The Effect of the Infection Rate on Oncolytic Virotherapy

Dongwook Kim¹*, Haeyoung Kim², Hui Wu³, Dong-Hoon Shin⁴

¹Department of Mathematics, Texas A&M University-Kingsville, Kingsville, United States of America
²Department of Biological and Health Sciences, Texas A&M University-Kingsville, Kingsville, United States of America
³Department of Mathematics, Clark Atlanta University, Atlanta, United States of America
⁴Department of Global Finance and Banking, Inha University, Incheon, South Korea

Email address:
Dongwook.Kim@tamuk.edu (D. Kim)
*Corresponding author

To cite this article:

Received: June 10, 2020; Accepted: June 23, 2020; Published: July 4, 2020

Abstract: Oncolytic viruses have become a novel therapeutic tool for various cancer treatments. Several naturally occurring oncolytic viruses and engineered oncolytic viruses are developed for oncolytic virotherapies. Although we have a good understanding on molecular mechanisms of viral replication and virus-induced cell lysis at the cellular level, it is unclear how oncolytic viruses and cancer cells interact as a population. Several mathematical models of oncolytic virotherapy have been developed to advance the understanding of dynamic interaction between oncolytic viruses and cancer cells. Many authors investigated the effect of the virus replication on dynamics of cancer cell population and proposed that the bursting rate of viruses is an important factor for successful oncolytic virotherapy. In this study, we investigate the effect of infection rate of oncolytic viruses on an oncolytic virotherapy model. Particularly, we focused on studying the relationship between two control parameters, bursting rate and infection rate of the virus, to generate the patterns from equilibrium steady state to periodic solutions. Based on the model, the interaction between cancer cells and oncolytic viruses shows an intriguing two-dimensional bifurcation, showing three parameter regions (equilibrium steady state, damped oscillations and oscillations). Our result suggests that both infection rate and bursting rate are crucial properties of oncolytic viruses to design a successful oncolytic virotherapy.

Keywords: Computational Biology, Oncolytic Virotherapy, Bifurcation, Dynamical System

1. Introduction

Oncolytic virotherapy (OV) is a novel treatment by which oncolytic replication competent viruses are either inoculated to a cancer patient or directly injected into the tumor to kill cancer cells without causing harm to healthy normal cells. Upon introduction, viruses selectively infect cancer cells by binding to a specific receptor protein present in cancer cells. The antitumor activity of OV can be achieved either by direct lysis of cancer cells or by triggering a host immune response [1-3]. Upon the lysis of infected cancer cell, new virus particles burst out and infect adjacent cancer cells. Recent studies have shown that measles, adenovirus, Newcastle-disease virus (NDV), reovirus, herpes simplex virus 1 (HSV-1), and coxsackievirus are relatively non-toxic and tumor specific [4, 5]. In addition to naturally occurring viruses, viruses have been engineered to increase the specificity. For instance, an adenovirus H101 was the first engineered virus that have been approved by a government regulatory agency [6].

Many mathematical models of OV have been developed to understand complex and dynamic interactions between tumor cells and viruses. In addition, mathematical models can give an insight to design treatment regimens for a cancer intervention with minimal adverse effects. Many studies have focused on examining how variation in viral and host parameters influence the outcomes of treatments [7-14]. Wodarz et al. proposed a mathematical model to investigate the effect of virus-specific lytic cytotoxic T lymphocytes on the dynamics of OV [8]. The ordinary differential equations (ODEs) were used to describe the interactions between two types of cancer cells populations (uninfected cancer cells and infected cancer cells) according to predator-prey model including virus...
population. There are also mathematical models using partial differential equations (PDEs) [12-14].

It has been suggested that the ability of the viral replication, represented as a bursting rate, is a key factor for successful OV. In addition, the oscillations of cancer cell and virus populations are an intrinsic property of OV [11]. Compared to the ability of virus replication, little attentions have been given to the effect of infectivity, represented as an infection rate, of the virus. Furthermore, it will be crucial to understand the relationship between bursting rate and infection rate of viruses for the development of optimal OVs. In this paper, we investigated how the variable -infection rates impact- on the cancer cell population using dynamical system (phase space analysis) and bifurcation approaches. We also analyzed the population dynamics of oncolytic viruses and cancer cells while considering (or applying) both bursting rate and infection rate in a simple OV model. Our results suggest that 1) the infection rate coefficient is a bifurcation parameter; existence of two threshold values, 2) these threshold values of infection rate are dependent on the bursting rate values and 3) there are a set of values or parameter region in choosing the bursting rate and infection rate to the therapeutic way to succeed or fail.

