Optimization of Newcastle Disease Virus Production in T-flask

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Abstract—In this work, the propagation of Asplin F strain of Newcastle disease virus (NDV) in DF-1 cells was investigated. Experiments were carried out in T-flask to investigate the effects of serum concentration in the culture medium during virus replication phase and multiplicity of infection (MOI) on virus production. Virus infectivity titre of $6.62 \times 10^7$ TCID₅₀/ml was achieved when serum concentration of 0.5% and MOI of 20 was used.

Keywords-component; Newcastle disease virus; Asplin F strain; DF-1 cell; T-flask

I. INTRODUCTION

Newcastle disease (ND) is regarded as one of the most important disease in the poultry industry [1][25]. The disease which had its first outbreaks in 1926, in Java, Indonesia [12], and in Newcastle-upon-Tyne, England [7] is highly contagious affecting 27 of the 50 orders of birds [27]. Chickens are the most susceptible host, in which the severity of the disease may vary from mild infection with no apparent clinical signs to a severe form causing 100% mortality. The causative agent responsible for the disease is the Newcastle disease virus (NDV) [2]. NDV which is also known as avian paramyxovirus 1 is a member of the genus Avulavirus of the family Paramyxoviridae, in the order Mononegavirales [19]. This family also includes other important pathogens such as the mumps virus, human parainfluenza virus, sendai virus, simian virus 5 and recently emerging nipah and hendra viruses [13].

NDV strains are classified into three main pathotypes; lentogenic, mesogenic, and velogenic, based on the clinical signs and severity of the disease. Lentogenic strains cause mild or avirulent infections that are largely limited to the respiratory. Mesogenic strains are of intermediate virulence causing respiratory infection with moderate mortality while velogenic strains are highly virulent causing 100% mortality in chickens. Velogenic strains can be further categorized into two types: viscerotropic and neurotropic. Viscerotrophic velogenic strains produce hemorrhage in the digestive tract whereas neurotropic velogenic strains produce neurological and respiratory disorders [2][25].

The disease caused by NDV remains a potential threat to commercial or backyard production until today. In 2002, an outbreak caused by a virulent strain of NDV occurred in California, USA. The outbreak has caused $200 million worth of losses from the depopulation of birds [11]. Although the disease is not yet eradicated, it can be controlled either by importing birds from disease free flocks or through vaccination that must continue throughout the life of the bird [19].

Currently, vaccines for ND are produced by using embryonated chicken eggs, a technology that has remained almost unchanged since the late 1930s. This traditional method however poses some drawbacks, such as the need for high amounts of specific pathogen-free eggs, high labor-intensity, time consuming and requires big area for the incubation of eggs. Besides the process is slow and difficult to scale-up, so large strategic stocks must be kept to respond in cases of epidemics [24]. An alternative to this method is the propagation of NDV in cell culture systems. It has been reported that Newcastle disease virus strains are able to replicate in an enormous range of cells [14]. Among cell substrate systems that have been identified are Vero cell, CEF cell and DF-1 cell [5][6][22].

In the present work, the production of Newcastle disease virus (lentogenic Asplin F strain) in T-flasks using DF-1 cells as hosts was studied. The influence of factors such as serum concentration during virus replication phase and multiplicity of infection, MOI on virus yield was investigated. This study serves as the foundation for future live Newcastle disease vaccine production in larger cell culture systems such as the bioreactor.

II. METHODOLOGY

A. Cell line and virus strain

Established DF-1 cell line (ATCC-CRL-12203™) was purchased from the American Type Culture Collection (ATCC). Lentogenic Asplin F strain of NDV was obtained from Malaysia Vaccine Pharmaceuticals (MVP) Sdn. Bhd.
B. Culture medium and chemicals

Dulbecco’s Modification of Eagle’s Medium, DMEM (with glucose and L-glutamine) in powder form and fetal bovine serum (FBS) were supplied by Gibco®.

C. Cell infection with NDV in T-flask

Confluent monolayers of cells were used for infection with NDV as described by Hussain and Rasool [10]. Spent medium was removed and cell monolayer was washed with 5 ml of PBS. Later cells were infected with NDV according to the designed levels by inoculation of virus in DMEM supplemented with trypsin. The virus inoculums was spread uniformly and incubated in humidified CO\textsubscript{2} incubator for 1 hour with intermittent rotation to allow adsorption. Five milliliters of maintenance medium with designed concentration of serum was added to flask. The flask was later returned to humidified CO\textsubscript{2} incubator and monolayers were examined daily under inverted microscope for evidence of CPE.

