A Luciferase-EGFP Reporter System for the Evaluation of DNA Methylation in Mammalian Cells

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Abstract—DNA methylation is an essential epigenetic modification involved in numerous biological processes. Here, we present a cell-based system pLTR-Luc2P-EGFP for evaluation of DNA methylation in mammalian cells. In this system, the expression of reporter gene luciferase2P (Luc2P)-EGFP is under the control of HIV-1 promoter 5' long terminal repeat (LTR), which contains multiple CpG sites. Once these sites are methylated, the expression of Luc2P-EGFP is turned off, which may be visualized under fluorescence microscopy, with quantification performed in luciferase activity assay. As a proof of principle, pLTR-Luc2P-EGFP was methylated in vitro, and transfected into 293T cells, where the reduction of Luc2P-EGFP expression was confirmed. Premixed reporter DNA samples with the methylation levels varying from 0 to 100% were used for quantitative measurements of DNA methylation. The resulting standard curves indicated the accuracy of luciferase activity exceeding that of the Western blotting against EGFP. The Bland–Altman analysis showed that data from luciferase activity assay were in good agreement with the actual DNA methylation levels. In summary, we have established a reporter system coupled with reliable detection technique capable of efficient quantifying the changes in methylation in mammalian cells. This system may be utilized as a high throughput screening tool for identifying molecules that modulate DNA methylation.

Keywords: Luc2P firefly luciferase, cell-based reporter system, HIV-1 promoter 5’ LTR, DNA methylation

INTRODUCTION

Epigenetics describes the phenotypic changes that alter gene expression without disturbing the primary DNA sequences [1]. Epigenetic regulations are heritable and reversible; their list includes DNA methylation, histone modifications and small noncoding microRNAs (miRNA) [1, 2]. Since Riggs [3] and Holliday et al. [4] proposed a convincing model of molecular mechanism of inheritance more than 40 years ago, DNA methylation had been considered as a paradigm of epigenetic information transfer [5]. Currently, DNA methylation is a common epigenetic modifications in eukaryotes, which plays an important regulatory role in biological processes such as transposable element silencing, genomic imprinting, X chromosome inactivation and developmental processes [6, 7], and abnormal methylation patterns are often associated with the incidence of diseases [8].

In mammals, most of the methylations occur at the carbon-5 position of cytosine (5 mC) [9–11]. The methylcytosine is mainly found in cytosine-guanine (CpG) dinucleotides. Although the CpG dinucleotides constitute only 1% of the human genome, CpG-rich stretches, so-called CpG islands, are located in the promoter regions of more than 70% of all known human genes [12–14]. 5mCs, especially when clustered at CpG sites, are important transcriptional silencers at gene promoters and endogenous retrotransposons in the genome [15–17]. Many studies have shown that, the epigenetic silencing of a variety of genes by hyper methylation of promoter-associated CpG islands is often related with particular diseases [18–20]. Therefore, a simple, reliable and sensitive method for detecting DNA methylation and its changes (e.g. hyper- or hypomethylation) is of great interest.

The traditional DNA methylation assays are mainly based on sodium bisulfite treatment, which converts non-methylated cytosine into uracil, while methylated cytosine is resistant to bisulfite and remains unchanged [21]. This allows discrimination between methylated DNA and non-methylated DNA, and usually followed by methylation-specific PCR, DNA sequencing or combined bisulfite restriction
assays. Due to their reliability and accuracy, they are widely used to quantify the site specific DNA methylation. However, these assays require complex procedures such as cloning and sequencing, which limits their usage in high-throughput analysis [22, 23]. Meanwhile, techniques based on high-performance liquid chromatography (HPLC) [24], restriction enzyme PCR [25] and gas chromatography/mass spectrometry (GC/MS) [26] have been developed. These techniques are complex, time consuming and expensive. Therefore, the more convenient and easy to use assay assessing DNA methylation warrants development.

