Characterization of SARS-CoV-2 Dynamics in the Host

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Abstract

While many epidemiological models have being proposed to understand and handle COVID-19, too little has been invested to understand how the virus replicates in the human body and potential antiviral can be used to control the replication cycle. In this work, using a control theoretical approach, validated mathematical models of SARS-CoV-2 in humans are properly characterized. A complete analysis of the main dynamic characteristic is developed based on the reproduction number. The equilibrium regions of the system are fully characterized, and the stability of such a regions, formally established. Mathematical analysis highlights critical conditions to decrease monotonically SARS-CoV-2 in the host, such conditions are relevant to tailor future antiviral treatments. Simulation results show the potential benefits of the aforementioned system characterization.

Keywords: SARS-CoV-2 infection, In-host model, Equilibrium sets characterization, Stability analysis.

1. Introduction

By December 2019, an outbreak of cases of pneumonia of unknown etiology was reported in Wuhan, Hubei province, China \cite{1}. On January 7, a novel betacoronavirus was identified as the etiological agent by the Chinese Center of Disease Control and Prevention (CDC), and subsequently named as Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) \cite{2}. On February 11, the World Health Organization (WHO) named the disease as Coronavirus disease 2019 (COVID-19) \cite{3}. Although, prevention and control measures were implemented rapidly, from the early stages in Wuhan and other key areas of Hubei \cite{4}, the first reporting of cases outside of China, 2 in Thailand and 1 in Japan \cite{5}, showed that the disease was starting to spread around the world. On March 11, with more than 111,800 cases in 114 countries, and 4921 fatality cases, the COVID-19 is declared pandemic by the WHO \cite{5}. So far, with more than 7,000,000 total cases confirmed in 213 countries and territories \cite{6,7}, and an estimated case-fatality rate (CFR) of 5.7\% (H1N1 pandemic, CFR<1\%) \cite{8}, the potential risks associated with this disease are evident.

Facing this situation, and taking into account the nonexistence of vaccines or specific therapeutically treatments, preventive measures such as social and physical distancing, hand washing, cleaning and disinfection of surfaces and use of face masks, among others, have been implemented in order to decrease the transmission of the virus, which is spread mainly from person-to-person through respiratory droplets produced when an infected person coughs, sneezes or talks \cite{9}. Furthermore, this infection prevention and control measures, helps avoid healthcare systems from becoming overwhelmed.

Many epidemiological mathematical models \cite{10,11,12} have been proposed to predict the spread of the disease and evaluate the potential impact of infection prevention and control measures in outbreak management \cite{13}. However, mathematical models at within-host level that could be useful to understand SARS-CoV-2 replication cycle and interaction with immune system as well as pharmacological effect of potential drug therapies \cite{14,15} are needed. So far, there are approximately 109 trials (including those not yet recruiting, recruiting, active, or completed) to assess pharmacological therapy for the treatment of COVID-19 in adult patients\cite{16}, including antiviral drugs (i.e. Hydroxychloroquine, Remdesivir, Favipiravir, Lopinavir/Ritonavir, Ribavirin), immunomodulatory agents (i.e. Tocilizumab) and immunoglobulin therapy, among others.
Recently, Hernandez-Vargas et al. [17] proposed different within-host mathematical models (2 based on target cell-limited model, with and without latent phase, and 1 considering immune response) for 9 infected patients with COVID-19. Numerical results in [17] showed a mean infecting time between susceptible cells of 30 days (about 3 times slower than Ebola and 60 times slower than influenza), which could explain the slow recuperation rate (12-22 days post symptom onset, pso) showed in COVID-19 infected patients. Furthermore, they informed within-host reproductive number values consistent to influenza infection (1.7-5.35).

Although models in [17] have been fitted to COVID-19 patients data, a control theoretical approach is needed to characterize the model dynamics. Even when the equilibrium states are known, a formal stability analysis is needed to properly understand the model behavior and, mainly, to properly design efficient control strategies. Note that the target cell model has been employed previously taking into account pharmacodynamic (PD) and pharmacokinetics (PK) models of antiviral therapies [18, 19], and this can be potentially done also for COVID-19.

