Effective inhibition of HCMV UL49 gene expression and viral replication by oligonucleotide external guide sequences and RNase P

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Abstract
Background: Human cytomegalovirus (HCMV) is a ubiquitous herpesvirus that typically causes asymptomatic infections in healthy individuals but may lead to serious complications in newborns and immunodeficient individuals. The emergence of drug-resistant strains of HCMV has posed a need for the development of new drugs and treatment strategies. Antisense molecules are promising gene-targeting agents for specific regulation of gene expression. External guide sequences (EGSs) are oligonucleotides that consist of a sequence complementary to a target mRNA and recruit intracellular RNase P for specific degradation of the target RNA. The UL49-deletion BAC of HCMV was significantly defective in growth in human foreskin fibroblasts. Therefore, UL49 gene may serve as a potential target for novel drug development to combat HCMV infection. In this study, DNA-based EGS molecules were synthesized to target the UL49 mRNA of human cytomegalovirus (HCMV).

Results: By cleavage activity assessing in vitro, the EGS aimed to the cleavage site 324 nt downstream from the translational initiation codon of UL49 mRNA (i.e. EGS324) was confirmed be efficient to direct human RNase P to cleave the target mRNA sequence. When EGS324 was exogenously administered into HCMV-infected human foreskin fibroblasts (HFFs), a significant reduction of ~76% in the mRNA and ~80% in the protein expression of UL49 gene, comparing with the cells transfected with control EGSs. Furthermore, a reduction of about 330-fold in HCMV growth were observed in HCMV-infected HFFs treated with the EGS.

Conclusions: These results indicated that UL49 gene was essential for replication of HCMV. Moreover, our study provides evidence that exogenous administration of a DNA-based EGS can be used as a potential therapeutic approach for inhibiting gene expression and replication of a human virus.

Background
Human cytomegalovirus is a ubiquitous herpesvirus that typically causes asymptomatic infections in healthy individuals but may lead to serious complications in newborns and immunodeficient individuals. For example, this virus accounts for one of the most common opportunistic infections in AIDS patients (i.e., CMV retinitis). Moreover, HCMV infection is the leading viral cause of birth defects in newborns and a major cause of morbidity and mortality in bone marrow and solid organ transplant recipients [1]. The emergence of drug-resistant strains of HCMV has posed a need for the development of new drugs and treatment strategies [2]. Antisense molecules are promising gene-targeting agents for specific regulation of gene expression [3]. Conventional antisense oligonucleotides have been used as anti-HCMV agents to inhibit the expression of HCMV-essential genes and abolished viral replication [4,5]. External guide sequences (EGSs) are antisense oligonucleotides that can be used in conjunction with ribonuclease P (RNase P) or the catalytic RNA subunit of RNase P from Escherichia coli (M1 RNA) to diminish gene expression [6-9]. RNase P is one of the most abundant and active enzymes in cells and is responsible for 5’ termini maturation of tRNAs [10]. This enzyme catalyzes a hydrolysis reaction to remove the 5’ leader sequence of tRNA precursors (ptRNA) by recognizing the common...
structure shared among all tRNAs (Fig. 1A). The EGS-based technology is unique in inducing endogenous RNase P to cleave a target mRNA when the EGS hybridizes to the mRNA to form a structure resembling a tRNA substrate (Fig. 1B). This approach is highly specific and does not generate nonspecific "irrelevant cleavage" that is observed in RNase H-mediated cleavage induced by conventional antisense phosphothioate molecules [11]. Thus, EGSs represent a new class of agents that may lead to highly effective and specific inhibition of gene expression.

RNA-based EGSs have been expressed endogenously as transgenes in both bacteria and mammalian cells [12], and were effective in inhibiting the gene expression of herpes simplex and influenza virus and in abolishing the replication of influenza virus in human cells [13,14]. In vitro studies have also shown that DNA-based EGSs, as well as EGS molecules with modified nucleotides, can direct M1 RNA or human RNase P to cleave a mRNA sequence, although their targeting efficiencies are lower than those of unmodified RNA-based EGSs [15]. However, little is known about whether DNA-based EGSs are functionally active in cultured cells. Also, whether DNA-based EGSs can be exogenously administered into human cells to inhibit gene expression and growth of human viruses is yet to be determined.