Luciferase is a type of bioluminescence producing enzymes isolated from several animal species. Due to the high sensitivity, robust signal and assay convenience, it is often used as reporter to monitor gene activity/gene promoter activity. For example, Sanchez et al. used the luciferase reporter gene to analyze the relationship between the structure and biological activity of strigolactones [27]. Solberg et al. used a luciferase assay to characterize the activity of the 5'-flanking promoter of mouse Tcf3 [28]. Moreover, the combination of two luciferases: firefly luciferase and Renilla luciferase, improved the quantitative accuracy and assay reproducibility, where Renilla luciferase serves as an internal control to monitor cell numbers and viability. So the luciferase reporter assay is a powerful tool used to study the regulatory elements of genes of interest. For example, Xiao et al. established the 293-Sox2-Luciferase cell line as a luciferase reporter system to study transcriptional regulation of the human Sox2 gene [29]. And recently, a luciferase reporter virus icSARS-CoV-2-nLuc-GFP was established to test the cross–CoV neutralization of sera from SARS and COVID–19 patients [30].

The epigenetic regulation theory suggests that the promoter activity is consistent with the level of DNA methylation [31, 32]. Therefore, the luciferase was further used as a reporter to reflect the methylation status and/or methylation changes of gene promoter. For example, Li et al. used a cell-based firefly luciferase reporter assay to test the effect of Gadd45a on DNA methylation [33]. And the evaluation of CpG methylation by using HIV LTR-luciferase plasmid was mentioned in a short report [34]. However, the quantitative accuracy and reliability of the detection, including the dose-response relationship between methylation levels and luciferase activity, remain to be assessed.

In order to fully characterize the properties of luciferase-based methylation reporter system, here we used a modified firefly luciferase gene Luc2P fused with EGFP as the reporter to represent the promoter activity, and established a cell-based system to measure the changes of DNA methylation. The result from luciferase activity assay was validated by comparing it with the data obtained by HpaII sensitivity assay [35] and Western blotting analysis [36, 37]. After carefully analysis, we proved that this is a reliable reporter system for accurate measuring the changes of DNA methylation in living cells.

**EXPERIMENTAL**

**DNA constructions.** The EGFP expression plasmid pEGFP-N1 was purchased from BD Biosciences Clontech, and used as backbone for pLTR-Luc2P-EGFP construction. The HIV-1 5' LTR was PCR amplified from HIV-1 pNL4.3, and used to replace the CMV promoter of pEGFP-N1 by AseI and NheI digestion. Luc2P was amplified from pGL4.32 [luc2P/NFkB-RE/Hygro] (Promega), and cloned into pEGFP-N1 within HindIII and BamHI sites. The constructed plasmid pLTR-Luc2P-EGFP was verified by sequencing. Primers used for PCR amplification are: 5'-AseI-LTR, TCG-TATTAATTGGAGGGGCATAATTGTGC, 3'-NheI-LTR, CTAGCTAGCTGCTAGAGATTTTCCACTGAC, 5'-HindIII-EGFP, CGGATCCTAGGAAAGTGGAGGCGAAGATGCCAAAAACATTA, 3'-BamHI-luc2P, CGGGATCCCGGCTGTCGCTGG.

**In vitro methylated of plasmid DNA.** The plasmid pLTR-Luc2P-EGFP was treated with CpG methylase M.SssI (Zymo). Briefly, 2 μg plasmid was incubated with 0.6 mM S-adenosylmethionine and 4 units M.SssI at 30°C for 6 h, then 2 units of M.SssI were added to the reaction system and continuing incubated at 30°C for 6–8 h. The treated plasmid was recycled by 3 M sodium acetate. Complete CpG methylation of the LTR–Luc2P-EGFP was confirmed by endonucleases (HpaII and MspI, Thermo) digestion and bisulfite-mediated methylcytosine mapping.

**Bisulfite-mediated methylcytosine mapping.** After bisulfite conversion (Zymo), regions of interest in M.SssI-treated plasmid pmLETR–Luc2P-EGFP and non-methylated plasmid pLTR–Luc2P-EGFP were amplified by PCR (primers targeting LTR: LTR-BSP-F, 5'-TATGAGTTAATGAGGAGGTGGAGGCT‘-3’, and LTR-BSP-R, 5'-AATCTAACAAAAAACCACATA'-3’). The PCR products were gel-purified by using DNA Gel Extraction Kit (Axygen) and cloned into pJET1.2 for sequencing. Data were analyzed by online software QUantification tool for Methylation Analysis (http://quma.cdb.riken.jp/) [38].

**Cell culture and transient transfection.** Human embryonic kidney (HEK) 293T cells and HeLa cells were maintained in DMEM (Gibco) supplemented with 100 units/mL penicillin, 0.1 mg/mL streptomycin, and 10% fetal bovine serum (FBS, Gibco). Cells were cultured at 37°C in 5% CO₂ incubator.

