The Self-Interaction of a Nodavirus Replicase Is Enhanced by Mitochondrial Membrane Lipids

Yang Qiu1,2, Zhaowei Wang1, Yongxiang Liu1, Yajuan Han1, Meng Miao1, Nan Qi1, Jie Yang1, Hongjie Xia1, Xiaofeng Li2, Cheng-Feng Qin2, Yuanyang Hu1*, Xi Zhou1*

1 State Key Laboratory of Virology, College of Life Sciences, Wuhan University, Wuhan, Hubei, China, 2 State Key Laboratory of Pathogen and Biosecurity, Beijing Institute of Microbiology and Epidemiology, Beijing, China

Abstract

RNA replication of positive-strand (+)RNA viruses requires the protein-protein interactions among viral replicases and the association of viral replicases with intracellular membranes. Protein A from Wuhan nodavirus (WhNV), which closely associate with mitochondrial membranes, is the sole replicase required for viral RNA replication. Here, we studied the direct effects of mitochondrial membrane lipids (MMLs) on WhNV protein A activity in vitro. Our investigations revealed the self-interaction of WhNV protein A is accomplished via two different patterns (i.e., homotypic and heterotypic self-interactions via different interfaces). MMLs stimulated the protein A self-interaction, and this stimulation exhibited selectivity for specific phospholipids. Moreover, we found that specific phospholipids differently favor the two self-interaction patterns. Furthermore, manipulating specific phospholipid metabolism affected protein A self-interaction and the activity of protein A to replicate RNA in cells. Taken together, our findings reveal the direct effects of membrane lipids on a nodaviral RNA replicase.

Introduction

One universal feature of positive-strand (+)RNA viruses is the assembly of their viral RNA replication complexes (vRCs), including viral replicase proteins, viral RNA, and host proteins, on host intracellular membranes [1–3]. During viral RNA replication, these viruses often induce specific intracellular membrane remodeling and lipid biosynthesis modifications via viral replicates [4–6]. On the other hand, lipids are major components of intracellular membranes, as they control membrane fluidity and plasticity [6,7], and virus-induced modifications of lipid biosynthesis are closely linked to the formation and function of vRCs [2].

The viral protein-protein interaction is important for (+)RNA viruses replication [1]. Most (+)RNA viruses encode multiple viral proteins, which work together for the vRCs formation and function [8–13]. Although some replication proteins from many viruses have activity as individual units, they still require self-interaction/oligomerization for the complete functionality [14–17].

Nodaviruses (family Nodaviridae) are (+)RNA viruses that contain a bipartite genome consisting of two nonpolyadenylated RNAs, RNA1 (~3.1 kb) and RNA2 (~1.4 kb), which encode protein A, the RNA-dependent RNA polymerase (RdRP) [18] and capsid precursor protein α [19], respectively. A subgenomic RNA3 (sgRNA3), which is not encapsidated into virion, is synthesized during RNA1 replication and encodes nonstructural protein B2, a viral suppressor of RNA silencing [20].

In contrast to many (+)RNA viruses such as bromovirus, flavivirus, picornavirus and tombusvirus, in which a set of viral RNA replicase proteins synthesizes their RNA genomes, nodaviruses encode a sole RNA replicase protein, protein A, for viral RNA replication [1,21]. This feature renders nodaviruses such as Flock House virus (FHV) and Wuhan nodavirus (WhNV) well-recognized and simplified models for studying viral RNA replication [22–27]. Previous studies of FHV, the most extensively studied member of the Nodaviridae family, revealed that FHV protein A contains multiple activities including synthesizes RNA, mitochondrial membrane association and self-interaction [22]. Disrupting the self-interaction of FHV protein A by the point mutations revealed that FHV protein A self-interaction is important for its function [29]. Moreover, the domains responsible for FHV protein A self-interaction include the trans-membrane regions, implying the correlation between membranes and protein A self-interaction [28,29].

Multiple lines of evidence indicated that intracellular membranes, particularly membrane lipids, mediate FHV RNA protein A function. FHV protein A is a lipid-binding protein with particular affinity for specific anionic phospholipids, which may mediate the protein A-membrane interactions required for vRCs assembly [30]. The in vitro study showed that complete replication activity of FHV vRCs isolated in membrane fraction is disrupted by membrane-disrupting detergents, and can be augmented by the
addition of exogenous phospholipids [31,32]. Moreover, the genes involved in the synthesis of phosphatidylcholine play an important role in FHV RNA replication in *Drosophila* cells [33]. Inhibition of fatty acid synthesis using cerulenin resulted in the block of FHV RNA replication in *Drosophila* cells [34]. However, whether membrane lipids directly mediate nodaviral RNA protein A self-interaction is not well understood.

As a virus closely related to FHV, WhNV has been well characterized and provides novel insights for nodaviral subgenomic RNA replication [26] and RNA silencing suppression [35,36]. Moreover, WhNV protein A can initiate RNA synthesis via *de novo* mechanism and contains a terminal nucleosidyl transferase activity [37]. Previous study showed that the activity of WhNV protein A to associate with mitochondrial membranes is closely linked with its activity for recruitment and stabilization of viral genomic RNA templates [38], suggesting the direct role of membrane lipids in WhNV protein A function. In this study, we focused on the effects of membrane lipids on WhNV protein A self-interaction. We expressed WhNV protein A in *vitro*, and isolated mitochondrial membrane lipids (MMLs) from mitochondrial outer membrane. Our study reveals that WhNV protein A is self-interacted and MMLs directly mediate protein A self-interaction in many aspects.