2. Materials & Methods

2.1. Model

The OV model is a three dimensional [11]

\[
\frac{dx}{dt} = k_1x \left(1 - \frac{x+y}{K}\right) - k_2xz
\]

\[
\frac{dy}{dt} = k_2xz - \delta_1y
\]

\[
\frac{dz}{dt} = k_3y - k_2xz - \delta_2z
\]

(1)

Where \(x(t)\), \(y(t)\) and \(z(t)\) represent populations of uninfected cancer cells, infected cancer cells and free viruses, respectively. The \(k_1\) is the growth rate of cancer cell and \(K\) being the carrying capacity of a tumor. The term \(k_1x \left(1 - \frac{x+y}{K}\right)\) describes the logistic growth rate of an uninfected cancer cell population \(x(t)\). The constant value \(k_2\) represents the infection rate of the virus in the infected cancer cells so that the term \(k_2xz\) describes the rate of infected cancer cells by free viruses \(z(t)\). \(\delta_1\) represents the death rate of infected cancer cells. The \(k_3\) is the bursting rate of free virus particles. The term \(\delta_2z\) is the rate of elimination of free virus particles by various causes including nonspecific binding, generation of defective interfering particles, and host immune system.

For non-dimensionalization, we set \(\tau = \delta_1 t\), \(x = K \hat{x}\), \(y = K \hat{y}\), \(z = K \hat{z}\). Then Equation (1) become

\[
\frac{d\hat{x}}{d\tau} = \frac{k_1}{\delta_1} \hat{x}(1 - \hat{x} - \hat{y}) - \frac{k_2K}{\delta_1} \hat{z} \hat{x}
\]

\[
\frac{d\hat{y}}{d\tau} = \frac{k_2}{\delta_1} \hat{y} - \frac{k_2K}{\delta_1} \hat{z} \hat{x} - \frac{\delta_2}{\delta_1} \hat{z}
\]

(2)

We have the following model by setting the parameters;

\[
a = \frac{k_1}{\delta_1}, \quad b = \frac{k_2K}{\delta_1}, \quad c = \frac{k_3}{\delta_1} \quad \text{and} \quad d = \frac{\delta_2}{\delta_1}
\]

\[
\frac{dx}{dt} = ax(1 - x - y) - bxz
\]

\[
\frac{dy}{dt} = bxz - y
\]

\[
\frac{dz}{dt} = cy - bxz - dz
\]

(3)

All parameters are described in Table 1.

### Table 1. Non-dimensionalized model parameters. All parameters are assumed to be non-negative.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a)</td>
<td>Growth rate of cancer cell</td>
<td>(\frac{k_1}{\delta_1})</td>
</tr>
<tr>
<td>(b)</td>
<td>Infection rate of virus into cancer cell</td>
<td>(\frac{k_2K}{\delta_1})</td>
</tr>
<tr>
<td>(c)</td>
<td>Bursting rate of virus</td>
<td>(\frac{k_3}{\delta_1})</td>
</tr>
<tr>
<td>(d)</td>
<td>Death rate of free virus particles</td>
<td>(\frac{\delta_2}{\delta_1})</td>
</tr>
</tbody>
</table>

2.2. Analysis and Stability of Equilibrium

There are three fixed points (equilibrium points); two equilibrium solutions \(E_1(0,0,0)\) and \(E_2(1,0,0)\) in the positive invariant domain \(D\) and the other one \(E_3(x^*, y^*, z^*)\) in either the negative or the positive domain depending on the parameter values.

\[
E_3(x^*, y^*, z^*) = \left(\frac{d}{b(c-1)(bc-b+ad)}, \frac{a(bc-b-d)}{b(c-1)(bc-b+ad)}\right)
\]

The Jacobian matrix \(J\) of the Equation (3) is given by

\[
J = \begin{bmatrix}
  a - 2ax - ay - bx & -ax & -bx \\
  bz & -1 & bx \\
  -bz & c & -bx - d
\end{bmatrix}
\]

The Jacobian matrix at \(E_1(0,0,0)\) is

\[
\begin{bmatrix}
  a & 0 & 0 \\
  0 & -1 & 0 \\
  0 & c & -d
\end{bmatrix}
\]

