Protein Phosphatase 2A B55 and A Regulatory Subunits Interact with Nitrate Reductase and Are Essential for Nitrate Reductase Activation¹[W]²[OA]

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Posttranslational activation of nitrate reductase (NR) in Arabidopsis (Arabidopsis thaliana) and other higher plants is mediated by dephosphorylation at a specific Ser residue in the hinge between the molybdenum cofactor and heme-binding domains. The activation of NR in green leaves takes place after dark/light shifts, and is dependent on photosynthesis. Previous studies using various inhibitors pointed to protein phosphatases sensitive to okadaic acid, including protein phosphatase 2A (PP2A), as candidates for activation of NR. PP2As are heteromeric enzymes consisting of a catalytic (C), structural (A), and regulatory (B) subunit. In Arabidopsis there are five, three, and 18 of these subunits, respectively. By using inducible artificial microRNA to simultaneously knock down the three structural subunits we show that PP2A is necessary for NR activation. The structural subunits revealed overlapping functions in the activation process of NR. Bimolecular fluorescence complementation was used to identify PP2A regulatory subunits interacting with NR, and the two B55 subunits were positive. Interactions of NR and B55 were further confirmed by the yeast two-hybrid assay. In Arabidopsis the B55 group consists of the close homologs B55α and B55β. Interestingly, the homozygous double mutant (b55α × b55β) appeared to be lethal, which shows that the B55 group has essential functions that cannot be replaced by other regulatory subunits. Mutants homozygous for mutation in Bf and heterozygous for mutation in Bα revealed a slower activation rate for NR than wild-type plants, pointing to these subunits as part of a PP2A complex responsible for NR dephosphorylation.

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divalent cations to extracts for activity (MacKintosh, 1992). This makes PP6 an unlikely candidate for regulation of NR. Arabidopsis (Arabidopsis thaliana) has two, weakly expressed, genes encoding PP4 (Farkas et al., 2007). Furthermore, the two PP4 phosphatases (PPX1 and PPX2) are localized inside plastids (Pujol et al., 2000), therefore not likely to dephosphorylate NR, which is localized in the cytosol. Protein phosphatases with Kelch-repeat domains (BSU1, BSL1, 2, and 3) are also inhibited by okadaic acid, although higher concentrations are needed than for PP2A. BSU1 is localized in the nucleus and present in young tissue. BSL1, 2, and 3 are expressed in leaves and older tissue (Mora-García et al., 2004), hence can be considered candidates for dephosphorylating NR. PP5 is known to promote expression of the light-regulated genes, CHALCONE SYNTHASE, CAB2, and RBCS (Ryu et al., 2005), and is a putative candidate for regulating also NR.

In Arabidopsis, as well as in other eukaryotes, PP2As are oligomeric proteins composed of a 36- to 38-kD catalytic subunit (C) bound to a scaffolding subunit (A) of 65 kD. A number of variable regulatory subunits of molecular mass 48 to 74 kD associate with the core dimer to form a trimeric PP2A. Presently three B families are identified in plants (B/B35, B’, B”; Farkas et al., 2007; Wang et al., 2007). The A subunit is required as a scaffold for the formation of the heterotrimeric PP2A complex. In mammals, two different A subunits are present, whereas Arabidopsis has three such scaffolding subunits, and yeast (Saccharomyces cerevisiae) has one. The A subunits are composed of 15 imperfect HEAT repeats, each of 39 amino acids, that form a hook-shaped molecule. The repeats consist of two α-helices connected by an intrarepeat loop. B subunits bind to repeats 1 to 10, whereas C subunits bind to repeats 11 to 15 (Li and Virshup, 2002). The Arabidopsis A subunits have been studied by different research groups, and are known to be important for root morphology (Zhou et al., 2004) and auxin transport (Michniewicz et al., 2007). They show overlapping functions, and the triple knockout is most likely lethal (Zhou et al., 2004; Michniewicz et al., 2007). One of the A subunits, ROOTS CURL IN NAPHTHYLPHTHALAMIC ACID1 (RCN1), was suggested to have a cardinal role in regulation of phosphatase activity, whereas functions of the other two, PP2AA2 and PP2AA3, were only unmasked when RCN1 was absent (Zhou et al., 2004). By using these different loss-of-function mutants and a transgenic line with tamoxifen-inducible expression of artificial microRNAs (amiRNAs) targeting all three A subunits of PP2A (Michniewicz et al., 2007), we have investigated the importance of A subunits for activation of NR. A clear effect on NR by inducible knockdown of all three A subunits established the involvement of PP2A for activation of NR in vivo.

