Viral Envelope Protein 53R Gene Highly Specific Silencing and Iridovirus Resistance in Fish Cells by AmiRNA

Yu-Sin Kim¹²,*, Fei Ke¹*, Xiao-Ying Lei¹, Rong Zhu¹, Qi-Ya Zhang¹*

¹ State Key Laboratory of Freshwater Ecology and Biotechnology, Institute of Hydrobiology, Chinese Academy of Sciences, Wuhan, China, ² College of Life Science, Kim Il Sung University, Pyongyang, Democratic People's Republic of Korea

Abstract

Background: Envelope protein 53R was identified from frog Rana grylio virus (RGV), a member of the family Iridoviridae, and it plays an important role in the virus assembly. Although inhibition of iridovirus major capsid protein (MCP) by small hairpin RNAs (shRNAs) has been shown to cause resistance to viral infection in vitro, RNA interference (RNAi) to inhibit aquatic animal virus envelope protein gene product has not been reported.

Methodology: We devised artificial microRNAs (amiRNAs) that target a viral envelope protein gene RGV 53R. By incorporating sequences encoding amiRNAs specific to 53R of RGV into pre-miRNA155 (pSM155) vectors, which use the backbone of natural miR-155 sequence and could intracellularly express 53R-targeted pre-amiRNAs. The pre-amiRNAs could be processed by the RNase III-like enzyme Dicer into 21–25 nt amiRNAs (amiR-53Rs) in fish cell lines. The levels of 53R expression were analyzed through real-time PCR and RGV virions assembly were observed by electron microscopic in fish cells transfected with or without amiR-53Rs at 72 h of RGV infection.

Conclusion/Significance: The results argue that viral envelope protein RGV 53R can be silenced and the virions assembly was deficient by amiR-53R-1, and further identified the first amiRNA of envelope protein gene from iridovirus that was able to cause resistance to virus infection in fish cells. The data demonstrate that the viral infection is efficiently suppressed (58%) by amiR-53R-1 targeting position 36–57 of RGV 53R. Moreover, electron microscopic observations revealed virion assembly defect or reduced virions assembly capacity was closely correlated to expression of amiR-53R-1. Based on real time PCR of the Mx gene, we found no evidence of activation of IFN by amiR-53R-1.

Introduction

Iridoviruses are one of the significant viral pathogens for aquatic animals that have caused great economic losses in the aquaculture industry worldwide [1]. Rana grylio virus (RGV), a member of iridoviruses, causes systemic infectious iridovirus disease in cultured pig frog (Rana grylio) [2]. Some RGV genes and their functions have been identified, including deoxyuridine triphosphatase (dUTPase), thymidine kinase (TK), 3β-hydroxysteroid dehydrogenase (3β-HSD) and ERV1 (essential for respiration and vegetative growth 1) [3–6]. Recently, an envelope protein that may play an important role in virus assembly, RGV 53R, was identified and characterized from RGV. RGV 53R has homologues in all sequenced iridoviruses. The structural conserved features shared by iridovirus envelope proteins were also found in RGV 53R. The intracellular distribution and dynamic changes of RGV 53R revealed that it co-localized with the ER components at early stage of post transfection[7]. Therefore, RGV 53R was selected for further characterization to understand the molecular pathogenesis and gene silencing by artificial microRNAs (amiRNAs) in RGV infection.

Discovery of the RNA interference (RNAi) pathway has led to exciting new strategies for developing treatment for viral diseases [8]. To date, siRNAs and miRNAs have been used for silencing expression of the major capsid protein (MCP) encoded by tiger frog virus (TFV) and red seabream iridovirus (RSIV), two iridovirus causing severe disease in aquatic animals [9–11]. But the envelope protein of iridovirus has not been demonstrated as a target for RNAi as well as MCP. RNA interference is a process of post-transcriptional sequence specific gene silencing in many eukaryotes. The use of RNAi to inhibit virus, including siRNAs and miRNAs, offers a new approach for controlling viral infections [12,13]. Artificial miRNAs (amiRNAs) expression plasmid vectors such as pre-miRNA 155-designed shRNAs vectors (pSM155) which were designed to target on ORF of endogenous and exogenous genes have been developed and used for RNA
interference [14–18]. The plasmid vector pSM155 used in this study to construct plasmids that express amiRNAs was based on the backbone of natural miR-155. And the natural miRNA: miRNA duplex sequence was replaced by the artificial one [19]. The appearance of plasmid-based expression systems that is effective and inexpensive for amiRNAs generation present rational way for the design and expression of 53R targeted amiRNA.

