Growth kinetics of a Vero cells adapted Bangladeshi strain of *peste des petits ruminants* (PPR) virus in cell culture

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Abstract

Growth kinetics of a Vero cells adapted Bangladeshi strain of *peste des petits ruminants* virus was studied in Vero cells to determine maximum virus yield. One-step growth curve was formulated after determining virus in both supernatant (CFV) and cell lysate (CAV) at different time categories by microtitre plate titration in Vero cells and the viral presence was confirmed by real-time RT-PCR. The virus was first detected in both the supernatants and cell pellets at 12 hpi. The virus titre reached its plateau at 72 hpi. Maximum virus titre of CAV was 6.2 log_{10} TCID_{50}/ml and that of CFV was 5.2 log_{10} TCID_{50}/ml at 72 hpi. After that, the titer gradually declined, but maintained at 4.5 log_{10} TCID_{50}/ml in case of CAV and 4.2 log_{10} TCID_{50}/ml in case of CFV at 96 hpi. It was concluded that the optimum time point for harvesting Vero cell culture is 72 hpi.

Keywords Vero cell · PPR virus · Virus titre · Growth curve

Introduction

*Peste des petits ruminants* (PPR) virus belongs to genus Morbillivirus under the family Paramyxoviridae, popularly known as Goat plaque of small ruminant as characterized by fever, anorexia, ulcerative necrotic stomatitis, diarrhea, purulent ocular and nasal discharges, pneumonia and respiratory distress, and death (Yongqiang et al. 2015; Chowdhury et al. 2014). Mortality and morbidity may reach up to 90 and 100%, respectively (Chowdhury et al. 2014). In Bangladesh, PPR is endemic since 1993. PPR is considered as the major hindrance of goat rearing as economic loss due to PPR in small ruminants 1000 USD annually (DLS 2010). The viral replication occurs in the host cell cytoplasm and the virus is released by budding (Cromeans et al. 1989; Takimoto and Portner, 2004). Virus use the machinery and metabolism of a host cell to produce multiple copies of themselves, and they assemble in the cell (Yin and Redovich 2018). When infected, the host cell is forced to rapidly produce thousands of identical copies of the original virus. However, these mechanisms are not very efficient in the viruses of Paramyxoviridae family and a substantial amount of the produced viruses is kept associated with the host cell membranes. During cell culture, the viral RNA can stay free in the cell and be replicated as such, or it can be incorporated into the host chromosome and be replicated simultaneously with it. Viral proteins are next synthesized with the host’s machinery under the direction of viral RNA and the new virus particles are assembled mechanically. These particles can find their way out of the cell or lysate, and be released into the medium, ready to infect new cells (Más and Meloro 2013). It is essential to measure the viral load in both supernatant (cellfree virus—CFV) and cell lysate (cellassociated virus—CAV).

To reduce the burden due to PPR an effective, potent vaccine production is essential; therefore, adaptation of virus
in cell culture, viral replication, as well as high virus yield are equally important in vaccine production strategy. Classic study of this kind defined the one-step growth curve, in which cells in a culture are infected simultaneously using a high multiplicity of infection and the increase in infection virus over time is followed by sequential sampling and titration. Virus that is free in the medium can be titrated separately from viruses that remain in cell lysate (Frederick et al. 1999).

The titre of PPR virus increases significantly if the virus is passaged in Vero cells serially. Adaptation of PPR virus on Vero cells may be suitable for large-scale production of antigen. The field isolates of PPRV could be passaged in Vero monolayer cell culture, which showed characteristic CPE and were readily adapted between 5 and 7th passages. The CPE usually develops within 48 h. Production of PPRV vaccine in tissue culture is widely used all over the world. To get maximum yield of virus, which is essential for potent vaccine formulation, the virus has to be harvested at the proper time following cell infection. To determine such optimum time, the growth curve of the virus in infected tissue culture, the development of complement-fixing antigen has to be studied. Therefore, the present study was designed to determine the multiplication profile of infectious viral particles in PPR-infected Vero cells at 60th passage level.

Materials and methods

Place of work

The research work was done at Gene laboratory and Cell culture and Virology laboratory of the Department of Pathology, Faculty of Veterinary Science, Bangladesh Agricultural University, Bangladesh.

Virus

Locally isolated Bangladeshi strain of peste des petits ruminants (PPR) virus (BD/PPR/08) had been attenuated by 60 serial passages in Vero cell culture was used in this study.

Tissue cultures

Vero cells (CLS, Germany; order no. 605372) were brought and used in this study.

