flash-frozen in liquid nitrogen. 550 For the co-immunoprecipitation (CoIP) 8 µl affinity-purified anti-CP31A antibody (Kupsch et al., 2012) 551 was bound to 50 µl Dynabeads ProteinG (Invitrogen) under rotation (15 rpm). The plant extract was thawed 552 and centrifuged for 10 min at 20,000xg and 4°C. 350 µl of the supernatant were diluted 1:1 with CoIP buffer 553 (150 mM NaCl, 20 mM Tris-HCl pH 7.5, 2 mM MgCl₂, 0.5% Nonidet P-40, 5µg/ml Aprotinin) and 554 incubated with the antibody-coated magnetic beads for 75 min at 15 rpm and 4°C. An aliquot of the 555 antibody-bead solution was taken to serve as the input control. The beads were washed four times in CoIP 556 buffer and resuspended in Proteinase K buffer (100 mM NaCl, 10 mM Tris-HCl pH 7.0, 1 mM EDTA, 0.5% 557 SDS). 558 The crosslink was reversed with 0.1 mg/ml Proteinase K (ThermoFisher Scientific) at 50°C for 1 h and RNA 559 was extracted from input and pellet fractions using TRIzol and RNA Clean and Concentrator Columns 560 (Zymo Research) according to the manufacturer's instructions.

Library preparation was performed with the NEBNext® Multiplex Small RNA Library Prep Set for Illumina (New England BioLabs) according to the manufacturer's instructions with few deviations. Library preparation was performed for half the volume. Additionally, a 5' adaptor including unique molecular identifiers (5'-(UMI) was used rGrUrUrCrArGrArGrUrUrCrUrArCrArGrUrCrCrGrArCrGrArUrCGATCNNNNNNNN-3´). **PCR** amplification was performed using the KAPA HiFi HotStart ReadyMix with the cycling protocol for library amplification for Illumina platforms and an annealing temperature of 62°C. The PCR amplified cDNA construct was purified using the GeneJET PCR Purification Kit (ThermoFisher Scientific) according to the manufacturer's instructions and then separated on 6% polyacrylamide gel. Library fragments between 160bp and 190bp were extracted from the gel according to the NEBNext® protocol and subjected to Illumina sequencing. Read numbers and mapping results are summarized for all libraries in Supple. Table 3.

- For the bioinformatic identification of the CP31A binding events the following steps were performed:
- 573 1) Mapping of the Illumina reads. First, reads were split depending on its barcode using 574 fastx_barcode_splitter (version 11sep2008) to identify to which sample they correspond, next the UMI 575 barcode was extracted for each read using umi tools, later, reads were trimmed using Trimmomatic version 576 0.36 with the next parameters: LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:10 in order to 577 eliminate Illumina adapters and bad quality sequence regions. The obtained trimmed reads were aligned to 578 the Arabidopsis thaliana genomes (Araport11) using STAR version 2.5.2a with parameter: --579 outFilterMultimapNmax 2 --outSAMtype BAM SortedByCoordinate --alignIntronMax 3000 580 outFilterIntronMotifs RemoveNoncanonical --outSAMstrandField intronMotif --outSAMattrIHstart 0. At 581 this point, duplicated reads were removed from the BAM file using umi_tools. Only reads that mapped to 582 the chloroplast genome with a length bigger than 35 bp were used for downstream analysis.
 - Remaining reads were used to identify BSs using CSAR (version 1.31.0; parameters: b=10, w=70, nper=100, test="Ratio") for each strand and sample/replicate independently. Only those regions that showed a significant enrichment in mapped reads when comparing IP versus corresponding input samples at a false discovery rate (FDR) < 0.05 were kept (Muino et al., 2011), which led to the discovery of 371 candidate BSs across all samples. In order to obtain a common set of candidate BSs for all samples, the list

2) Identification of candidate BSs in each condition studied. Reads mapped in rRNA were eliminated.

- candidate BSs across all samples. In order to obtain a common set of candidate BSs for all samples, the list of BSs for each sample/replicate were merged using the software *mergeBed* from the package *bedtools*
- 590 (v2.26.0). This resulted in 75 reference BSs.

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- 3) Quantification of BSs. The program *featurecounts* (v1.6.0; parameters: -s 1 -M) was used to count the
- number of reads strand-specifically mapping to the common set of candidate BSs obtained from the previous
- step. DESeq2 (v1.14.1) was used with defaults parameters except fitType="local" to obtain normalized

number of mapped read in each sample, next it was used to calculate enrichment and significance of number

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of the candidate BSs comparing IP vs control (two biological replicates). The normalized number of reads were used to calculate Pearson correlation coefficients. SSMART analysis of CP31A RIP-seq data SSMART analysis was carried out as described (Munteanu et al., 2018). We used all significant BSs identified in wild type plants grown in cold conditions as input, as in this sequence set most binding events were detected. The input sequences of the BSs were scored using their adjusted p-value according to the actual binding site analysis of the RIP-seq data. RNA extraction and editing analysis Total RNA was extracted from fully developed leafs (0.1 g) powdered in liquid nitrogen using Trizol (Thermo Fisher) according to the manufacturer's protocol. DNA was removed from RNA samples by three consecutive DNase I treatments and Phenol/Chloroform/Isoamyl alcohol extractions. DNA removal was checked by PCR with the chloroplast-specific rps14 primer set. cDNA synthesis was performed with 2 µg RNA using SuperScript III first strand synthesis system (Invitrogen) according to the manufacturer's protocol. A one-to-ten dilution of cDNA was used as a template for amplifying ten cDNAs encompassing 16 editing sites (Fig. 2A) on nine plastid genes using primer sets previously reported (Tillich et al., 2009). RT-PCR products were purified with MinElute PCR Purification Kit (Qiagen). Bulk RT-PCR products were cloned into the barcoded cloning vector and sequenced using the Illumina MiSeq machine. Sequenced data were sorted according to barcoding of vectors and analyzed with the CLC Genomics Workbench (CLC bio). Selected RT-PCR products were directly analyzed by Sanger Sequencing and analyzed using the Geneious software. RNA gel blot analysis Total RNA (4 µg) was fractioned on 1.2% agarose gels containing 1.2% formaldehyde, blotted and hybridized with radiolabeled RNA probes produced by T7 in vitro transcription from PCR products generated with primer combinations described in Suppl. Tab. 2. Acknowledgement We wish to acknowledge expert technical support by Irina Passow. This work was supported by grants of the Deutsche Forschungsgemeinschaft to CSL (A02 of TRR175), DL (C05 of TRR175) and TR (FOR 2092). Support of MKL by the IRI Life Sciences and the "Frauenförderung des Instituts für Biologie" of the Humboldt University is gratefully acknowledged. AO was supported

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- 625 discussion of the data.
- 626 AO, MKL, and TR performed the research and analyzed the data. JMM performed the
- 627 computational analysis of the RNA seq data. BL performed the analysis of the CP31A BS
- 628 consensus. DL, UO and CSL designed the research and analyzed the data. CSL wrote the paper
- with contributions of all authors to the final version.
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