Identification of candidates for interacting partners of the tail domain of DcNMCP1, a major component of the *Daucus carota* nuclear lamina-like structure

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**ABSTRACT**

NMCP/CRWN (NUCLEAR MATRIX CONSTITUENT PROTEIN/CROWDED NUCLEI) is a major component of a protein fibrous meshwork (lamina-like structure) on the plant inner nuclear membrane. NMCP/CRWN contributes to regulating nuclear shape and nuclear functions. An NMCP/CRWN protein in *Daucus carota* (DcNMCP1) is localized to the nuclear periphery in interphase cells, and surrounds chromosomes in cells in metaphase and anaphase. The N-terminal region and the C-terminal region of DcNMCP1 are both necessary for localizing DcNMCP1 to the nuclear periphery. Here candidate interacting partners of the amino acid position 975–1053 of DcNMCP1 (T975–1053), which is present in the C-terminal region and contains a conserved sequence that plays a role in localizing DcNMCP1 to the nuclear periphery, are screened for. *Arabidopsis thaliana* nuclear proteins were subjected to far-Western blotting with GST-fused T975–1053 as a probe, and signals were detected at the positions corresponding to ~70, ~40, and ~18 kDa. These ~70, ~40, and ~18 kDa nuclear proteins were identified by mass spectrometry, and subjected to a yeast 2-hybrid (Y2H) analysis with T975–1053 as bait. In this analysis, the ~40 kDa protein ARP7, which is a nuclear actin-related protein possibly involved in regulating chromatin structures, was confirmed to interact with T975–1053. Independently of the far-Western blotting, a Y2H screen was performed using T975–1053 as bait. Targeted Y2H assays confirmed that 3 proteins identified in the screen, MYB3, SINAT1, and BIM1, interact with T975–1053. These proteins might have roles in NMCP/CRWN protein-mediated biologic processes.

**INTRODUCTION**

Animal inner nuclear membrane (INM) is lined with a protein fibrous meshwork called the nuclear lamina, and plant INM is also lined with a lamina-like structure. The animal lamina consists of lamin proteins, while the plant lamina-like structure consists of NMCP/CRWN (NUCLEAR MATRIX CONSTITUENT PROTEIN/CROWDED NUCLEI) family proteins. The amino acid sequence similarities between lamin proteins and NMCP/CRWN proteins are low, thus they are thought to have different origins. However, they are predicted to have a similar secondary structure, which consists of the N-terminal head domain, the central coiled-coil domain, and the C-terminal tail domain. Lamin proteins are involved in regulating chromosome positioning in the nucleus, gene expression, nuclear mechanical properties, and nuclear shape. Some mutations in lamin proteins cause genetic diseases called laminopathies (for a review see ref. [14]). NMCP/CRWN proteins also regulate nuclear shape, chromatin structure, and plant growth.

The model plant *Arabidopsis thaliana* has 4 CRWN/NMCP proteins (CRWN1–4). CRWN1 physically interacts with 2 SUN (Sad1/UNC-84) proteins, SUN1 and SUN2. SUN proteins are present in the INM, and interact with WIP (WPP domain-interacting protein) outer nuclear membrane (ONM) proteins. WIP proteins interact with WIT (WIPP domain-interacting tail-anchored protein) ONM proteins, and WIT proteins interact with myosin XI-i, which would interact with cytosolic actin.
filaments.\textsuperscript{22} The complexes formed by the SUN, WIP, WIT, and myosin XI-i proteins are thought to act as the plant LINC (linker of nucleoskeleton and cytoskeleton) complexes, which transmit the force from the cytoplasmic actin filaments to the nuclear membrane, and thereby regulate nuclear shape. The NMCP/CRWN proteins may affect the structures and/or the functions of the LINC complexes via the interaction with the SUN proteins, and vice versa, although it has also been proposed that the NMCP/CRWN proteins and the LINC complexes have distinct roles in regulating nuclear shape.\textsuperscript{23} KAKU4, another regulator of nuclear shape, is localized to the nuclear periphery and interacts with CRWN1,\textsuperscript{24} but it is still unclear how the CRWN1-KAKU4 complex functions.