The eigenvalues are \(a\), -1, and -d. Here all parameters are positive values, so that the fixed point \(E_1(0,0,0)\) is a saddle point (unstable) with a stable manifold space on \(y-z\) plane, and the unstable manifold space on the \(x\)-axis.
The Jacobian matrix at $E_2(1,0,0)$ is

$$\begin{bmatrix}
-a & -a & b \\
0 & -1 & b \\
0 & c & -b - d
\end{bmatrix}$$

The eigenvalues are $\lambda_i = -a$ and $\lambda_{2,3} = -\frac{(b+d+1)\pm\sqrt{(b+d+1)^2+4bc}}{2}$. There is no complex number in eigenvalue since $(b + d - 1)^2 + 4bc$ is positive. However, $E_2(1,0,0)$ becomes stable if $\sqrt{(b + d - 1)^2 + 4bc} < b + d + 1$ i.e., $a < \frac{d}{c-1}$. Otherwise, $E_2(1,0,0)$ is a saddle point (unstable).

The Jacobian matrix at $E_3(x^*,y^*,z^*)$ is

$$\begin{bmatrix}
-\frac{ad}{b(c-1)} & -\frac{ad}{b(c-1)} & -\frac{d}{c-1} \\
-\frac{b(c-b-d)}{b(c-1)} & 1 & \frac{d}{c-1} \\
-\frac{a(bc-b-d)}{b(c-1)} & \frac{c}{c-1} & \frac{cd}{c-1}
\end{bmatrix}$$

The characteristic equation is given by

$$P = \lambda^3 + p_1\lambda^2 + p_2\lambda + p_3$$  \hspace{1cm} (4)

where

$$p_1 = \frac{bc + ad + bcd - b}{b(c-1)}$$

$$p_2 = \frac{ad(cd + c - 1) + ad(bc - b - d)(a-d)}{b(c-1)^2} + \frac{ad(bc-b-d)(a-d)}{b(c-1)(bc-b+ad)}$$

$$p_3 = \frac{ad(bc-b-d)}{b(c-1)}$$

It is hard to find the roots of Equation (4) explicitly. We will discuss the stability of $E_3(x^*,y^*,z^*)$ with our numerical results.

Theorem $E_3$ is stable if and only if $p_i > 0$, $i=1, 2, 3$ and $p_1 \cdot p_2 > p_3$.

Proof: Let us consider that the cubic characteristic equation (Equation 4) can be factored completely in the complex plane such as $(\lambda - r_1)(\lambda - r_2)(\lambda - r_3) = 0$ where $r_1, r_2$ and $r_3$ are complex roots of Equation (4). Then

$$r_1 + r_2 + r_3 = -p_1$$

$$r_1 r_2 + r_1 r_3 + r_2 r_3 = p_2$$

$$r_1 r_2 r_3 = -p_3$$  \hspace{1cm} (5)

$\rightarrow$ If $E_3$ is stable, there are two possibilities: (a) There are three real roots, $r_1 < 0$, $i=1, 2, 3$. According to Equation (5), $p_1 > 0$, $p_2 > 0$, $p_3 > 0$ and $p_1 \cdot p_2 > p_3$. (b) There is one real root $r_1 < 0$ and two complex conjugate pair roots $r_2 = p + qi$, $r_3 = p - qi$, $p > 0$. According to Equation (5), it is easy to verify $p_1 > 0$, $p_2 > 0$, $p_3 > 0$ and $p_1 \cdot p_2 > p_3$.

$\rightarrow$ If $p_1 > 0$, $p_2 > 0$, $p_3 > 0$ and $p_1 \cdot p_2 > p_3$. Consider the eigen function $f(\lambda) = \lambda^3 + p_1\lambda^2 + p_2\lambda + p_3$. There is no change of signs between all coefficients. According to Descartes’ Rule of signs, $f(\lambda)$ has zero number of positive roots. Therefore, there are two possibilities: (a) If there are three negative real roots, $E_3$ is stable. (b) If there are on negative real root $-r$ ($r > 0$) and two complex conjugate roots $-p + qi$ and $-p - qi$,

$$f(\lambda) = (\lambda + r)(\lambda + p + qi)(\lambda + p - qi)$$

$$= \lambda^3 + (r+2p)\lambda^2 + (p^2 + q^2 + 2rp)\lambda + r(p^2 + q^2)$$

Therefore, $p_1 \cdot p_2 > p_3$, $2p((r+p)^2 + q^2 > 0$, which implies $p > 0$. Thus, the complex roots have negative real parts, which means $E_3$ is stable.