The PP2A regulatory B subunits play an important role in conferring substrate specificity and determine subcellular localization of PP2As in mammals (Sontag, 2001; Eichhorn et al., 2009). It is assumed that also in plants the different B subunits have such functions (Matre et al., 2009). In Arabidopsis, the TON2/FASS is the only B (a B’) regulatory subunit that has been firmly characterized so far, and was found to be involved in cortical microtubule organization and important for cell shape (Camilleri et al., 2002; Wang et al., 2007). Very recently it was shown that silencing of TAP46 in Arabidopsis, the ortholog of the yeast TAP42 protein, a PP2A catalytic subunit interacting protein, affects NR expression (Ahn et al., 2011). When we tested different B subunits for interaction with NR, positive results were found for the B55 group. In other eukaryotes B55 has a regulatory role in mitosis (Mochida et al., 2009; Schmitz et al., 2010). Little is known about the functions of the two 55-kD B isoforms in plants. This type of subunit is almost entirely composed of β-sheets and turns, whereas B’ and B” are mostly α-helical. B55 subunits contain four to seven degenerate WD40 repeats. In mammals the B55 subunit family consists of four members, α, β, δ, and γ (Eichhorn et al., 2009). The mammalian Bγ is the form most similar to the plant B55. Evolutionarily, plants and mammals divided before the various isoforms were developed, and their isoforms are named independently (Teron et al., 2002). A necessary step for dissecting the signal transduction chain connecting processes in the chloroplasts and activation of NR in the cytosol is to identify phosphatase(s) of physiological importance for NR activation, and this is the subject of the present report.

RESULTS

PP2A-Like Protein Phosphatases

PP5 is a protein phosphatase susceptible to okadaic acid inhibition, and a possible candidate for activation of NR. NR activity state following dark-to-light transitions was tested for a PP5 loss-of-function mutant (pp5ko) and also a PP5 overexpressor (pp5ox; papp5-1 and PAPP5-OX1 in Ryu et al., 2005). No effects on NR activation during dark-to-light transitions were revealed for these mutants (see Supplemental Fig. S2). Mutants homozygous for T-DNA insert in PPP-KELCH motif protein phosphatase genes BSL1, BSL2, and BSL3 were tested, however no indications of effects on NR activation were seen. From these experiments it can be concluded that none of these phosphatases had a major effect on NR activation (Supplemental Fig. S2), however they cannot be excluded from having some effect on NR, that would be overtaken by another phosphatase in the mutants.

PP2A A1, A2, and RCN1 Subunits and NR Activity

A triple knockout for all three A subunits is to our knowledge not available, and this is most likely due to
lethality of such a mutant (Michniewicz et al., 2007). Knockdown of all three A subunits of PP2A was achieved by using tamoxifen-inducible microRNA overexpression with amiRNA-1 and amiRNA-2 lines (Michniewicz et al., 2007). Three-week-old plants were treated with tamoxifen every day for 1 week, which led to a strong decrease (90%) in expression of the A subunit examined (Fig. 1A). A marked decrease in NR activation was observed in these amiRNA plants (Fig. 1B). The NR activity state gives the amount of non-phosphorylated (active) NR as a percentage of the total amount of NR (the nonphosphorylated plus phosphorylated forms; MacKintosh et al., 1995). Only the phosphorylated form of NR is inhibited by Mg++. When Mg++ is present in the assay only nonphosphorylated NR is active, but when EDTA is added to the assay also the phosphorylated form becomes active because of sequestering of cations and release of the 14-3-3 inhibitor (Huber et al., 2002). The differential Mg++/EDTA assays, therefore, enable an efficient and precise way of calculation of activity state and also adjust for possible variations in total NR in each extract (Supplemental Fig. S3). Within 40 min of illumination, NR activity state increased from 10% to 66% in mock-treated amiRNA-1 plants, and from 11% to 28% in tamoxifen-treated amiRNA-1 plants (Fig. 1). Wild-type (Columbia [Col]) plants treated with tamoxifen confirmed that NR was still activated as in nontreated wild-type plants (Fig. 1B). The low PP2AA expression seemed to have no effect on NR inactivation during the dark period since the NR activation state was the same in mock- and tamoxifen-treated darkened plants (time zero in Fig. 1B). NR activity was also measured in A-subunit loss-of-function mutants (Zhou et al., 2004), i.e. rcn1 (Wassiljevskaja [WS] background) and double mutant pp2aa2 pp2aa3 (Col background; Fig. 2). Activity state for samples harvested during the photoperiod was 47% ± 3% for the pp2aa2 pp2aa3 double mutant and significantly higher, 62% ± 4%, for the wild-type Col control (Fig. 2A). NR activity state in the rcn1 mutant was not significantly different from the control (WS; Fig. 2A). Samples harvested after 2 h of darkness showed activity state between 11% and 16%, but no significant differences between mutants and controls (Fig. 2B). Change in activity state in response to light on/off was also tested, and showed that for the pp2aa2 pp2aa3 double-mutant activation of NR was lower than in wild type. After 20 min of light, wild type showed activity state of 38% ± 7% and the double mutant showed activity state of 26% ± 5% (Fig. 2C). For the rcn1 mutant the difference from the WS control was not significant (Fig. 2D). In summary these experiments showed that PP2A is required for rapid NR activation in vivo, and that the structural A subunits have overlapping functions in this process.