D. Virus harvesting

The virus was harvested by combination of freeze thawing, sonication and centrifugation process. Four days post inoculation; flasks were transferred to -80°C freezer for 1 hour and later thawed at room temperature. This process was then resumed with sonication at 15°C for 1.5 minutes. Subsequently, virus suspensions were poured into centrifuge tubes and centrifuged at 300g for 10 minutes at 4°C to pellet the cell debris. The clear supernatant fluid containing virus was collected carefully, labeled and kept in -80°C freezer.

E. TCID\textsubscript{50} assay

Fifty percent tissue culture infectious dose (TCID\textsubscript{50}) assay was carried out in 96 well flat shaped microtiter plates. Plates were seeded with DF-1 cells such that confluence was reached in 2–3 days. DF-1 cells were grown in DMEM media containing 10% FBS and grown at 37 °C, 5% CO\textsubscript{2}. Appropriate dilutions of each virus were prepared in DMEM medium and 100μl of each dilution was added to each well. Plates were incubated at 37°C for 30–60 min before 100μl of DMEM was added to each well. The plates were then incubated for 4 days until the CPE effect was appeared. The log TCID\textsubscript{50}/ml calculation was based on the Reed and Muench [23] formula.

F. Experimental design

Optimization of NDV propagation in T-flask was carried out according to 3***(2-0) full factorial design (FD) generated by STATISTICA®. Two selected parameters; serum concentration and MOI were varied in this experiment to investigate their optimum value in response to virus infectivity titre produced. The design consists of 9 main runs, 4 replicates at the centre point and each parameter have 3 levels. The layout of the design is shown in Table 1.

| Table 1. 3***(2-0) Full factorial design for ndv propagation in T-flask |
|---|---|---|
| Serum concentration (%), A | 0.5 | 1.5 | 2.5 |
| MOI, B | 0.2 | 2 | 20 |

III. RESULTS AND DISCUSSION

A. Determination of time of infection (TOI)

To propagate viruses in cell culture, a suitable time of infection (TOI) is required. To this regard, the growth profile of DF-1 cell in the T-flask was studied and the appropriate time to infect with the virus was thereof determined.

As shown in Figure 1, culture of DF-1 was started by inoculation of 1.00 x 10\textsuperscript{7} cells/ml into the flask. Growth of DF-1 cells were in lag phase from 0 hour until 48 hours. Starting from 48 hours, the culture entered exponential phase and reached the maximum cell concentration of 1.29 x 10\textsuperscript{5} cells/ml at 64 hours. Once reached the maximum, cell concentration dropped and entered the death phase.

Based on this, TOI was determined to be between 56 hours and 64 hours where the culture will be in the exponential phase. During this period, cells which will serve as the virus replication hosts are highly available and this will contribute to higher virus production.

Figure 2 shows the morphology of DF-1 cells before and after the infection by NDV. As can be observed, characteristic changes in the appearances of DF-1 cells were very obvious in which formation of giant multinucleated cell or syncytia, formation of plaques and also elongation of cytoplasmic tails has occurred after the infection. These changes are known as cytopathic effect or CPE. CPE is defined as pronounced morphologic changes induced in individual cells or groups of cells by virus infection, which are easily recognizable under a light microscope. CPE are very reproducible and can precisely character the virus type providing significant clues to the identity of a virus [4].

B. Optimization of NDV production in T-flask

Virus infection parameters, serum concentration during virus replication phase and multiplicity of
infection (MOI) as independent factors were optimized to yield high virus infectivity titre. Experiments were carried out as per the design matrix of the 3**2 full factorial design (FD) (Table II), and the final virus titre produced by the culture was used as the response. For predicting the optimal values of virus titre obtained within the experimental constrains, a second order polynomial model was fitted to the experimental results by using the STATISTICA® software. The model developed is as follows:

\[
Y = \text{Virus infectivity titre, TCID}_{50}/\text{ml} = -8222414 + 20301437(A) - 2781034(A^2) + 19951268(B) - 815773(B^2) + 920045(AB)
\]

where the virus infectivity titre as yield (Y) is a serum concentration (A), and MOI (B).