For transient transfection, 293T cells were plated onto 12-well plates. When about 80% confluent, cells were transfected with unmethylated pLTR–Luc2P-EGFP or methylated pmLETR–Luc2P-EGFP plasmids using FuGENE® HD reagent (Promega), following manufacturer’s protocol. The *Renilla* luciferase
control vector pGL4.74 [hRLuc/TK] (Promega) was cotransfected with reporter vector at the ratio of 1 : 10.

**Immunofluorescence assay.** For the detection of 5mC, HeLa cells grown on cover slips were transfected with the unmethylated and methylated pLTR-Luc2P-EGFP (0.2 μg/well) for 36 hours and fixed in 4% paraformaldehyde in PBS for 15 min at room temperature. The cells were rinsed with PBS thrice and incubated in PBS containing 5% fetal bovine serum and 0.3% Triton X-100 for blocking, and then incubated overnight at 4°C with 5-methylcytosine monoclonal antibodies (1 : 100, EpiGentek). The next day after reworking at RT for 1 h, the cells were washed thrice with PBS, and incubated with goat anti-mouse IgG (H + L) coupled with Alexa Fluor 555 dyes antibody (1 : 500, Thermo) for 1 h at 37°C. After three washes with PBS, the cells were incubated with DAPI staining solution (Beyotime) for 5–5 min. Following three washes with PBS, the samples were mounted on slides with anti-fade Fluorescence Mounting Medium (Leica Inc, Germany).

For the detection of EGFP, 36–48 hrs post-transfection, 293 T cells with EGFP fluorescence signal were observed under Nikon fluorescence inverted microscope (Nikon ECLIPSE Ti), images were captured and processed with NIS-Elements D.

**Western blotting.** 48 hrs post transfection, 293T cells were lysed on ice in RIPA buffer plus protease inhibitors (Beyotime, China). Lysates were fractionated by SDS–PAGE and transferred to nitrocellulose (NC) membrane (Millipore). Membranes were blocked with 5% nonfat dry milk in PBS, incubated with rabbit anti-GFP antibody (1/400 dilution in PBS, Santa Cruz), and again incubated with secondary antibody goat anti-rabbit IgG-HRP (1/10000 dilution in PBS, Elabscience), GAPDH immunoblotting was performed with anti-human-GAPDH antibody (1 : 10000, Sigma) as control. Detection was performed using enhanced chemiluminescence (Bio–Rad).

**Luciferase activity assay.** The Dual-Glo® Luciferase Reporter Assay System (Promega) was used for the firefly and Renilla luciferase activity measurement. Briefly, 48-h post transfection, Dual-Glo® Reagent equal to the volume of culture medium was added to the plate wells. The plate was subjected to end-over-end rotation for 20 min to achieve complete lysis. And the firefly luminescence (Fluc) was measured in a GloMax®-96 microplate lumimeter (Promega). Then, equal amount of Dual-Glo® Stop & Glo® Reagent was added, and the Renilla luminescence (Rluc) was measured 20 min later. The Relative Luciferase Activity (RLA) was calculated by dividing the Fluc by Rluc. Relative Response Ratio (RRR) was calculated by [(experimental RLA) – (RLA of cells transfected with 100% methylated DNA)]/[RLA of cells transfected with 0% methylated DNA) – (RLA of cells transfected with 100% methylated DNA)].

**HpaII sensitivity assay.** The CpG methylation levels of 5’-LTR in pLTR-Luc2P-EGFP were quantified using the quantitative real-time PCR(qPCR) after HpaII digestion, which is blocked by CpG methylation [35]. Total DNA was extracted from transfected cells with Quick-DNA™ Miniprep Plus Kit (Zymo). 0.5 μg DNA was incubated with 10 units HpaII or in a mock reaction without HpaII at 37°C for 4 h, and inactivated for 20 min at 80°C. Equal amounts from both the mock reaction and the HpaII reaction were used in qPCR using the SYBR Premix Ex Taq (Tli RNaseH Plus, Takara). 5’-LTR flanking two HpaII digestion sites was amplified. Housekeeping gene GAPDH was used as the internal reference. PCR reaction was run on an Applied Biosystems 7500 Real-Time PCR system under the following conditions: 95°C for 30 s, followed by 40 cycles of 95°C for 5 s and 60°C for 30 s. Primer sequences for 5’-LTR are: forward-GCCAATGGAAGGAGAACAACA; reverse-AAGCGGAAAAGTCCCTTGTA; for GAPDH are: forward-GAAGGTGAAGGTCGGAGTCAAC and reverse-CAGAGTTAAAAAGCAGCCCTGTG. Reaction was run in triplicate. HpaII sensitivity of the CCGG site was calculated by \[1–2^{−ΔΔCt}\text{(mock)}−ΔCt\text{(HpaII)}\] × 100%.

