Simultaneous introduction of multiple biomacromolecules into plant cells using a cell-penetrating peptide nanocarrier†

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Introduction

Plant cells contain groups of biomolecules that participate together in a particular biological process. Exogenous codelivery of multiple biomolecules is an essential step for elucidation of the biological significance of these molecules and enables various biotechnological applications in plants. However, the currently existing biomolecule delivery methods face difficulties in delivering multiple components into plant cells, mediating transgene expression, and maintaining the stability of the numerous components and lead to delays in biomolecular function. Cell-penetrating peptides (CPPs) have demonstrated remarkable abilities to introduce diverse biomolecules into various plant species. Here, we employed the engineered CPP KH9-BP100 as a carrier to deliver multiple biomolecules into plant cells and performed a bimolecular fluorescence complementation assay to assess the simultaneous introduction of multiple biomolecules. We demonstrate that multiple biomolecule/CPP cargos can be simultaneously internalized by a particular plant cell, albeit with different efficiencies. We present a cutting-edge technique for codelivery of multiple biomolecules into plant cells that can be used for elucidation of functional correlations and for metabolic engineering.

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functions of interacting proteins, advanced delivery systems for simultaneous introduction of multiple biomolecules into plant cells must be developed.

Cell-penetrating peptides (CPPs) are short chains of amino acid residues that have the ability to traverse cellular lipid membrane bilayers. CPPs are typically positively charged or amphipathic polypeptides that are derived from viral proteins as well as peptide toxins. These short functional peptides can efficiently deliver biocromolecules such as double-stranded DNA (dsDNA) and double-stranded RNA (dsRNA) and high-molecular weight proteins into animal cells as well as several types of plant cells. The constituent amino acid residues of CPPs interact with various biomolecules to form biomolecule/CPP complexes. Combination of CPPs with cationic domains, e.g., lysine-rich, arginine-rich, and poly(lysine/histidine) domains, has been reported to improve the efficiency of biomolecule transport into different plant species. Among the CPPs commonly used for biomolecule delivery into plant cells, BP100, a CPP originally used as a plant antimicrobial peptide, and its cationic domain-fused derivatives have exhibited an exceptional ability to deliver biomolecules into intact plants. The efficiency of functionalized CPPs in transporting multiple components into plant cells remains to be explored.

To contribute to advances in the biomolecule delivery techniques for plants, we utilized N-terminal poly(lysine/histidine) domain-fused BP100 (KH9-BP100) as a CPP carrier to transport multiple biomolecules into plant cells. We used a superfolder GFP (sfGFP)-based BiFC assay with a cytosolic homodimer of the *Coffee arabica* 7-methylxanthine methyltransferase 1 (MxMT) protein as a model to examine the effects of simultaneous introduction of two molecules with functional complementarity into plant cells. The biomolecules paired for protein–protein interactions, namely, plasmid DNA (pDNA) molecules harbouring transgene expression cassettes, single-stranded RNA (ssRNA) molecules, and fusion proteins, formed biomolecule/CPP complexes with KH9-BP100. These biomolecule/CPP complexes were cointroduced into intact plant leaves, and the formation of MxMT homodimers was notably observed in the cytosol by sfGFP-based BiFC. The efficiency of KH9-BP100 in delivering multiple biocomponents into plant cells was evaluated and compared among pDNA, ssRNA and proteins. Our CPP-mediated multiple-biomolecule delivery system provides a versatile platform with which to elucidate the functions of numerous components that control particular plant metabolic processes and cellular mechanisms.

**Results**

**Multiple pDNA transfer into plant cells via KH9-BP100 peptide carriers**

Nucleic acids such as DNA and RNA form nucleic acid/CPP complexes with cationic CPPs by electrostatic interaction.

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Fig. 1 Direct introduction of multiple biomolecules into plant cells using cell-penetrating peptide KH9-BP100 nanocarrier. (a) Two expression vectors harboring the expression cassettes of two independent non-fluorescent sfGFP fragments-fused *Coffee arabica* 7-methylxanthine methyltransferase (MxMT) proteins (psfGN155-MxMT and psfGC155-MxMT), (b) two single-stranded RNA (ssRNA) molecules of sfGN155-MxMT and sfGC155-MxMT transcripts, and (c) two cell-free synthesized non-fluorescent sfGFP fragment-fused MxMT proteins (sfGN155-MxMT and sfGC155-MxMT) were formed biomolecule/CPP complexes with KH9-BP100. A mixture containing paired biomolecule/CPP complexes was infiltrated into plant leaves using needle-less syringe. Successful introduction of multiple biomolecules into a single plant cell mediated by KH9-BP100 was elucidated from the formation of sfGFP-visualized MxMT homodimer using fluorescence microscopy.
To produce cargo molecules, we first introduced two different pDNA molecules (psfGN155-MxMT and psfGC155-MxMT) encoding the nonfluorescent N-terminal and C-terminal sfGFP fragment-fused MxMT proteins26,27 (sfGN155-MxMT and sfGC155-MxMT, respectively) into plant cells using KH9-BP100 (Fig. S1a†). Different amounts of KH9-BP100 were added into solutions containing a consistent amount of pDNA at various N/P ratios (the molar ratios of the positively-charged NH₃⁺ groups of basic side chains of lysine and histidine in the KH9-BP100 to the PO₄⁻ groups of the pDNA).28 Sub-micron-sized, negative-to-positive surface charge-transitioned pDNA/CPP complexes were formed with increasing N/P ratios (Fig. 2a, b, Fig. S1b, c and Tables S1, S2†). Positively surface-charged pDNA/CPP complexes with particle sizes of approximately 100–150 nm efficiently transfected various plant cell types.15,19 We rationally selected globular-shaped, ∼120 nm pDNA/CPP complexes that formed at an N/P ratio = 2.5 (Fig. 2a, b and Fig. S1d, e) to monitor the successful delivery of the two different pDNA molecules into plant cells. We then infiltrated a mixture of the psfGN155-MxMT/KH9-BP100 and psfGC155-MxMT/KH9-BP100 complexes into tobacco (Nicotiana benthamiana) leaves. Via confocal laser scanning microscopy (CLSM) imaging analysis, the formation of sfGN155-MxMT/sfGC155-MxMT homodimers was gradually detected as complemented sfGFP fluorescence in the cytosol of leaf cells at different times post cotransfection (Fig. 2c, d and Fig. S2†). The sfGFP intensities in both epidermal and spongy mesophyll cells peaked at 24 hours before declining significantly by 48 hours post transfection (Fig. 2c, d and Fig. S2, S3a†). sfGN155-MxMT and sfGC155-MxMT mRNA accumulated to high levels in the cotransfected leaves by 24 hours post transfection (Fig. 2e and f). Immunoblotting confirmed the high levels of accumulation of both sfGN155-MxMT and sfGC155-MxMT proteins in the cotransfected leaves at 24–48 hours post transfection with pDNA/CPP complexes (Fig. 2g and Fig. S3b†). These results indicate that two pDNA molecules harbouring different gene expression cassettes delivered by a CPP can be coordinately transported into a plant cell and function as templates for the expression of two corresponding proteins that participate in protein–protein interactions.