The UL49-deletion BAC of HCMV was significantly defective in growth in human foreskin fibroblasts [16]. Furthermore, the expression of HCMV UL49 protein has been shown in our mock infected human foreskin fibroblasts (unpublished data, doctoral dissertation). Therefore, UL49 gene may serve as a potential target for novel drug development to combat HCMV infection. In this study, we provide direct evidence that exogenous administration of chemically synthesized DNA-based EGS is highly effective in inhibiting HCMV gene expression and growth in human cell culture and, furthermore, demonstrate the feasibility of using DNA-based EGSs for the studies and treatment of infections caused by human viruses including HCMV.

Materials and methods

Construction of the RNA substrate and EGSs

The DNA sequence that encodes HCMV UL49 gene was constructed by PCR using AD169 genomic DNA as a template and oligonucleotides 49F: 5'-AGTTGGATCCTGCGCTCTC-3' and 49R: 5'-ATCTAAGCTTGTAGACATGGGGCAGGCCGT-3' as 5' and 3' primers, respectively. The PCR fragments were digested with BamHI and Hind III, and then inserted into vector pGEM3z (Invitrogen) to form a recombinant plasmid, UL49-pGEM3z. The RNA substrates were synthesized by bacteriophage T7 RNA polymerase using the Hind III-linearized UL49-pGEM3z plasmid as transcriptional templates and labeled with 32P-UTP. The restriction endonucleases, LA Taq polymerase, T4 DNA ligase and T7 RNA polymerase were purchased from Takara (Dalian, China). All oligonucleotides used as PCR primers were purchased from Invitrogen (Guangzhou, China). EGS oligonucleotides used in this report including EGS324(5' CGACACACTTGGGGTTTCCCCACGCAGGT TCGAATCTCGCCATCCCA3'), EGS1539(5'GTCGCGGT TGGGGTTTCCCCACGCAGGT TCGAATCTCGCCACGTCCCA3'), C-EGS324(5' CGACACACTTGGGGTTTCCCCACGCAGGAATCCTGCG CCA TTCCCA3'), C-EGS1539(5'GTCGCGGT TGGGGTTTCCCCACGCAGGAATCCTGCG CCA TTCCCA3'), and TK-EGS(5'TACGTCGGTG GTCTCCGCGCGCAGGTTCAAATCCTGCCGCAGAC CA3') were chemically synthesized directly by using a DNA synthesizer.

Purification of human RNase P

RNase P was purified from HeLa cells according to the method as described [17] with the following modifications. The DEAE-Sepharose Fast-flow column was equilibrated and washed with buffer A containing 10 mM Tris-
were incubated with human RNase P (5 units). The cleavage reactions were carried out at 37°C for 30 min in buffer A (50 mM Tris-HCl, pH 7.5, 100 mM NH₄Cl, 10 mM MgCl₂). To disrupt aggregates, which might form during storage, EGS and substrate together were heated to 90°C for 1 min and then cooled to room temperature before the addition of other components. Substrate and cleavage products were separated under denaturing conditions on a 15% acrylamide gel containing 8 M urea. The amount of radioactivity per band was quantified using a Typhoon 9200 phosphorimager. The procedures to measure the equilibrium dissociation constants (K_d) of the EGS-UL49 mRNA complexes were carried out as described [18]. In brief, various concentrations of EGSs were preincubated in buffer B (50 mM Tris, pH 7.5, 100 mM NH₄Cl, 20 mM MgCl₂, 3% glycerol, 0.1% xylene cyanol, 0.1% bromphenol blue) at 37°C for 10 min before mixing with an equal volume of different concentrations of ³²P-labeled substrate RNA preheated under identical conditions. The samples were incubated for 15 min to allow binding, loaded on a 5% polyacrylamide gel, and run at 10 watts. The electrophoresis running buffer contained 100 mM Tris-Hepes, pH 7.5, and 10 mM MgCl₂. Each band was quantitated with a Typhoon 9200 phosphorimager. The value of K_d was then extrapolated from a graph plotting percentage of product bound versus EGS concentration. The values were the average of three experiments.