In this context, the main contribution of this article is twofold. First, a full characterization of equilibrium and stability proprieties is performed for the COVID-19 target cell-limited model [17]. Then, formal properties concerning the state variables behavior before convergence - including an analysis of the virus peak times - are given. A key aspect in the target cell model for acute infections shows some particularities such as it has a minimal non-punctual stable equilibrium set, whose stability does not depends on the reproduction number. On the other side, assuming a basic reproduction number greater than 1, the virus would not be cleared before the target cells goes under a given critical value, which is independent of the initial conditions.

After the introduction given in Section 1 the article is organized as follows. Section 2 presents the general in-host target cell-limited model used to represent SARS-CoV-2 infection dynamic. Section 3 characterizes the equilibrium sets of the system, and establishes their formal asymptotic stability, by stating both, the attractivity of the equilibrium set in a given domain, and its $\epsilon - \delta$ (Lyapunov) local stability. Then, in Section 4 some dynamical properties of the system are stated, concerning the values of the states at the infection time $t = 0$. In Section 5 the general model for the SARS-CoV-2 infection is identified according to patient data, and the general characteristics of the infection are analyzed. Finally, Section 6 gives the conclusion of the work, while several mathematical formalism - necessary to support the results of Sections 3 and 4- are given in Appendices 7, 8 and 9.

1.1. Notation

$\mathbb{R}$ and $\mathbb{I}$ denote the real and integer numbers, respectively. The real vector space of dimension $n$ is denoted as $\mathbb{R}^n$. $\mathbb{R}_0^n$ represents the vectors of dimension $n$ whose components are equal or greater than zero. The distance from a point $x \in \mathbb{R}^n$ to a set $\mathcal{X} \subset \mathbb{R}^n$ is defined by $\|x\|_\mathcal{X} := \inf_{z \in \mathcal{X}} \|x - z\|_2$, where $\| \cdot \|_2$ denotes the norm-2. The open ball of radius $\epsilon$ around a point $x \in \mathbb{R}^n$, with respect to set $\mathcal{X}$, is defined as $\mathbb{B}_\epsilon(x) := \{z \in \mathcal{X} \mid \|x - z\|_2 < \epsilon\}$. Let us consider the real function $y(z) = ze^z$, then, the so-called Lambert function is defined as the inverse of $y(\cdot)$, i.e., $W(z) := f^{-1}(z)$ in such a way that $W(f(z)) = z$.

2. SARS-CoV-2 Within-Host Mathematical Model

Although incomplete by definition, mathematical models of in-host virus dynamic improves the understanding of the interactions that govern infections and, more important, permits the human intervention to moderate their effects [20]. Basic in-host infection dynamic models usually include the susceptible cells, infected cells, and the pathogen particles [21]. Among the most used mathematical models, the target cell-limited model has been employed to represent and control HIV infection [22, 23, 24], influenza [25, 26, 27, 18], Ebola [28], dengue [29, 30] among others.

In this work, we consider the mathematical model proposed by Hernandez-Vargas [17] given by the following set of differential equations (ODEs):

\begin{align}
\dot{U}(t) &= -\beta U(t)V(t), \quad U(0) = U_0, \tag{2.1a} \\
\dot{I}(t) &= \beta U(t)V(t) - \delta I(t), \quad I(0) = I_0 = 0, \tag{2.1b} \\
\dot{V}(t) &= pI(t) - cV(t), \quad V(0) = V_0, \tag{2.1c}
\end{align}

where $U$ [cell], $I$ [cell] and $V$ [copies/mL] represent the susceptible cells, the infected cells, and the virus load, respectively. The parameter $\beta$ [(copies/mL)$^{-1}$day$^{-1}$] is the infection rate of susceptible cells by the
responses are not explicitly described in this model, but they are implicitly included in the death rate of infected cells.

The model (2.1) is positive, which means that \( U(t) \geq 0 \), \( I(t) \geq 0 \) and \( V(t) \geq 0 \), for all \( t \geq 0 \). If we denote \( x(t) := (U(t), I(t), V(t)) \), then the states are constrained to belong to:

\[
X := \{ x \in \mathbb{R}^3_{\geq 0} \}. \tag{2.2}
\]

Another meaningful set is the one consisting in all the states in \( X \) with strictly positive amount of virus and susceptible cells, i.e.,

\[
\mathcal{X} := \{ x \in X : U > 0, V > 0 \}. \tag{2.3}
\]

Note that the set \( \mathcal{X} \) is an open set.