To localize an NMCP1/CRWN homolog in Daucus carota (DcNMCP1) to the nuclear periphery, its N-terminal region (position 1–141) and its C-terminal region (position 908–1053) are both necessary. The C-terminal region of DcNMCP1 contains the conserved sequence motif RYNLRRHK, and mutating it to either RYNLAAAA or RAAARRHK disrupts the nuclear periphery localization of DcNMCP1.\textsuperscript{25} At least 3 other D. carota NMCP/CRWN proteins (DcNMCP2–4) have been identified (Fig. 1). NMCP/CRWN proteins could be classified into 2 major clades.\textsuperscript{17} DcNMCP1, DcNMCP3, and DcNMCP4 belong to a clade that includes A. thaliana CRWN1–3, while DcNMCP2 belongs to the other clade with CRWN4. Among DcNMCP1–4, DcNMCP1 is the best characterized.

Here, candidates for interacting partners of the C-terminal region of DcNMCP1 are screened for using far-Western blotting (FWB) and a yeast 2-hybrid (Y2H) system. The potential roles of the identified candidate interactors in regulating nuclear functions are also discussed.

Results

Far-Western blotting for detecting interacting partners of the partial tail domain of DcNMCP1

The partial tail domain corresponding to the amino acid position 975–1053 of DcNMCP1 (T975–1053) was expressed in Escherichia coli as glutathione S-transferase (GST)-fused protein, purified (Fig. S1), and used as the probe for FWB to detect T975–1053-binding nuclear proteins of A. thaliana suspension-cultured cells. The A. thaliana nuclear proteins were fractionated into hydrochloric acid (HCl)-soluble proteins and HCl-insoluble proteins. The HCl-soluble proteins should include histone,\textsuperscript{26} which is known to directly interact with a lamin protein in Drosophila melanogaster.\textsuperscript{27} The nuclear proteins were separated by SDS-PAGE (sodium dodecyl sulfate-polyacrylamide gel electrophoresis), transferred to a membrane, and reacted with the probe and antibodies. In the FWB, signals were detected at the positions corresponding to \( \sim 40 \), \( \sim 38 \), \( \sim 33 \), \( \sim 19 \), \( \sim 17 \) kDa for the HCl-soluble proteins, while at \( \sim 70 \) and \( \sim 40 \) kDa for the HCl-insoluble proteins (Fig. 2A). Similar banding patterns were observed when nuclear proteins were prepared using phenol extraction-based methods, which could reduce DNA contamination in protein samples (Fig. S2). When GST alone was used instead of the GST-T975–1053 fusion protein for the FWB, no clear signals were observed. When the blots were reacted with untagged T975–1053 before being reacted with GST-T975–1053 in the FWB, no clear signals were observed, either (Fig. 2B). These results suggest that the signals detected in the FWB are specific to the interaction between T975–1053 and other proteins. An in vitro pull-down assay was also performed using the same nuclear proteins and GST-T975–1053, but reproducibility of results was low (data not shown).

The signals of \( \sim 70\)- and \( \sim 40\)-kDa proteins were strong in the FWB even when a low concentration (40 ng/ml) of GST-T975–1053 was used (Fig. 2A, right panel), and the \( \sim 19\)- and \( \sim 17\)-kDa nuclear proteins

![Figure 1. Schematic representation of DcNMCP1 (accession number: BAA20407.1), DcNMCP2 (BAI67718.1), DcNMCP3 (BAN14787.1), and DcNMCP4 (KX828842). Their predicted coiled-coil regions are shown as boxes.](image-url)
should include histones. Such proteins detected in the FWB (Fig. 2A, arrows, and Fig. S2, arrowheads) were identified by the liquid chromatography-mass spectrometry (LC-MS). Some of the identified proteins were chosen for a Y2H analysis on the basis of their functional annotations and their coverage scores, which indicate what percentages of the full-length protein sequences were covered by the peptide sequences identified by the LC-MS (Table S1). In the targeted Y2H analysis, in summary, most of the tested proteins failed to enable yeast cells to grow on the selection medium when co-expressed with T975–1053, but the actin-related protein ARP7 (Arabidopsis Genome Initiative (AGI) code: AT3G60830) did (Fig. 2 and Table 1, upper 18 rows), supporting the idea that T975–1053 interacts with ARP7 in yeast.