Fig. 2 Simultaneous transfection of two plasmid DNA molecules into plant cells mediated by cationic CPP KH9-BP100. (a) and (b) Physicochemical properties of plasmid DNA (pDNA)/KH9-BP100 complexes formed at different N/P ratios. pDNA indicates a solution containing only pDNA molecules. Error bars represent standard deviation of the average value from three independent measurements. (c) Cytoplasmic localization of sfGFP-visualized MxMT homodimers in plant cells co-transfected by psfGN155-MxMT- and psfGC155-MxMT/KH9-BP100 complexes at different times after transfection. Scale bars = 50 µm. (d) Fluorescent intensity in plant cells transfected by psfGN155-MxMT- and psfGC155-MxMT/KH9-BP100 complexes at different time points. The distributions of fluorescent intensity were presented as the box plot. Black bars indicate means of distributed values. Different letters show significant differences of mean fluorescent intensity in the different times analyzed by one-way ANOVA with Tukey’s HSD test at p = 0.05 (n = 9 for 0 h and 12 h, n = 12 for 24 h and 48 h). (e) and (f) Transcript abundances of sfGN155-MxMT and sfGC155-MxMT in plant leaves transfected with pDNA/KH9-BP100 complexes. Transcript levels were analyzed by qRT-PCR. Error bars represent the standard deviation of the average transcript number per 100 ng of total RNA in 4 biological independent samples (n = 4). Different letters indicate the significant differences of mean fluorescent intensity analyzed by one-way ANOVA with Tukey’s HSD test at p = 0.05. Same letter represents no significant difference among compared samples. (g) Accumulation of non-fluorescent sfGFP fragment-fused MxMT proteins in plant leaves transfected with two different pDNA/KH9-BP100 complexes at various incubation times. The accumulation of recombinant proteins in total plant leaf proteins was detected by western blotting (top panel). WT and P are total proteins from wild-type and pDNA only-infiltrated leaf samples. N = sfGN155-MxMT protein. C = sfGC155-MxMT protein. R = plant cell abundant protein Rubisco large subunit stained with Ponceau S diazo dye (lower panel).
Delivery of multiple ssRNA molecules into plant cells

Although RNA molecules are less stable biochemically than DNA, double-stranded interfering RNA molecules delivered to plant cells by CPPs can efficiently suppress the expression of target genes in plants.\(^{18}\) We therefore examined whether two different RNA molecules could be coordinately transferred into plant cells with a CPP delivery platform (Fig. S4a†). ssRNA molecules of both sfGN155-MxMT and sfGC155-MxMT were synthesized by in vitro transcription (Fig. S4b and S5a, b†). Adding different amounts of KH\(_{9}\)-BP100 to solutions containing a constant amount of individual ssRNA molecules changed the physicochemical properties of the ssRNA/CPP complexes at different N/P ratios (Fig. S5c–f and Tables S3, S4†). Taking into consideration the hydrodynamic diameters (Fig. S5c†), surface charges (Fig. S5d†), RNA release characteristics (Fig. S5e and ff) and sphere-shaped appearances of the ssRNA/CPP complexes, a positively charged ssRNA/CPP complex that formed at an N/P ratio of 2.5 (Fig. 3a and b) was chosen for cotransfection into plant cells. Because incorporating dsRNA molecules with peptides can enhance the ability of dsRNA to resist degradation by extra- and intracellular RNases,\(^{18}\) we subsequently monitored the degradation of ssRNA molecules in plant tissues by quantitative reverse transcriptase-PCR (qRT-PCR). Notably, the ssRNA abundances of sfGN155-MxMT and sfGC155-MxMT in the leaves cotransfected with the two ssRNA/CPP complexes were significantly higher than the ssRNA levels in free ssRNA-infiltrated samples (Fig. 3c). These ssRNA molecules accumulated at high levels for up to 4 hours before gradually declining in abundance to ~30% at 24 hours post transfection (Fig. 3c). Based on the qRT-PCR analysis, the half-lives of the ssRNA molecules in plant leaves cotransfected with ssRNA/CPP complexes were 2.8 ± 1.6 hours for sfGN155-MxMT and 3.2 ± 2.7 hours for sfGC155-MxMT. These results imply that KH\(_{9}\)-BP100 protects ssRNA molecules from the RNA degradation process in plant cells.

