Selection of reference genes for quantitative real-time PCR analysis in chicken embryo fibroblasts infected with avian leukosis virus subgroup J

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Abstract

Background: The selection of stably expressed reference genes is a prerequisite when evaluating gene expression, via real-time PCR, in cells in response to viral infections. The objective of our study was to identify suitable reference genes for mRNA expression analysis in chicken embryonic fibroblasts (CEF) after infection with avian leukosis virus subgroup J (ALV-J).

Findings: The expression levels of 11 potential reference genes in CEF infected with ALV-J were determined by real-time PCR. The expression stability of these genes were analyzed and ranked using the geNorm tool. Analysis indicated that the genes RPL30 (ribosomal protein L30) and SDHA (succinate dehydrogenase complex, subunit A) were the most stably expressed genes in the ALV-J infected CEF.

Conclusions: The RPL30 and SDHA were deemed suitable for use as reference genes for real-time PCR analysis of mRNA gene expression during ALV-J infection, whereas commonly used ACTB and GAPDH are unsuitable to be reference genes.

Keywords: Avian leukemia virus, Reference gene, Real-time polymerase chain reaction

Background

Avian leukosis virus subgroup J (ALV-J) is an avian oncogenic retrovirus that induces myeloid leukemia in poultry, a disease that causes significant economic losses in the broiler breeder industry worldwide. Disease severity of ALV-J is linked to an increase incidence of tumor formation, immunosuppression, with ensuing high mortality rates [1]. To elucidate the molecular pathogenesis of ALV-J infection and tumor development, many studies have focused on analyzing host gene expression profiles following ALV-J infection [2-4]. Assessment of mRNA expression levels via real-time PCR has been a standard approach with high accuracy, sensitivity and reproducibility, and thus it has been widely used to infer host gene expression in response to virus infection [5]. To ensure a reliable result in gene expression analysis, the selection of stably expressed reference gene (or genes) is an important technical prerequisite for each individual experimental setting [6]. This is especially important because even the expression stability of candidate reference housekeeping genes varies across host tissue cells, and virus strains [7,8]. Therefore, the selection of reference genes should ideally be determined for each specific cell type and virus. Chicken embryonic fibroblasts (CEF) are frequently used for propagation of ALV-J. Those fibroblasts have also been used as model cells for the study of host-virus interaction. To identify suitable reference genes for mRNA analysis in poultry, here we present the expression stability of 11 housekeeping genes in CEF after ALV-J infection, and propose stably expressed genes for use as reference genes in ALV-J/CEF settings.

Findings

The ranking of the 11 candidate reference genes according to their expression stability values (M) is show in
Figure 1. From the most stable (lowest M-value) to the least stable (highest M value) genes: RPL30/ SDHA < HPRT1 < YHWAZ < TBP < ALB < GAPDH < TUBB < ACTB < B2M. RPL30 and SDHA have the lowest M value (0.41) and therefore are the most stably expressed genes.

To determine the optimal number of reference genes for accurate normalization, geNorm calculated the pairwise variation value $V_{n/n+1}$ (Figure 2). It was showed that a combination of the three most stable genes RPL30, SDHA and RPL4 has a lowest pair-wise variation value ($V_{3/4}$) of 0.117, lower than cut-off value of 0.15. If RPL4 is excluded, $V_{2/3} = 0.129$ is a bit higher than $V_{3/4}$, but still lower than cut-off value of 0.15. Therefore, two reference genes (RPL30 and SDHA) are sufficient for normalization.

Discussion
Since the introduction of real-time PCR in virology, RT-PCR has been extremely useful to document host cell responses to virus infection. Although there are several advantages, many factors can affect the performance of the test when quantitating mRNA expression levels. Validation
of a given quantitative PCR assay, and that of a normal- 
ilization process of CT values, is critical to obtain reliable 
results. The use of “reference genes” as markers of stability 
has served to normalize variations arisen from differences 
in nucleic acid integrity, the efficiency of the reverse tran- 
scription, and the amount of sample loaded onto a PCR 
master mix. An ideal reference gene should be one which 
is stably expressed in cells and unaffected by the experi-
mental treatments.