**In vitro cleavage studies**

The EGSs (50 nM) and ³²P-labeled UL49 mRNA (50 nM) were incubated with human RNase P (5 units). The cleavage reactions were carried out at 37°C for 30 min in buffer A (50 mM Tris-HCl, pH 7.5, 100 mM NH₄Cl, 10 mM MgCl₂). The EGS-lipid mixtures were prepared according to the manufacturer’s recommendation (GIBCO) and incubated with cells for 7 hrs. HCMV (strain AD169) was propagated in human foreskin fibroblasts in DMEM supplemented with 10% FBS (GIBCO). Cells (1 × 10⁶) were treated with liposome complexes in the absence and presence of EGS, followed by HCMV infection or mock infection at a MOI of 1 in an inoculum of 1.5 ml DMEM supplemented with 1% fetal calf serum (FCS). Total cellular RNAs and proteins were isolated from cells as described 72 hrs postinfection [19].

**Fluorescence Quantitative RT-PCR**

Fluorescent Quantitative RT-PCR (FQ-RT-PCR) was performed using standard protocols on an Applied Biosystem’s 7300 Sequence Detection System. Briefly, total RNAs were extracted using Trizol, then 5 ml of a 1/100 dilution of cDNA in water was added into 12.5 ml of the 2 × SYBR green PCR master mix (Takara), with 800 nM of each primer in a total volume of 25 ml. All reactions were run in triplicate using Applied Biosystem’s 7300 Sequence Detection System. UL49 gene sense primer: 5’-CGTCTCTTCGTCCTTCATCTC’-3’, anti-sense primer: 5’-CACAAAGTGAGGCTTGGTGCAT-3’. As internal standard, a fragment of human endogenous β actin was amplified simultaneously in each PCR. β actin sense primer: 5’-TCGTCCACCGAAATGCTTCTAG-3’, anti-sense primer: 5’-ACTGCTGTCACTTTCACCGTTC-3’. Quantitative RT-PCR conditions: 95°C for 10 s, 1 cycle, followed by 95°C for 5 s, 60°C for 20 s, 40 cycles for amplifying UL49 and β actin.

**Western blotting**

For Western analyses, the denatured, solubilized poly-peptides were separated on 12% (vol/vol) SDS polyacrylamide gels cross-linked with N, N”-methylenebisacylamide, and then transferred onto PVDF membranes (Amersham, Freiburg, Germany). The membranes were incubated in the primary antibody solution (including antibody against human β actin, ul49, IE1/IE2, pp28, or gB), then were incubated in a goat anti-mouse HRP-conjugated antibody solution. After washing with PBS containing 0.05% Tween 20, the membranes were subsequently stained with a chemiluminescent substrate with the aid of a Western chemiluminescent substrate kit (Pierce, Rockford) and quantitated with a Typhoon 9200 phosphorimager [20]. The monoclonal antibodies against HCMV UL49 protein were kindly provided by Doctor Feng Zhu (BOAOSEN, Beijing). The monoclonal antibodies against human β actin was purchased from SIGMA.

**Assay of viral growth in EGS-treated cells**

To determine the level of the inhibition of viral growth, 5 × 10⁵ human foreskin fibroblasts were first incubated with liposome complexes in the absence and presence of EGSs, and then mock-infected or infected with HCMV.
AD169 at a MOI of 1. The cells and medium were harvested at 1, 2, 3, 4, 5, 6, and 7 days postinfection. The viral stocks were prepared by adding an equal volume of 10% skim milk, followed by sonication. The titers of the viral stocks were determined by infecting 2 × 10^5 foreskin fibroblasts in 6-well plates and counting the number of plaques 7~10 days postinfection. The values obtained were the average from triplicate experiments.

**Results**

**Design of EGSs**

To achieve optimal cleavage, it is critical to choose the target regions of the UL49 mRNA. According to general principles of designing EGSs [21], two positions, 324 nt and 1539 nt downstream from the translational initiation codon, were chosen as the cleavage sites for EGSs. The flanking sequence of these cleavage sites exhibited several sequence features that need to be present in order to interact with an EGS and RNase P to achieve efficient cleavage. These features include that ? the nucleotides 3′ and 5′ adjacent to the site of cleavage are a guanosine and a pyrimidine, respectively, and ? a U is 8 nt downstream from this cleavage site. The interactions of these sequence elements with the EGS facilitate the formation of the mRNA-EGS complex into a tRNA-like structure while those with RNase P are critical for recognition and cleavage by the enzyme [22]. On the basis of the target sites, two EGSs (EGS324 and EGS1539) were constructed respectively, which resemble a part of the tRNA structure, containing a T-loop, a stem, and a variable region but not the anticodon region (Fig. 1C and 1E). The anticodon domain has been shown to be dispensable for EGS activity. Binding of EGSs to the target mRNA results in a helix structure of 7 bp that is equivalent to the D stem of a tRNA, which usually has 4 bp. Our choice of a D stem-equivalent structure of 7 bp instead of 4 bp is to increase the targeting sequence specificity of the EGS (Fig. 1C and 1E). The two control (C-EGS324 and C-EGS1539) were derived respectively from EGS324 and EGS1539 by introducing base substitution mutations in three positions of the T-loop (Fig. 1D and 1F). The nucleotides in these three positions are highly conserved among tRNA molecules and are important for the recognition of tRNA molecules by RNase P [23].

