Development of potent promoters that drive the efficient expression of genes in apple protoplasts

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Abstract
Protoplast transient expression is a powerful strategy for gene functional characterization, especially in biochemical mechanism studies. We herein developed a highly efficient transient expression system for apple protoplasts. The abilities of the Arabidopsis thaliana and Malus domestica ubiquitin-10 (AtUBQ10 and MdUBQ10) promoters to drive the expression of multiple genes were compared with that of the CaMV 35S promoter, and the results revealed that the AtUBQ10 and MdUBQ10 promoters were more efficient in apple protoplasts. With this system, we demonstrated that active AtMKK7ac could activate MAPK6/3/4 signaling cascades, which further regulated MdWRKY33 phosphorylation and stability in apple. Furthermore, the ligand-induced interaction between the immune receptor AtFLS2 and the coreceptor AtBAK1 was reconstituted in apple protoplasts. We also found that the stability of the bacterial effector AvrRpt2 was regulated by feedback involving auxin and the immune regulator RIN4. The system established herein will serve as a useful tool for the molecular and biochemical analyses of apple genes.

Introduction
Apple is a rosaceous fruit tree plant that is cultivated worldwide, and researchers are becoming increasingly interested in the functionally important genes and molecular mechanisms involved in controlling apple fruit and plant growth and development1–3, fruit quality4–8, resistance9–12, and other physiological activities13,14. The expression of genes in plant materials is of critical importance for gene functional characterization and signal transduction pathway identification. Currently, apple genes are transiently expressed in onion epidermal cells14, apple and maize protoplasts7,8,13,14, pollen tubes13, tobacco and apple leaves3,10, and apple fruit surface cells5 and are stably expressed in transgenic apple plants3,7,12, callus6–8,11, and Arabidopsis8 for gene functional characterization. A protocol for transient expression in apple fruit cells was also developed by Spolaore et al.15. All these methods contribute to elucidating the molecular mechanisms controlling the physiological processes in apple.

Protoplasts are cell wall-free cells isolated from plant tissues and are capable of perceiving external stimuli. Transient expression in protoplasts has been proven to be a powerful strategy for gene functional characterization and signal pathway identification16, as it provides valuable information for understanding the molecular mechanisms controlling plant immune responses, hormone signaling, growth and development, epigenetic gene expression regulation and other physiological processes17–20. The expression of target genes in protoplast cells can be suppressed by the overexpression of artificial microRNAs or by modulation of their regulatory elements20,21. Mutagenesis in protoplast cells can be performed by CRISPR/Cas9 gene editing22,23. Protoplast transient expression has been found to efficiently, precisely, accurately, and consistently reveal molecular mechanisms. In apple, the use of an apple protoplast transient expression strategy is of special and critical importance. It can realize homologous expression without technical obstacles or the long time required to obtain transgenic apple plants and callus. Importantly, it is unaffected by the problems of heterologous expression. For example, certain Arabidopsis proteins were mislocalized after their expression in...
A precisely controlled expression strategy with an accurate evaluation method will be a powerful tool for apple gene functional characterization and signal transduction pathway identification, which will substantially deepen the understanding of the gene functions and signaling processes involved. In a recent report, apple protoplasts were subjected to CRISPR/Cas9 modification to generate genome-edited apple plants23. Despite the application of this method in several studies on subcellular localization and protein interactions8,13, it has not been extensively employed in studies on apple gene functional characterization and signal transduction pathway identification, especially in biochemical mechanism studies. Low gene expression in apple protoplast cells might be an important factor impeding the use of apple protoplasts.

In this study, target genes were stably and efficiently expressed in apple protoplasts by using the AtUBQ10 orMdUBQ10 promoter. The data obtained herein using these promoters provide insight into the novel mechanisms regulating immune and disease resistance in apple. This research paves the way for the application of this method in several studies on subcellular localization and protein interactions.