Subcellular Localization of B55 Subunits and Interactions with NR

NR1 and NR2 linked by their C terminus to EYFP were transiently expressed in protoplasts of Arabidopsis suspension culture and mesophyll cells. The fusion proteins localized to the cytosol, with a tendency to form speckles (representative pictures are shown in Fig. 3, A and B). The B55α and B55β subunits both localized to the cytosol regardless of N- (Fig. 5, C and D) or C-terminal (not shown) fusion to the EYFP. Interaction of NR1 and B55α or B55β was tested in planta by bimolecular fluorescence complementation (BiFC). This assay is based on the reconstitution of

Figure 1. The scaffolding A subunits of PP2A are required for light activation of NR. A, Relative transcript levels for the PP2AA subunit in tamoxifen-treated wild-type and amiRNA-1 plants, and mock-treated plants. Expression levels were tested with RT-PCR using SYBR green with plant samples also used for testing NR activation. SE for the RT-PCR assay is given, n = 3. B, Activity state of NR following a dark-to-light transition of wild-type (dots) and amiRNA-1 (squares) plants treated with tamoxifen (black symbols), and mock-treated plants (white symbols). Plants were grown in 12-h light/12-h darkness for 3 weeks. Two hours into the photoperiod, when NR activity was at a high level, plants were transferred to darkness and samples harvested after 1 h of darkness (time zero), then after 20 and 40 min of white light (150 μmol m⁻² s⁻¹). Data are means of three samples, SE is given.
EYFP when nonfluorescent N-terminal (NFP) and C-terminal (CFP) fragments of the EYFP are brought together by two interacting proteins fused to NFP and CFP (Hu et al., 2002; Maple et al., 2005). The NFP was fused to the C-terminal end of full-length NR1, and CFP was fused to the C-terminal end of full-length B55\(\alpha\) and B55\(\beta\). The results firmly established that NR1 and B\(\alpha\) interact in planta (Fig. 4, A and B). In experiments performed with B\(\beta\), some interactions were detected, however, with lower occurrence. Several other B’ and B” subunits (At5g03470/B’\(\alpha\), At3g09880/B’\(\beta\), At3g26030/B’\(\delta\), At1g13460/B’\(\theta\), At4g15410/B’\(\gamma\), At3g26020/B’\(\eta\), At3g54930/B’\(\epsilon\), At3g21650/B’\(\zeta\), At5g18580/TON2, and At5g53000/TAP46) were also tested for interaction with NR1 without positive results. Combined these results showed that specifically B55, but not other B subunits tested, interacted with NR1. To confirm NR interactions with B55\(\alpha\) and to determine the interacting domains, we furthered the BiFC experiments by testing interaction between NR protein fragments and B55-type subunits. The experiments showed that B55\(\alpha\) interacted with the domain containing the hinge1 flanked by dimerization domain and heme-binding domains (amino acid 340-621; Fig. 4, C and D). The N-terminal 100 amino acids containing an acidic domain, and the flavin-binding domain linked with hinge2 (amino acid 621–917), gave the same type of positive images (not shown). The molybdenum cofactor factor binding domain (amino acid 100–340) did not interact with B\(\alpha\). For scheme of interacting domains see Supplemental Figure S4 (Buchanan et al., 2000).