**In vitro studies of targeting activities of the EGSs**

The EGSs were subsequently incubated with human RNase P and substrate RNA, containing the targeted UL49 mRNA sequence of 1713 nt. In the absence of any EGS, no cleavage of the UL49 mRNA sequence was observed (Fig. 2, lane 1). Efficient cleavage of this substrate by RNase P was observed in the presence of EGS324 (Fig. 2, lane 2). In contrast, cleavage of the same substrate by RNase P was barely detected in the presence of C-EGS324 (Fig. 2, lane 3). It is possible that the differential cleavage efficiencies observed with EGS324 and C-EGS324 were due to their different binding affinities to the UL49 mRNA sequence. To investigate whether this was the case, the binding between the EGSs and substrate UL49 was studied using a gel-shift assay. In this assay, the EGSs were incubated with the substrate to allow for binding, and the EGS-UL49 mRNA complexes were separated in polyacrylamide gels under non-denaturing conditions. Similar amounts of complexes formed between these two EGSs and the UL49 mRNA sequence were observed when the same amount of EGSs was used (Fig. 3, lane 1 & lane 2). Further detailed assays under different concentrations of the EGS324 and UL49 mRNA indicated that the binding affinity of C-EGS324 to UL49 mRNA (K_d = 0.83 ± 0.12 μM) is similar to that of EGS324 (K_d = 0.82 ± 0.10 μM). Meanwhile, a very little amount of cleavage products was observed in the presence of C-EGS324 even under high concentrations of the EGS and RNase P and a prolonged incubation period (data not shown). These observations suggest that the mutations in C-EGS324 do not significantly affect the binding affinity of the EGS to the mRNA sequence but abolish its targeting activity to induce RNase P cleavage, possibly by disrupting the recognition of EGS-UL49 mRNA complex by RNase P. Thus, C-EGS324 may be used as a control for the antisense effect in our experiments in cultured cells (see below). Nevertheless, both EGS1539 and C-EGS1539 showed no activity in guiding RNase P to cleave the target mRNA (Fig. 2, lane 4 & lane 5). An additional EGS, TK-EGS, which was targeting the HSV-1 TK mRNA, was also constructed. This EGS was used to determine whether an EGS with an incorrect targeting sequence can direct human RNase P to cleave UL49 mRNA. No cleavage of UL49 mRNA by RNase P was observed in vitro in the presence of TK-EGS (Fig. 2, lane 6).

**Inhibition of HCMV UL49 expression in EGS-treated cells**

The chemically synthesized EGS molecules were complexed with Lipofectamine 2000 liposomes and delivered into human foreskin fibroblasts. Treatment of cells with the EGS-liposome complexes by using our transfection protocol consistently yielded a transfection efficiency of about 90% (data not shown). To investigate whether the internalized EGSs inhibit viral UL49 expression, the cells were treated with EGSs and then infected with HCMV at a moi of 1. The levels of viral UL49 mRNA were determined by quantitative RT-PCR analyses (Fig. 4). A reduction of 75.57 ± 3% in the level of UL49 mRNA expression was observed in cells treated with EGS324, whereas a reduction of less than 10% was observed in the cells treated with C-EGS324. No reduction in the expression...
lel of UL49 mRNA was observed in cells that were treated with liposome complexes in the absence of EGSs or in the presence of TK-EGS. These results suggest that the significant reduction of UL49 mRNA expression in the EGS324-treated cells was due to targeted cleavage by RNase P. The low level of inhibition observed in the C-EGS324 treated cells was presumably due to the antisense effects of the EGS.