Materials and Methods

Plasmids

Standard procedures were used for restriction nuclease digestion and plasmid construction and purification. To analyze WhNV protein A activity in cells, protein A ORF and RNA1 was inserted into pAC5.1/V5-His B vector (Invitrogen, Carlsbad, CA). Plasmids for the purification of MBP fusion protein A were constructed by inserting protein A ORF into pMAL-c2X (New England BioLabs, Ipswich, MA). For *in vitro* translation, WhNV and FHV protein A ORF was inserted into pET-28a (Novagen, Germany), respectively. Mutations were introduced into protein A ORF via PCR-mediated mutagenesis as described previously [26,38]. The oligonucleotides used in this study are shown in Table 1.

Cells and Transfection

Pr-E cells, which is derived from *Pieris rapae* larvae, the natural host of WhNV, and was successfully utilized to study WhNV RNA replication previously [Qiu et al., 2011; Qiu et al., 2013], were maintained at 27°C in Grace’s medium (Gibco, Carlsbad, CA, USA) supplemented with 10% fetal bovine serum (Gibco). DNA plasmids were transfected into cells using FuGENE HD transfection reagent (Roche, Basel, Switzerland) according to the manufacturer’s protocol. All subsequent assays were performed 36 hrs after transfection except where indicated otherwise.

WhNV *trans*-replication System

*Whnv* *trans*-replication system was previously established to study WhNV RNA replication [26,38]. Briefly, we constructed two WhNV RNA1 mutants based on pAC1E plasmid, in which an EGFP open reading frame (ORF) is inserted at the 3’ end of RNA1 sequence [38]. The plasmid pAC1E is functional template for RNA1 replication (the transcribed and repeated products are labeled as “RNA1E”), but the ORF of protein A is closed by the mutation of the start codon [38]. WhNV protein A is provided by the plasmid pA, which the RdRp activity is remained but the ability for replication as RNA template is destroyed by deleting the 5’ and 3’ untranslated regions [38]. This WhNV *trans*-replication system, in which the RNA1 template and protein A mRNA are separately provided by two plasmids, was successfully used to study WhNV RNA replication and RNA recruitment/stabilization [38]. The assay was tested in Pr-E cells, 36 hrs after transfecting with the indicated plasmids, cells were collected and total RNA was separated, and 2 μg of total RNA was analyzed by Northern blotting.

Western Blot Analysis and Antibodies

The proteins extracted from cells were subjected to 10% SDS-PAGE and Western blot analysis as previously described [26,38]. Unless otherwise indicated, the anti-MBP polyclonal antibody was purchased from New England BioLabs, and the other primary and secondary antibodies were purchased from Proteintech, Chicago, IL, USA.

RNA Extraction and Northern Blot Analysis

Total RNA was extracted from cells using TRIzol reagent (Invitrogen) and digested with RQI RNase-free DNase I (Promega, Madison, WI, USA) as previously described [26,38]. For Northern blot analysis, 2 μg of each RNA sample was analyzed via Northern blot analysis as previously described [26,38]. The probes for (+) and (−) EGFP were complementary to the entire EGFP sequences. All probes were labeled with DIG-UTP (Roche) for *in vitro* transcription; the corresponding oligonucleotides are shown in Table 1.

Purification of Protein A and its Derivatives

The expression and purification of recombinant WhNV protein A and its derivatives were carried out as previously described [35–37,39]. Briefly, to obtain soluble recombinant protein, Maltose-binding protein (MBP)-tagged full-length protein A and its mutants as well as the negative control protein MBP were expressed in *Escherichia coli* strain TB1 at 20°C in the presence of 0.2 mM IPTG. Cell pellets were resuspended in binding buffer (20 mM Tris-HCl [pH 7.4], 200 mM NaCl, 1 mM EDTA, 10 mM 2-Mercaptoethanol) supplemented with 1.5% Triton-X 100 and protease inhibitors cocktail (Sigma, St. Louis, Mo, USA). Cells were lysed by sonication and then debris was removed by centrifugation for 30 min at 11,000 xg. The proteins in the supernatant were purified using amylose resin (New England BioLabs) according to the manufacturer’s protocol and concentrated using Amicon Ultra-15 filters (Millipore). Lipids were obtained from Sigma in the highest purity grades available: 1,1’,2,2’-tetraoleoyl cardiolipin (CL), 1,2-dioleoylsn-glycero-3-phosphoethanolamine (sphingomyelin), 1,2-dioleoylsn-glycero-3-phosphocholine (PC).

Mitochondrial Membrane Lipids and Liposomes

Mitochondrial outer membranes were isolated from Pr-E cells by mechanical disruption and differential centrifugation as previously described [40,41], and then determined by immunodetections (Fig. S1). Subsequently, the purified outer mitochondrial membranes were treated with 0.1 mg/ml protease K (Sigma) for 10 min in hypotonic buffer supplemented with 1.5% Triton-X 100 to dissolve integral membrane proteins. MMLs were then reisolated by centrifugation at 12,000 x g for 20 min and resuspended in hypotonic buffer. MMLs were further purified and concentrated by using Amicon Ultra-15 filters (Millipore). Lipids were obtained from Sigma in the highest purity grades available: 1,1’,2,2’-tetraoleoyl cardiolipin (CL), 1,2-dioleoylsn-glycero-3-phosphoethanolamine (sphingomyelin), 1,2-dioleoylsn-glycero-3-phosphocholine (PC).
phosphate (PA), 1,2-dioleoyl-sn-glycero-3-[phospho-rac-(1-glycerol)] (PG), 1,2-dioleoyl-sn-glycero-3-phospho-L-serine (PS), 1,2-dioleoyl-sn-glycero-3-phosphocholine (PC) and 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (PE). The liposomes were prepared as described [30–32,42]. Briefly, the purchased lipids were dissolved and mixed in chloroform/methanol (2:1) at 10 mg lipid per 1 ml organic solvent. The mixture was dried under nitrogen and lyophilized to remove any traces of solvent. The dry film was hydrated with 20 mM HEPES buffer at pH 7.4 by vortexing overnight at 4°C. The purified MMLs and liposomes were quantified by Bradford protein assay (Bio-Rad) using a UV-visible spectrophotometer (Shimadzu).