Viruses, as obligate intracellular parasites, replicate in-
side cells and use various strategies to induce cell apopto-
sis, cell transformation, cell death or other dysfunctions 
by shifting host gene expression on different scales. There-
fore, it is reasonable to think that the expression of so-
called “housekeeping” genes is probably unstable in virus-infected cells, and that it depends on the virus and host cell types. This highlights the necessity of ref-
ence gene selection during real-time PCR analysis for 
virus-infected cells. Radonić et al. [7] compared the ex-
pression of 10 candidate reference genes in cell lines 
infected with 6 human viruses: cytomegalovirus, human 
herpesvirus-6, camelpox virus, SARS coronavirus or yellow 
fever virus, and found that a commonly used gene ACTB, 
is unsuitable as reference gene, whereas TATA-Box binding 
protein (TBP) and peptidyl-prolyl-isomerase A (PPI) were 
stable genes for use as reference genes in expression 
studies in virus infected cells. However, Yin et al. [9] 
found that ACTB is a stable reference gene in CEF 
infected with Newcastle disease virus (NDV). Li et al. 
[10], after comparing 6 housekeeping genes, also showed 
that ACTB is the most stably expressed gene in infectious 
bursal disease virus (IBDV) infected CEF. Our previous 
study [11] with CEF infected with H5N1 avian influenza 
virus demonstrated that ACTB and RPL4 were the most 
suitable genes for use as endogenous reference genes in 
avian influenza virus H5N1/CEF settings. In the study 
carried out by Watson et al. [8], PPIA (peptidylprolyl 
isomerase A), GAPDH and SDHA were recommended as 
the best reference genes for host gene expression analysis 
in cells infection with immunodeficiency virus and her-
pes viruses.

Although identifying the best reference genes for each 
type of study may be time and resource-intensive, the 
studies listed above highlight the need to identify the 
most stable gene markers for each host virus assay to 
ensure reliable data. In the present study, RPL30 and 
SDHA were found to be the most stable reference genes 
in CEFs infected with ALV-J.

| Table 1 Primers used in this study [11] |
|-----------------------------|-----------------------------|-----------------------------|
| Symbol | Gene name | Accession no. | Primer sequences(5′-3′) | Amplicon(bp) |
| ALB | Albumin | NM_205261 | F: CCTGGACACCAAGGAAAT  
R: TGTGGACGCCGATAGAAT | 197 |
| B2M | Beta-2-microglobulin | NM_001001750 | F: CGTCCTCTAAGCTCTGTTG  
R: TCTCGTGCTCCACCTTGC | 194 |
| GAPDH | Glyceraldehyde-3-phosphate dehydrogenase | NM_204305 | F: AGCACCCGCATCAAAAGG  
R: CATCATCCAGGTCGGA | 283 |
| HRPT1 | Hypoxanthine phosphorribosyltransferase 1 | NM_204848 | F: ACTGCGCTGTCCTTGTTG  
R: GGTTGGTGTGCTTGT | 245 |
| RPL30 | Ribosomal protein L30 | NM_001007967 | F: GAGTCACCTGGGTCATAAA  
R: CCAAACACTGCTGGTCTT | 160 |
| RPL4 | Ribosomal protein L4 | NM_001007967 | F: TTATGCCATCTGCTTGGCC  
R: GCGATCTCCTACCCACTCCT | 235 |
| SDHA | Succinate dehydrogenase complex, subunit A | XM_419054 | F: CAGGGATGTAGTGTCTGTTG  
R: GGGAATAGGCTCCTTAGTG | 187 |
| TBP | TATA box binding protein | NM_205103 | F: CGTCAGGGAAATAGGCA  
R: ACTGCGCTGTCCTTGTTG | 470 |
| TUJ2B | Beta-tubulin | NM_205315 | F: AAAACAAGTTATCAGGGTCTGA  
R: ATGCCGACCAAAATCG | 243 |
| YWHAZ | Tyrosine 3-monooxygenase/tryptophan S-monooxygenase activation protein, zeta polypeptide | NM_001031343 | F: TCCACACCGACAGACCA  
R: CCAGCCTCCTCAACTC | 358 |
| ACTB | Beta-actin | NM_205518 | F: CCTGTCGCCATCTGATAGGCTA  
R: ATTTCCTCTCGGGCTTGT | 139 |
Conclusions
In conclusion, RPL30 and SDHA could be used as reference genes for the standardization of in CEF gene response to the infection with ALV-J, whereas commonly used ACTB and GAPDH are unsuitable to be reference genes in ALV-J/CEF settings.