### Results and discussion

Intact apple protoplast cells that were spherical in shape and did not exhibit bursting or clustering were isolated from ‘Orin’ and ‘Zhong’ apple callus cells cultured in MS medium for different durations (Fig. 1a). The highest protoplast amounts were obtained from both ‘Orin’ and ‘Zhong’ apple callus cells cultured in MS medium for 10 days, yielding 3.68 × 10^6 and 4.07 × 10^6 protoplasts per gram of fresh weight callus (per g FW callus), respectively (Fig. 1b). Although apple callus cells cultured in MS medium for 6 days yielded only 1.3 × 10^6 protoplasts per g FW, this condition yielded the strongest target protein expression (Supplementary File 1: Fig. S3a) and was therefore used for protoplast transfection in our research.

**Pro-BIUTNT is a potent promoter that enables the efficient expression of target genes in apple protoplast cells**

Ethylene response factors (ERFs) are a large family of transcription factors that are extensively involved in controlling fruit ripening, disease resistance and other physiological activities or traits. MdERF1 and MdERF2 are two ERF transcription factors involved in controlling fruit ripening28. In our apple protoplast transient expression system driven by the cauliflower mosaic virus (CaMV) 35S promoter, MdERF1 expression was not detectable, and a weak MdERF2 signal was detected by western blot. However, a clear and specific MdERF1 signal and a strong MdERF2 signal were detected when the promoter of *Arabidopsis ubiquitin-ten* (Pro-BIUTNT) was utilized (Fig. 2b), which is the 1307 nucleotide sequence upstream of the ATG translational start codon of the *Arabidopsis ubiquitin-10* (At4g05320) gene. Further studies revealed that two of the three independent MdERF2 clones under the control of CaMV 35S generated a dramatically weaker signal than that generated using Pro-BIUTNT (Fig. 2c). Only trace amounts of MdERF1 were detected in two MdERF1 clones under the control of CaMV 35S compared with the significantly stronger and more specific signal of the gene driven by Pro-BIUTNT (Fig. 2c).

MdERF3 and MdERF6 are two ERF transcription factors that are responsive to pathogen infection26. Homologs of MdERF3 and MdERF6, MdERF3a and MdERF6a were cloned and transfected into apple protoplast cells. Strong MdERF3a and MdERF6a signals were detected when the genes were driven by Pro-BIUTNT, but little or no signal was detected when the genes were driven by CaMV 35S (Fig. 2c).

ERF98 (At3g23230) is an ERF transcription factor that is responsive to treatment with flg22, a conserved 22-amino acid peptide from eubacterial flagellin that induces immune responses in plants27, and MdERF98 is the ortholog of ERF98 in the apple genome. No expression was found when driven by CaMV 35S, but a strong signal was detected for MdERF98 driven by Pro-BIUTNT (Fig. 2b). Analysis of a different CaMV 35S::MdERF98 clone also revealed that the gene under the control of CaMV 35S was expressed at lower levels in protoplast cells than the gene driven by the Pro-BIUTNT promoter (Supplementary File 1: Fig. S1). This result suggests that CaMV 35S was not suitable for driving the expression of the target genes in apple protoplast cells and was consistent with the expression of three independent MdERF2 and MdERF1 clones driven by the CaMV 35S promoter in further assays.

MdEIL2 is the ortholog of EIN3 and regulates the expression of *MdPGI*28. A strong and clear MdEIL2 signal was achieved using Pro-BIUTNT, in contrast to the weak signal achieved with CaMV 35S (Fig. 2d).