2.3. Numerical Simulation

We used the Runge-Kutta 2nd order method [15] to compute the numerical solutions in MATLAB (The Mathworks, Natick, MA). The small-time step $\Delta t = 0.05$ was used to check the accuracy of the numerical method. For numerical simulations, we set the parameters $a = 0.31$ (growth rate of cancer cell from the experimental data [16]) and $d = 0.44$ with initial conditions $x(0) = 0.5, y(0) = 0$, and $z(0) = 1.5$.

3. Results

3.1. Infection Rate Coefficient Is a Control Parameter in Changing the Structure of Cancer Cell Population

We investigated the effect of the infection rate of the virus on the cancer cell population. We set the bursting rate ($c = 5$; different values of $c$ will be discussed later) and increased the infection rate constant from 0 to 1 with step size 0.01. Our numerical result shows that the increased infection rate ($b$) results in the altered pattern of cancer cell population over time. The cancer cell population exhibits three patterns over the relative time depending on the infection rate coefficient (Figure 1A). For the infection rate $b = 0.1$, the cancer cell population approaches to the free virus equilibrium $E_2(1,0,0)$ in phase portrait (Figure 1B) or maximal capacity (green curve in Figure 1A). For $b = 0.4$, it shows fluctuations over short time period and then converges to the relative population less than the maximal capacity (damped oscillation, Figure 1C). For $b = 0.64$, the cancer cell population converges to the minimum population 0.172 (data not shown). For $b = 0.7$, the population starts to oscillate over time (red curve in Figure 1A) and the trajectory of populations circulates around a stable limit cycle in a dynamical system (Figure 1D). As the infection rate increases, the amplitude of oscillation of the cancer cell population also increases. This result indicates that the infection rate coefficient should be considered as a control parameter that can change the dynamic structure of the relative cancer cell population over time as it changes.

3.2. Transcritical and Hopf Bifurcations

Qualitative changes in the stability of fixed points
depending on a parameter are called bifurcation in the dynamics. The control parameter values where bifurcations occurred are called bifurcation points. A bifurcation diagram of cancer cell population can be obtained by changing the parameter of virus infection rate. We computed the relative cancer cell population density with different values of \( b \) over the relative time. The bifurcation diagram for the cancer cell population over the infection rate is shown in Figure 2. There are two bifurcations at \( b = b_1 \) and \( b_2 \) where the shape of the graph changes. In our numerical result, we calculated two values \( b_1 = 0.12 \) and \( b_2 = 0.64 \) (Figure 2A). We also calculated the Lyapunov exponent to determine whether cancer cell population for \( b > b_2 \) shows periodic solutions or non-periodic solutions (Figure 2B). The cancer cell population density in this parameter interval shows periodic patterns since the Lyapunov exponent is less than 0 for \( b > b_2 \).

For \( 0 < b \leq 0.12 \), the cancer cell population reaches its maximum population (capacity). This indicates that the viral population diminished due to low strength of infectivity of the virus so that the cancer cell population approaches to the maximum. In this case, it can be concluded that the virotherapy fails in that parameter regions. For \( 0.12 < b \leq 0.64 \), the cancer cell population shows oscillations for a short time period and then approaches to the values less than 1. Particularly, the population decreases to the relative population of 0.172 at \( b = 0.64 \), which means that the virotherapy is partially successful in the interval and the cancer cell population can be reduced to minimum population of 0.172. For \( b > 0.64 \), the population start to show oscillations over the relative time. Populations of the infected cancer cells and viruses also oscillate (data not shown). Oscillations of cancer cells and viruses suggests that the interaction between cancer cells and viruses are the same dynamics as the prey-predator system (coexistence in dynamical system).