Methods
All animal experiments were done in accordance with institutional and national ethical guidelines. The protocol was approved by the Ethical Committee for animal experiments of Southwest University for Nationalities.

Infection of CEF with ALV-J virus
Primary cultures of chicken embryo fibroblasts (CEF) were prepared from 10-day-old specific-pathogen-free (SPF) chicken embryos (Yebio Bioengineering Co. Ltd, Qindao, China) as described previously [12], and maintained in DMEM supplemented with 10% fetal bovine serum (FBS). The cells were seeded (approximately 5 x 10^6 cells/well) in 24-well culture plates. Then the cells were infected with 100 TCID50 (50% tissue culture infective dose) ALV-J strain NX0101, obtained from the China Animal Health and Epidemiology Centre in Qindao, China. After 1 h incubation, the cells were washed and further incubated in media with 2% FBS. Host-virus interactions were tested in triplicate wells, which were harvested at 0, 24, 120 and 192 hours after infection for RNA extraction.

RNA extraction and cDNA synthesis
Total RNA was extracted from each sample using commercial RNAiso Reagent (TaKaRa, Japan) and purified with RNase-free DNase (TakaRa, Japan) following the manufacturer’s instructions. Complementary DNA (cDNA) was synthesized using the PrimeScript™ RT reagent (TaKaRa, Japan) with random primers following the product instructions. cDNA products were treated with RNase H to remove remnants of RNA, and stored at −20°C until the time of real-time PCR testing.

Real-time PCR
A total of 11 housekeeping genes, previously used as references for the evaluation of expression stability in CEF infected with avian influenza virus [11], were also used as candidate reference genes in the present study (Table 1).

Real-time PCR reactions were performed on an ABI 7300 Real-Time PCR Detection System (Applied Biosystems, USA) with SYBR® Premix Ex Taq™ Kit (TaKaRa, Japan). The 20 μl reaction volume consisted of 10 μl SYBR® Premix Ex Taq™, 2 μl cDNA, 0.4 μl ROX reference dye and 0.25 mM of each primer. The following PCR cycling profile was used: one single step at 95°C for 5 min, followed by 45 cycles of 95°C for 30 sec, 56°C for 30 sec, and 72°C for 30 sec, ending with a melting curve analysis from 65°C to 95°C.

Determination of gene expression stability
To determine the expression stability of 11 candidate genes over CEF cells collected from different time points, the geNorm software [13] was used to calculate gene expression stability measure (M value), which is the mean pair-wise variation for a gene compared with that of all the other tested candidate genes. Genes with higher M values are more variable and therefore have low expression stability. The stepwise exclusion of any given gene with the highest M value allows the ranking of the remaining tested genes based on their increasing expression stability. Candidate genes with the lowest M values have the most expression stability and thus should preferentially be used as reference genes for gene expression and normalization of other genes.

The optimal number of reference genes required for accurate normalization was also determined using geNorm by calculating the pairwise variation value Vn/n+1 between two sequential normalization factors containing an increasing number of reference genes. A large variation means that the added gene has a significant effect and should preferably be included for calculation of a reliable normalization factor. A Vn/n+1 value of 0.15 has been proposed as a cut-off value, below which the inclusion of an additional reference gene is not required.

Abbreviations
CEF: Chicken embryonic fibroblasts; ALV-J: Avian leukemia virus subgroup J (ALV-J); FBS: Fetal bovine serum; TCID50: 50% tissue culture infective dose.

Competing interests
The authors declare that they have no competing interests. This work was supported by the “863” National High-tech Development Research Project (2012AA101304) and Veterinary Medicine Discipline Program of Southwest University for Nationals (2011WWD-50906).

Authors’ contributions
CT and HY created the idea of this study and participated in the design of study. FY designed the primers, participated in data analysis and drafted the manuscript. XL carried out the cell culture, the virus infection and performed the real-time PCR study. AR participated in analysis and interpretation of the data and revised the manuscript. All authors read and approved the final manuscript.

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