To verify the BiFC interactions of NR and B55 the yeast two-hybrid assay was employed. To this end we decided to assay the specific PP2A regulatory B55 subunits by coexpression with NR1 in yeast. On selection medium lacking His, the yeast cells are able to grow only when BD-NR1 and an AD-B subunit interact and reconstitute the transcription factor needed for growth in the absence of His. His auxotrophy was restored in cells cotransformed with BD-NR1 and AD-B55\(\alpha\). Growth experiments were repeated in triplicate and showed reconstituted growth when the BD-NR1 and AD-B55\(\alpha\) were coexpressed, whereas controls with empty AD vector showed nearly no growth, and controls with empty BD vector showed some retarded growth only. Relative growth was measured by incubating 5 \(\mu\)L of yeast cultures on the agar medium for 48 h, the colonies were resuspended in synthetic dropout medium lacking His, Trp, Leu (SD/-HTL), and the absorbance was read at 600 nm (Fig. 5). Similar results were obtained for BD-NR1 and AD-B55\(\beta\), however, due to high autoactivating for the BD-empty control plus AD-B55\(\beta\) coexpression, the assay was nonconclusive for B55\(\beta\).

**Mutations in B55 Subunits Affect NR Activity State**

The T-DNA single-mutant lines of B55\(\alpha\) or B55\(\beta\) were not different from wild type regarding NR activation (Supplemental Fig. S5). We did not succeed in obtaining a homozygous double mutant, most likely due to lethality. An additional complicating factor was that the B\(\alpha\) and B\(\beta\) genes are situated on the same chromosome.
The homozygous single-mutant lines were crossed, and the F1 progeny was selfed. When two genes are linked it is easier to find double mutants in the F3 generation after suitable selection of crossover between the two mutated alleles (Weigel and Glazebrook, 2002). First two plants homozygous for the \( B_b \) mutation and heterozygous for the \( B_a \) mutation were selected in the F2 progeny. In these two plants a crossing over had taken place, leading to mutated \( B_b \) and \( B_a \) on one chromosome, whereas the sister chromosome was mutated in \( B_b \) only. Selfing of such plant should give 25% homozygous double-mutant plants in the F3 generation (if viable), but none appeared among 30 plants first tested. Therefore, plants heterozygous for T-DNA insert in \( B_55a \) and homozygous for T-DNA insert in \( B_55b \) were propagated, and identified in each generation by PCR. In total approximately 100 plants were tested, but a homozygous double mutant never appeared, which confirmed the assumption that such homozygous mutations would be lethal. See Supplemental Figure S1 for details. All seeds from the selfed plants germinated and seedlings were heterozygous, showing that lethality would have occurred prior to seed formation, i.e. a homozygous double mutant would likely be embryo lethal. To test for a possible dose effect of \( B_55 \) on NR activation, plants heterozygous for \( B_55a \) insert and homozygous for \( B_b \) insert were identified and used to test for NR activation (Fig. 6; Supplemental Fig. S6). In these plants a reproducible and significant decreased NR activation rate was found. NR activity state increased from 13% to 30% in wild type (four intact \( B_55 \) alleles), and from 13% to 21% in the mutant (one intact \( B_55 \) allele).

**DISCUSSION**

Based on previous studies with inhibitors, it was anticipated that PP2A would be important for dephosphorylation and hence activation of NR in response to dark/light transitions of plants (MacKintosh, 1992; Huber et al., 2002). In agreement with this hypothesis knockdown of all three A (scaffolding) subunits by amiRNA strongly impaired NR activation hence the involvement of PP2A in NR activation in situ was firmly established (Fig. 1). Complete abolishment of expression is, however, difficult to achieve by inducible amiRNA, and a low background activity of subunit A expression could therefore possibly be responsible for creating the slow NR activation observed (Fig. 1). Some residual activity due to a complex of the catalytic PP2A subunits with \( B_55 \), but devoid of A subunits, as has been found for yeast PP2A (Koren et al., 2004) cannot be excluded. Alternatively, a different protein phosphatase, not yet identified, may additionally be involved. It was previously found that PP2A activity, as measured by phospho-