To test the function of KH\(_{9}\)-BP100 in transporting multiple ssRNA molecules into plant cells, we coinfiltrated a mixture of sfGN155-MxMT/KH\(_{9}\)-BP100 and sfGC155-MxMT/KH\(_{9}\)-BP100 complexes into N. benthamiana leaves. Successful internalization of these two ssRNA/CPP complexes by the same cells was confirmed via observation of complemented sfGFP fluorescence of an sfGN155-MxMT/sfGC155-MxMT homodimer in the plant cells. In CLSM analysis, significant increases in fluorescence intensity and a uniform distribution of the complemented sfGFP-visualized MxMT homodimers in the cytosol of plant cells were detected in the leaves (Fig. 3e). The localization of sfGFP-visualized MxMT homodimers in the plant cells after transformation with sfGN155/sfGC155 complex was monitored by polyclonal anti-GFP antibody (Fig. 3f). Equivalent protein application (indicated by R; rubisco large subunit) on the membrane was confirmed by Ponceau S staining (lower panel). N = sfGN155-MxMT protein; C = sfGC155-MxMT protein. WT = wild type leaves, RNA = ssRNA only-treated leaves.
were observed at 12 and 24 hours after infiltration (Fig. 3d, e and Fig. S6†). Immunoblotting revealed that the accumulation of the sfGN155-MxMT and sfGC155-MxMT proteins in transfected plant cells gradually increased over the incubation periods and reached maximum levels at 24 hours post transfection (Fig. 3f). The CLSM imaging and immunoblotting results suggest that cationic KH₇₆BP100 can effectively deliver multiple ssRNA molecules for synchronous expression of multiple functional proteins in the same plant cell.

**Translocation of multiple proteins into the cytoplasm by KH₇₆BP100**

Some functional proteins can be translocated across plant cell boundaries by their CPPs. Negatively charged proteins such as BSA can form complexes with C-terminal-fused cationic (BP100)ₓ-KH₇₆ peptides, and BSA/CPP complexes can be delivered into plant cell cytoplasm. We analyzed the ability of KH₇₆BP100 to form protein/CPP complexes with two nonfluorescent N-terminal and C-terminal sfGFP fragment-fused MxMT proteins (sfGN155-MxMT and sfGC155-MxMT, respectively) for translocation into plant cells (Fig. S7a†). Recombinant sfGN155-MxMT and sfGC155-MxMT proteins with predicted molecular weights of 60.5 kDa and 52.4 kDa, respectively, were synthesized by cell-free protein synthesis (Fig. S7b†). Both proteins were calculated to show net charges of −15.2 mV in solution at pH 7.0. Equal amounts of both proteins were added to solutions containing various amounts of KH₇₆BP100 to form protein/CPP complexes at different molar ratios.

Increasing the amounts of KH₇₆BP100 in the solutions increased the hydrodynamic diameters of both the sfGN155-MxMT/KH₇₆BP100 and sfGC155-MxMT/KH₇₆BP100 complexes (molar ratios of 0.1 and 0.5; Fig. 4a and Tables S5, S6†). Unexpectedly, both protein/CPP complexes that formed at molar ratios of 1.0 and 2.5 exhibited hydrodynamic diameters larger than 1 µm (Fig. 4a and Tables S5, S6†). The surface charges of the protein/KH₇₆BP100 complexes formed at various molar ratios increased progressively with increasing amounts of KH₇₆BP100 (Fig. 4b and Tables S5, S6†). Native-PAGE analysis revealed that protein mobility gradually decreased with increasing molar ratios (Fig. S8†). Based on the hydrodynamic diameters, surface charges and release of proteins from the protein/CPP complexes in the electrophoretic field, the sfGN155-MxMT/KH₇₆BP100 and sfGC155-MxMT/KH₇₆BP100 complexes that formed at molar ratios of 0.5, 1.0, and 10 were chosen for in vitro and in vivo protein-protein interaction assays by sfGFP-based BiFC with the MxMT homodimer.

We subsequently investigated the in vitro interaction of two nonfluorescent sfGFP fragment-fused MxMT proteins (sfGN155-MxMT and sfGC155-MxMT) after they formed complexes with KH₇₆BP100 at molar ratios of 0.5, 1.0, and 10 by dynamic light scattering (DLS) analysis and fluorospectrometry. The hydrodynamic diameters of protein/KH₇₆BP100 complexes that formed at molar ratios of 0.5, 1.0 and 10 in the in vitro interaction assay buffer showed significant increases at 24 hours post incubation (Fig. 4c). The fluorospectrometric analysis showed that the in vitro BiFC reaction between sfGN155-MxMT and sfGC155-MxMT was reduced by complexation with KH₇₆BP100 at these molar ratios (Fig. 4d). We further investigated whether the incorporation of KH₇₆BP100 into sfGN155-MxMT and sfGC155-MxMT could inhibit the spontaneous dimerization of sfGN155-MxMT and sfGC155-MxMT in plant cells. Mixtures of sfGN155-MxMT/KH₇₆BP100 and sfGC155-MxMT/KH₇₆BP100 complexes at molar ratios of 0.5, 1.0, and 10 were infiltrated into the tobacco leaves. Complemented sfGFP fluorescence in the cytoplasm was observed in all samples transfected with a mixture of the two protein/KH₇₆BP100 complexes at 2 hours post infiltration (Fig. S9†). However, sfGFP signals were not detectable in the apoplasts of plant cells transfected with a solution containing free proteins (Fig. S9†). On the basis of the results regarding the inhibition of the in vitro BiFC reaction by KH₇₆BP100 and the ability to deliver multiple proteins into plant cells, the protein/KH₇₆BP100 complex that formed at a molar ratio of 10 was chosen for atomic force microscopy (AFM) imaging. The morphologies of the sfGN155-MxMT/KH₇₆BP100 and sfGC155-MxMT/KH₇₆BP100 complexes that formed at a molar ratio of 10 were globular in shape with an average size of ∼200 nm (Fig. 4e and f).