Protein flotation Assays

For MML-binding assays, MBP-tagged protein A or its derivatives (10 pmol each) was incubated with MMLs (50 μg per 100 μl reaction mixture) in flotation buffer (50 mM HEPES [pH 7.4], 50 mM KCl, 2 mM MgCl2, 1 mM DTT) for 1 h at room temperature. After the incubation, the reaction mixtures were diluted with 4 volumes of flotation buffer, and Nycondenz (Sigma) was added to the mixtures to a final concentration of 37.5% (wt/vol), and samples were loaded under a 5% to 25% discontinuous Nycondenz gradient and centrifuged to equilibrium at 100,000 g for 20 hrs at 4°C in a Beckman Coulter SW40 rotor. After centrifugation, the gradient was divided into two fractions including the upper half of the gradient (low-density fraction, LD) and the lower half of the gradient (high-density fraction, HD). Protein samples were isolated from each fraction via centrifugation at 180,000 g in a Beckman Coulter SW40 rotor for 3 hrs and then analyzed via Western blotting.

Pull-down Assays

MBP pull-down assays were performed with recombinant MBP fusion proteins and His-tagged proteins as previously described.

Table 1. Oligonucleotides used in this work.

| Primers       | Sequences (5’ to 3’)                                             |
|---------------|-----------------------------------------------------------------|
| pA GAA-His-R1 | CTAACTGTAATCTGGGAACATCGTATGGTAGCTAAGGAACATTTCTTAAAGACTGGT       |
| pA GAA-His-R2 | GCCGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTGGT |
| pA GAA-HA-R1  | CTAATGATGATGATGATGATGGCTTAAGGAACATTTCTTAAAGACTAGAG             |
| pA GAA-HA-R2  | GCCGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTGGT |
| 1–1014 MBP/protA-F | GGATCCATGAGTGTCAGATACCAACAGAGGTCTTAATCTTAAAGACTAGAG             |
| 1–1014 MBP/protA-R | GCCGCGAGGTACCTTAACTTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–254 MBP/protA-F | GCGGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–254 MBP/protA-R | GCCGCGAGGTACCTTAACTTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–480 MBP/protA-F | GTCGACCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–480 MBP/protA-R | GCCGCGAGGTACCTTAACTTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–659 MBP/protA-F | GTCGACCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–659 MBP/protA-R | GCCGCGAGGTACCTTAACTTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 255–1014 MBP/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 255–1014 MBP/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 481–1014 MBP/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 481–1014 MBP/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 660–1014 MBP/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 660–1014 MBP/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 840–1014 MBP/protA-F | GCGGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 840–1014 MBP/protA-R | GCGGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| His/control-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| His/control-R | GCGGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–1014 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–1014 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–254 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–254 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–480 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–480 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–659 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–659 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 255–1014 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 255–1014 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 481–1014 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 481–1014 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 660–1014 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 660–1014 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 840–1014 His/protA-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 840–1014 His/protA-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–254/M1-F | GGATCCATGAGTGTCAGATACCAACAGAGGTCTTAATCTTAAAGACTAGAG             |
| 1–254/M1-R | GCCGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |
| 1–254/M2-F | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| 1–254/M2-R | GATATTCTTAAAGACTAGATTCGGTAGCTAAGGAACATTTCTTAAAGACTAGAG             |
| FHV-protein A-F | GGATCCATGAGTGTCAGATACCAACAGAGGTCTTAATCTTAAAGACTAGAG             |
| FHV-protein A-R | GCCGCCCTCCCTAAGCTGAAACCTGATATGGTGGTAGCTAAGGAACATTTCTTAAAGACTAGAG |

Sequence specific primers are designed according to Genbank no. AY962576 (WhNV RNA1). Characters in bold indicate restriction endonuclease sites, and the types are shown in brackets.
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[35,36]. Briefly, amylose resin (New England Biolabs) was added to the reaction mixtures containing MBP fusion and His fusion proteins and then incubated at room temperature for 4 hrs. To test the stimulating effects of MMLs on protein A self-interaction, MMLs were added (0.1 to 10 μg MMLs per 1 μl reaction mixture) to the reaction mixtures. After subsequent washing and collection, proteins that bound to the amylose resin were subjected to Western blot analysis. Immunoblotting signals were quantified and plotting results against standard curves from immunoblotting of serially diluted samples (data not shown).

Chemical Cross-linking Assays

Chemical cross-linking assays were performed as previously described [36]. MBP-tagged proteins were cross-linked in cross-linking buffer (10 mM HEPES [pH 7.4], 50 mM MgCl2, 1 mM DTT, 1% glycerol, and 0.03% [vol/vol] glutaraldehyde) for 30 min. The complexes were then analyzed via 10% SDS-PAGE.

PA Inhibitor and Cells Viability Assays

5-fluoro-2-indolyled-chloroalopine (FIPI) (Sigma) was used to inhibit PA production as previously described [43]. Briefly, 12 hrs after transfection, cells were treated with 75 nM FIPI in DMSO and incubated for another 24 hrs. Then cells were collected, and divided into two equal fractions. One fraction was used for Co-IP experiments as described below, and the other fraction was used for total RNA extraction and following analyzing by Northern blot as described above. Cell viability assays were performed using MTT (Sigma) as previously described [33].