In the present study, we devised amiRNAs of structured 64-nucleotides that targets different positions of RGV 53R as pre-miRNA, which could be processed by the RNase III-like enzyme Dicer into 21–25 nt miRNAs (amiR-53R-1, amiR-53R-2 and amiR-53R-3) target 36–57, 476–498 or 37–58 oligo position, respectively) [20], to investigate whether viral envelope protein gene silencing and iridovirus resistance mediated by the amiRNAs in fish cells.

Results

Expression of pSM155-amiR-53Rs

Three pairs of oligonucleotides encoding 53R-specific amiRNAs of RGV (referred as amiR-53R-1, amiR-53R-2 and amiR-53R-3) (Table 1), and a pairs of oligonucleotides corresponding to PB2 gene of avian influenza virus, AIV (referred as amiR-PB2), were annealed and ligated into the pre-miRNA155 (pSM155) vector to create plasmids (pSM155-amiR-53Rs and pSM155-amiR-PB2) capable of producing 53R or PB2 gene encoded pre-amiRNAs in plasmid-transfected cells. The predicted structures of the engineered pre-amiRNAs incorporated into the pSM155 backbone are shown in Fig. 1. When pSM155-amiR-53Rs were transfected into grass carp ovary (GCO) cells, they allowed co-cistronic expression of pre-amiRNAs with GFP in cells under the control of the Pol II human CMV promoter. The co-cistronic expression of the pre-amiRNAs was monitored microscopically under a fluorescence microscope.

Inhibitory effect of amiR-53Rs on 53R expression

When GCO cells were co-transfected with pSM155-amiR-53R-1/pEGFP-N3-53R, pSM155-amiR-53R-2/pEGFP-N3-53R or pSM155-amiR-53R-3/pEGFP-N3-53R, pSM155-amiR-53Rs silenced the expression of the 53R gene in different levels compared with the control that were co-transfected with pSM155-amiR-PB2/pEGFP-N3-53R. 53R mRNA levels were evaluated in different groups using the real-time quantitative RT-PCR assay. The results indicated that expression of the 53R gene were reduced by 74% (FC = 0.26±0.02), 56% (FC = 0.44±0.04) and 35% (FC = 0.65±0.08) in cells transfected with pSM155-amiR-53R-1, pSM155-amiR-53R-2 and pSM155-amiR-53R-3 at 72 h, respectively (Fig. 2). The pSM155-amiR-53R-1 was more efficient in inhibiting 53R gene expression than pSM155-amiR-53R-2 and pSM155-amiR-53R-3. Thus, pSM155-amiR-53R-1 was chosen in our further studies on amiRNAs-mediated inhibition of RGV infection.

Delayed emergence of CPE of RGV in host cells by amiR-53R-1

Cytopathic effect (CPE) was the most intuitive parameter that reflected the viral quantity of virus accumulation [21]. In GCO cells that were infected with RGV after transfected with pSM155-amiR-53R-1, CPE were markedly delayed than that were transfected with pSM155-amiR-PB2, as well as only infected with RGV.

Reduction of RGV titer and virions by amiR-53R-1

To test whether pSM155-amiR-53R-1 could impede the packaging and production of infectious RGV, pSM155-amiR-53R-1 transfected cells were infected with RGV (MOI of 3) at 24 h of transfection. Cell supernatants were collected at 24, 48, 72 and 96 h post infection (p.i.) to determine the production of virions. In TCID50 assays, the average titer of RGV in pSM155-amiR-53R-1 transfected samples were about 17-fold, 12-fold, 14-fold and 6-fold lower than those of the transfected with pSM155-amiR-PB2 at different time intervals, respectively (Fig. 3). At the same time, the average titers in pSM155-amiR-53R-3 transfected cells were higher than those in pSM155-amiR-53R-1 transfected cells.