WRKY33 and WRKY29 are involved in regulating ethylene and phytoalexin biosynthesis and pathogen-associated molecular pattern (PAMP)-triggered immune responses, respectively29,30. The orthologs of the two genes in apple, MdWRKY33 and MdWRKY29 (Table S1), were cloned and transfected into apple protoplast cells. MdWRKY33 was hardly expressed under the control of CaMV 35S; however, a strong signal was obtained when it was driven by Pro-BIUTNT (Fig. 2b, Supplementary File 1: Fig. S3a, b). To further explore and compare the capabilities of Pro-BIUTNT and CaMV 35S to drive the expression of target genes in apple protoplast cells, the C-terminal epitope HA tag of MdWRKY33 was replaced with GFP. Strong MdWRKY33-GFP expression was detected when the gene was driven by Pro-BIUTNT, while
Fig. 1 (See legend on next page.)
no expression was found under the control of \textit{CaMV 35S} (Supplementary File 1: Fig. S4a). \textit{CaMV 35S} has numerous variants, and the variant we herein named \textit{CaMV 35S-2} was subcloned from the pBI121 binary vector\textsuperscript{31} and analyzed for its ability to drive the expression of the target genes. Both of the \textit{CaMV 35S} promoters contained the core sequence of the \textit{CaMV 35S} promoter (Supplementary File 1: Sequences 1 and 2), and their abilities to drive the expression of target genes in apple have been proven\textsuperscript{11} (Fig. 4f). This variant of \textit{CaMV 35S} resulted in weak MdERF98 expression and noMdWRKY33 expression (Supplementary File 1: Fig. S4b, c).

The above results suggested that \textit{Pro-BIUTNT} is a strong promoter that enables the stable expression of target genes in apple protoplast cells.

Previous research showed that the \textit{AtUBQ10} promoter was more stable and persistent than \textit{CaMV 35S} in \textit{Arabidopsis} and tobacco\textsuperscript{32}, and the target gene was expressed at higher levels under the control of the \textit{AtUBQ10} promoter in both dicot (\textit{Arabidopsis}) and monocot (rice)
target genes in apple protoplast cells significantly better than CaMV 35S (Fig. 2c). Notably, the MdUBQ10 promoter sequence was not highly similar to that of Pro-BIUTNT (Supplementary File 1: Table S2).

Our research demonstrated that the Pro-BIUTNT and MdUBQ10 promoter enable the efficient expression of target genes in apple protoplast cells.

**AtMKK7ac overexpression activates MAPK signaling in apple**

MAPK signaling is an ancient and conserved signaling pathway in eukaryotes\(^3\) that helps to control plant growth, development, and disease resistance\(^30,35\). However, its upstream activators have not been identified in apple.

AtMKK7 (At1g18350) is thought to be involved in regulating both basal and systemic acquired resistance in plants\(^36\). The constitutive expression of AtMKK7 in its active form (AtMKK7ac: AtMKK7 with S193ES199D mutation) induces hypersensitive reactions in tobacco leaves (Fig. 4c). The in vivo detection of MdMAPK phosphorylation using an anti-pERK antibody showed that the constitutive expression of AtMKK7ac in apple protoplast cells markedly activated the phosphorylation of MdMAPK3/6/4, particularly MdMAPK4 (Fig. 4d). AtMKK7ac was then expressed in apple callus cells via Agrobacterium-mediated transient expression, and AtMKK7ac overexpression strongly activated MdMAPK phosphorylation (Fig. 4g). An in vitro kinase assay showed that MBP-AtMKK7ac could phosphorylate MdMAPK6, and Thr-231 was important for this process (Fig. 4f).

In this study, apple protoplasts were isolated from genetically transformed ‘Orin’ apple callus cells expressing MdMAPK6-FLAG driven by CaMV 35S and from wild-type ‘Orin’ apple callus cells. The coexpression of AtMKK7ac and MdWRKY33 induced a band shift of MdWRKY33 in apple protoplast cells overexpressing MdMAPK6-FLAG, which mimicked the band shift of Botrytis-induced kinase 1 (BIK1) observed in PAMP-triggered immune responses\(^37\), whereas a similar band shift was not observed in protoplast cells expressing only MdWRKY33. This result suggested that the expression of MdWRKY33 in apple protoplast cells was modified, potentially by phosphorylation, by upstream activators. The level of MdWRKY33 in the presence of MdMAPK6-FLAG was markedly higher than that obtained without MdMAPK6-FLAG overexpression (Fig. 4e), which suggests that the stability of MdWRKY33 is influenced by the status and degree of its phosphorylation upon activation by upstream factors.