Comununoprecipitation Assays

Coimmunoprecipitation (co-IP) assays were performed as previously described [35,36]. Briefly, 36 hrs after transfection and FIPI treatment, cells were lysed with NETN buffer [20 mM Tris-HCl (pH 7.4), 150 mM NaCl, 1 mM EDTA, and 0.5% NP-40] for 20 min at 4°C in the presence of protease inhibitors cocktail (Sigma). Lysates were clarified at 12,000 rpm for 10 min at 4°C, and then postnuclear lysates were precleared via incubation with protein-G agarose beads (Roche) coupled to goat anti-mouse IgG and then incubated with antibodies (mouse anti-His antibody, mouse anti-HA antibody, or control mouse anti-FLAG antibody) at room temperature for 4 hrs. The antibody-bound complexes were captured, washed, and then subjected to SDS-PAGE and Western blotting analysis with rabbit anti-His antibody or rabbit anti-HA antibody.

PA Determination

Total PA content was determined using a modified phospholipase D-based enzymatic method [44]. Briefly, cells were detached by treatment with PBS plus 5 mM EDTA and resuspended in 1 × PBS buffer containing 1 mM MgCl2, 1 mM CaCl2, 5 mM glucose, and 0.2% BSA. Cells were then immediately frozen and the total cellular lipids were incubated first with lipoprotein lipase of E. coli and purified (Fig. 1A). MBP-protA was expressed at its expected molecular weight (around 150 kDa; Fig. 1B, lane 3), and MBP protein alone was expressed as the negative control under the same condition (Fig. 1B, lane 2).

To determine whether WhNV protein A can self-interact in vitro, we used 1 μM of MBP-protA to pull down 1 μM of His-tagged protein A (His-protA) that was expressed in nucleasen-treated rabbit reticulocyte lysates (RRLs) in vitro. The bound complexes were analyzed by Western blots using anti-His antibody, and the input lysates were also detected using anti-His and anti-MBP antibodies, respectively. His-protA was efficiently pulled down by MBP-protA (Fig. 2A, lane 4), but it did not interact with negative control MBP protein alone (Fig. 2A, lane 2). The dimerization of protein A was further confirmed using MBP-protA cross-linking assay in vitro (Fig. 2B). MBP-protA was incubated in the chemical cross-linking for 20 min, and then the samples were analyzed via SDS-PAGE, revealing one band with molecular weight about 330 kDa (Fig. 2B, lane 4), indicating that protein A can form homodimer.

The protein A self-interaction was further confirmed via co-IP in Pr-E cells. As shown in Fig. 2C, Pr-E cells were transfected with either empty vector (pAC) (lane 1), a plasmid expressing protein A with C-terminal HA tag (pHA-HA) (lane 2), a plasmid expressing protein A with a C-terminal His tag (pA-His) (lane 3), or with both pA-HA and pA-His (lane 4). After 36 h of transfection, cells were harvested, and protein complexes were immunoprecipitated with anti-HA antibody and following by Western blots with anti-His antibody. Protein A self-interaction was present in cells (Fig. 2C, lane 4), whereas no protein was immunoprecipitated with a control antibody anti-Flag (Fig. 2C, lane 5). Taken together, these results show that protein A can be self-interacted (homodimerized) in vitro and in cells.

Characterization of the Fragments Responsible for WhNV Protein A Self-interaction and the Homotypic and Heterotypic Interactions among these Fragments

We sought to determine the fragments required for protein A self-interaction. Thus, a series of MBP-protA fragments were produced according to the hydrophobicity of protein A amino acid (aa) sequences (Fig. 3A) and then used to pull down His-protA (Fig. 3B). The self-interaction efficiency of these protein A fragments was measured as the percentages of the self-interaction of FL to FL protein A (i.e., MBP-protA pulls down His-protA; Fig. 3B, lane 1; Fig. 3C, right, “FL”). We found that multiple fragments were required for protein A self-interaction. MBP-protA fragments aa 1–254, aa 255–480, and aa 1200–1314 did not contribute to protein A self-interaction (Fig. 3B, lanes 13–15; Fig. 3C, right). The elongation of aa 1–254 to aa 480, aa 659, or aa 839 resulted in an increase in self-interaction to a level comparable to that of FL to FL protein A self-interaction (94%, 96%, and 108%, respectively; Fig. 3B, lanes 3–5; Fig. 3C, right). On the other hand, the elongation of aa 255–480 to aa 659, aa 839, or aa 1014 did not further affect protein A self-interaction (Fig. 3B, lanes 7–9; Fig. 3C, right). Similar results were observed
when aa 481–659 was elongated to aa 839 or aa 1014 (Fig. 3B, lanes 11 and 12; Fig. 3C, right). Taken together, we conclude that aa 1–254 and aa 255–480 are sufficient to mediate protein A self-interaction.

Given that protein A self-interaction is mediated at least by two distinct fragments, we also sought to determine whether the self-interaction is formed by homotypic (i.e., aa 1–254/1–254 and aa 255–480/255–480) and/or heterotypic (i.e., aa 1–254/255–480) interactions of these two fragments of protein A. To that end, we assessed the potential homotypic and heterotypic interactions using MBP-protA fragments aa 1–254, aa 255–480, and aa 481–659 to pull down in vitro translated His-protA fragments aa 1–254, aa 255–480, and aa 481–659, respectively. Various homotypic and heterotypic interactions were detected (Fig. 3D), and the results were graphed as the percentages of the FL to FL protein A self-interaction (Fig. 3E). The homotypic interactions of aa 1–254 and aa 255–480 were 73% and 55%, respectively, of the level of FL protein A self-interaction, whereas the homotypic interaction of aa 481–659 was very weak (2%). Heterotypic interactions between aa 1–254 and aa 255–480 were also detected (50–52%). Interestingly, although the homotypic self-interaction of aa 481–659 was minimal, the heterotypic interactions of this fragment with aa 1–254 and aa 255–480 were relatively substantial (30% and 22%, respectively), thereby implying that the aa 481–659 fragment mediates protein A self-interaction via facilitating heterotypic interactions (Fig. 3B). Taken together, these results show that both homotypic and heterotypic interactions of protein A fragments exist and act together to mediate protein A self-interaction.