Observation of RGV virions in ultrathin sections of epithelioila papulosum cyprini (EPC) cells by electron microscopy showed that larger numbers of RGV virions were assembled and arranged orderly in the cytoplasm of the host cells only infected with RGV (Fig. 4-a). In pSM155-amiR-53R-1 transfected EPC cells, several sporadic RGV virions were present in the cytoplasm with irregular arrangement and the number of virions were significantly decreased (Fig. 4-b).

RGV resistance by amiR-53R-1

It has been shown in above that 53R silencing and RGV inhibition by pSM155-amiR-53R-1 delayed emergence of CPE, or reduced RGV titer and assembled virions in transfected fish cells. So 53R expression was assayed as the RGV inhibition efficiency in RGV infected GCO cells that transiently transfected with pSM155-amiR-53R-1 or pSM155-amiR-PB2 by real-time PCR. The data indicated that the 53R expression was reduced by 58% (FC = 0.42±0.05) at 72 h.p.i. of RGV in pSM155-amiR-53R-1 transfected cells with compared to the pSM155-amiR-PB2 transfected cells (Fig. 5).

Table 1. Oligonucleotides sequence encoding 53R-specific pre-miRNAs.

| Name    | Strand | Oligo sequence   | Position of target in gene |
|---------|--------|------------------|---------------------------|
| amiR-53R-1 | Top    | TGCTGTGACAGTTGTATAGATTCCGGTTTTGGCCACTGACCGGAATCTAAACACTGTCT | 36-57                     |
|         | Bottom | CCTGAGACAGTTGTATAGTTCCGGTGACCTGACAGTGGCACAAAACCGGAAATCTTACACACTGTCA |                     |
| amiR-53R-2 | Top    | TGCTGTAGCTGGAGTTTTGCAAAGTCCCTGAGCTGACAGTCCAAAAACTCCTAGTT | 476-498                   |
|         | Bottom | GCTAAGCTCATAGTTTTGCAAAGCTCGTAGCTGGACAAAAACGTTGTTGCAAACATCCTGAGTCA |                     |
| amiR-53R-3 | Top    | TGCTGTGACAGTTGTATAGTTCCGGTGACCTGACAGTGGCACAAAACCGGAAATCTTACACACTGTCA | 37-58                     |
|         | Bottom | CCTGAGACAGTTGTATAGTTCCGGTGACCTGACAGTGGCACAAAACCGGAAATCTTACACACTGTCA |                     |

Bold letters represent sense sequences of engineered amiRNAs derived from the target gene.

doi:10.1371/journal.pone.0010308.t001
No Mx response by amiR-53Rs

Mx is a key component of the interferon response. To determinate whether transfection of GCO cells with plasmids expressing 53R-targeted amiR-53Rs and pEGFP-N3-53R resulted in induction of IFN response, mRNA levels of Mx were detected. There were no significant changes in expression of Mx gene in GCO cells that were transfected with different pSM155-amiR-53Rs (Fig. 6). These results indicated that the antiviral activity was not induced by Mx expression but by RNAi.

Figure 1. Schematic presentation of predicted stem-loop sequences of pSM155-amiR-53Rs. +1, 11, 12 and 21 are corresponded to positions in stem structure. The antisense of the target sequences for the ORF of 53R and the 3' UTR of PB2 are underlined.
doi:10.1371/journal.pone.0010308.g001

Figure 2. Quantitative analysis of 53R mRNA levels in cells co-transfected with different plasmids. GCO cells co-transfected with pSM155-amiR-53R-1/pEGFP-N3-53R, pSM155-amiR-53R-2/pEGFP-N3-53R and pSM155-amiR-53R-3/pEGFP-N3-53R, respectively. 53R mRNA level from each group was measured by real-time PCR analysis 72 h post transfection. Group co-transfected with pSM155-amiR-PB2/pEGFP-N3-53R was used as negative control. The value of negative control was designated as 1.0 (n = 3). The values represent averages of three independent experiments, with the range indicated (±S.D.). *P<0.05 versus control.
doi:10.1371/journal.pone.0010308.g002
Discussion

Envelope protein gene 53R has been reported as one of the twenty six core genes existed in all iridoviruses [22]. Present study was the first time that the envelope protein targeted amiRNA was used for silencing an iridovirus efficiently. It revealed that envelope protein was also good target for iridovirus inhibition besides the MCP and offers a new approach for controlling viral infections.