The initial conditions of (2.1) must be carefully established in order to properly represent the host body evolution from the beginning of the infection. So, it is assumed that the system is at a healthy steady state before the infection time \( t = 0 \), i.e., \( V(t) = 0 \), \( I(t) = 0 \), and \( U(t) = U_0 \), for \( t < 0 \). At time \( t = 0 \), a small quantity of virions enters to the host body and, so, a discontinuity occurs in \( V(t) \). Indeed, \( V(t) \) jumps from 0 to a small positive value \( V_0 \) at \( t_0 = 0 \) (formally, \( V(t) \) has a discontinuity of the first kind at \( t_0 \), i.e., \( \lim_{t \to 0^-} V(t) = 0 \) while \( \lim_{t \to 0^+} V(t) = V_0 > 0 \). The same scenario arises, for instance, when an antiviral treatment affects either parameter (say \( p \) or \( \beta \)).

The jump of \( p \) or \( \beta \) can be considered as a discontinuity of the first kind. In any case, for the time after the discontinuity, the virus may spread or be cleared in the body, depending on its infection effectiveness. To properly determine what such a spread means, the following (mathematical) definition is given

**Definition 1** (Spreadability of the virus in the host body). Consider system (2.1), constrained by the positive set \( X \), at some time \( t_0 \), with \( U(t_0) > 0 \), \( I(t_0) \geq 0 \) and \( V(t_0) > 0 \) (i.e., \( x(t_0) = (U(t_0), I(t_0), V(t_0)) \in X \)). Then, it is said that the virus spreads (in some degree) in the body host for \( t > t_0 \) if there exists at least one \( t^* > t_0 \) such that \( V(t^*) > 0 \).

The latter definition states that the virus spreads in the body host if \( V(t) \) has at least one local maximum. On the other hand, the virus does not spread if \( V(t) \) is strictly decreasing for all \( t > t_0 \), which means that \( V(t) \) has neither local minima nor local maxima. As it will be stated later on (Property[1], \( \lim_{t \to \infty} V(t) = 0 \) for system (2.1), independently of the fact that the virus reaches or not a maximum (this is a key difference between acute and chronic infection models[20, 21]).

The infection severity could be related with the virus spreadability established in Definition1. Liu et.al. [31] have shown that patients with severe COVID-19 tend to have a high viral load and a long virus shedding period. The mean viral load of severe cases was around 60 times higher than that of mild cases, suggesting that higher viral loads might be associated with severe clinical outcomes. Furthermore, they found that the viral load of severe cases remained significantly higher for the first 12 days after the appearance of the symptoms than those of corresponding mild cases. Mild cases were also found to have an early viral clearance, with 90% of these patients repeatedly testing negative on reverse transcription polymerase chain reaction (RT-PCR) by day 10 post symptoms onset (pso). By contrast, all severe cases still tested positive at or beyond day 10 pso. In addition, Zheng et.al. [32] from an study with 96 SARS-CoV-2 positive patients (22 with mild disease and 74 with severe disease) reported a longer duration of SARS-CoV-2 in lower respiratory samples of severe patients, such as, for patients with severe disease (21 days, 14-30 days) was significantly longer than in patients with mild disease (14 days, 10-21 days; p=0.04). Moreover, higher viral loads were detected in respiratory samples, although no differences were found in stool and serum samples. Although, these findings suggest that reducing the viral load through clinical means and strengthening management should help to prevent the spread of the virus, they are preliminary and it remains controversial whether virus persistence is necessary to drive the dysfunctional immune response characteristic of COVID-19 patients [33].

**Remark 1.** Note that the virus spreadability may or may not cause a severe infection (a disease that eventually cause the host death) depending on how much time the virus is above a given value.