The statistical model was checked by Fischer variance ratio, the F-value and the analysis of variance (ANOVA) for the response surface quadratic model is summarized in Table III. In Table III, the model F-value of 36.7793 implies that the quadratic regression model was significant. F-value is a statistically valid measure of how well the factors describe the variation in the mean of data. The greater the F-value from 1 the better the factors explain the variation in the data about its mean, and the estimated factor effects are real [17]. Model was further approved to be significant by having a very low probability value \((P_{\text{model}} > F) = 0.0001\).

At the model level, the correlation measures for the estimation of the regression equation are the correlation coefficient R and the determination coefficient R². The correlation between the experimental and predicted values is better when the value of R is closer to 1 [9][26]. In this experiment, the value of R and R² were 0.98149 and 0.96333 respectively. These values indicate a high degree of correlation between the experimental and the predicted values. The value of R² indicates that 96.333% of the factors: serum concentration and MOI contribute very positively to the response. The value of R² is also a measure of fit of the model and it can be mentioned that only 3.667% of the total variation were not explained by the virus infectivity titre. The value of the adjusted determination of coefficient was also very high (0.93714) which indicates high significance of the model.

The P values are used as a tool to check the significance of each of the factors which, in turn, are necessary to understand the pattern of the mutual interactions between the factors. The lesser the P value than 0.05, the bigger the significance of the corresponding factor [16]. Based on this, the factors that were highly significant in the model were the MOI concentration (B), square terms of MOI (B²) and the interactive term between serum concentration and MOI (AB). Serum concentration (A) and its square term (A²) were not significant because their P values was larger than 0.05. However, in this study, all terms were included due to the significance of overall model.

**Table 2.** Experimental design using 3**2 full factorial design (FD) with experimental and predicted (using model equation) values of virus infectivity titre.

<table>
<thead>
<tr>
<th>Standard run</th>
<th>Serum conc. (%)</th>
<th>MOI</th>
<th>Virus infectivity titre (TCID₉₀/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Experimental</td>
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<td></td>
</tr>
<tr>
<td>Predicted</td>
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<td></td>
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<tr>
<td>1</td>
<td>2.50000</td>
<td>2.00000</td>
<td>5.76 x 10⁻¹</td>
</tr>
<tr>
<td>2</td>
<td>1.50000</td>
<td>20.00000</td>
<td>5.89 x 10⁻¹</td>
</tr>
<tr>
<td>3</td>
<td>0.50000</td>
<td>0.20000</td>
<td>1.11 x 10⁻¹</td>
</tr>
<tr>
<td>4</td>
<td>0.50000</td>
<td>20.00000</td>
<td>6.62 x 10⁻¹</td>
</tr>
<tr>
<td>5</td>
<td>2.50000</td>
<td>0.20000</td>
<td>2.75 x 10⁻¹</td>
</tr>
<tr>
<td>6</td>
<td>1.50000</td>
<td>0.20000</td>
<td>1.48 x 10⁻¹</td>
</tr>
<tr>
<td>7</td>
<td>1.50000</td>
<td>2.00000</td>
<td>5.13 x 10⁻¹</td>
</tr>
<tr>
<td>8</td>
<td>0.50000</td>
<td>2.00000</td>
<td>2.95 x 10⁻¹</td>
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<tr>
<td>9</td>
<td>1.50000</td>
<td>2.00000</td>
<td>5.14 x 10⁻¹</td>
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</tr>
<tr>
<td>11</td>
<td>2.50000</td>
<td>20.00000</td>
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</tr>
<tr>
<td>12</td>
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<td>2.00000</td>
<td>5.12 x 10⁻¹</td>
</tr>
<tr>
<td>13</td>
<td>1.50000</td>
<td>2.00000</td>
<td>5.13 x 10⁻¹</td>
</tr>
</tbody>
</table>