Figure 3. Comparison of virus titers. GCO cells that transfected with pSM155-amiR-53R-1, pSM155-amiR-53R-3 and pSM155-amiR-PB2, respectively, were infected with RGV at 24 h post transfection (MOI of 3). Cell supernatants were collected at 24, 48, 72 and 96 h.p.i. and assayed for virus titration. The titers of RGV in pSM-amiR-53R-1 transfected samples were about 17-fold, 12-fold, 14-fold and 6-fold lower than those of the transfected with pSM155-amiR-PB2 at different time intervals, respectively. The data show the average titers of three independent experiments in lgTCID50 plus S.D. doi:10.1371/journal.pone.0010308.g003

Figure 4. Electron micrograph of RGV. EPC cells were transfected with pSM155-amiR-53R-1 and infected with RGV 24 h post transfection (MOI of 3). After 72 h of RGV infection, cells were collected for electron microscopy. (a) Crystalline aggregation contained large number of regular arranged virions in host cells only infected with RGV. (b) Several sporadic RGV virions with irregular arrangement in host cells transfected with pSM155-amiR-53R-1 and then infected with RGV. doi:10.1371/journal.pone.0010308.g004
Three amiRNAs targeted to different positions were constructed and used in this study. Two of these, amiR-53R-1 and amiR-53R-3, had the diversity in just one nucleotide in the targeted positions. AmiR-53R-1 targeted sequence was located in positions 36-57 of RGV 53R whereas amiR-53R-3 targeted 37-58. But the inhibition efficiency of these two amiRNAs was distinctly different. AmiR-53R-1 reduced 53R expression in 74% compared to 35% inhibition efficiency of these two amiRNAs was distinctly different. AmiRNAs device and plasmids construction

Three pairs of oligonucleotides encoding 53R-specific amiRNAs of RGV (referred as amiR-53R-1, amiR-53R-2 and amiR-53R-3) (Table 1), and a pair of oligonucleotides corresponding to PB2 gene of avian influenza virus, AIV (referred as amiR-PB2), were designed using the BLOCK-iTTM RNAi Designer/miR RNAi gene of avian influenza virus, AIV (referred as amiR-PB2), were designed using the BLOCK-iTTM RNAi Designer/miR RNAi (http://rnadesigner.invitrogen.com). Each oligonucleotide pair (“top strand” and “bottom strand” oligos) was annealed and ligated into the pre-miRNA155 (pSM155) vector to create plasmids (pSM155-amiR-PB2 and pSM155-amiR-53Rs) capable of producing pre-amiRNAs in plasmid transfected cells. The pSM155 vector was kindly provided by Prof. De-Yin Guo and Dr. On-Sam Sin. It was based on the backbone of natural miR-155 in which the natural miRNA/miRNA duplex sequence was replaced by the artificial one. Briefly, the natural miR-155 sequence was replaced by the artificial one. Briefly, the natural miR-155 sequence was inserted into the eGFP-N3 vector under a CMV promoter to create pSM155 vector. The oligonucleotides encoding amiRNA was inserted into the BsmBI site of miR-155 to replace the nature miRNA duplex sequence [19]. The 53R-expressing plasmid (pEFGP-N3-53R) constructed in previous studies [7] was used in co-transfection experiments to express the target 53R.

Transfection

GCO and EPC cells were seeded into 12-well or 6-well cell culture plates using 199 medium containing 5% of FBS 24 h before transfection. Cells were transfected with plasmids using Lipofectamine 2000 (Invitrogen, U.S.A.) following the manufacturer’s protocol. The transfection mixtures were removed at 6 h post transfection, and transfected cells were maintained for further processing.