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**Figure 3.** Fusion proteins of NR and B55 subunits localize to the cytoplasm in Arabidopsis protoplasts. A, NR1 (At1g77760) and B, NR2 (At1g37130) were fused at their C terminus to EYFP (NR-EYFP) and transiently expressed in protoplasts from Arabidopsis cell suspension culture. C, B55a (At1g51690) protein fused at its N terminus to EYFP and transiently expressed in protoplasts localized to cytosol. D, B55B (At1g17720) protein fused at its N terminus to EYFP and transiently expressed in protoplasts localized to cytosol.

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**Figure 4.** Full-length and truncated forms of NR interact with \( B_a \) as tested by BiFC. A and B, Interaction of full-length NR1 and \( B_a \) in cytosol of Arabidopsis mesophyll protoplasts. C and D, Interaction of the NR1 fragment comprising the dimerization domain, hinge1, and heme domain with \( B_a \) in Arabidopsis mesophyll protoplasts. Scale bar = 10 \( \mu \)m.
histone dephosphorylation, in extracts from the rcn1 mutant was reduced by about 50% (Zhou et al., 2004). However, NR activation was not impaired in the rcn1 mutant. NR activation was lowered in the pp2aa2 pp2aa3 double mutant, showing that the PP2AA2 and PP2AA3 subunits were necessary for full NR activation, but since activation of NR was still taking place to a high degree, RCN1 could apparently substitute for PP2AA2 and PP2AA3. In conclusion the results are in agreement with NR being activated by PP2A complexes containing different structural A subunits.

Clearly, the B55α subunit interacted with NR as shown by BiFC and the yeast two-hybrid assay, and is thereby a candidate for targeting a PP2A complex to NR. B55α interacted with different domains of NR. Two of the domains tested, i.e. the N-terminal 100 amino acids, and the FAD-binding domain, do not contain the known regulatory, phosphorylation site. This implies that interaction between B55 and NR does not require the conserved motif in hinge1. This is in agreement with the knowledge that short motifs are recognized by protein kinases, whereas phosphatases do not appear to recognize consensus motifs within their substrates (Virshup and Shenolikar, 2009). Interestingly, the N-terminal end, which interacted with B55α in the BiFC test, was previously pointed out as important for regulation of NR from Nicotiana plumbaginifolia (Nussaume et al., 1995; Provan et al., 2000).

We did not find any effect on NR activity when comparing B55α or B55β single mutant with wild-type plants. Since B55α and B55β are close homologs, i.e. 87% similar, they may have overlapping functions. The single mutants were crossed, but interestingly a mutant homozygous for T-DNA inserts in both genes could not be obtained and is most likely lethal. This showed that the B55 subunits are essential and not replaceable by the other B (B’ and B”) subunits. By propagating the double mutant through plants homozygous for insertion in Bβ and heterozygous for insertion in Ba, and identifying these plants in every generation a dosage effect of the B55 genes on NR activation was shown, confirming that B55 is indeed involved in activation of NR. Single mutants of several B’ and B” subunits had been tested without finding effects on NR activity, thereby excluding each for being solely responsible for NR activation. However, a contribution from some of the B’ and B” subunits in targeting PP2A to NR cannot be excluded. By testing several PP2A-like phosphatases it was also recognized that none of these were exclusively responsible for PP2A interaction with NR, but again some contribution to NR activation is possible.

In conclusion, the results showed that PP2A structural genes are essential for activation of NR in situ after a dark-to-light transition. Furthermore, the regulatory B55 subunit genes are required for full activation rate of NR. Further work will include analysis of the posttranslational modifications of the subunits in response to light and photosynthesis.

**MATERIALS AND METHODS**

**Plant Growth**

Seeds for inducible knockdown of all three A subunits of PP2A were provided by Professor Jiri Friml (Ghent University, Belgium). Two different amiRNAs had been used to construct lines with inducible overexpression of...
amirNA (amiRNA-1, amiRNA-2) that simultaneously targeted all three A subunit genes of PP2A (Schwab et al., 2006; Michniewicz et al., 2007). Seeds with loss-of-function mutations in the RCN1, PP2AA2, and PP2AA3 genes were provided by Professor Alison DeLong (Brown University, RI). The rcn1 and pp2aa2-1 pp2aa3-1 double mutants were used (Zhou et al., 2004).