To determine whether the inhibition of UL49 expression directed by EGS324 is dose-dependent, the EGS324 were transfected into HFFs at different concentrations (2.5, 5, 10, 20, 40, 80, and 160 nM), followed by infection with HCMV at a moi of 1. At 72 hrs postinfection, total RNAs were extracted using Trizol reagent, and then quantitative RT-PCR were performed. The quantitative RT-PCR showed that when the concentration of EGS324 was 2.5 nM, the inhibition efficiency was 15%; and then 5 nM was 22%, 10 nM was 31%, 20 nM was 43%, 40 nM was 51%, 80 nM was 75%, and 160 nM was 76.6% (Fig. 5). The results showed that the inhibition efficiency of EGS324 was dose-dependent.

The expression level of UL49 protein was also determined by Western analyses (Fig. 6). A reduction of 80% ± 3% in the expression level of UL49 protein was observed in cells treated with EGS324, whereas a reduction of less than 10% was found in the C-EGS324 treated cells.

Inhibition of UL49 expression is not expected to affect the expression of other viral genes

It is possible that the observed reduction of UL49 expression in the EGS324-transfected cells is not necessarily due to specific EGS-directed RNase P cleavage of the target mRNA, but is due to other effects of the EGS on viral lytic replication, such as blocking the expression of viral immediate-early genes. To exclude these possibilities and further determine the antiviral mechanism of the EGS-directed cleavage, we examined the expression of other viral genes in the EGS324-treated cells. Relative RT-PCR analyses were carried out to determine the mRNA levels

Figure 2 Cleavage of the 32P-labeled substrate by human RNase P in the presence of different EGSs. No EGS was added to the reaction mixture in lane 1. Cleavage reactions were carried out in the presence of EGS324 (lane 2), C-EGS324 (lane 3), EGS1539 (lane 4), C-EGS1539 (lane 5) or TK-EGS (lane 6).

Figure 3 Binding of UL49 mRNA substrate by EGS324 and C-EGS324. Substrate (10 nM) was either in the presence of 10 nM of EGS324 (lane 1), C-EGS324 (lane 2), or TK-EGS (lane 3) to allow binding and then loaded on a 5% polyacrylamide gel.
of an immediate-early gene (IE) and an early-late gene (UL44). Moreover, Western analyses were performed to determine the levels of viral protein IE2, a viral immediate-early (α) protein, gB, a viral early (β) protein, and pp28, a viral late (γ) protein. No significant differences in the expression levels of these genes were observed in cells that were treated with liposome complexes in the absence of EGS or in the presence of TK-EGS, EGS324, or C-EGS324 (Table 1). These results suggest that EGS324 specifically inhibits the expression of UL49 and does not affect overall viral gene expression.

Inhibition of HCMV growth in the EGS-treated cells

The impact of UL49 inhibition by EGS on viral growth was further investigated. Cells were treated with liposome complexes in the absence and presence of EGSs and then infected by HCMV at an moi of 1. Virus stocks were prepared from the infected cultures at 1-day intervals through 7 days postinfection. The count of plaque-forming units (PFU) was determined by measurement of the viral titer on cells. After 4 days postinfection, a reduction of about 330-fold in viral yield was observed in the EGS324-treated cells, whereas no significant reduction was found in those that were either treated with C-EGS324 or TK-EGS (Fig. 7). Thus, EGS324 is effective in inhibiting HCMV infection and blocking viral growth.

Discussion

Compared to other nucleic acid-based gene interference approaches, the EGS technology with the use of endogenous human RNase P exhibits several unique and attractive features as a gene targeting tool. First, the mechanism of the EGS technology for degradation of a specific mRNA is different from other RNA- or DNA-based gene-targeting approaches. It uses the endogenous RNase P, which is one of the most ubiquitous, abundant, stable and efficient enzymes in all type of cells. This essential enzyme is highly expressed (5 × 10⁴ copies per cell) and is responsible for the processing of all tRNA precursors that account for approximately 2% of total cellular RNA[24]. The action of RNase P with the EGS will result in irreversible cleavage of the target mRNA in a highly efficient catalytic fashion. Second, the sequence specificity of the EGS technology is governed by two different types of interactions between the EGS and the target mRNA: the base-pairing interactions in which the sequence of 12 nucleotides in the EGS hybridizes with the target mRNA, and the interactions between the target mRNA and the other part of the EGS sequence (equivalent to the T-stem and T-loop, and variable regions of a tRNA) which are required for folding of the RNase P-recognizable tertiary structure. Thus, the EGS-based technology is highly specific and does not generate nonspecific "irrelevant cleavage" that is observed in RNase H-mediated cleavage induced by conventional antisense phosphorothioate molecules [25].