Materials and methods

Cell lines and virus

A strain of frog iridovirus, *Rana grylio* virus (RGV), and two fish cell lines, grass carp ovaries (GCO) and *epitheloma papaolusum* cyprinid (EPC) cells, were used in these studies (GCO cell and EPC cell were maintained in our laboratory, please see the references) [29,30]. Cell culture, virus propagation and virus titer determination were performed as described previously [31,32].
AmiRNAs expression and anti-RGV activity detection

To select the amiR-53R that has the best inhibition efficiency, the amiR-53Rs were initially tested for sequence-specific silencing on the target 53R gene by employing transient transfection of a plasmid expressing 53R (pEGFP-N3-53R). GCO cells were co-transfected with pSM155-amiR-53R-1/pEGFP-N3-53R, pSM155-amiR-53R-2/pEGFP-N3-53R and pSM155-amiR-53R-3/pEGFP-N3-53R, respectively. At 72 h of transfection, total RNA was extracted from transfected cells and reverse transcribed to cDNA for real-time PCR analysis. Cells transfected with pSM155-amiR-PB2/pEGFP-N3-53R were used as negative control.

To elucidate antiviral effect of amiR-53R-1 on RGV replication, the expression of 53R gene was monitored in GCO cells transfected with pSM155-amiR-53R-1 and infected with RGV at an MOI of 3 after 24 h of transfection. The pSM155-amiR-PB2 was used for negative control. At 72 h of p.i, cDNA obtained by method above were used for real-time PCR analysis.

To assess inhibitory effect of amiR-53R-1 on RGV replication in terms of the production of viral particles, GCO cells transfected with pSM155-amiR-53R-1 and infected with RGV at an MOI of 3 after 24 h of transfection. Cell supernatants were collected at 24 h, 48 h, 72 h and 96 h p.i., respectively. The pSM155-amiR-PB2 was used as comparison and pSM155-amiR-53R-3 was used as negative control. Monolayer of GCO cells seeded in 96-well plates was inoculated with serial 10-fold dilutions of RGV samples for TCID$_{50}$ detection.

Electron microscope observation

EPC cells were transfected with pSM155-amiR-53R-1 and infected with RGV at an MOI of 3. Cells were centrifuged at 2000 x  g  for 72 h p.i. and the pellets were fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH = 7.4) for 1 day, rinsed in 0.1 M phosphate buffer containing 1% osmium tetroxide for 1 h at 4°C, dehydrated in a graded ethanol series and embedded in Spurr’s resin. The cell pellets were post-fixed with 2% osmium tetroxide in sodium cacodylate buffer for 1 to 2 h, dehydrated in a graded acetone series, and embedded in LR White acrylic resin. Ultra-thin sections were post-stained with 2% uranyl acetate and 3% lead citrate and observed at 100 kV with a JEOL 1230 transmission electron microscope.

Mx expression detection

The expression level of Mx was investigated by real time PCR in cells co-transfected with poly (I:C)/pEGFP-N3-53R, pSM155-amiR-PB2/pEGFP-N3-53R, pSM155-amiR-53Rs/pEGFP-N3-53R, respectively. Control cells were transfected only with Lipofectamine 2000 as well as in cells transfected with pEGFP-N3-53R.

Real-time PCR

RNA extracted by Trizol reagent (Invitrogen, USA) was treated with DNase I and added to the reverse transcription reaction containing 200 U M-MLV reverse transcriptase (promega, USA), 0.5 mM each of dNTP, 1×M-MLV reaction buffer, 500 ng of oligo(dT)$_{15}$ Primer and 25 U Rnasin (TOYOBO).

Real-time PCR was performed with 2 μl of cDNA in a final volume of 25 μl containing 11.25 μl of 2.5×RealMasterMix/20×SYBR solution(TIANGEN BIOTECH), 300 nM of sense primer and antisense primer. PCR was carried out in the 7300 Fast Real-time PCR System (Applied Biosystems, USA) using the following thermal cycling profile: 95°C for 3 min, followed by 40 cycles of amplification (95°C for 20 s, 58°C for 30 s, 68°C for 45 s). The absorption values of the SYBR Green I in each tube were detected at the end of each cycle. The melting curve analysis...
of PCR products from 55 to 95°C were also performed after PCR amplification. PCR was performed with the 53R forward primer (5'-CATCAGAAGGGAGGACAGA-3'), the 53R reverse primer (5'-GGTGGTAACTCATGGGAACTGCAG-3'), the β-actin forward primer (5'-GTGATGGAAATTCCGCGACCTG-3'), the β-actin reverse primer (5'-ACCAACATGAACACTGATGCT-3'), the Mx forward primer (5'-GCTGGAAGCGGAGGAGGAT-3'), and the Mx reverse primer (5'-CCCGGTGTGCCTTCTGTAG-3'). Fragments were 244 bp (53R), 120 bp (β-actin) and 250 bp (Mx), respectively. The Fold Change (FC = 2^ΔΔCt) in 53R and Mx mRNA expression levels were normalized to those of the same cDNAs. The relative 53R and Mx mRNA expression values in each group were calculated by the mathematical delta–delta method [33] and all samples were run in triplicate.