The revelation that AtMKK7ac activates MdMAPK6 provides an effective method for activating MAPK signaling in apple cells and for elucidating the elements controlling MdMAPK6 activation in apple.
Pro-BIUTNT enables the reconstitution of BAK1 and FLS2 interaction in apple protoplast cells

The interaction of AtBAK1 and AtFLS2 is an important and fundamental signaling event in plant immune responses triggered by PAMPs. Immune complex formation was observed in apple protoplast cells after flg22 elicitation, and the phenotypes were characterized 2 days later. AtMKK7 overexpression activated MAPK cascades in apple protoplast cells. AtMKK7 overexpression induced hypersensitive responses in tobacco leaves. The white arrows denote the phenotype of hypersensitive reactions.

This study elucidated the paradigmatic signaling event in apple protoplast cells and provides a direct method for identifying novel signaling components of PAMP-triggered immune responses in highly developed perennial woody plants. This investigation also provides an accurate and convenient method for comparative studies of immune signaling in model and highly developed plants.
Consistent with previous findings, AXR2 expression was significantly reduced in the presence of AvrRpt2 and 1-naphthalacetic acid (NAA), and the P87S mutation in AXR2 blocked NAA- and AvrRpt2-mediated degradation (Fig. 6b). Interestingly, the protein level of AvrRpt2 was markedly decreased to undetectable levels in the presence of 1 mM NAA (Fig. 6b). The same results were found in tobacco leaves when AvrRpt2 was expressed or coexpressed together with AXR2 or AXR2 P87S (Fig. 6c).

The degradation of AvrRpt2 was also identified in its interactions with RIN4. The functional processing of RIN4 by AvrRpt2 was confirmed by the disappearance of full-length RIN4-GFP [indicated as RIN4 (FL) in Fig. 6d] and the detection of the GFP-tagged RIN4 C-terminus [indicated as RIN4 (AMC) in Fig. 6d] upon the coexpression of RIN4-GFP with AvrRpt2-HA. We also repeatedly found that the AvrRpt2 level was decreased or even undetectable in the presence of RIN4 (Fig. 6d, Supplementary File 1: Fig. S5a, b).

These results suggest that the microbial effector is negatively regulated by auxin and the RIN4-associated complex in the plant. This phenomenon might represent two important disease resistance mechanisms in plant basal and effector-triggered resistance. The strengthened auxin signaling might provide feedback to remove the virulence source inside cells and thereby enhance plant disease resistance. In AvrRpt2-activated disease resistance mediated by the nucleotide-binding domain leucine-rich repeat (NLR) protein RPS2, AvrRpt2 might be negatively controlled to prevent the excessive induction of immune responses triggered by the causal agent itself. The detailed mechanisms by which auxin and the RIN4 complex control the microbial effector in plant cells will be elucidated in the future.

**Materials and methods**

**Plasmid construction**

*Pro-BIUTNT*, an abbreviation for ‘promoter of *ubiquitin-ten*’, the 1307-base pair (bp) sequence upstream from the translational start codon ATG of *Arabidopsis*
ubiquitin-10 (At4g05320), was amplified from Arabidopsis genomic DNA.

pHBT-CaMV 35S::AvrRpm1-2HA was used as the backbone for the construction of the protoplast transient expression vectors used in this study. The vector driven by CaMV 35S was constructed by directly replacing AvrRpm1 with the coding sequence of the target gene. The vector driven by the Pro-BIUTNT or MdUBQ10 (MDP00000820500) promoter was constructed by substituting Pro-BIUTNT or nucleotide sequences of different lengths upstream from the ATG translational start codon of MdUBQ10 for CaMV 35S.