A brief bifurcation diagram in terms of fixed points, discussed in section 2.2, is shown in Figure 2C. A horizontal line is used to describe the fixed points of \( E_1(0,0,0) \) and \( E_2(1,0,0) \) and a curve is used to describe the parameter dependent fixed point \( E_3(x^*, y^*, z^*) \). Here, solid and dotted
lines represent stable and unstable, respectively. For $0 < b \leq b_1$, $E_2$ is a stable fixed point and both $E_1$ and $E_3$ are saddle (unstable), so that their populations approach to the free virus equilibrium value $E_2$ in dynamical system. Particularly, $E_3$ is in the negative invariant domain that is outside of our interest region. However, $E_3$ is in the positive invariant domain $D$ and it is in the same position as $E_2$ at the bifurcation point $b = b_1$. In other words, $E_3$ moves closely to $E_2$ from the negative invariant domain to the positive invariant domain $D$ as the $b$ value increases from 0 and then $E_3$ is in the position of $E_2$ at $b=b_1$. For $b > b_1$, the stability of $E_2$ and $E_3$ changed from stable to unstable and from unstable to stable, respectively. This indicates a transcritical bifurcation occurs at $b=b_1$. For $b_1 < b \leq b_2$, both $E_1$ and $E_2$ become unstable fixed (saddle) points but $E_3$ becomes a stable spiral fixed point, so that the trajectories circulate about $E_3$ and then gradually converge to $E_3$. For $b > b_2$, all fixed points are unstable fixed points and the eigenvalues of Jacobian matrix at $E_3$ has a family of complex numbers with positive real part. The trajectories rotate around a stable limit cycle near the fixed point $E_3$ and the populations show oscillations, which implies that there is a Hopf bifurcation occurs at $b=b_2$.

**Figure 2.** Bifurcation diagram and stability of fixed points. A. Bifurcation diagram of cancer cell population density with respect to the infection rate coefficient. B. Lyapunov exponent of cancer cell population with respect to infection rate. $b_1$ and $b_2$ are two bifurcation values. C. The stability of three fixed points; $E_1(0,0,0), E_2(1,0,0)$ and $E_3(x^*,y^*,z^*)$. The system undergoes two bifurcations: transcritical at $b=b_1$, Hopf bifurcation at $b=b_2$. The dotted blue line represents the fixed point is unstable and the solid red line represents the fixed point is stable. For $0 < b \leq b_1$, $E_2$ is stable and the others, $E_1$ and $E_3$ are unstable (saddle points). For $b_1 < b \leq b_2$, $E_3$ becomes a stable fixed point but both $E_1$ and $E_2$ are unstable fixed points. For $b > b_2$, there exist periodic solutions. The bifurcation values of $b_1=0.12$ and $b_2=0.64$ are computed in MATLAB with parameters $a = 0.31, c = 5$ and $d = 0.44$.

### 3.3. Bursting Rate-dependent Bifurcation Values

We investigated the effect of the correlation between the bursting rate and the infection rate of the virus on the cancer cell population. We simulated the bifurcation diagrams for the infection rate coefficient with various fixed bursting rate
coefficients to examine how the bifurcations points \((b_1 \text{ and } b_2)\) change (Figure 3). For the fixed value of \(c=4\), the cancer cell population undergoes the first (transcritical) bifurcation at \(b_1=0.146\) and the second (hopf) bifurcation at \(b_2=0.82\) (Figure 3A). The other values are given by \((c, b_1, b_2) = (7, 0.073, 0.45), (15, 0.031, 0.205), \text{ and } (30, 0.015, 0.102)\) (Figure 3B-3D). These results provide increasing the bursting rate of the virus results in decreasing the bifurcation points \((b_1 \text{ and } b_2)\). Those bifurcation values are dependent on the bursting rate values, which implies that different values of bursting rate induce different bifurcation points in the infection rate. These results leave questions about how the correlation between the bursting rate of the virus and the infection rate of the virus affects the success or failure of the oncolytic virotherapy.

Figure 3. Bifurcation values depend on the bursting rate. (A)-(D) are bifurcation diagrams for relative cancer cell population density with respect to the infection rate with bursting rate values \(c = 4, 7, 15 \text{ and } 30\), respectively. Increasing the bursting values result in decreasing values of two bifurcation values \(b_1 \text{ and } b_2\). Parameters used were \(a = 0.31\) and \(d = 0.44\).

3.4. Parameter Regions from Equilibrium Steady State to Periodic Solutions

The oncolytic model undergoes two bifurcations with respect to two parameters; the bursting rate and the infection rate of the virus. The transcritical bifurcation occurs at \((b, c) = (b_1, c_1)\) where the equation form of the bifurcation point clearly can be formulated from the stability of the fixed point \(E_2(1,0,0)\) and is given by \(b_1 = \frac{d}{c-1}\). In other words, the set of bifurcation points \((b_1, c_1)\) are the set of points on the graph of \(c = \frac{b+d}{b}\) in the \(b-c\) plane. Therefore, the set of parameters \((b, c)\), which satisfies the inequality \(c \leq \frac{b+d}{b}\), represents the region where the cancer cell population converges to the maximum carrying capacity and the virotherapy is a failure. Otherwise, the therapy is partially successful, or all populations coexist showing oscillations over the relative time. Figure 4A shows the two-dimensional bifurcation diagram. The diagram shows three different colored regions which represent the following; 1) The blue region is the set of parameters \((b, c)\) at which the relative cancer cell population converges to the maximal capacity, 2) the
yellow one is the region where the population exhibits damped oscillations and converges to the population less than the maximal capacity and 3) the red one is the region where the population shows oscillations over the time. Our numerical result provides the critical set of values of bifurcation points \((b_1, c_1)\) and \((b_2, c_2)\) which are located on the border between two different colored regions. For example, the transcritical bifurcation occurs at the border between blue and yellow color and the Hopf bifurcation occurs on the border between yellow and red colored regions.