Cell culture media and reagents

Cell culture medium-M-199 (Gibco-Invitrogen, cat no. 11825) and calf serum-FBS (Gibco-Invitrogen, cat no. 10437) were used for culture and maintenance of Vero cells. Cell culture working medium was made by adding l-glutamine, HEPES buffer, Sodi-bi-carb, antifungal solution (Fungizone), antibiotic (gentamicin), and distilled water with commercially available M-199 solution. 0.25% Trypsin with EDTA (Gibco-Life technologies 20,367, C13) were used in this study for dissociation and detachment of cells from cell culture flask during subculture.

Growth curve design

Cells were grown to approximately 90% confluency in petridish (Nunc, Denmark) and infected with PPRV at a fixed multiplicity of infection (MOI) of 0.01. All petridishes are marked and grouped into different time category e.g., 12, 24, 36... 96 h. After incubation at 37 °C for 1 h with rocking every 10 min, the virus suspension was discarded and cells were washed thrice with 1x phosphate-buffered saline (PBS) to remove unadsorbed virus. The end of the adsorption period was considered zero time of post-infection (p.i.). Cells were fed with M-199 containing 5% FBS and returned to the incubator. Cell lysate (CAV) and corresponding supernatants were harvested at different time interval as described below. The medium from one petridish, selected randomly, was harvested using a pipette and stored immediately at – 70 °C as cell-free virus (CFV). Attached cells from the petridish were harvested with a cell scraper (Nunc, Denmark), washed four times with 1x PBS, and suspended in 1.0 ml M-199 supplemented with 5% FBS. The cell suspension was freeze-thawed thrice, centrifuged at 2790xg for 10 min, discard supernatant, and stored the cell suspension at – 70 °C as cell lysate or cell-associated virus (CAV). The same procedure was followed for all the subsequent samples.

Infectivity titration

After completion of the experiment, samples (both CFV and CAV) were thawed at 4 °C, diluted serially (tenfold dilution), and titrated in Vero cells. Cells were observed daily for CPE and 50% tissue culture infective dose (TCID₅₀/ml) was determined by microtitre assay as described by Reed and Muench (1938)

Confirmation of PPR viral RNA by real-time RT-PCR

The presence of virus in the harvested samples (both CFV and CAV) was first confirmed by real-time RT-PCR. For this, one-Step RevTrans-qRT-PCR Evagreen (No Rox) kit (Bio sell Inc., Germany) and AB 7500 Fast Real Time PCR were used. Briefly, the methods of real-time RT-PCR was as follows-

Syber green real-time RT-PCR for F& N genes were performed in AB 7500 Fast Real-Time PCR Machine using One-Step RevTrans-qRT-PCR Evagreen (No Rox) kit (Bio Sell, Germany). PPRV F1b, PPRV F2d primer for F gene
and NP3 and NP4 primer for N gene were used in this study (Ozkul 2002; Couacy-Hymann et al. 2002). The master mix for syber green real-time RT-PCR was prepared as reaction volume 25.0 μl comprising of both forward and reverse primer (100 pmol/μl) 1.0 μl, 2X RT-qPCR EvGr-Reaction Mix 12.5 μl, Rnase free water 8.5 μl, and template RNA 2.0 μl.

The syber green real-time RT-PCR was performed in MicroAmp Optical Fast 0.1 ml 8-tube strips. First, 20 μl master mix was dispensed to each tube of the strip. Then, 5 μl RNA of the samples, positive control, and negative control was added to respective wells and mixed with the help of pipette tips. The tubes were closed with cap strip and the tube strip was placed to the tube rack in the machine. The optical setting was selected for syber green detection and the test was run with the preset thermal profile includes reverse transcription stage at 50 °C for 15 min, Initial denaturation at 95 °C for 5 min and 40 cycles of PCR (denaturation at 95 °C for 15 s, annealing at 60 °C for 20 s, and extension at 72 °C for 30 s). The results were presented as the C_{t} value for each sample, which was determined from the amplification plot.

Results

One-step growth curve was performed on Vero cells to determine the growth characteristics of the PPRV isolates. One-step growth curve is a plot typical of the rapid growth of a virus in cell culture when all cells are infected simultaneously. Released (supernatant or extracellular virus or cellfree virus—CFV) and lysates or cell-associated viruses (CAV) were harvested at different time intervals in between 12 and 96 hpi and their titres were determined. Viruses were detected in both culture supernatants and cell lysates at 12 hpi. The latent phase in both lysates and supernatant was found short, but the exponential phase was rapid because of using Vero cell adapted PPRV tissue culture fluid, i.e., 60th passaged PPR virus tissue culture fluid (PPRV- TCF) used in growth curve study (Fig. 1).