### Y2H screen for interacting partners of T975–1053

Independently of the FWB, a Y2H screen was performed using T975–1053 as bait and *A. thaliana* cDNA library. Approximately 200 thousand cDNA clones were screened, and 122 colonies survived on selection media. The majority of the plasmid inserts of these colonies were the genes encoding the MYB3 (AGI code: AT1G22640) and BIM1 (AT5G08130) transcription factors (TFs) and the putative E3 ubiquitin ligase SINAT1 (AT2G41980). The full-length coding sequences (CDSs) of MYB3, BIM1, and SINAT1 were recloned into the Y2H vector for a targeted Y2H analysis. In this analysis, MYB3, BIM1, and SINAT1 all enabled yeast cells when co-expressed with T975–1053 (Fig. 3 and Table 1, lower 6 rows), confirming that those proteins interact with T975–1053 in yeast.

### Table 1. The numbers of the colonies that could survive on DDO/X/A in the targeted Y2H analyses.

| AD-fused protein | BD-T975–1053 | Number of colonies assessed | Number of colonies surviving on DDO/X/A | \(^{3}\)P value in the chi-square test |
|------------------|-------------|----------------------------|-----------------------------------------|--------------------------------------|
| ARP7             | +           | 34                         | 32                                      | \(< 10^{-10}\)                        |
| MED37F           | +           | 34                         | 2                                       | 0.888                                |
| PDI              | +           | 30                         | 6                                       | 0.958                                |
| HDT2             | +           | 30                         | 4                                       | 0.816                                |
| RPS14A           | +           | 38                         | 5                                       | 0.78                                 |
| HisH3.3          | +           | 33                         | 2                                       | 0.949                                |
| HisH2AXa         | +           | 33                         | 3                                       | 0.187                                |
| HisH2B.11        | +           | 31                         | 5                                       | 0.42                                 |
| HAP1             | +           | 33                         | 2                                       | 0.348                                |
| MYB3             | +           | 35                         | 31                                      | 0.00178                              |
| BIM1             | +           | 38                         | 20                                      | \(< 10^{-4}\)                        |
| SINAT1           | +           | 34                         | 33                                      | \(< 10^{-10}\)                       |

\(^{1}\)AD: GAL4 activation domain.

\(^{2}\)BD: GAL4 DNA-binding domain. The presence of BD-fused T-975–1053 and its absence are indicated as + and −, respectively.

\(^{3}\)The chi-square test was performed for each AD-fused protein to examine whether the presence of BD-T-975–1053 is independent of the percentages of the colonies that could survive on DDO/X/A.
Subcellular localization of the candidates for T975–1053-interacting partners

Full-length DcNMCP1 cannot be expressed as a GFP-fused protein in a transient expression system with Apium graveolens, but a DcNMCP1 variant that lacks the central coiled-coil domain can be, allowing to visualize positions where its tail domain is present.25 Such a GFP-fused DcNMCP1 variant (GFP-DcNMCP1HT) was co-expressed with m Cherr y-fused ARP7, MYB3, BIM1, or SINAT1 (ARP7-mCherry, MYB3-mCherry, BIM1-mCherry, or SINAT1-mCherry, respectively) to examine their subcellular localization. In agreement with the previous study,25 in all the cases studied, GFP-DcNMCP1HT was detected as either uniform or dotted signals at the nuclear periphery. Signals of ARP7-mCherry were detected in the nucleus and the cytosol, but in agreement with the previous finding that ARP7 is localized to the nucleus in interphase cells,28 the signal intensity of ARP7-mCherry was higher in the nucleus than in the cytosol (Fig. 4, top panels). Signals of MYB3-mCherry and BIM1-mCherry were also detected mainly in the nucleus (Fig. 4, second and third panels from the top). The signals of ARP7-mCherry, MYB3-mCherry, and BIM1-mCherry were not enriched in the nuclear periphery, but this does not rule out the possibility that these proteins could interact with the tail domain of DcNMCP1. Signals of SINAT1-mCherry was often detected as large dots (Fig. 4, fourth from the top), which may be protein aggregates, but they were detected at peripheral regions of the nucleus in some cells (Fig. 4, bottom panel). This difference in the localization of the SINAT1-mCherry signals might be due to the difference in their expression levels in the cells, although other factors might also be involved.