The efficiency of KH₇₆BP100 in transporting multiple proteins into plant cells was evaluated by time-course CLSM imaging. Cellular distribution of complemented sfGFP-visualized MxMT homodimers in the cytoplasm could be observed at 30 minutes post infiltration (Fig. 4g, h and Fig. S10†). The complemented sfGFP signals reached maximum intensity at 2 hours post infiltration and subsequently declined with time (Fig. 4g, h and Fig. S10†). Immunoblotting of the total proteins recovered from leaves transfected with the two protein/CPP complexes showed that up to 50% of the initial amounts of the sfGN155-MxMT and sfGC155-MxMT proteins remained in the plant leaves at 2 hours post transfection (Fig. 4i and Fig. S11†). However, the amounts of these proteins decreased drastically to only 3% of the initial amounts by 6 hours post infiltration (Fig. 4i and Fig. S11†). These results imply that significant degradation of the sfGN155-MxMT and sfGC155-MxMT proteins occurred in plant tissues after transfection with the protein/KH₇₆BP100 complexes.

**KH₇₆BP100-mediated codelivery of multiple biomolecules into plant cells is an endomembrane-dependent event**

The positively charged helical CPP BP100 induces the formation of endocytic vesicles for transport of biomolecules through plant plasma membranes. We thus investigated the cellular uptake of multiple biomolecule/CPP cargos into plant cells. We observed the plant cell internalization of pDNA/KH₇₆BP100 complexes as representative biomolecules by detecting vesicle formation and colocalization with the plasma membrane-staining fluorescent dye FM4-64. Elevated numbers of large FM4-64-stained vesicles (∼1 µm) were found on the plasma membranes of plant cells at 2 hours post transfection (Fig. S12†). This result shows that the pDNA/KH₇₆BP100 com-
plexes induced the formation of large membrane vesicles in the transfected plant cells.

In our experiments, two different pDNA/CPP complexes were codelivered into plant cells. We performed fluorescence microscopy and colocalization analysis of fluorescent dye-labeled psfGN155-MxMT/KH$_2$-BP100 and psfGC155-MxMT/KH$_2$-BP100 complexes in plant cells at 2 hours post transfection. Fluorescent cyanine-3 (Cy3)-labeled psfGN155-MxMT/KH$_2$-BP100 complexes were found to colocalize with fluorescent cyanine-5 (Cy5)-labeled psfGC155-MxMT/KH$_2$-BP100 complexes inside plant cells (Fig. 5a–c). We also observed colocalization of both Cy3-psfGN155-MxMT/KH$_2$-BP100 and Cy5-psfGC155-MxMT/KH$_2$-BP100 complexes with intracellular vesicles stained by FM4-64 (∼1 µm) on the plasma membranes of plant cells at 2 hours post transfection (Fig. 5d and Fig. S13†). Moreover, we surprisingly detected colocalization of the fluorescent dye-labeled pDNA/KH$_2$-BP100 complexes with the intracellular vesicles and observed trafficking to the nuclear membrane via the cellular network (Fig. 5d, e, Fig. S13 and Movie S1†). These results suggest that multiple nucleic acid cargos can be simultaneously internalized by plant cells via large endocytic vesicles and translocated together to the nucleus via intracellular vesicle trafficking.

To investigate how multiple protein/CPP complexes are transported into plant cells, we coinfiltrated intact leaves with

Fig. 4 Direct co-delivery of two different proteins into plant cytoplasmic region by cationic CPP KH$_2$-BP100. (a) and (b) Hydrodynamic diameter and surface charge of sfGN155-MxMT and sfGC155-MxMT protein/KH$_2$-BP100 cargos formed at various protein/peptide molar ratios. Data were presented as the average value of three independent replicates. Error bars represent standard deviation. (c) DLS measurement of combined protein/CPP complexes formed at molar ratio = 0.5, 1.0, and 10 after incubation in in vitro interaction assay condition. The sfGN155-MxMT and sfGC155-MxMT proteins without KH$_2$-BP100 (free) were added to the in vitro interaction buffer as the control. Data were presented as mean of 4 biological replicates ± SD. Different letters indicate the significant differences of mean analyzed by one-way ANOVA with Tukey’s HSD test at $p = 0.05$. (d) In vitro interaction assays of sfGFP-visualized MxMT homodimers. The spectra show the average emissions of complemented sfGFP-visualized MxMT homodimers in complex with KH$_2$-BP100 at molar ratios = 0.5, 1.0, and 10 at 24 hours post-incubation (n = 4). (e) and (f) Morphologies of sfGN155-MxMT/CPP complexes and sfGC155-MxMT/CPP complexes at molar ratio = 10 under AFM imaging. Scale bars = 500 nm. The heated map represents the different height of particle on the mica surface. (g) Localization of sfGFP-visualized MxMT homodimers in plant cell cytoplasm after transfection with two protein/CPP complexes. Scale bars = 20 µm. (h) Fluorescent intensity in the plant cells transfected with protein/KH$_2$-BP100 cargos. The fluorescent intensities from 15 cells (n = 15) were represented as the box plot. Black bar indicates the mean of fluorescent intensities. Different letters indicate the significant differences of mean analyzed by one-way ANOVA with Tukey’s HSD test at $p = 0.05$. (i) Degradation of non-fluorescent sfGFP fragment-fused MxMT proteins in plant leaves after transfection with two protein/KH$_2$-BP100 complexes. Total protein recovered from plant leaves were analyzed by western blotting. The band intensities of sfGN155-MxMT and sfGC155-MxMT proteins were converted to the percentage of protein recovery. The bar graph represents the mean of protein recovery ± SD of 3 independent samples (n = 3). Asterisks indicate significant difference analyzed by Student’s t-test at $p < 0.05$. 