**Statistical analysis.** Data are shown as the mean ± SD of at least two independent experiments; each replicate has at least three technical replicates. The two-tailed Student’s t-test and one-way analysis of variance (ANOVA) were performed using SPSS (version 21, IBM). A value of \(P < 0.05\) was considered statistically significant.

Histogram was obtained by Graphpad Prism 6 to show the data in the most intuitive way. The standard curve was obtained by Excel 2013 to show correlation between three DNA methylation assays and standard. Bland–Altman plot was obtained by MedCalc software to analysis the consistency of three DNA methylation assays.

**RESULTS**

**Construction and Methylation of the Reporter Vector pLTR-Luc2P-EGFP**

To construct the report vector, pEGFP-N1 was used as the backbone. The modified firefly luciferase Luc2P was inserted into multiple cloning site, and the fusion protein Luc2P-EGFP was used as the reporter. The original promoter CMV was removed and replaced with HIV-1 5’LTR, which contains eight CpG sites and is easily methylated [34, 39] (Fig. 1a). After methylated in vitro by M.sssI, the pmeLTR-Luc2P-EGFP can be delivered into mammalian cells. Then the methylation status of the promoter 5’LTR can be monitored by HpaII sensitivity assay (real time–PCR) directly [35], or indirectly reflected by the expression of the reporter gene Luc2P-EGFP. In transfected cells, the expression of Luc2P-EGFP can be qualitatively monitored by fluorescence micros-
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After M.sssI treatment, pmeLTR-Luc2P-EGFP was analyzed by HpaII/MspI digestion. Both enzymes recognize the same site-CCGG, but the HpaII digestion can be blocked by CpG methylation [40]. As shown in Fig. 2a, both methylated and non-methylated DNA could be cleaved by MspI, which confirmed the presence of CCGG sites in pLTR-Luc2P-EGFP. But only M.sssI treated DNA was resistant to the HpaII digestion, indicating the CpG sites in pmeLTR-Luc2P-EGFP were fully methylated by M.sssI treatment. Also, the methylation efficiency was confirmed by the bisulfite sequencing of LTR region. As shown in Fig. 2b, among the six M.sssI treated plasmid clones (M1–M6), the methylation rate of CpG sites was 87.5% in four clones, and 75% in the other two clones. In contrast, 0% of CpG sites were methylated in all six non-methylated clones (U1–U6). Then pmeLTR-Luc2P-EGFP was transfected into HeLa cells, the presence of 5mC was measured by immunofluorescence assay. In the cells transfected with non-methylated DNA, the presence of 5mC was not detected (Fig. 2c, lower panel). In pmeLTR-Luc2P-EGFP transfected cells, 5mC was detected and mainly located in the nucleus (Fig. 2c, upper panel), suggesting that, pmeLTR-Luc2P-EGFP remains methylated and stable after delivered into living cells.

Validation of the Reporter System

In order to evaluate the reporter system, equal amounts of M.sssI treated or untreated reporter plasmids were delivered into 293T cells. 48-h following transfection, the cells are assayed by methods listed in Fig. 1b. Compared with cells transfected with non-methylated plasmid, strong silencing of Luc2P-GFP expression was observed in pmeLTR-Luc2P-EGFP transfected cells, as examined by fluorescence microscopy and Western blotting (Figs. 3a, 3b). Meanwhile, similar result was obtained by luciferase activity assay, that is, the relative luciferase activity of the pmeLTR-Luc2P-EGFP transfected cells was significantly reduced (Fig. 3c, P < 0.01). Moreover, the HpaII sensitivity of DNA isolated from pmeLTR-Luc2P-EGFP

Fig. 1. Overview of reporter vector and experimental procedures. (a) Schematic representation of the pLTR-Luc2P-GFP construction, key components were shown, in which the expression of Luc2P-EGFP fusion protein was under the control of HIV-1 5’LTR promoter. (b) A schematic illustration of the experimental procedure.
transfected cells was significantly decreased (Fig. 3d, P < 0.01), confirmed that the gene silencing is in consistency with the promoter methylation. These data showed that these three methods (by detecting EGFP, Luc2P, 5mC in LTR region) are capable of efficient distinguishing methylated DNA samples from non-methylated ones.