Discussion

In this study, ARP7, MYB3, BIM1, and SINAT1 were identified as the candidate interactors of T975–1053. The idea that T975–1053 possibly interacts with ARP7 is consistent with a previous finding that the tail domain of human lamin A interacts with purified actin.29 ARP7 can be phylogenetically classified into a plant-specific clade in the actin-related protein family. ARP7 is localized in the cytosol in cells in metaphase and anaphase, and localized to the nucleus in cells in the interphase.28
Figure 5. Subcellular localization of the possible interacting partners of the C-terminal region of DcNMCP1. A GFP-fused DcNMCP1 variant lacking the central coiled-coil domain (GFP-DcNMCP1HT) was co-expressed with mCherry-fused ARP7, MYB3, BIM1, or SINAT1 (ARP7-mCherry, MYB3-mCherry, BIM1-mCherry, or SINAT1-mCherry, respectively) in epidermal cells of *Apium graveolens*. For each combination of constructs, more than 10 transformed cells were observed, and representative results are shown. For SINAT1-mCherry, 2 patterns of signals, which might reflect different expression levels of SINAT1-mCherry in transformed cells, are shown (fourth and fifth panels from the top). Dotted lines indicate cell boundaries. Bars = 25 μm.
suspension-cultured cells, an NMCP/CRWN homolog, AgNMCP1, is associated to chromosomes in the metaphase, anaphase, and telophase. Thus, the nuclear localization of ARP7 is likely to depend on the formation of the nuclear membrane. It would be interesting to visualize the subcellular localization of both an NMCP1/CRWN protein and ARP7 at each phase of mitosis. Knockdown of ARP7 causes growth defects such as dwarfism and reduced fertility, and knockout of ARP7 causes embryonic lethality. Another nuclear actin-related protein, ARP6, is a component of the SWR1 complex, which catalyzes replacement of histone variants and thus regulates gene expression. It would be interesting to examine whether the regulation of chromatin structures by NMCP/CRWN proteins are mediated by nuclear actin-related proteins.

MYB3 is a MYB TF possibly involved in phenylpropanoid metabolism, and BIM1 is a basic helix-loop-helix (bHLH) TF involved in regulating the gene expression mediated by the phytohormone brassinosteroid, and SINAT1 is a putative E3 ubiquitin ligase functions of which have not been characterized. No TF or ubiquitin ligase subunit has been characterized. No TF or ubiquitin ligase subunit has been reported to interact with NMCP/CRWN proteins in plants. However, in animal cells, lamin A/C interacts with several TFs and DNA-binding proteins such as the c-Fos TF, the SREBP1 bHLH-leucine zipper TF, and the MYB telomere repeat-binding factor TRF2. Lamin A affects the ubiquitin-proteasome-dependent degradation of heterochromatin proteins, which promote heterochromatinization. These interactions are thought to facilitate the processes mediated by these proteins. A double knockout of CRWN1 and one of CRWN2–4 causes dwarfism, and further knockouts of them cause more severe dwarfism or lethality in A. thaliana. A double knockout of CRWN1 and CRWN3 stabilizes the ABI5 (ABSCISIC ACID INSENSITIVE 5) TF and increases sensitivity of plants to the stress-related phytohormone abscisic acid at the seed germination stage in A. thaliana. It would be interesting to examine whether NMCP/CRWN proteins regulate the subcellular localization, the protein stability, and/or the physiologic functions of MYB3, BIM1, and SINAT1. It would also be interesting to examine whether MYB3, BIM1, or SINAT1 regulates the physiologic functions of NMCP/CRWN proteins.

Materials and methods

Plant materials

Hypocotyls of D. carota and A. thaliana were incubated on the medium 1 and the medium 2 (Table S2), respectively, in the darkness at 25°C for approximately one month to obtain their calli. To develop suspension-cultured cells, the calli of D. carota and A. thaliana were liquid-cultured in the medium 1 and the medium 2, respectively, at 82 rpm at 25°C in the darkness. Every 7–10 days, 10 ml of the suspension cultures of D. carota and A. thaliana was transferred to 90 ml of the medium 1 and the medium 2, respectively, and cultured at 82 rpm at 25°C in the darkness to maintain these cells. These cells were used to prepare RNA and nuclear proteins.