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a solution containing sfGN155-MxMT/KH9-BP100, sfGC155-MxMT/KH9-BP100, and FM4-64. In a control experiment without KH9-BP100, free proteins could neither be internalized by plant cells nor form complemented sfGFP-visualized MxMT homodimers in the intercellular spaces and apoplasts (Fig. S14f). We further detected the colocalization of complemented sfGN155-MxMT/sfGC155-MxMT homodimers with FM4-64-stained vesicles in the cytoplasm (Fig. 5f and Fig. S14f), and the results indicated that the two protein/CPP complexes colocalized with vesicles when the colocalization coefficient ($R$) was higher than 0.75, associated with vesicles when $0.25 \leq R \leq 0.75$, or were distinctly released when $R$ was lower than 0.25 (Fig. S14f). At 30 minutes post transfection, 63% of the complemented sfGFP-visualized MxMT homodimers were colocalized or associated with the membrane-bound vesicles with a median $R$-value of 0.41 (Fig. 5g and h). Fifty-nine percent of these complemented sfGFP-visualized MxMT homodimers then dissociated from the vesicles and were distinctly located in the cytoplasm at 2 hours post transfection (median $R = 0.23$) (Fig. 5f–h). Furthermore, 31% of the complemented sfGFP-visualized MxMT homodimers colocalized with large endomembrane vesicles ($\sim$5 µm) bound to the plasma membrane at 6 hours post transfection (Fig. 5f–h). These results imply that the two nonfluorescent sfGFP fragment-fused MxMT proteins delivered by KH9-BP100 were internalized by plant cells via membrane vesicle formation and movement. Additionally, the complemented sfGFP-visualized MxMT homodimers were later released into the cytoplasm (Fig. 5f, 2 hours post transfection).

### Discussion

Introduction of multiple biomolecules across plant cell boundaries is a challenging step in investigations regarding the integrative functions of these biomolecules in particular cellular
Nanoscale Paper

processes.8,10 For example, reprogramming of a significant biosynthesis pathway of plant metabolites requires efficient cotransfection of numerous independent vectors into plant cells.31–33 The translocation efficiencies for miscellaneous biomolecules depend mainly on the availability of the biomolecules after their introduction into and trafficking and accretion within target cells.34 Multiple pDNA molecules that each harbor an individual gene expression cassette can be mechanically and biologically transformed into the nuclei of plant cells.30 In addition, electroporation and PEG-mediated protoplast transfection have been used for direct delivery of short RNA molecules.35,36 Recent advances in the delivery of Cas9/gRNA ribonucleoprotein complexes into protoplasts have also been accomplished via PEG-mediated transfection.37 The applicability of and effort associated with these techniques are decidedly dependent on several factors. These include tedious material preparation and handling steps, the formulation of biomolecule/carrier complexes and the availability of expensive instruments. Our data suggest that multiple pDNA molecules, ssRNA molecules and proteins can be concurrently introduced into plant cells of intact plant tissues using CPP carriers, although with different efficiencies. These biomolecule/CPP cargos can penetrate plant cell barriers without external force or cumbersome plant material preparation, thus overcoming the limitations of conventional plant cell transfection techniques that require mechanical or chemical treatment. The CPP-mediated method also requires fewer biomolecules than gold particle-based microprojectile bombardment and polymeric PEG-mediated protoplast transformation methods.30,35–37

Generally, integration of CPPs into biomacromolecules can fortify the molecules against cellular degradation processes or suppress their biological functions.38,39 However, our results suggest that incorporating CPPs into negatively charged pDNA and ssRNA molecules greatly enhances the stability of these polynucleotides without disrupting their biological functions. The functional KH₃-BP100 peptide was rationally designed for CPP-based gene delivery.19 Constituent poly(llysine/histidine) (KH₃) residues of KH₃-BP100 are responsible for the condensation of the pDNA or ssRNA molecule, thus enabling the BP100 region to be freely exposed on the outer surface of the complex.18,19,40,41 This arrangement protects the polynucleotide from plant nucleases while the complex is transported into plant cells and finally to the nucleus. Cationic KH₃-BP100 also interacts with the sfGN155-MxMT and sfGC155-MxMT proteins via electrostatic interactions.17 However, our in vitro interaction study results suggested that integration of CPP into the proteins significantly inhibited physical interaction between sfGN155-MxMT and sfGC155-MxMT. Moreover, KH₃-BP100 did not contribute a protective effect against protein degradation in plant cells. Unexpectedly, this disorderly protein/peptide complexation could lead to contact of the proteins with proteases, priming the proteins for undesirable degradation before and after entry into plant cells.

The ability of CPPs to enter plant cells depends on the biological and physicochemical properties of both the CPPs and their cargos.20,21,12 The CPP BP100 and its cationic derivatives, KH₃-BP100 and BP100-KH₃, can destabilize the plasma membrane as well as induce the formation of membrane vesicles.17–19,23 Our results regarding the internalization of multiple biomolecule/KH₃-BP100 complexes into plant cells are in accordance with these findings. Consistent with a previous report,23 our study revealed that the numbers of large membrane vesicles increased during the transfection of plant cells with biomolecule/CPP complexes. We also found that the biomolecule/CPP complexes relocated rapidly from the plasma membrane to the cytoplasm. Moreover, two different biomolecule/CPP complexes colocalized inside conspicuous large membrane vesicles. These vesicles subsequently trafficked the biomolecules and released them into the nucleus or cytosol. The two expression vectors were stable and coexisted in the plant cells transfected with pDNA molecules (Fig. S15†). However, the transcript levels of sfGN155-MxMT and sfGC155-MxMT and the complemented sfGFP-visualized MxMT homodimers were significantly decreased in pDNA/CPP complex-infiltrated leaves. Moreover, sfGN155-MxMT and sfGC155-MxMT RNA was rapidly degraded within 3 hours after transfection of the plant leaves with the ssRNA/CPP complexes, indicating that these RNA molecules were relatively unstable. In addition, the complemented sfGFP-visualized MxMT homodimers only showed significant interactions with the protein/CPP complexes within 2–6 hours post infiltration. The reductions in sfGFP fluorescence in the plant cells transfected with the biomolecule/CPP complexes could have been a consequence of the consecutive degradation of mRNA and the recombinant MxMT proteins in plant tissues. Since the trafficking of biomolecule/CPP complexes to the cytoplasm and nucleus occurs via intracellular vesicle movement, the biomolecular cargos could also have been translocated to vacuoles for nonspecific biomolecule degradation.43 It is also possible that the RNA molecules and proteins distributed in the cytoplasm were degraded by cytosolic RNases and proteases upon release from the CPP-based carriers.16