**Table 3.** Analysis of Variance (ANOVA) for Response Surface Quadratic Model for optimization

<table>
<thead>
<tr>
<th>Source</th>
<th>Sum of squares</th>
<th>F value</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>3.520427 x 10⁻⁷</td>
<td>36.7793</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Serum concentration, A</td>
<td>3.907699 x 10⁻⁷</td>
<td>2.0413</td>
<td>0.196144</td>
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<tr>
<td>MOI, B</td>
<td>2.575082 x 10⁻⁷</td>
<td>134.5168</td>
<td>0.000008</td>
</tr>
<tr>
<td>A²</td>
<td>2.136099 x 10⁻⁷</td>
<td>1.1159</td>
<td>0.325900</td>
</tr>
<tr>
<td>B²</td>
<td>1.474961 x 10⁻⁷</td>
<td>77.0489</td>
<td>0.000050</td>
</tr>
<tr>
<td>AB</td>
<td>4.059055 x 10⁻⁷</td>
<td>21.2036</td>
<td>0.002470</td>
</tr>
</tbody>
</table>

Figure 3 shows the virus infectivity titre predicted by the model and the real values obtained from the experiment. It can be observed that the point’s distribution around the line for the response fitted to the linear model.
Figure 3. Plot of observed versus predicted values for virus infectivity titre

The 3D response surface plots described by the regression model were drawn to illustrate the effects of the independent factors and the interactive effects of each independent factor on the targeted response. The shape of the corresponding 2D contour plots indicates whether the mutual interactions between the independent variables are significant or not. An elliptical contour plots indicates that the interactions between the independent factors are significant. While a circular contour plot indicates that the interactions between related factors are negligible [17]. By analyzing the 3D response surface plots and the corresponding 2D contour plots, the optimal values of the independent factors could be observed, and the interaction between each independent factor can be easily understood [15]. Figure 2 shows elliptical contour plots since the interactive term between serum concentration and MOI, AB was significant (refer to Table 2). The maximum virus infectivity titre can be obtained at the point of intersection of the major and minor axes of the ellipse or by solving the inverse matrix of Eq. 1. By using STATISTICA®, the optimum values for each factor were determined; 1.79% for serum concentration and 11.22 for MOI which will yield the maximum virus infectivity titre of $1.22 \times 10^8$ TCID$_{50}$/ml.

Figure 4: 3D response surface and 2D contour plots shows the effect of serum concentration (%) and MOI on virus infectivity titre

Genzel et al. [8] has reported that the multiplicity of infection (MOI) influences virus growth dynamics but not final virus yield. In the study of propagation of influenza virus in Vero and MDCK cells conducted by Audsley and Tannock [3], it has been observed that cells infected with higher MOIs maximum titers were attained earlier than in cultures infected at lower MOIs. While according to Maranga et al. [18] in his study of production of virus-like particles with a baculovirus insect cell system, he noticed that at high MOIs the specific productivity decreased when cells were infected at late growth phase.

Different theories have encouraged us to investigate the effects of MOI on Newcastle disease virus production. As shown in Table II, three different MOIs, 0.2, 2.0 and 20 were used. Based on the result, the maximum virus infectivity titre of $6.62 \times 10^7$ TCID$_{50}$/ml was achieved when we used high MOI, 20 (while serum concentration used was 0.5%). High MOI probably better suited NDV production in T-flasks. While for serum concentration during viral replication phase, it has been analyzed by STATISTICA® that the factor was not significant towards virus infectivity titre. In addition, serum was suggested to be removed from the culture medium to facilitate the growth of virus in the host cells [21].

Apart from serum concentration and MOI, other factors such as type of cell, type of culture medium and time of infection also may partially contribute to the results we
obtained in this experiment. Investigation on their effects on virus production is much recommended.

IV. CONCLUSION

In the present work, two factors of interest for the establishment of a new platform for the production of Newcastle disease virus were investigated. It was verified by statistical analysis using STATISTICA®, that multiplicity of infection, MOI has substantial effect on the virus infectivity titre. High virus infectivity titre of 6.62 x 10⁷ TCID₅₀/ml was achieved when high MOI of 20 was used. Serum concentration during virus replication however was analyzed to have minor effect on the virus titre. Also it has been analyzed that the optimum conditions to produce Newcastle disease virus using DF-1 cell culture are serum concentration of 1.79% and MOI of 11.22. These results constituted important information when large scale production of Newcastle disease virus is considered.

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