RNA isolation and cDNA synthesis

Approximately 80 mg of the A. thaliana suspension-cultured cells was frozen in liquid nitrogen, ground with a motor and pestle to a fine powder, and resuspended in 1 ml TRIzol Reagent (Thermo Fisher Scientific, 15596026). The suspension was incubated for 5 min at room temperature, mixed with 200 μl chloroform, incubated for 2 min at room temperature, and centrifuged at 12000 × g for 15 min. The supernatant was removed and the pellet was washed with 750 μl solution containing 1 mM DTT, 1 mM RNase-free water, 4 μl DcNMCP1, and 1 ml 100 mM DTT, 1 μl RNase Inhibitor, and 1 μl SCRIPT RT. The solution was incubated at 42°C for 10 min, 50°C for 40 min, 75°C for 15 min, and used as the RNA sample.

First strand cDNA was synthesized using Transcriptor First Strand cDNA Synthesis Kit (Roche, 04379012001). A 10 μl solution containing 1 μg of the total RNA and 12 μM Random Hexamer was incubated at 65°C for 5 min, immediately put on ice, and mixed with 2 μl RNase-free water, 4 μl SCRIPT RT Buffer, 1 μl dNTP Mix, 1 μl of 100 mM DTT, 1 μl RNase Inhibitor, and 1 μl SCRIPT RT. The solution was incubated at 42°C for 10 min, 50°C for 40 min, 75°C for 15 min, and used as the cDNA sample.

Plasmid preparation

For FWB, the CDS of T975–1053 was amplified by PCR using the construct with the full-length DcNMCP1 in the pKD0330 background as the
template and the primer pair shown in Table S3. The PCR products were digested by SalI, and inserted into the SalI site of the pGEX-6P-1 vector (GE Healthcare, 28954648), generating pGEX-6P-T975–1053.

For Y2H experiments, the same PCR products were digested by SalI and inserted into the SalI site of pGBK-T7 DNA-BD Vector (Clontech, 630443), generating pGBK-T975–1053. To clone the other genes for the Y2H experiments, their full-length CDSs were amplified by PCR using the A. thaliana cDNA as template and the primer pairs shown in Table S3. The PCR products were digested and inserted into pGADT7 AD Vector (Clontech, 630442) using the restriction enzyme sites shown in Table S3. The pGAD vectors containing the full-length CDSs of MYB3, BIM1, and SINAT1 were extracted from the corresponding colonies identified in the Y2H screen (see the subsection “Y2H”), transformed into the E. coli strain DH5α, and purified from the transformed E. coli cells for further analyses.

To express GFP-DcNMCP1HT, a construct to express GFP-fused full-length DcNMCP1 was digested with BglII. The resulting larger DNA fragment was self-ligated, generating p35S-GFP-DcNMCP1HT. To express mCherry-fused proteins, the full-length CDSs of ARP7, MYB3, BIM1, and SINAT1 were amplified by PCR using the above primer pairs shown in Table S3. The PCR products were digested and inserted into pBS-35SMCS-mCherry using the restriction enzyme sites shown in Table S3.

**Nuclear protein preparation**

To prepare nuclear proteins of the D. carota and A. thaliana suspension-cultured cells, the cells were collected from 25 ml culture by leaving them without rotation, washed with the Man-MES-1/2N4 solution (0.35 M mannitol, 10 mM NaCl, 15 mM KCl, 1.5 mM MgCl2, 1 mM CaCl2, 20 mM 2-(N-morpholino)ethanesulfonic acid (MES)-KOH, pH 5.8), resuspended in 10 ml of the ice-cold homogenization buffer (20% (v/v) glycerol, 0.3 M sucrose, 1% (v/v) Triton X-100, 10 mM NaCl, 15 mM KCl, 1.5 mM MgCl2, 1 mM CaCl2, 2 mM 2-mercaptoethanol, 1× Protease Inhibitor Cocktail for General Use (Nacalai Tesque, 04080–24), 20 mM MES-KOH, pH 5.8), homogenized with a Potter glass homogenizer on ice, filtered through stainless steel mesh with 37 μm pores, and then through stainless steel mesh with 22 μm pores. The filtered suspensions were centrifuged at 2500 × g at 4°C for 15 min. The pellets were resuspended in 5 ml of the ice-cold homogenization buffer, and the suspensions were centrifuged at 2500 × g at 4°C for 15 min. The pellets were resuspended with Potter Teflon homogenizer in the ice-cold Man-MES-1/2N4 solution containing 25% (v/v) glycerol, and stored as the isolated nucleus sample at -40°C until used.