The two MxMT proteins form a homodimeric complex that localizes in the cytosolic fraction to catalyze the second methylation steps in caffeine biosynthesis from xanthosine.24,26,44 Notably, mRNA levels and protein accumulation are not definitively related to the transcriptional and translational activity of the corresponding genes.45,46 Small oscillations in protein accumulation can occur due to variations in the stability of mRNA and protein molecules that are tightly regulated posttranscriptionally and posttranslationally.15,16,45–48 Our findings showed that the transcript (or synthetic RNA) and protein abundances of the two nonfluorescent sfGFP fragment-fused MxMT components in plant leaves transfected with pDNA and ssRNA/CPP cargos were comparable. Additionally, the sfGFP-visualized MxMT homodimers were homogenously distributed in the cytoplasm. However, in plant cells transfected with ssRNA/CPP cargos, sfGFP fluorescence could be detected within 30 minutes post transfection, much faster than in the pDNA/CPP complex-transfected cells. Our results also showed that the sfGFP-visu-
alized MxMT homodimers matured earlier in the endomembrane vesicles after entering the cells via multiple protein/CPP cargos than after entering via pDNA and ssRNA cargos. However, the sfGFP fluorescence intensity and protein accumulation in cells transfected using multiple protein/CPP cargos were much lower than those in cells transfected using multiple pDNA and ssRNA cargos. Moreover, the protein levels in protein/CPP complex-transfected plant cells drastically decreased within 6 hours after transfection, suggesting that protein degradation occurred. Sub-micron-sized (200 to 1000 nm) positively charged protein/CPP complexes can enter cells by membrane-associated macropinocytosis or heparan sulfate-dependent endocytosis. There is a strong possibility that two BiFC proteins can dissociate from their respective CPP-based carriers and interact with each other inside the endomembrane vesicles. BiFC complexes with MxMT homodimers may then be intermittently released from the vesicles into the cytoplasm, resulting in an uneven distribution of sfGFP signals within the plant cells. These sfGFP-visualized MxMT homodimers may then be intermittently released from the trans-Golgi networks and early endosomes to be recycled back to the plasma membrane or to the vacuoles for complete degradation, as drastic reductions in recombinant MxMT protein levels have been observed after transfection.

Biogenesis of plant metabolites requires multistep enzymatic reactions. The involved enzymes are strictly controlled not only at the expression level but also at the protein level. Therefore, comprehensive engineering of targeted metabolic processes can be accomplished by introducing numerous gene expression regulators and corresponding enzymes as well as their activation components into plant cells. Multiple genetic modifications at key reaction steps are possible via classic gene transfer techniques. However, alteration of multiple genes in a single transgenic line can occasionally cause transgene silencing and unbalanced protein activity as well as cryptic phenotypic defects. Uncontrolled expression ratios and protein levels in transgenic plants can strongly influence the bifurcated reactions of target metabolic pathways.

Conclusions

Our results show that pairs of molecules that act as interaction partners, i.e., pDNA, ssRNA, and protein pairs, can be codelivered into the same plant cell using cationic CPP-based carriers. The CPP-based multiple-biomolecule delivery system is compatible with different plant species and various biomolecules of interest but does not require specialized equipment or complicated plant material preparation. Furthermore, the CPP can be combined with selected OTPs, i.e., peptides that designate biomolecule translocation to specific organelles, for targeted delivery of multiple components to various plant organelles. Our peptide-mediated multiple-biomolecule delivery platform provides a robust tool for comprehensive engineering of plant metabolic processes and quality trait improvement.

Experimental

Cationic KH_{10\times}BP100 peptide

The cationic CPP KH_{10\times}BP100 (NH_{2-KHKHKHKHKHKHKHKHKHKKLLFKKILKYL-COO^-}) was synthesized as previously described. The peptide was prepared in a 1.0 mg ml^{-1} stock solution with water.

Plant cultivation

Seeds of *N. benthamiana* were germinated on soil (Promix, Rivière-du-Loup, Canada) supplemented with vermiculite (ratio of 2 : 1) at 25 °C under 12/12-hour light/dark periods with 100 μmol photons per m² per s in a plant growth chamber for 7 days. The seedlings were then transferred to individual pots and cultured at 22 °C under 8/16-hour light/dark periods with 100 μmol photons per m² per s in a plant growth chamber. Fully expanded leaves from 4–5-week-old plants were used for transfection.

Preparation of pDNA, ssRNA, and protein molecules

pDNA molecules were purified from liquid cultures of *Escherichia coli* DH5α-harbouring psfGN155-MxMT or psfGC155-MxMT using a Qiagen Plasmid Giga Kit (Qiagen, Hilden, Germany). The concentration of pDNA was adjusted to 1.0 mg ml^{-1} with water and then kept at −20 °C.

ssRNA molecules of sfGN155-MxMT and sfGC155-MxMT were synthesized by in vitro transcription. The coding sequences of the sfGN155-MxMT and sfGC155-MxMT genes were synthesized.
were amplified from psfGN155-MxMT and psfGC155-MxMT plasmids, respectively, with the primers shown in Table S7.† The gene fragments were subcloned into pET-30b(+) vectors at XbaI/BamHI restriction sites. The ssRNA molecules were synthesized from a pET-30b(+) vector harbouring a codon sequence of the sfGN155-MxMT or sfGC155-MxMT gene with a T7 Ribomax Express System (Promega, Madison, WI, USA) and purified with a QIAquick Nucleotide Removal Kit (Qiagen). The purified ssRNA molecules were analyzed by 20% polyacrylamide gel electrophoresis to determine the integrity of the ssRNA (Fig. S4b†). The ssRNA samples were kept in small aliquots at −80 °C (1.0 mg ml⁻¹ stock solution).