Arabidopsis (Arabidopsis thaliana) mutant lines SALK_062514 (B55b), SALK_095004 (B55a), SALK_055355 (B5L2), SALK_088260C (B5L1), and SALK_071689C (B5L3) were from the Ecker collection (Alonso et al., 2003) and obtained through the European Arabidopsis Stock Centre (Nottingham).

Homologous mutant selection was done by PCR using primers for T-DNA insertion lines recommended by the SALK institute Web site SIGnALL (http://signal.salk.edu/tdnampriamers2.html). For the B55a × B55b double mutant, plants heterozygous for the B55a mutation and homozygous for the B55b mutation were identified (Supplemental Fig. S1). To confirm that B55 mutant lines are true knockouts they were verified with reverse transcription (RT)-PCR using gene-specific primers (Supplemental Fig. S1).

Seeds from Pp5 overexpression plants (PpPp5-0X1) and loss-of-function mutant (papp5-1) were provided by Professor Hong Gil Nam (Pohang University, South Korea).

Small (2 weeks after sowing) amiRNA plants were induced by tamoxifen/ estradiol treatment. Plants were sprayed every day for 1 week with 10 μM tamoxifen and 50 μM estradiol (Sigma-Aldrich).

The seeds were stratified at 4°C for 1 to 3 d and then transferred to the green house. During germination and growth plants were placed at 20°C under artificial light in short days (8-h light/16-h dark) or long days (16-h light/8-h dark) or 12-h light/12-h dark regimens.

**Cell Culture**

Arabidopsis (Col) cell suspension was cultured in JPL medium as described by Droillard et al. (2000) at 22°C in constant light. Cells were subcultured every week and harvested 3 to 4 d after subculture. Protoplasts were prepared using 1% w/v cellulase RS (Yakult) and 0.2% Macerozyme R-10 (Yakult; Boudsocq et al., 2007).

**Assay of NR Activity**

Plants were grown in soil. Rosette leaves (0.2–0.3 g) were homogenized in a mortar with cold, 800 to 1,000 μL extraction buffer 100 mM HEPES-KOH, 1 mM EDTA, 7 mM cystein. The extract was centrifuged for 2 min, and the supernatant, 50 μL, was assayed in 700 μL assay volume with 200 μM NADH and generally 2 mM EDTA to assay total NR activity (nonphosphorylated plus phosphorylated NR). To measure actual activity (nonphosphorylated NR) NR was tested in the presence of 5 mM MgCl2. The assay was run at 25°C for 10 min. Nitrite formed was determined by addition of 700 μL 1% sulfanilamide and 0.02% N-(naphthyl)-ethylendiamine dihydrochloride in 1.5 N HCl, and read spectrophotometrically at 540 nm. Absorbance of 1 corresponds with 27 μmol NO2 formed per g fresh weight per h. NR activity state was defined by the actual activity (Mg assay) to the total NR activity (EDTA assay) multiplied by 100 (Mackintosh et al., 1995).

**Gene Cloning for in Planta Expression**

All cDNAs were cloned into pWEN18, pWEN 25, pWEN-NY, and pWEN-CY in the same way: Full-length coding sequences of NRI, NR2, Bo, Bj, and other B and B’ subunits were PCR amplified from the cDNAs in pCRScript using the primer pairs listed in Supplemental Table S1. The PCR fragments were then cloned into pCRScript (Stratagene) before digestion with the restriction enzymes of pWEN18, pWEN-CY, or pWEN-NY as N-terminal fusions or pWEN25 as a C-terminal fusion. The vectors pWEN18 and pWEN25 have previously been described by Kost et al. (1998), and these vectors were modified by Fujisawa et al. (2004). pWEN18 provided in planta expression of B-EYFP (free N terminus of B), and pWEN25 provided expression of EYFP-B (free C terminus of B). The vectors pWEN-CY or pWEN-NY used for the BiFC have previously been described by Maple et al. (2005).