To extract proteins from the nuclei using HCl, the suspensions containing the isolated nuclei were mixed with an equal volume of ice-cold double-deionized water, and centrifuged at 2800 × g at 4°C for 10 min. The pellets were resuspended in 1.5 ml ice-cold double-deionized water. The suspensions were mixed with 1.5 ml of 0.8 M HCl, incubated at 4°C for 1 h with gentle shaking, and centrifuged at 9500 × g at 4°C for 15 min. The supernatants were transferred to a new tube. The pellets were resuspended in 3 ml of ice-cold 0.4 M HCl, and centrifuged 9500 × g at 4°C for 15 min. The supernatants were combined with those collected above. The resulting solutions were mixed with 36 ml ice-cold acetone, incubated at 4°C overnight, and centrifuged at 3600 × g at 4°C for 15 min. The pellets were dissolved in ice-cold 0.4 M HCl, and the solutions were dialyzed with Dialysis Membrane, Size 20, and double-deionized water. The resulting solutions were stored as the nuclear protein samples at -40°C until used. The HCl-insoluble pellets obtained above were dissolved in the denaturation solution (7 M urea, 3 mM EDTA, 6 mM 2-mercaptoethanol, 10 mM Tris-acetate, pH 7.6), and used for FWB.

Nuclear proteins were also prepared using SDS and phenol as described previously. Briefly, the solutions containing the isolated nuclei were centrifuged at 2000 × g at 4°C for 10 min. The pellets were washed twice with
pre-chilled (-20 °C) 70% (v/v) ethanol, resuspended in the resuspension solution (2% (w/v) SDS, 0.1% (v/v) 2-mercaptoethanol, 20 mM EDTA, 20 mM Tris-HCl, pH 7.6), incubated at 100°C for 3 min, and incubated on ice for 1 min. The suspensions were mixed with an equal volume of phenol. The phenol phase was collected and dialyzed with Dialysis Membrane, Size 20, and the dialysis solution (0.1% (w/v) SDS, 0.1% (v/v) 2-mercaptoethanol, 10 mM Tris-HCl, pH 7.6) to obtain the nuclear protein samples. Nuclear protein samples were also prepared using a phenol- and guanidine isothiocyanate-based reagent, TriPure Isolation Reagent (Roche, 11667157001), according to the manufacturer’s instructions. Protein concentrations in the protein samples were determined using the Pierce Coomassie Plus (Bradford) Protein Assay kit (Thermo Fisher Scientific, 23236).

**FWB and LC-MS**

BL21-Gold (DE3) Competent Cells (Agilent Technologies, 230132) were transformed with either pGEX-6P-1 or pGEX-6P-T975–1053, and liquid-cultured in 3 ml LBA (Luria-Bertani medium supplemented with 50 mg/l ampicillin) at 37°C for 4 h. The cultured cells were transferred to 90 ml LBA, further cultured at 37°C overnight, transferred to 360 ml LBA, and further cultured at 37°C for 1 h. Isopropyl β-D-1-thiogalactopyranoside was added to the culture at the 0.5 mM final concentration, and the cells were further cultured at 30°C for 5 h. The cells were collected by centrifugation at 9500 × g for 10 min, washed with PBS (phosphate-buffered saline: 137 mM NaCl, 2.7mM KCl, 10 mM NaH2PO4, 1.76 mM KH2PO4, pH 7.4), resuspended in 10 ml cell lysis buffer (150 mM NaCl, 10 mM Tris-HCl, 0.2 mg/ml lysozyme, 1× Protease Inhibitor Cocktail for General Use, pH 8.0), incubated at room temperature for 10 min, mixed with 1 μl (250 units) of TurboNuclease (Accelagen, N0103P), and further incubated at room temperature for 20 min. The suspension was centrifuged at 33000 × g at 4°C for 10 min. The supernatant was transferred to a new tube, and ammonium sulfate was dissolved in it at the 0.56 g/ml final concentration. The solution was then incubated at 4°C overnight, and centrifuged at 33000 × g at 4°C for 10 min. The pellet was dissolved in PBS and used as the crude protein sample containing GST or GST-T975–1053.