The nonfluorescent sfGFP fragment-fused MxMT proteins sfGN155-MxMT and sfGC155-MxMT were synthesized with a dialysis-mode cell-free protein synthesis system.17 Briefly, pDNA for in vitro transcription was added to a solution containing the necessary substrates, buffer solutions and enzymes for in vitro transcription and translation. His-tagged proteins were purified from total protein in a cell-free synthesis reaction with TALON® Cell Thru HisTrap resin using the manufacturer’s protocol (GE Healthcare, Little Chalfont, UK). The polyhistidine affinity tags were removed by in-column TEV protease digestion and purification (Fig. S3a†). The resulting proteins were dialyzed through a membrane with a 15 kDa molecular weight cutoff (Pierce Biotechnology, Rockford, IL, USA) in dialysis buffer at 30 °C for 16 hours. The protein solution recovered from dialysis was analyzed via 4–15% SDS-PAGE (Bio-Rad Laboratories, Hercules, CA, USA) (Fig. S5b†). The proteins were diluted to a 1.0 mg ml⁻¹ concentration and stored at 4 °C.

**Formation of biomolecule/KH₇-BP100 complexes**

Two different pDNA/KH₇-BP100 complexes were formed at different N/P ratios (0, 0.1, 0.5, 1.0, 2.5, 5.0, 10 and 50) by adding different amounts of KH₇-BP100 peptide solution (1.0 mg ml⁻¹ stock solution) into 100 µl of water containing 10 µg of pDNA. The solutions were mixed thoroughly and incubated at ambient temperature for 30 minutes without shaking. After incubation, the solutions were diluted with 700 µl of water. The particle size of the complex was determined by DLS with a Zeta Nanosizer using a 633 nm He–Ne laser at 25 °C with a backscatter detection angle of 173°. The surface charge of the complex in solution was measured with a Zeta potentiometer. The averaged data from three replicates were obtained using Zetasizer software ver. 6.20 (Malvern Instruments, Ltd, Worchestershire, UK). Release of pDNA molecules from the complexes was measured by resolved 20 µl of complex solution via 1.0% (w/v) agarose gel retardation assays in 1× TAE buffer at 100 V for 30 minutes.

Complexes of ssRNA/KH₇-BP100 were prepared by mixing 1.0 µg of ssRNA in 100 µl of RNase-free water containing different amounts of KH₇-BP100 peptide (N/P ratios = 0, 0.1, 0.5, 1.0, 2.5, 5.0, 10 and 50). After mixing, each solution was incubated at ambient temperature without shaking for 30 minutes. The solution was diluted with 700 µl of RNase-free water before being subjected to measurement of complex size and surface charge as previously described. Release of ssRNA molecules from the complexes was analyzed by resolving 20 µl of complex solution via 2.0% (w/v) agarose gel retardation assays in 0.5× TBE buffer at 80 V for 60 minutes.

To form nonfluorescent sfGFP fragment-fused MxMT protein/KH₇-BP100 complexes at various molar ratios (0, 0.1, 0.5, 1.0, 2.5, 5.0, 10 and 25), different volumes of 26.3 µM KH₇-BP100 were added to 100 µl of water containing 82.5 µM sfGN155-MxMT protein or 95.5 µM sfGC155-MxMT protein. The complex solutions were incubated at ambient temperature for 30 minutes before dilution with 700 µl of water. The complex size and surface charge were determined with a Zetasizer. The electrostatic interactions of the proteins with the KH₇-BP100 peptide were analyzed by protein gel-shift assays on 4–16% Invitrogen NativePAGE Bis-Tris Gels (Life Technologies, Carlsbad, CA, USA).

**N. benthamiana leaf infiltration**

After complex formation, two biomolecule/KH₇-BP100 complexes formed separately were combined in 1.5 ml microcentrifuge tubes at a ratio of 1 : 1 and gently mixed by pipetting. One hundred microliters of the mixture was infiltrated into the abaxial side of an N. benthamiana leaf with a 1 ml needleless syringe. The transfected plants were incubated at 22 °C under 12/12-hour light/dark periods prior to the expression analyses. Only leaves with complete liquid dispersion through a specific area were used in the subsequent experiments.

**CLSM observation and image analysis**

Approximately 0.5 × 0.5 cm² sections of leaves that had been infiltrated with complex solution were cut from the region around the syringe-attached area (indicated by leaf tissue necrosis). The leaf segments were washed twice and deaerated in water prior to observation with a Zeiss LSM 880 confocal microscope (Carl Zeiss, Oberkochen, Germany) with 488 nm/490–535 nm excitation/emission wavelengths for the detection of sfGFP. A 640–700 nm emission wavelength was used to detect chlorophyll autofluorescence in plant cells. Plant cell images with high fluorescence intensity and low background fluorescence were chosen for image analysis with Fiji ImageJ.62 The means of the relative green fluorescence intensity were measured from at least 9 different regions of interest (ROIs) taken from three independent experiments only in the cytoplasm (the nucleus was excluded). The average background signal measured from the 5 ROIs surrounding the cell was subtracted from all fluorescence signals.

**Transcript analysis**

qRT-PCR was used to determine the transcript abundance in pDNA- or ssRNA/KH₇-BP100 complex-transfected leaves. Total RNA was extracted from transfected plant leaves with an RNasy Plant Mini Kit (Qiagen) and treated with DNase I (Qiagen) to eliminate contaminating genomic DNA. Complementary DNA (cDNA) molecules were synthesized from 0.5 µg of purified RNA with a QuantiTect Reverse Transcription Kit (Qiagen). Known concentrations of ssRNA molecules from
in vitro transcription were converted to cDNA, which was used as the template in qRT-PCR. qRT-PCR was performed using SYBR™ Green RealTime Master Mix Plus (Toyobo, Osaka, Japan) with the primer sequences provided in Table S7.† A linear regression equation for each mRNA molecule was generated by plotting the threshold cycle of the amplification value (ΔCt) against the amount of transcript (Fig. S16†). The transcript amount per 100 ng of total RNA was determined by fitting the ΔCt value to the linear equation.