Mitochondrial Membrane Lipids Stimulate WhNV Protein A Self-interaction

Subsequently, we examined the direct effect of MMLs on the self-interaction of protein A under the same conditions described in Fig. 2A with the addition of 2 μg/μl MMLs. In the presence of...
MMLs, the ability of protein A to self-interact was substantially increased (Fig. 4A, lane 4), whereas MMLs had no effect on MBP alone (Fig. 4A, lane 2). To further confirm the stimulating effect of MMLs on protein A dimerization, we conducted a dose-response assay (Fig. 4B). As the concentration of MMLs increased, the self-interaction of protein A was gradually enhanced (Fig. 4B, “Bound”). Protein A self-interaction was stimulated about 4.6-fold at an MML concentration of 1 μg/ml, about 9-fold at an MML concentration of 2 μg/ml, about 12-fold at an MML concentration of 5 μg/ml, and then plateaued at an MML concentration of 10 μg/ml (Fig. 4C). Together, these results confirmed that MMLs promoted protein A self-interaction.

Mitochondrial Membrane Lipids Stimulate WhNV Protein A Self-interaction by Promoting the Homotypic and Heterotypic Interactions of Protein A

After identifying the stimulating effects of MMLs on protein A self-interaction and the fragments responsible for protein A self-interaction, we next attempted to determine whether these fragments are responsible for protein A’s binding to MMLs. To this end, we incubated various protein A fragments in the presence or absence of MMLs and then subjected them to Nycodenz gradient flotation assays to examine their MML association (Fig. 5A). MBP alone was used as the negative control, thereby ruling out the possibility that MBP induces protein-MML interaction. The flotation gradients were divided into two

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**Figure 3.** Characterization of the fragments responsible for WhNV protein A self-interaction and the homotypic and heterotypic interactions among these fragments. (A) Potential hydrophobic regions of WhNV protein A. (B) MBP-tagged protein A fragments (1 μM each) were used to pull-down FL His-protA (1 μM). The sizes of the molecular weight markers are indicated on the left in thousandths. (C) Summary of MBP fusion proteins and their activities to interact with His-protA, representing the results shown in (B). The self-interaction efficiency of protein A fragments was measured as the percentage of protein A FL self-interaction. ND, not detected. FL, full-length. (D) The self-interacting fragments form homotypic and heterotypic interactions. MBP-tagged protein A fragments (1 μM each) were used to pull-down His-tagged protein A fragments (1 μM each). (E) Summary of the homotypic and heterotypic interactions of protein A, representing the results shown in (D).

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interaction Activity

Characterization of the Stimulating Effect of Mitochondrial Membrane Lipids on Protein A Self-interaction Activity

Because WhNV protein A is a membrane binding protein, it is possible that protein A interacts with another protein A via a common lipid “bridge”. To test this possibility, we sought to define the sites critical for protein A self-interaction, mutate these regions without affecting the MML binding of protein A, and then determine the self-interaction of these mutants in the absence or presence of MMLs. To this end, amino acid substitutions were introduced into the aa 1–254 and were expressed in nuclease-treated RRLs; and in each mutant, the original amino acid was changed to alanine. We constructed multiple single-site mutations spanning aa 1–254; however, the triple-sites mutations completely lost their self-interacting activities. Then two aa 1–254 mutants, aa 1–254/M1 (K91A, W92A, and R93A) and aa 1–254/M2 (S163A, R165A and Y169A) were used to test their abilities to binding to MMLs and self-interact in the presence of MMLs, respectively. As shown in Fig. 6, aa 1–254/M1 and aa 1–254/M2 still contain the MML-binding property as being determined by Nycodenz gradient centrifugation (Fig. 6A), but completely lost their self-interacting activities (Fig. 6B, lanes 2 and 6). Furthermore, our result showed that the self-interactions of these mutants were unable to be stimulated by MMLs at various concentrations (Fig. 6B, lanes 3–5 and 7–9).

Furthermore, we used another MML binding protein, FHV protein A, to test if FHV protein A could also interact with WhNV protein A via the possible “bridging” effect of MMLs. MBP-protA was used to pull-down His-tagged FHV protein A (His-protA_FHV) that was expressed in nuclease-treated RRLs in vitro. The interactions between WhNV protein A with MBP and His tags were used as the positive control (Fig. 6C, lane 1). As shown in Fig. 6C, MBP-protA can not interact with His-protA_FHV in the absence of MMLs (lane 2), showing that WhNV protein A and FHV protein A have no direct protein-protein interaction. The presence of 2 μg/μl MMLs resulted in a very weak interaction of these two proteins (compared lane 3 to lane 1, the positive control). However, after increasing the MML concentrations (Fig. 6C, lane 3–5) or increasing the His-protA_FHV concentrations (Fig. 6C, lanes 6–8) showed no stimulating effect on the weak indirect protein A WhNV-protein A_FHV interaction. These results indicate that binding to common lipid may contribute to but could not be the major cause for the stimulation on protein A self-interaction, since the indirect interaction through binding to MMLs is much weaker than the protein-protein

Figure 4. MMLs stimulate WhNV protein A self-interaction. (A) Protein A self-interaction is increased by MMLs. MBP-protA (lanes 3 and 4, 1 μM) or MBP alone (lanes 1 and 2) was used to pull down the His-protA (1 μM) in the absence (lanes 1 and 3) or in the presence (lanes 2 and 4) of 2 μg/μl MMLs, and then subjected the pull-down products to Western blotting with anti-His antibody. (B–C) MMLs stimulate protein A self-interaction in a dose-response manner. Increasing concentrations (wt/vol) of MMLs were incubated with MBP-protA and His-protA (1 μM each). The concentrations of MMLs are indicated above each lane. The self-interaction of protein A in the absence of MMLs is used as the control (1-fold). The increases in the self-interaction of protein A at each point concentration of MMLs are graphed as the fold of the control as shown in (C). Error bars represent S.D. values from at least three independently repeated experiments and the represent results were shown in (B).