Quantification of Methylation Status by the Reporter System

To determine whether this system allows quantitation of DNA methylation, we performed a dilution experiment. Fully methylated plasmid (pmeLTR-Luc2P-EGFP) was set as 100% and non-methylated plasmid (pLTR-Luc2P-EGFP) as 0%. The plasmids were mixed by different ratios (3:1, 2:2, 1:3) to yield 75, 50 and 25% methylated plasmid. DNA mix with different methylation gradients (100, 75, 50, 25, 0%) were transfected 293T cells, the methylation status were detected by corresponding methods. As shown in Fig. 4, decreasing of methylation levels result in a gradually increase of Luc2P-GFP expression, as indicated by fluorescence intensity, Western blotting, relative response ratio (RRR) of luciferase assay. Consistently with protein expression, HpaII sensitivity assay showed the same trend. Hereby, we confirmed that
presented reporter system may find its use in quantitative measurements of DNA methylation.

**Accuracy and Reliability Assessment**

To evaluate quantitative accuracy of the detection methods, data from Fig. 4, plotted against their respective methylation percentage, were subjected to linear regression analysis. Figure 5 revealed a strong linear regression between normalized data points and the percentage of methylated DNA ($R^2 = 0.9156$ for Western blotting; $R^2 = 0.9613$ for luciferase assay; $R^2 = 0.9867$ for HpaII sensitivity assay), suggesting a good correlation between the measured value and actual methylation level.

The consistency of the measurement was further analyzed by Bland—Altman plot [41, 42]. All data points collected by Western blotting, luciferase assay and HpaII sensitivity assay are within the consistency limit (Fig. 6), namely 100% differences are within the consistency limit. It is generally believed that, if more than 95% of the differences are within the consistency limit, the consistency is considered to be good [42]. Therefore, the consistency of three assays was all in good agreement with the actual 5mC level of pLTR-Luc2P-EGFP.

**DISCUSSION**

Here we present a reporter system for assessing the 5mC modification in mammalian cells coupled with accurate and reliable assay for visualization and quantitation of these methylation events including real time-PCR, Western blotting and luciferase activity assay. Despite that the first two methods only require resources commonly available in a laboratory of molecular biology, they are more cumbersome and time-consuming than luciferase activity assay. Luciferase activity assay requires specialized equipment and reagents, but does not require the sample preparation step. Therefore, it is rapid in operation, plus very sensitive in detection and easy to analysis, which enables
Fig. 4. Quantitative assessment of the reporter system accuracy. 293T cells transfected with pLTR-Luc2P-EGFP methylated at different levels (100, 75, 50, 25, 0%) were observed under fluorescence microscopy (a) and analyzed by Western blotting (b), luciferase activity assay as indicated by relative response ratio (c) and HpaII sensitivity assay (d). The experiments were done in triplicate, with cultures repeated three times in total. Data of the histogram are presented as mean ± SD. The fluorescence image and blot plot is a representative of the three experiments. Open circles in (d) represent data from individual experiments and lines represent mean values. *P < 0.05, **P < 0.01, one-way analysis of variance (ANOVA).

Fig. 5. Linear regression analysis between the methylated level of transfected plasmid and normalized data points from Western blotting (a), luciferase activity assay (b) and HpaII sensitivity assay (c) of Fig. 4. Solid circle represents mean values, straight line represents the corresponding standard curve, broken line represents the connection between actual values. The equation of the linear regression curve and the correlation coefficient ($R^2$) are indicated.
researchers to obtain accurate results with minimal effort.