**Transformation and Fluorescence Microscopy**

Arabidopsis (Col) protoplasts, prepared from suspension culture, were transiently transformed using the polyethylene glycol transformation protocol as described by Droillard et al. (2003). For the BiFC in Arabidopsis mesophyll protoplasts plasmids for interacting proteins as well as negative control plasmids were cotransformed into protoplasts using polyethylene glycol transformation protocols described by Sheen (http://genetics.mgh.harvard.edu/sheenweb/). Microscopy was carried out on a Nikon TE-2000U fluorescence microscope equipped with an Exo X-cite 120 fluorescence illumination system and filters and a special red chlorophyll autofluorescence filter set (exciter HQ630/39, emitter HQ680/40; Chroma Technologies). Images were captured using a Hamamatsu Orca ER 1394 cooled CCD camera and using Openlab and Velocity II (Improvision). Images were subsequently processed for optimal presentation with Adobe Photoshop version 7.0 (Adobe Systems).

**Yeast Two-Hybrid Assay**

The assay was performed according to Maple et al. (2005). Yeast (Saccharomyces cerevisiae) strain AH109 (MATa, trp-1, leu2-112, ura3-52, i, gal4D, gal80D), and plasmids pGADT7 and pGBK7 encoding the Gal4 activation domain and the Gal4 DNA-binding domain, respectively, were derived from MATCHMAKER two-hybrid system version 3 (Clontech Laboratories).

The plasmids pGADT7 and pGBK7 encoding Gal4 AD (activating domain) and Gal4 DNA-BD (binding domain), were used for protein-protein interaction studies in yeast (MATCHMAKER two-hybrid system version 3, Clontech Laboratories). The NR1 construct was cloned into pGBK7 by the primers: forward ATCATAATGCCACCTCCGCTGAAACC, reverse ATCCCGGGTCGCTACAGGATCGGTAGT, and reverse ATCCCGGGTCGCTAACAGGATCGGTAG.

Yeast H7c cells were cotransformed with the resulting plasmids according to the manufacturer’s instructions (Clontech). After 5 d of incubation at 30°C single colonies were inoculated in SD/–TL and grown at 30°C in a shaking incubator for one night. Cultures were grown to OD600 of 1.0 and 5 μL spotted onto SD/–TL plates, or SD/–HTL to monitor HIS3 reporter expression by cell growth. Plates were incubated at 30°C for 48 h, each spot was resuspended in SD/–HTL medium, and the OD600 measured. This OD was taken as a measure of relative cell growth.

**RT-PCR**

Total RNA was isolated using RNeasy plant mini kit (Qiagen), and cDNA synthesized using the high-capacity cDNA archive kit (Applied Biosystems). Real-time PCR reactions were assayed using an ABI 7300 fast real-time PCR system. SYBR green real-time RT-PCR was carried out in triplicate with SYBR green PCR master mix (Applied Biosystems Inc.) using primers, forward TACATATGAACGGTGGTGATGAGGTCGTC, and reverse ATCCCGGGTCGCTACAGGATCGGTAGT, and reverse ATCCCGGGTAAGCAGATACAGAACGC. Yeast H7c cells were cotransformed with the resulting plasmids according to the manufacturer’s instructions (Clontech). After 5 d of incubation at 30°C single colonies were inoculated in SD/–TL and grown at 30°C in a shaking incubator for one night. Cultures were grown to OD600 of 1.0 and 5 μL spotted onto SD/–TL plates, or SD/–HTL to monitor HIS3 reporter expression by cell growth. Plates were incubated at 30°C for 48 h, each spot was resuspended in SD/–HTL medium, and the OD600 measured. This OD was taken as a measure of relative cell growth.

**Supplemental Data**

The following materials are available in the online version of this article.

**Supplemental Figure S1.** Selecting double mutant A1g51690 × A1g17720.

**Supplemental Figure S2.** NR activity state after a darkness to light transition of plants with mutations in various PP2A-like protein phosphatases.

**Supplemental Figure S3.** Assays of NR in the presence of EDTA or Mg2+ used to calculate NR activity state.

**Supplemental Figure S4.** NR domains and B55 tested for interaction by bimolecular fluorescence complementation.

**Supplemental Figure S5.** NR activation rate in the single mutants b55a and b55b.

**Supplemental Figure S6.** Assays of NR in the presence of EDTA or Mg2+ used to calculate NR activity state for wild type and b55.

**Supplemental Table S1.** Primers used for cloning and RT-PCR.

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