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Figure 5. MMLs stimulate WhNV protein A self-interaction by promoting the homotypic and heterotypic interactions of protein A. (A) MBP-tagged protein A fragments were incubated without (left) or with the MMLs (right) and subjected to Nycodenz flotation. The LD and HD fractions were analyzed via Western blotting with anti-MBP antibody. (B–C) The effects of MMLs on different homotypic (B) and heterotypic (C) interactions of protein A.
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interaction and can not be further enhanced by increasing the concentrations of MMLs or protein.

**Specific Anionic Phospholipids Stimulate Protein A Self-interaction**

MMLs are composed of various specific phospholipids [7]. The various phospholipid compositions of intracellular membranes are the key determinants of the activities of membranes as well as membrane-associated proteins [7]. Thus, we further analyzed the self-interaction activity of protein A with liposomes that made of individual major outer mitochondria membrane phospholipids. A series of dose-response assays were performed to determine the effect of distinct liposomes on the self-interaction of protein A (Fig. 7A). And the data was graphed as the fold of the self-interaction of protein A without lipids. As shown in Fig. 7B, protein A self-interaction was substantially stimulated in the presence of increasing concentrations of CL and PA. PG and PS stimulated protein A self-interaction moderately, whereas PC and PE did not affect protein A self-interaction. These results indicate that protein A self-interaction is selectively stimulated by specific anionic phospholipids.

Specific Anionic Phospholipids Favor Different Types of Self-interactions of Protein A aa 1–254 and aa 255–480

Having shown that homotypic and heterotypic interactions exist during protein A self-interaction and that specific anionic phospholipids stimulate protein A self-interaction at various levels, we hypothesized that the homotypic and heterotypic interactions of protein A could be differentially mediated by specific anionic phospholipids. To test this hypothesis, we assessed the effects of various anionic phospholipids on the homotypic interactions of aa 1–254 and the heterotypic interactions of aa 1–254 and aa 255–480. MBP pull-down assays were conducted in the presence of liposomes containing increasing concentrations of CL, PG, or PS (Fig. 8A and B). Increases in the levels of homotypic or heterotypic interactions at different concentrations of various liposomes were measured and graphed as the fold of interactions without lipids (Fig. 8C). Interestingly, our results revealed the different levels of homotypic and heterotypic interactions at different concentrations of various liposomes. As shown in Fig. 8C, as the liposome concentration increased, CL gradually favored the homotypic interactions (black bar is gradually higher than gray bar at each point concentrations of CL). According to PG, the homotypic and heterotypic interactions show no significant differences.

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*Figure 6. Characterization of the stimulating effect of MMLs on protein A self-interaction activity.* (A) The in vitro translation His-protA fragments aa 1–254/M1 (K91A, W92A, and R93A) and aa 1–254/M2 (S163A, R165A and Y169A) were incubated without (left) or with the MMLs (right) and subjected to Nycodenz flotation. (B) MBP-protA fragment aa 1–254 was used to pull-down His-protA fragments aa 1–254/M1 (lanes 2–5) and aa 1–254/M2 (lanes 3–8) in the increasing concentrations of MMLs (lanes 3–5 and 7–8). The concentrations of MMLs are indicated above each lane. The wt protein A fragment aa 1–254 was used as the control (lane 1). (C) MBP-protA was used to pull-down 1 μM His-protArev at the increasing concentrations of CL (lanes 3–5), or increasing concentrations of His-protArev at the 2 μg/μl MMLs (lanes 6–8). The concentrations of His-protArev and MMLs are indicated above. The WNV protein A self-interaction was used as the control (lane 1).

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Figure 7. Specific anionic phospholipids stimulate protein A self-interaction. (A–B) MBP pull-down was performed on the increasing concentrations (wt/vol) of liposomes generated from specific purified phospholipids with MBP-protA-His-protA complex. The concentrations of liposomes are indicated above each lane. The self-interaction of protein A in the absence of liposomes is used as the control (1-fold). The increases in the self-interaction of protein A at each point concentration of liposomes are graphed as the fold of the control as shown in (E). Error bars represent S.D. values from at least three independently repeated experiments.

Manipulation of Phospholipid Metabolism Affects Protein A-induced RNA Replication and Self-interaction in Cells

To further investigate the effects of MMLs, particularly changes in MMLs, on the functions of protein A in cells, we aimed to manipulate phospholipid synthesis in Pr-E cells to assess protein A activity with regard to membrane association, self-interaction, and RNA1/sgrRNA3 replication in cells.