Luciferase was first found from firefly (Photinus pyralis) by de Wet J.R. et al. in 1985, and advanced in detection technology across academia and industry [43]. Currently, luciferase has been widely used in reporter system for functional genomics (such as RNAi screening), signaling pathways, well-defined molecular mechanism, and biological activity studies. While searching for the term “luciferase reporter assay”, over 20000 publications since 1987 appeared [44]. As a reporter, luciferase genes have the following important features: exceptional sensitivity (100- to 1000-fold higher than fluorescent reporters such as GFP), wide dynamic range, typically no endogenous activity in host cells to interfere with quantitation, and, the measurements are almost instantaneous [45].

Therefore, here we used a dual-luciferase assay to evaluate the methylation of HIV-1 promoter 5' LTR in transfected 293T cells. Firefly luciferase Luc2P tracks the transcriptional activity of LTR promoter, while the Renilla luciferase acts as an internal control to minimize experimental variability due to pipetting errors, cell viability and transfection efficiency [46]. The ratio of two luciferase signals (relative luciferase activity) represents the relative expression the Luc2P-EGFP, and indirectly reflects the methylation status of the promoter. In our study, the expression of Luc2P-EGFP was also detected by Western blotting. The results revealed that, both methods could monitor protein expression with the linear range from 0 to 100% methylated DNA. While compared with Western blotting ($R^2 = 0.9156$), luciferase assay has a higher accuracy ($R^2 = 0.9613$). Meanwhile, we used the HpaII sensitivity assay to directly examine the methylation of promoter LTR, which is based on the combination of methylation sensitive restriction enzyme HpaII and qPCR [35, 47]. The results showed that, the HpaII sensitivity was also negatively correlated with the methylation level of pLTR-Luc2P-EGFP.

And the Bland-Altman analysis revealed that, the consistency of luciferase activity assay was similar to HpaII sensitivity assay, and, in good agreement with the actual DNA methylation level. Thus, these data proved that, the luciferase activity assay is a valid substitute for detection of DNA methylation in living cells.

Moreover, the simplicity and sensitivity of luciferase activity assay makes it a high throughput screening tool for identification of novel compounds that modulate DNA methylation. In fact, luciferase assays have been used for screening of antimicrobial agents against Mycobacterium tuberculosis and immunosuppressive drugs as early as 1990s [48, 49]. After that, increasing studies proved the potential usefulness of luciferase assay for the screening. Since this method could not only reduce the cost of drug screening, but also improve the reliability and predictability of the results [50].

In terms of our reporter system, where luciferase is driven by HIV-1 promoter, it can be used to screen compounds targeting HIV-1 latency reactivation. HIV-1 latency is the major barrier for HIV-1 eradication in infected individuals. And reactivation of latent virus is the first step in the “kick and kill” strategy, a novel direction in HIV-1 cure [51]. DNA methylation in LTR region has been shown to be highly associated with latency regulation [52]. Hypermethylation suppresses viral gene expression and stabilizes HIV-1 latency. Compounds demethylating LTR reactivate latent virus, and can be further used in eliminating or reducing viral reservoirs [51, 53]. Therefore, instead of using HIV-infected cell lines or primary cells, where latency may be achieved by more than one mechanism, compounds reactivating virus solely via DNA demethylation may be screened in our reporter system, which is also applicable to the studies of LTR methylation in HIV-1 latency. Hence, the screens for therapeutic compounds against diseases caused by abnormal methylation changes may be broadened [54].
In conclusion, a luciferase-based reporter system pLTR–Luc2P-EGFP was developed and validated. Feasibility and reliably of this system for assessment of methylation changes is proven. This study promotes the use of bioluminescence in the field of epigenetics, and provides new method for screening compounds targeting DNA methylation/demethylation.

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COMPLIANCE WITH ETHICAL STANDARDS
This article does not contain any research involving humans or animals as subjects of research.

The authors declare no conflicts of interest.

AUTHOR CONTRIBUTIONS
Xiaoxia Wang and Huaijie Jia conceived and designed the study. Xiaoxia Wang, Huijijie Jia and Xiaole Wei performed the experiments and wrote the paper. Yanrong Lv, Huihui Sun and Jiying Tan analyzed the data. Zhizhong formed the experiments and wrote the paper. Yanrong Lv, the study. Xiaoxia Wang, Huaijie Jia and Xiaole Wei performed the experiments and wrote the paper. Zhizhong Jing reviewed and edited the manuscript. All authors read and approved the final manuscript.

ADDITIONAL INFORMATION
The text was submitted by the author(s) in English.

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