**Protoplast isolation, transfection, and western blot analysis**

Apple protoplast cells were isolated from 'Orin' and 'Zihong' apple callus cells that had been cultured in MS medium for different durations according to the method described by Gao et al. One gram of apple callus cells was dispersed evenly in 10 mL of enzyme digestion solution consisting of 1.5% cellulase, 0.4% macerozyme, and 0.05% pectinase dissolved in 20 mM MES buffer containing 20 mM KCl, 10 mM CaCl₂, 2% sucrose, and 0.4 M mannitol (pH adjusted to 5.7). After vacuum infiltration for 30 min, the digestion solution was maintained undisturbed at room temperature for 8 h. The solution was then passed through a nylon mesh with a diameter of 100 μm and collected in a 50 mL Eppendorf tube, to which an equal volume of W5 solution was added. The nylon mesh was washed twice with W5 solution, and the solution was collected and added to the protoplast cell suspension. After the suspension was centrifuged at 650 × g for 2 min, the protoplast cells were collected, resuspended in 15 mL of W5 solution, and maintained on ice for 30 min. The supernatant was then discarded, and the protoplasts were suspended in 1 mL of MMG solution and later used for protoplast cell number calculations and plasmid DNA transfection.

Protoplasts were transfected as described by He et al. Protein expression in protoplast cells was detected by western blot using anti-HA, anti-FLAG, or anti-GFP antibodies.

**Luciferase activity assay**

AvrRpm1 in pHBT-CaMV 35S::AvrRpm1-2HA was replaced with a luciferase (Luc) coding sequence to generate pHBT-CaMV 35S::Luc. CaMV 35S was then replaced to generate luciferase vectors under the control of the promoters investigated in the research. The luciferase activity in apple or Arabidopsis protoplast cells was measured according to He et al.

**Co-immunoprecipitation (Co-IP)**

Driven by Pro-BIUTNT, AtBAK1-HA and AtFLS2-FLAG were expressed in apple protoplast cells for 6 h. After treatment with 100 nM flg22 or the equivalent amount of solvent for an additional 10 min, the protoplast cells were harvested. Co-IP was conducted using anti-FLAG beads according to Lu et al.

Cells expressing AtBAK1 that were treated or not treated with only flg22 were used as controls.

**In vitro kinase assay**

MdMAPK6 (MDP0000340624) or its mutant MdMAPK6 T231A, MdMAPK6 T231D, or MdMAPK6 T236A was cloned into a pBlI121 binary vector under the control of CaMV 35S with a C-terminal FLAG epitope tag and transformed into apple callus cells according to He et al.

The recombinant protein MBP-AtMKK7ac was induced and purified from E. coli BL21.

In the kinase activity assay, MdMAPK6 or one of its mutants was purified from one gram of transgenic apple callus cells and mixed with two micrograms of MBP-AtMKK7ac in 25 μL of kinase reaction buffer (20 mM Tris-HCl [pH 7.5], 5 mM EDTA [pH 8.0], 1 mM DTT, 20 mM MgCl₂, 0.1 mM ATP and 5 μCi [γ-32P] ATP). After reacting for 2 h at 23 °C, the proteins were denatured and separated by SDS-PAGE. Phosphorylation was detected by autoradiography.

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Author contributions
X.W., X.C., and S.W. conceived the project and designed the research; X.W., LXu., X.L., and L.X. performed all the experiments; X.W., X.C., and S.W. analyzed the experimental data and discussed the results; and X.W. and S.W. wrote the manuscript.

Data availability
All data supporting the findings of this study are available within the paper or within its supplementary information published online. The materials used in this study are available from the corresponding author upon reasonable request.

Conflict of interest
The authors declare no competing interests.

Supplementary information
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