We used PA inhibitor FIPI to down-regulate PA in cells [43] because PA is a precursor in the CDP-DAG pathway [7]. Pr-E cells were treated with 75 nM FIPI, which inhibits PA production efficiently and show little negative effect on cells [43]. FIPI treatment yielded a 40% reduction in cellular levels of PA (Fig. 9A). The incomplete blockage of PA production was likely due to the presence of a de novo PA synthesis pathway [6,7]. Moreover, we also assessed cell viability and found that FIPI minimally affected cell viability (~10% reduction), which was comparable with the effect of the vehicle DMSO (Fig. 9B). The effect of FIPI treatment on mitochondrial associated protein was also assessed via the detection of porin protein, which is an integral membrane protein associated with mitochondria. Our results show that FIPI minimally affected the porin expression (Fig. 9C, left). Moreover, we further determined that FIPI treatment was unable to alter the membrane association of porin via Nycodenz flotation assay (Fig. 9C, right), thereby ruling out the possibility that FIPI treatment can damage the property of mitochondrial membranes to associate with membrane-bound proteins.

Furthermore, we assessed the membrane association of protein A via Nycodenz flotation assay. WhNV protein A was expressed via transfection with plasmid Pa. As shown in Fig. 9D, left, FIPI treatment did not alter the activity of protein A to associate with membranes. Moreover, we examined whether inhibiting PA affects the initial transcription from input plasmid. As shown in Fig. 9D, right, the initial transcription from the input plasmid pAC1E was almost the same in cells with or without FIPI treatment.

We examined the effects of MML manipulation on WhNV RNA replication using WhNV trans-replication system (Material and Methods). To this end, the cells expressing protein A and RNA1E template were treated with or without FIPI. The accumulations of negative-strand (−)RNA1E, (+)RNA1E and (+)sgrRNA3E were determined by Northern blots. The accumulation of (−)RNA1E was only moderately reduced by about 15% in FIPI treated cells compared to that in non-treated cells (Fig. 9E, “(−)RNA1E”, compared lane 3 to lane 2 or 1; Fig. 9F), while the accumulation of (+)sgrRNA3E was reduced by about 60% (Fig. 9E, “(+sgrRNA3E)”, compared lane 3 to lane 2 or 1; Fig. 9F). Also, the FIPI treatment resulted in about 30% reduction in the accumulation of (+)RNA1E (Fig. 9E, “(+RNA1E)”, compared lane 3 to lane 2 or 1; Fig. 9F). Moreover, the FIPI treatment showed an apparent biased effect on sgrRNA3 production, suggesting that this step may be especially sensitive to inhibiting PA production. Besides, the self-interaction of protein A wt was also inhibited by about 60% via inhibiting PA production (Fig. 9E, “co-IP”, compared lane 3 to lane 2 or 1; Fig. 9F). Although (−)RNA synthesis was less affected, the levels of (+)RNA1 and (+)sgrRNA3 were still reduced by the FIPI treatment, indicating that at the similar level of the (−)RNA1 template, the activity of WhNV RdRBP to replicate (+)-stranded RNA products was indeed weakened.
Figure 8. Specific anionic phospholipids favor different types of self-interactions of protein A aa 1–254 and aa 255–480. (A–C) The homotypic interactions of aa 1–254 and the heterotypic interactions of aa 1–254 and aa 255–480 were examined in the presence of the increasing concentrations of liposomes generated from CL, PA, PG, or PS, respectively. The homotypic or heterotypic interactions in the presence of different liposomes at each concentration are graphed as shown in (C). The homotypic (gray bar) and heterotypic (black bar) interactions in the absence of liposomes are used as the control (1-fold). The increase in homotypic or heterotypic interactions in the presence of different liposomes at each concentration is graphed as the fold of control. Error bars represent S.D. values from at least three independent experiments and the represent results were shown in (A–B).

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Discussion

RNA replication of (+) RNA viruses requires the association of viral RNA and replicases with intracellular membranes to form vRCs [1–3]. To advance the understanding of the relationship between intracellular membranes and viral RNA replicases, we studied the direct effects of membranes, particularly membrane lipids, on the function of the replicase (protein A) from WHNV. We uncover the self-interaction of WHNV protein A and show that this activity of protein A could be stimulated by MMLs. Additional investigations show that MMLs interact with specific fragments of protein A, and this direct lipid-protein interaction may stimulate protein A self-interaction by promoting homotypic and heterotypic interactions of specific fragments. Moreover, the self-interaction of protein A could be selectively modulated by liposomes generated from specific anionic phospholipids, and specific anionic phospholipids favor different types of homotypic and heterotypic interactions. Furthermore, manipulating phospholipid metabolism via a PA inhibitor weakens protein A self-interaction and RNA replication in cells. Altogether, these findings demonstrate the direct role of membrane lipids in the activity of WHNV protein A.

Two mechanisms may be responsible for the stimulation on protein A self-interaction. One possibility is that MMLs directly mediate protein A activity. The changes in lipid composition may result in protein A’s property changes via altering protein A’s conformation. The other one is that MMLs partition protein A into liposome fraction and thus lead to the increase of protein A’s local density. Binding to common lipid may also contribute to the stimulation on WHNV protein A self-interaction (Fig. 6).

For many (+) RNA viruses, different patterns of protein-protein interactions of replicases are associated with the distinct functions for RNA replication. For example, the 3D polymerase of poliovirus was shown to homooligomerize via two interfaces, which may be related to different function [45,46]. Similarly, HCV RdRp changes its conformations to direct different function at the early stages of RNA replication [47,48]. The different homotypic and heterotypic interactions of WHNV protein A provide the direct evidence that the different protein-protein interaction interfaces exist and can be regulated by specific liposomes (Fig. 6). Such different patterns of self-interaction could also be seen from FHV protein A [28]. Although the function of the heterotypic and homotypic interactions is not known, it is possible that different interactions may serve to alter the structure.