GST or GST-T975–1053 in the crude sample was purified using Glutathione Sepharose 4B (GE Healthcare, 17075601) according to the manufacturer’s instructions. Briefly, 10 ml of the crude protein solution was incubated with 1 ml of the Glutathione Sepharose 4B slurry equilibrated with PBS at room temperature for 1 h. The Glutathione Sepharose 4B resin was then washed with the washing solution (50 mM NaCl, 2 mM EDTA, 20 mM PIPES-KOH, pH 6.8), and incubated in 9 ml elution solution (20 mM reduced glutathione, 5 mM EDTA, 100 mM Tris-HCl, pH 8.0) to elute GST or GST-T975–1053. The eluent was dialyzed against PBS, and the resulting solution containing purified GST or GST-T975–1053 was stored as the probe solution at 4°C until used. The concentration of GST-T975–1053 in the solution was estimated to be 1.6 mg/ml on the basis of the Coomassie Brilliant Blue (CBB) staining, where the signal intensity for GST-T975–1053 was compared with that of known amount of protein in a protein size marker (see the subsection “FWB”). To prepare untagged T975–1053, 1 ml of the solution containing purified GST-T975–1053 was mixed with 4 μl PreScission Protease (GE Healthcare, 27084301), and incubated at 25°C for 1 h. PreScission Protease and excised GST was removed from the solution using Glutathione Sepharose 4B according to the manufacturer’s instructions. The resulting solution was dialyzed against PBS, and centrifuged at 16000 × g at 4°C for 10 min. The supernatant was used as the solution containing untagged T975–1053.

The nuclear protein samples were separated by SDS-PAGE (1.5 mg protein/lane) as described previously, transferred to Immun-Blot PVDF Membrane (Bio-Rad, 1620177) according to the manufacturer’s instructions. The membrane was incubated at 30°C for 1 h in PBS containing 1% (w/v) Block Ace (DS Pharma Biomedical, UKB40) and 1% (w/v) bovine serum albumin, washed twice with PBST (PBS supplemented with 0.2% (v/v) Tween 20), incubated at 30°C for 3 h in PBST containing either GST or GST-T975–1053 as probe (probe concentrations are indicated in Fig. 1A), washed 3 times with PBST, incubated at 25°C for 3 h in PBST containing Anti-GST, Monoclonal Antibody, Peroxidase Conjugated (Wako, 011–21891), and washed 3 times with PBST. Signals were detected using the Chemi-Lumi One L chemiluminescent substrate (Nacalai Tesque, 07880), and the Lumivision Pro 400EX imager (AISIN). Images of FWB signals were processed using the Canvas X software (ACD Systems). To estimate the amounts and sizes of
proteins, XL-Western Marker color plus (Apro Science, SP-2170) and SDS-PAGE Molecular Weight Standards, Low Range (Bio-Rad, 161–0304) were subjected to SDS-PAGE as well as the nuclear proteins. Proteins in the gel or on the membrane were stained with Quick-CBB PLUS (Wako, 178–00551).

For LC-MS, 1.5 mg of the A. thaliana nuclear proteins prepared with HCl (see the “Nuclear protein preparation” subsection) and SDS-PAGE Molecular Weight Standards, Low Range, were separated by SDS-PAGE, and stained with Quick-CBB PLUS. The gel blocks containing ~70, ~40, ~19, ~17 kDa protein bands were excised, and the proteins in these blocks were digested using In-Gel Tryptic Digestion Kit (Thermo Fisher Scientific, 89871) to elute the peptides from the gel. The peptides were analyzed with the LTQ Orbitrap XL mass spectrometer (Thermo Fisher Scientific) and the Proteome Discoverer 1.4 software (Thermo Fisher Scientific) provided by Instrumental Analysis Services of Global Facility Center at Hokkaido University.