Acknowledgments

We thank Prof. De-Yin Guo and Dr. On-Sam Sin for kindly provided pSM155 vector and technical assistance.

Author Contributions

Conceived and designed the experiments: QYZ. Performed the experiments: YSK FK. Analyzed the data: YSK FK QYZ. Contributed reagents/materials/analysis tools: XYL RZ. Wrote the paper: YSK FK QYZ.

References

1. Williams T (1996) The iridoviruses. Adv Virus Res 46: 345–412.
2. Zhang QY, Xiao F, Li ZQ, Guo JF, Mao JH, et al. (2003) Characterization of an iridovirus from the cultured pig frog Rana grylio with lethal syndrome. Dis Aquat Org 48: 27–36.
3. Zhao Z, Ke F, Gui JF, Zhang QY (2007) Characterization of an early gene encoding for dUTPase in Rana grylio virus. Virus Res 123: 126–137.
4. Zhao Z, Ke F, Shi Y, Zhou GZ, Gui JF, et al. (2009) Rana grylio virus thymidine kinase gene: an early gene of iridovirus encoding for a cytolytic protein. Virus Genes 38: 345–352.
5. Sun Y, Huang YH, Zhao Z, Gui JF, Zhang QY (2006) Characterization of the Rana grylio virus 3β-hydroxysteroid dehydrogenase and its novel role in suppressing virus-induced cytopathic effect. Biochem Biophys Res Commun 351: 44–50.
6. Ke F, Zhao Z, Zhang QY (2009) Cloning, expression and subcellular distribution of a Rana grylio virus late gene encoding ERV1 homologue. Mol Biol Rep 36: 1653–1659.
7. Zhao Z, Ke F, Huang YH, Zhao JG, Guo JF, et al. (2008) Identification and characterization of a novel envelope protein in Rana grylio virus. J Gen Virol 89: 1866–1872.
8. Yeung ML, Bemasser Y, Le SY, Jeang KT (2005) siRNA, miRNA and HIV: promises and challenges. Cell Res 15: 935–946.
9. Xie JF, Lu L, Deng M, Weng S, Zhu JY, et al. (2005) Inhibition of reporter gene and iridovirus-tiger frog virus in fish cell by RNA interference. Virology 338: 41–52.
10. Dang LT, Kondo H, Aoki I, Hiroiro I (2006) Engineered virus-encoded pre-microRNA pre-miRNA induces sequence-specific antiviral response in addition to non-specific immunity in a fish cell line: convergence of RNA-related pathways and IFN-related pathways in antiviral response. Antiviral Rep 20: 316–323.
11. Dang LT, Kondo H, Hiroiro I, Aoki I (2008) Inhibition of red seabream iridovirus (RSIV) replication by small interfering RNA (siRNA) in a cell culture system. Antiviral Res 77: 142–149.
12. Denis AM, Tops BB, Plasser RH, Ketting RF, Hannon GJ (2004) Processing of primary microRNAs by the microprocessor complex. Nature 432: 231–235.
13. Sui HY, Zhao QY, Huang JD, Jin DY, Yuen KY, et al. (2009) Small Interfering RNA Targeting MJ Gene Induces Effective and Long Term Inhibition of Influenza A Virus Replication. PLoS One 4: e6571. doi:10.1371/journal.pone.0006571.
14. Du G, Yonenobu J, Zeng Y, Ouisse M, Frohman MA (2006) Design of expression vectors for RNA interference based on miRNAs and RNA splicing. FEBS Lett 57: 5421–5427.
15. Stegner F, Fu G, Rikkels RJ, Hannon GJ, Elledge SJ (2005) A lentiviral microRNA-based system for single-copy polymerase II-regulated RNA interference. Cell 121: 535–547.
16. Shin KJ, Wall EA, Zavazadjian JK, Santat LA, Lu J, et al. (2006) A single lentiviral vector platform for miRNA-based conditional RNA interference and coordinated transgene expression. Proc Natl Acad Sci USA 103: 13579–13574.