Figure 9. Phospholipids affect the proper functioning of protein A. (A) Measurement of PA content in Pr-E cells or cells treated with 75 nM FIP or with matching concentration of DMSO (vehicle). (B) Viability of cells treated with FIP or DMSO. (C) FIP treatment show less effect on the activity of mitochondrial membrane-binding protein porin to associate with membranes. Left, cells treated with or without FIP were harvested and then probed via Western blotting with anti-porin antibody. Right, total RNAs was isolated from FIP treated cells expressing (+)RNA1E templates and then probed via Northern blotting with EGFP and 18s rRNA probes, respectively. (D) RNA accumulation in cells treated with FIP or DMSO expressing protein A–His/HA treated with FIP or DMSO (F). GAPDH, glyceraldehyde-3-phosphate dehydrogenase. Error bars represent S.D. values from at least three independent experiments and the represent results were shown in (E). The accumulation of RNA and protein is normalized to 18s rRNA and GAPDH, respectively.

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dimerization, or function of protein A, in successive step such as replication complex assembly, RNA replication, and RNA capping. Indeed, when PA production was inhibited in cells, the activity of WhnV protein A to replicate (+)sgRNA3E was preferentially inhibited; however, the synthesis of (-)RNA1E template was minimally affected (Fig. 9E). Because the replication of (-)RNA1E and (+)sgRNA3E are all mediated by protein A, such selective regulation by reducing PA production may be induced by the different homotypic and heterotypic interactions of protein A in response to the changes of membrane lipids.

Membrane lipids are comprised of distinct phospholipids, and the composition of these phospholipids is different for different membranes [7]. It is possible that certain lipids have different effects on (+)RNA replicases. Semliki Forest virus (SFV) localizes to lysosomes and endosomes and the capping activity of SFV NSP1 protein requires association with negative phospholipids PS [42]. Hepatitis C virus (HCV) localizes to membrane lipid rafts and the activity of HCV RdRP requires association with sphingomyelin [49]. According to nodavirus, FHV protein A membrane association [30] and WhnV protein A self-interaction (Figs. 7 and 8) can be mediated by specific anionic phospholipids CL, PA and PG, which are enriched in mitochondroid membranes [7]. In these cases, particular phospholipids enriched in certain intracellular membranes, which are associated with these viruses, show preferential and direct effects on the activities of replicases. However, some universal phospholipids being enriched in many intracellular membranes [7], could also mediate (+)RNA virus replication. For example, PC show less direct impacts on FHV protein A’s membrane association but mediate protein A function in some other ways [30,33]. These results suggest that the regulations of phospholipids on (+)RNA virus replicase activities could be manifold.

Nonionic detergent (Triton X-100) is preferred for the isolation of membrane proteins, as it assists in the solubilization of proteins from lipids. Then, we used it for the purification of protein A and MMLs. Although we did our best to get rid of the detergent, we can not ensure that all detergents were completely removed. The transformation between liposomes and detergent/lipid mixed micelles is a reversible process that can be induced by the addition or reduction of the concentrations of detergent [50]. Our observation that increasing the concentrations of MMLs enhanced the protein A self-interaction (Fig. 4B) revealed that the concentrations of the remaining detergent is much low or even neglectable. However, it is still possible that the remaining detergents may affect the protein-MML interactions and subsequently weaken the enhanced protein A self-interaction in the presence of MMLs.

Although the in vitro data reveals the obvious effects of membrane lipids on WhnV protein A self-interaction (Figs. 4–6), the cellular experiment data shows relatively minor effects (Fig. 9). That may be partly due to that the simplified in vitro systems containing only one purified protein and one or a few kinds of lipids, and do not represent the whole behavior of the protein A in replication in cells. In addition, the PA content still remained at 60% level compared to that in cells without FIP1 treatment, probably due to the presence of a de novo PA synthesis pathway [6,7]. The production of other anionic phospholipids could be less affected by the FIP1 treatment, and may even compensate the lipid loss in mitochondria. Moreover, it is possible that inhibiting the self-interaction of protein A indirectly weaker the viral RNA replication by such as affecting the microenvironment of vRCS or the binding of host factor to vRCS, rather than directly weakens the ability of per unit protein A to synthesize RNA. Furthermore, inhibition of the total cellular PA content in cells may not reflect the real effects of membrane lipids on protein A function. The protein A function can be mediated by multiple factors, such as total cellular lipids content, membrane lipids, and the protein A microenvironnement. Our future studies will focus on the effects of other factors, including host proteins or/and other lipids, on WhnV protein A complete activity.

In summary, our findings further reveal the detailed mechanisms by which direct MML-protein interaction regulates the self-interaction of nodaviral replicase protein A. Nodavirus RNA replication is highly parallel with that of other (+)RNA viruses, with regard to the formation of vRCS on host intracellular membranes, the requirement of homono- and/or hetero-oligomerization of viral replicase components (for nodavirus, it is self-interaction of multiple fragments within the single replicase), and viral RNA replication-associated alterations in the composition of MMLs [28,29,33,51–53]. Considering the commonalities that exist between nodaviruses and other (+)RNA viruses in RNA replication, some of the principles revealed in this study may be generally applicable to a range of (+)RNA viruses.

Supporting Information

Figure S1 Detection of the purified outer mitochondrial membranes. The purified outer mitochondrial membranes (OMM) and intact mitochondrial (Mito) was subjected to Western blotting with anti-porin, anti-Tim 23 and anti-Calreticulin, respectively. Tim 23, an inner mitochondrial membrane protein. Calreticulin, an endoplasmic reticulum membrane protein. (TIF)

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Author Contributions

Conceived and designed the experiments: YQ XZ. Performed the experiments: YQ ZW YL YH MM NQ JY HX XL CQ. Analyzed the data: YQ ZW YL YH XZ. Contributed reagents/materials/analysis tools: YQ LX CQ. Wrote the paper: YQ XZ.

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