Y2H experiments

For the Y2H screen, Y2HGold Yeast (Saccharomyces cerevisiae) Strain (Clontech, 630498) was transformed with pGBK-T975–1053 using the lithium acetate method. Yeast mating was performed using the transformed Y2HGold cells and Mate & Plate Library - Universal Arabidopsis (Normalized) (Clontech, 630487), which is a yeast cell library constructed from mRNA isolated from A. thaliana seedlings, leaves, stems, flowers, and siliques, according to the manufacturer’s instructions. After the mating, the resulting cells were plated on the DDO/X/A selection medium (synthetic dextrose medium that lacks leucine and tryptophan) and contains X-a-Gal (Clontech, 630407) and the Aureobasidin A antibiotic (Clontech, 630466)), and grown for 5 d at 30°C. Colonies that could grow on DDO/X/A were transferred to the more stringent medium QDO/X/A (DDO/X/A lacking adenine and histidine), and grown for 5 d at 30°C. Plasmid inserts in the colonies that could grow on QDO/X/A were amplified by yeast colony PCR using KOD FX (TOYOBO, KFX-101) and Matchmaker AD LD-Insert Screening Amplimer Set (Clontech, 630433), and sequenced with 3130 Genetic Analyzer (Applied Biosystems) at DNA Sequencing Facility of Research Faculty of Agriculture at Hokkaido University.

For targeted Y2H analyses, pGBK-T975–1053 and one of the pGADT7 constructs with genes of interest (see the subsection “Plasmid preparation”) were co-transformed into the Y2HGold strain with the lithium acetate method. Transformed cells were selected on the DDO medium (synthetic dextrose medium that lacks leucine and tryptophan), transferred to DDO/X/A, and grown at 30°C for 5 d to examine the reporter gene activation mediated by the interaction between T975–1053 and the proteins of interest. To obtain percentages of the colonies that survived on DDO/X/A, more than 30 individual colonies that grew on DDO were grown on DDO/X/A at 30°C for 5 d for each combination of constructs. Tiny or almost invisible colonies were regarded as non-surviving colonies, and the numbers of surviving colonies and non-surviving colonies were used for the chi-square test.

Expression of GFP- and mCherry-fused proteins

One of the mCherry constructs and p35S-GFP-DcNMCP1HT (see the “Plasmid preparation” subsection) (0.5 µg each) were mixed, and co-introduced into celery (Apium graveolens) epidermal cells using the Biolistic PDS-1000/He particle delivery system (Bio-Rad) as described previously. Cells were incubated for 24 h at room temperature after being transformed, and signals were observed using the BX50 epifluorescence microscope (Olympus) equipped with the ORCA-ER-1394 digital camera (Hamamatsu Photonics, Hamamatsu, Japan). The fluorescence mirror units U-MGPHQ and U-MWIG2 (Olympus) were used to image GFP and mCherry, respectively. Images were processed with GIMP and Inkscape, respectively.

Accession numbers

Details regarding the sequences of the genes used in this study can be obtained with the following accession numbers (GenBank accession number for DcNMCP1 and AGI codes for A. thaliana genes): D64087 (DcNMCP1), AT5G42020 (MED37F), AT5G60640 (PDI), AT5G22650 (HD72), AT3G60830 (ARP7), AT2G36160 (PSI14A), AT1G08880 (HisH2-Axa), AT4G40030 (HisH3.3), AT5G59910 (HisH2B.11), AT1G02140 (HAP1), AT1G22640 (MYB3), AT5G08130 (BIM1), and AT2G41980 (SINAT1).
Abbreviations

2,4-D  2, 4-dichlorophenoxyacetic acid  
AGI  Arabidopsis Genome Initiative  
bHLH  basic helix-loop-helix  
CBB  Coomassie Brillant Blue  
CDS  coding sequence  
FWB  far-Western blotting  
GST  glutathione S-transferase  
INM  inner nuclear membrane  
LC-MS  liquid chromatography-mass spectrometry  
LINC  linker of nucleoskeleton and cytoskeleton  
MES  2-(N-morpholino)ethanesulfonic acid  
ONM  outer nuclear membrane  
SDS-PAGE  sodium dodecyl sulfate-PAGE  
TF  transcription factor  
Y2H  yeast 2-hybrid  

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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