17. Zeng Y, Cullen BR (2004) Structural requirements for pre-microRNA binding and nuclear export by Exportin 5. Nucleic Acids Res 32: 4776–4785.
18. Boudreau KL, Martini I, Davidson DL (2009) Artificial microRNAs as siRNA shuttles: improved safety as compared to shRNAs in vitro and in vivo. Mol Ther 17: 169–175.
19. Smit Onsma (2009) Relationship of gene silencing effects and the structure and targeting sites of intron-spliced artificial miRNAs. Wunan University Ph.D thesis. 45 p.
20. Zamore PD, Tuschl T, Sharp PA, Bartel DP (2000) RNA: double-stranded RNA directs the ATP-dependent cleavage of mRNA at 21 to 23 nucleotide intervals. Cell 101: 25–33.
21. Huang XH, Huang YH, Yuan XP, Zhang QY (2006) Electron microscopic examination of the viromatrix of Rana grylio virus in a fish cell line. J Virol Methods 133: 117–123.
22. Eaton HE, Metcalf J, Penny E, Tcherepanov V, Upton C, et al. (2007) Comparative genomic analysis of the family Iridoviridae: re-annotating and defining the core set of iridovirus genes. Virol J 4: 11.
23. Dykxhoorn DM, Novina CD, Sharp PA (2003) Killing the messenger: short RNAs that silence gene expression. Nat Rev Mol Cell Biol 4: 457–467.
24. Magee WE, Griffin MJ (1972) The liver as a site for interferon production in response to poly Ipoly C. Life Sci II 11: 1081–1086.
25. Manetti R, Amunziato F, Tomasovic L, Giannoni V, Paronchi P, et al. (1995) Polyinosinic acid: polycytidylic acid promoter T helper type 1-specific immuneresponses by stimulating macrophage production of interferon-alpha and interleukin-12. Eur J Immunol 25: 2656–2660.
26. Oso EL, Hiroiro I, Aoki T (2006) Functional characterization of the Japanese flounder, Paralichthys olivaceus, Mx promoter. Fish Shellfish Immunol 21: 292–304.
27. Jiang J, Zhang YB, Li S, Yu FF, Sun F, et al. (2009) Expression regulation and functional characterization of a novel interferon inducible gene Gig2 and its expression profile. Mol Immunol 46: 313–3140.
28. Robertson B (2006) The interferon system of teleost fish. Fish Shellfish Immunol 20: 172–191.
29. Zhang QY, Ruan HM, Li ZQ, Yuan XP, Guo JF (2003) Infection and propagation of lymphocystis virus isolated from the cultured flounder Paralichthys olivaceus in grass carp cell lines. Dis Aquat Org 57: 27–34.
30. Fijan N, Sulimanovic D, Berzotti M (1983) Some properties of the polyinosinic acid: polycytidylic acid promoter. In: Polyinosinic acid: polycytidylic acid promoters T helper type 1-specific immuneresponses by stimulating macrophage production of interferon-alpha and interleukin-12. Eur J Immunol 25: 2656–2660.
31. Zhang QY, Zhao Z, Xiao F, Li ZQ, Guo JF (2006) Molecular characterization of three Rana grylio virus (RGV) isolates and Paralichthys olivaceus lymphocystis disease virus (LCDV-C) in iridoviruses. Aquaculture 251: 1–10.
32. Zhang QY, Li ZQ, Guo JF (1999) Studies on morphogenesis and cellular interactions of Rana grylio virus in an infected fish cell line. Aquaculture 175: 182–197.
33. Pfaffl MW (2001) A new mathematical model for relative quantification in real-time RT-PCR. Nucleic Acids 29: e45.