The RNA decay in the ssRNA/CPP complex-treated plant leaves was determined from qRT-PCR analysis of the transcript abundances. The data points at 0, 0.5 and 4 hours post-transfection were used to determine the first-order decay kinetics.63 The transcript change over time was initially log-transformed, and the RNA decay rate constant (k) was determined through linear regression. The half-life (T(1/2)) was calculated from the RNA decay rate constant as described in eqn (1).

\[ T_{1/2} = \ln(2)/k \]  

(1)

In vitro protein–protein interaction assays

Ten-microgram samples of the nonfluorescent sfGFP fragment-fused sfGN155-MxMT and sfGC155-MxMT proteins were used to form protein/KH9-BP100 complexes at molar ratios of 0.5, 1.0 and 10. Equal amounts of the two proteins without KH9-BP100 were included in the experiment as controls. After complex formation, 25 µl of each protein/CPP complex was mixed together with 50 µl of 20 mM Tris-HCl, pH 7.5 + 0.2% Tween-20. The interaction reaction was incubated at 4 °C for 24 hours without shaking. Each solution was diluted with 800 µl of water and subjected to fluorescence measurement with an FP8500 spectrofluorometer (JASCO, Tokyo, Japan) with a 488 nm excitation wavelength and a scanning emission wavelength of 500 to 580 nm (5 nm bandwidth). The particle size of a 488 nm excitation wavelength and a scanning emission wavelength was resolved on 7.5% Mini PROTEAN® TGX Precast gels (Thermo Scientific, Waltham, MA, USA). The concentration of each protein was determined through DLS measurement with the parameters mentioned above.

Internalization of biomolecule/CPP complexes by plant cells

Internalization of biomolecule/KH9-BP100 complexes by plant cells was studied via assessment of colocalization of fluorescent dye-stained pDNA or fluorescent sfGFP with the plasma membrane-staining fluorescent dye FM4-64.29 pDNA molecules were labeled with Cy3 or Cy5 fluorescent dye using a Label IT® Nucleic Acid Labeling Kit (Mirus Bio, Madison, WI, USA). Solutions containing fluorescent dye-labeled pDNA/KH9-BP100 complexes or nonfluorescent sfGFP fragment-fused MxMT protein/KH9-BP100 complexes were introduced to the abaxial sides of fully expanded leaves of N. benthamiana by syringe infiltration. After incubation, the plant leaves were infiltrated with 10 µM FM4-64 solution and incubated for 10 minutes. The colocalization of the fluorescent dye-labeled pDNA/CPP complexes or sfGFP-visualized complemented MxMT homodimers with FM4-64 was observed on 0.5 × 0.5 cm² sections of leaf samples by CLSM with excitation/emission wavelengths of 555/560–580 nm for Cy3, 488/490–535 nm for sfGFP, and 488/620–700 nm for FM4-64 detection.

Immunoblotting

For immunoblot analysis of pDNA- and ssRNA/CPP cargo-transported plant leaves, total soluble proteins were extracted from at least 5 leaves with protein extraction buffer (100 mM HEPES, pH 7.0, 1 mM EDTA, pH 8.0, 1% (w/v) SDS, 200 mM DTT, 1% (v/v) Halt™ protease inhibitor cocktail (Pierce Biotechnology), and 10% (v/v) glycerol). The soluble proteins were concentrated through a filter membrane with a 10 kDa molecular weight cutoff (Pierce Biotechnology), and the retained proteins were used for immunoblotting. For protein/CPP complex-transfected leaves, total soluble proteins were extracted from one leaf with the protein extraction solution described above. Twenty micrograms of total protein was resolved on 7.5% Mini PROTEAN® TGX™ Precast gels (Bio-Rad Laboratories, Hercules, CA, USA), and the separated proteins were blotted onto Hybond-P PVDF membranes (GE Healthcare, Buckinghamshire, UK). The sfGN155-MxMT and sfGC155-MxMT proteins were detected with a primary antibody solution containing a 1 : 5000 dilution of a rabbit anti-GFP polyclonal antibody (NB600-308) (Novus Biologicals, Littleton, CO, USA) and then with a secondary antibody solution containing a 1 : 20 000 dilution of a horseradish peroxidase (HRP)-conjugated goat anti-rabbit IgG polyclonal antibody (ab6721) (Abcam, Tokyo, Japan). The signal of HRP activity on the membrane was detected after applying 1 ml of SuperSignal™ West Pico PLUS chemiluminescent substrate (Thermo Scientific, Waltham, MA, USA) onto the membrane using a LAS3000 imaging system (Fujifilm, Tokyo, Japan). The band intensities of sfGFP fusion proteins were quantified using Fiji ImageJ.62 Linear regressions of the purified sfGN155-MxMT- and sfGC155-MxMT proteins were performed as shown in Fig. S10b.† The abundance of the sfGFP fusion protein in each sample was calculated using the linear regression equation generated for each standard protein.

Statistical analysis

Multiple comparisons among groups of samples in the same experiment were performed using analysis of variance (ANOVA) with Tukey’s HSD test at p = 0.05. Significant differences in means between two samples were determined by Student’s t-test with StatPlus:mac statistical software (AnalystSoft, Walnut, CA, USA).

Live subject statement

This study was performed in strict accordance with the RIKEN guidelines for the genetically modified plants and was approved by the Wako safety center of RIKEN (Saitama, Japan).

Conflicts of interest

The authors declare no competing financial interests.
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