Our study used exogenous administration of DNA-based EGSs for antiviral applications and demonstrated that RNase P-mediated targeting directed by DNA-based EGS was highly effective in inhibiting HCMV gene expression and viral growth in cultured cells. We showed that these EGS molecules directed human RNase P to cleave the UL49 mRNA sequence efficiently in vitro. Moreover, we also showed that these EGSs were readily delivered in cultured cells. A reduction of about 76–80%
in the UL49 expression was achieved with a functional EGS, EGS324, whereas a reduction of less than 10% was observed in cells that were treated with C-EGS324 or TK-EGS. C-EGS324 bound to UL49 mRNA substrate \textit{in vitro} as well as EGS324 but contained nucleotide mutations that disrupted RNase P recognition. These results suggested that the overall observed inhibition with EGS324 was primarily due to targeted cleavage by RNase P as opposed to the antisense effect or other nonspecific effects of the EGSs. Moreover, the antiviral effect of the EGS treatment (inhibition of viral growth) appeared to be due to the reduction of the UL49 expression. Only the expression of UL49 mRNA and protein was found to be reduced in cells treated with EGS324. We found no significant change in the expression of IE2, UL44, pp28, and gB (Table 1). The observed level of inhibition of UL49 expression was consistent with the extent of the reduction of viral growth.

HCMV is a member of the human herpesvirus family, which includes seven other different viruses such as HSV and Epstein-Barr virus. UL49 is well conserved and necessary for the growth of this virus [16]. Therefore, UL49 is considered as an ideal target for antiviral therapy. EGSs containing modified oligonucleotides are required to increase their stability \textit{in vivo} [26,27]. It has been recently reported that chemically synthesized RNA-based EGSs with modified nucleotides can be administered exogenously into human cells and inhibit cellular gene expression [28]. Our study showed that exogenous administration of DNA-based EGSs is highly efficient in inhibiting gene expression and viral growth. Further understanding of how the functional groups in the nucleotides of an EGS interact with human RNase P and the mRNA substrate will lead to the construction of highly active and stable EGSs with either different bases or modifications at these nucleotide positions. Moreover, engineering different designs of EGSs [29] for increasing their targeting activity, as well as developing new means for improving their delivery, are needed to increase the efficacy of the EGSs \textit{in vivo}. These studies will facilitate the

![Figure 6](image6.png)  
**Figure 6** In Western analyses (A&B), protein samples were separated in two identical SDS-polyacrylamide gels and transferred electrically to two identical membranes. One membrane was allowed to react with an anti-ul49 antibody (A), whereas the other was stained with the monoclonal antibody (Anti-actin) against human actin (B).

![Table 1](image7.png)  
**Table 1:** Inhibiting rates of the mRNA and protein expression of different viral genes in cells.

| Viral genes     | HFFs | EGS324  | C-EGS324 | TK-EGS |
|-----------------|------|---------|----------|--------|
| UL49 mRNA       | 0%   | 76% ± 3% | 8%       | 3%     |
| IE2 mRNA        | 0%   | 0%      | 0%       | 0%     |
| UL44 mRNA       | 0%   | 0%      | 0%       | 0%     |
| UL49 protein    | 0%   | 80% ± 3% | 6%       | 5%     |
| IE2 protein     | 0%   | 0%      | 0%       | 1%     |
| Glycoprotein B  | 0%   | 2%      | 2%       | 1%     |
| pp28 protein    | 0%   | 1%      | 2%       | 1%     |

The values shown are the means from triplicate experiments, and values of standard deviation that were less than 5% are not shown.
development of the EGS-based technology for gene-targeting applications in both basic research and clinical therapy of HCMV infections.

Competing interests
The authors declare that they have no competing interests.

Authors' contributions
WJZ participated in gene cloning, sequence alignment, transfection, data analysis and drafting of the manuscript. YQL initially conceived of the study. ZF helped with PCR amplification on many of the sequences; XZ is responsible for the FQ-RT-PCR. SQ participated in western blotting. YZ revised the experiment design. THZ is the corresponding author. He assisted in the conception of data analyses, and in writing the manuscript. HJL is the supervising author. All authors read and approved the final manuscript.

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