The multiplication profile of infectious viral particles in PPR-infected Vero cells at 60th passage level was determined by growth kinetics. Figure 1 shows the virus growth pattern in the supernatant and cell lysate. Maximum virus titre observed in the lysates or cell pellet, i.e., cell-associated virus (CAV) was 6.2 log_{10} TCID_{50}/ml and in supernatant, i.e., cellfree virus (CFV) was 5.2 log_{10} TCID_{50}/ml on 72 hpi of 60th passage level, respectively. In this study, an increase in virus yield from 12 hpi to 72 hpi, with a maximum titer of 6.2 log_{10} TCID_{50}/ml was observed on 72 hpi, and beyond which the titer did not increase but was maintained as 4.5 log_{10} TCID_{50}/ml up to 96 h of observation in case of lysates (Fig. 1), and in case of supernatant, it was maintained as 4.2 log_{10} TCID_{50}/ml up to 96 hpi.

The commencement of CPE in Vero cell lines was observed at 12 hpi at 60th passage level and the pattern of induction of CPE resembled that observed in previous passages. The CPE was characterized by rounding, ballooning of cells followed by foci or small syncytia. During later
stage, large syncytia formation and disintegration of cells leading to total detachment of cells monolayer were found. 100% CPE was achieved at 72 hpi.

A real-time RT-PCR method was applied to estimate the presence of PPRV during 12–96 hpi (Fig. 2a, b). Melting curve was considered in real-time RT-PCR results. Minimum Ct value observed in the cell lysates or pellet, i.e., cell-llassociated virus (CAV) was 17.88 and in supernatant, i.e., cellfree virus (CFV) was 18.77 on 72 hpi of 60th passage level. In this study, a decrease in Ct value from 12 to 72 hpi was observed and beyond, which the Ct values did not decrease, would rather increase up to 25.05/25.74. Ct values with corresponding titres of all time categories of hpi for both supernatant and cell lysate in growth kinetics study are shown in Table 1.

Here, S1: 60th passage tissue culture fluid (TCF) of isolate BD_PPR_08, S2: cell lysate at 72 hpi, S3: cell lysate at 60 hpi, S4: tissue culture supernatant of 60th passage of isolate BD_PPR_08, S5: cell lysate at 84 hpi, S6: cell lysate at 48 hpi, S7: supernatant at 48 hpi, S8: supernatant at 60 hpi, S9: cell lysate at 96 hpi, S10: supernatant at 36 hpi, NC: threshold line, PC: positive control.

Discussion

To determine the proper time at which the harvested virus could contain the required amount of intact virus to produce a good and potent vaccine, maximum virus titer, the commencement of CPE, pattern of induction of CPE, the virus growth kinetics are still desirable (Jadi et al. 2010). A high virus yielding host system is equally important in large-scale propagation of viruses for diagnostic and vaccine development studies. In this study, single or one-step growth curves were undertaken on Vero cells to determine the growth characteristics of PPR viruses. Released (supernatant or extracellular virus or cellfree virus—CFV) and cell lysate virus or cell-llassociated viruses (CAV) were harvested at 12–96 hpi and their titers were determined. Viruses were detected in both the supernatant and cell lysate in 12 hpi. The latent phase or stages in both supernatant (CFV) and cell lysate (CAV) were found short, but the exponential phase is rapid. This might be due to using Vero cell adapted PPRV, i.e., 60th passed PPRV-TCF in growth curve study. On assay of the supernatant and cell extract collected at different time intervals as described above, a slow increasing pattern of PPR virus titre in culture supernatant was observed during the first 18 h of growth cycle (eclipse phase of virus multiplication was short as Vero cells adapted virus were used). The cellfree virus (CFV) gradually increased after 12 h and a maximum virus titre was achieved around 3 days (72 hpi), and it was 5.2 \( \log_{10} \) TCID\textsubscript{50}/ml at 72 hpi. Virus could be detected between 12 and 18 h in cell lysate, and a maximum virus titre was achieved around 3 days (72 hpi) and it was 6.2 \( \log_{10} \) TCID\textsubscript{50}/ml. Subsequently, the titre of cell lysate virus or intracellular or cell-llassociated virus (CAV) was always somewhat higher than that of the virus in supernatant, i.e., cellfree virus (CFV) during the entire phase virus of active replication of the virus (between 2 and 3 days) followed by a plateau. All the findings were found related closely to the findings of multistep growth kinetics study of two isolates of 5th passage level PPRV (PPR-Izatnagar/94 and PPR Sungri/96) in Vero and Marmoset B95a cells or B lymphoblastoid cells at an MOI of 0.001 by Sreenivasa et al. (2006).