![Figure 4](image-url)

**Figure 4.** Two-dimensional bifurcation diagram for the cancer cell population density; infection rate on the x-axis, bursting rate on the y-axis. A. Three colored regions represent the set of two parameters at which three different patterns of cancer cell population over time are observed; the steady-state equilibrium (blue), damped oscillations (yellow) and oscillations (red). Two- bifurcations occurs at the borders between two colors; (1) Transcritical bifurcation on the border between blue and yellow, (2) Hopf bifurcation on the border between (yellow and red). The examples of cancer cell population over time in each colored region are shown in B-C-D, blue-yellow-red, respectively. Parameters used were \(a = 0.31\) and \(d = 0.44\).

4. Discussion

OVs are promising new therapeutics for various types of cancers. There have been multiple clinical trials using oncolytic viruses [18]. For the OV, replication competent viruses are used such as reoviruses [19]. The infection oncolytic viruses can kill cancer cells by triggering a cytotoxicity directly in the infected cancer cells or antitumoral immune responses or both [18, 20]. For an effective and safe treatment with oncolytic viruses, it is important to understand the behaviors of cancer cells and viruses. Mathematical modelling is a useful method to study the dynamic interactions between cancer cells and viruses. Although it is a simplified model excluding many other variations that can happen in clinical settings, our model supports the idea that not only the bursting rate of viruses but also infection rate of viruses must be considered for OV. One of barriers in OV is finding the optimal number of viruses to exterminate cancer cells without adverse effects such as immune hypersensitivity or spontaneous infection of normal cells [21]. Our model can be useful to determine the optimal number of viruses to be used for OV based on the bursting rate and infection rate that can be expressed as a plaque-forming unit (pfu). Both the bursting rate and pfu of a specific virus can be determined by *in vitro* experiments.
In clinical setting, there can be many other factors to affect the interaction between cancer cells and viruses. One of factors have not been considered in our model is host immune response. The host immune system either can facilitate OVs by triggering cytotoxic response to kill infected cancer cells or can interfere OVs by eliminating viruses. Intrinsic properties of virus can also determine the lifespan or death rate of free virion. Based on figure 5, it will be important to develop advanced models to consider the death rate of viruses when assessing cancer cell-virus infections.

Figure 5. Bifurcation diagram of cancer cell population density with respect to the death rate of virus $d$ with infection rate $b=0.1$ (A) and $b=0.3$ (B). Parameters used were $a = 0.31$ and $c = 5$.

5. Conclusions

In this paper, we studied the effect of the infection rate of the virus on the cancer cell population using the bifurcation diagrams of the cancer cell population density and the dynamical system (phase-space analysis). Our results show that there are two bifurcation threshold values of infection rate, which are the bursting rate dependent. The first bifurcation threshold value ($b_1$) can be obtained from the stability of the fixed point at $E_0(1, 0, 0)$ and be expressed as $b_1 = \frac{d}{c-1}$, which has an inverse relationship with the bursting rate of the virus and is proportional to the death rate of free virus particles. However, it is difficult to compute the second threshold value explicitly and express it using parameter values. For this issue, we provided the two-dimensional bifurcation diagram to show the set of parameters where three different patterns are generated. Our result gives an insight into understanding the relationship between two control parameters at which the cancer cell population exhibits the patterns from equilibrium steady-state solution to periodic solutions. It provides the second threshold value ($b_2$) has the same functional inverse relationship with $c_2$. The two-dimensional bifurcation diagram provides a promising result in determining optimal parameters for the successful virotherapy.

Acknowledgements

The author would like to thank Dr. Chang K. Sung for useful discussion. This work was supported by the start-up fund at Texas A&M University-Kingsville and partially supported by NSF Grant HRD-1901028.

References