The replication kinetics and CPE development of viruses were found to depends upon the several factors such as (i) cell line used, (ii) the passage level or adaptation level, and (iii) method of selection of the virus population. The growth kinetics study of Vero cells adapted PPR virus in this study (at 60th passage level) revealed that the titres were 5.5 \( \log_{10} \) TCID\textsubscript{50}/ml on 48 hpi in case of cell lysate and 5.2 \( \log_{10} \) TCID\textsubscript{50}/ml on 72 hpi in case of supernatant. Both findings were found in accordance with the findings of Chandrahass et al., 2014 at 3\textsuperscript{rd} passage level where titre of tissue culture fluid ranges 3.5–4.5 \( \log_{10} \) TCID\textsubscript{50}/ml. In this study, the titre for cell lysate or cell-llassociated virus (CAV) reached 5.5 \( \log_{10} \) TCID\textsubscript{50}/ml and for released virus or supernatant or cellfree virus (CFV) 4.7 \( \log_{10} \) TCID\textsubscript{50}/ml on 84 hpi which were closely related the titre found in a study (Leonard et al. 2008) in regard to growth dynamics of another morbillivirus; measles virus in Vero/hSLAM cells which were 5.0 \( \log_{10} \) TCID\textsubscript{50}/ml on 84 hpi and 4.0 \( \log_{10} \) TCID\textsubscript{50}/ml on 108 hpi in case of cell lysate and supernatant respectively, though titres on 108 hpi in case of supernatant were not measured in this study. The titre observed in this study was found similar to the titre produced by the measles virus in Vero cells in another study which was approximately 5.0 \( \log_{10} \) TCID\textsubscript{50}/ml on day 4 of pi. at an MOI 0.01 (Takeuchi et al. 2002). In this study, it showed an increase in virus yield from 12 to 72 hpi, with a maximum titer observed at 72 hpi, and beyond which the titer did not increase. This finding was found close to the findings of Matumoto (1966) who first noted a rise in titer at 20 hpi, and the maximal titer was reached in 2–4 days post-infection (dpi) by measles virus in Monkey kidney cells and in 4 dpi by the same virus in Vero cells (Takeuchi et al. 2002) at an MOI 0.01. The infective titre of cell lysate virus in the culture increases more rapidly, is always much higher, and reaches a plateau earlier, than that in the fluid phase.
Fig. 2  a Confirmation of PPRV in both cell lysates and supernatant by real-time RT-PCR at different time categories. b: Confirmation of PPRV in both cell lysates and supernatant by real-time RT-PCR at different time categories.
The infective titres in both cells and fluid show increasingly earlier rise and greater rates of increase, and reach plateaus earlier, as the input multiplicity increases. This finding was in agreement with the research findings of other morbilli viruses as measles virus in Monkey kidney cell culture (Matumoto 1966), in mouse-derived cell cultures (Kohno et al. 1968).

Growth curve analysis of recombinant rinderpest virus (RPVs) containing chimeric H proteins was conducted by Parida et al. (2006) where Vero cells were infected with viruses such as RPV2C, PPRV, or RPV2C-PPRTm (at MOI 0.1) or virus RPV2C-PPRExt (at MOI 0.01), and the same titre was found on 72 hpi which was 3.5 log10 TCID50/ml. Marmoset B95a and Vero-SLAM cells were infected simultaneously with two rinderpest virus and titres were found between 5.0 log10 TCID50/ml and 6.0 log10 TCID50/ml by 96 hpi (Banyard et al. 2010).

In another study of rinderpest virus growth profile conducted by Nores and McCullough (1997) observed, the highest virus titres were obtained with the RPV-Saudi isolate at 10 dpi. in infected bovine monocytes. In another study by Aravind et al. (2015), the multiplication profile of infectious viral particles in Duck enteritis virus (DEV) infected Vero cells at 15th passage level was determined by growth kinetics. In that study, it showed an increase in virus yield from 6 to 48 hpi with a maximum titer of 5.6 log10 TCID50/ml observed at 48 hpi, and beyond which the titer did not increase, but was maintained as 4.0 log10 TCID50/ml up to 96 h of observation (Aravind et al. 2015). In growth kinetics study of Chandipura virus using Vero E6 cell line, maximum virus titre observed in the Vero cell pellet was 104.0 TCID50/ml at 6 hpi and in supernatant was 6.0 log10 TCID50/ml at 9 hpi. In this study, maximum titre was found on 72 hpi and more titre was found in cell lysate (CAV) than supernatant (CFV).

Based on these, to get maximum virus yield, cell lysate (CAV) along with supernatant (CFV) could be harvested on 72 hpi.

**Conclusion**

It has been concluded that, in this study, attenuated PPR virus in Vero cells showed high titre of progeny virus on 72 h of pi which could be the appropriate time of harvesting infected tissue culture fluid during production of an effective, potent vaccine.

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**Availability of data and materials** Available on request.

**Compliance with ethical standards**

**Conflicts of interest/competing interests** No conflict of interest.

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