To properly establish conditions under which the virus does not spread for \( t > 0 \) (i.e., after the infection time \( t = 0 \)) the so-called basic reproduction number within-host is defined next.
Definition 2. The basic reproduction number within-host $R$ is defined as the number of infected cells (or virus particles) that are produced by one infected cell (or virus particle), at a given time. Its mathematical expression is given by:

$$R(t) := U(t) \frac{\beta p}{c d}.$$  (2.4)

Particularly, for $t = 0$, this number describes the number of infected cells produced by one infected cell, when a small amount of virus, $V_0$, is introduced into a healthy stationary population of uninfected target cells, $U_0$.

$$R_0 := U_0 \frac{\beta p}{c d}.$$  (2.5)

A discussion about the way this value is obtained is given in Appendix 8. The relation between the basic reproduction number at the infection time ($R_0$) and the virus spreadibility is stated in the next theorem.

Theorem 2.1. Consider system (2.1), constrained by the positive set $\mathcal{X}$, at the beginning of the infection, i.e., $U(0) = U_0 > 0$, $I(0) = 0$ and $V(0) = V_0 > 0$ (i.e., $x(0) = (U(0), I(0), V(0)) \in \mathcal{X}$). Then, a sufficient condition (not necessary) for the virus not to spread in the host body is given by

$$R_0 < 1.$$

Proof: It is easy to see that the initial conditions are such that $R_V(0) := \frac{\beta I(0)}{c V(0)} = 0 < 1$. So, considering that $R_0 < 1$ by hypothesis, the result corresponds to Theorem 9.1. (i), when $t_0 = 0$. □

Remark 2. According to Theorem 9.1, in Appendix 9, there exists a positive value $\alpha(0) > 0$ (which is an implicit function of the initial conditions and the parameters, and it can be computed numerically) such that condition $R_0 < 1 + \alpha(0)$ is a necessary and sufficient condition for the virus not to spread in the host body, after the infection time $t = 0$. This means that for $R_0 > 1 + \alpha(0)$ the virus spreads in the host, as it is usually the case in real infections. In any case, the value of $\alpha(0)$ is generally close to zero (when $\beta$ is small), so to consider 1 as a threshold for the virus to spread seems to be a reasonable approximation.

Before proceeding with a full dynamic analysis of system (2.1), let us define first the so-called critical value of the susceptible cells, which shows to be an useful threshold value to properly understand the spread of the virus.

Definition 3. The critical value for $U$, $U_c$, is defined as

$$U_c := \frac{c \delta}{p \beta},$$  (2.6)

which, for fixed system parameters $\beta$, $p$, $\delta$ and $c$, is a constant.

Note that $U(t) < U_c$ if and only if $R(t) < 1$, for every $t \geq 0$.

2.1. Equilibrium set characterization

By equaling $\dot{U}$, $\dot{I}$ and $\dot{V}$ to zero, in (2.1), it can be shown that the system only has healthy equilibria of the form $x_s = (U_s, 0, 0)$, with $U_s$ being an arbitrary positive value, i.e., $U_s \in [0, \infty)$. Thus, there is only one equilibrium set, which is the healthy one, and it is defined by

$$\mathcal{X}_s := \{(U, I, V) \in \mathbb{R}^3 : U \in [0, \infty), I = 0, V = 0\}.$$  (2.7)

To have a first look on the stability of the equilibrium points in $\mathcal{X}_s$, system (2.1) can be linearized at a general state $x_s \in \mathcal{X}_s$. By simplifying (2.1) we have:

$$\dot{U} = f(U, I, V),$$
$$\dot{I} = g(U, I, V),$$
$$\dot{V} = h(U, I, V).$$

Then, the Jacobian matrix is given by
which means that

\[ \lambda \]

with

\[ \lambda = \begin{pmatrix} 0 & 0 & -\beta U_s \\ 0 & -\delta & \beta U_s \\ 0 & p & -c \end{pmatrix} \]

And the Jacobian evaluated at any point \( x_s \in X_s \) reads

\[ A_s = \begin{pmatrix} -\lambda & 0 & -\beta U_s \\ 0 & -\delta & \beta U_s \\ 0 & p & -c \end{pmatrix} \]

with \( U_s \in [0, \infty) \). Then, the eigenvalues \( (\lambda_1, \lambda_2, \lambda_3) \) are computed as the solution to \( \text{Det}(A_s - \lambda I) = 0 \), being the matrix \( A_s - \lambda I \) given by

\[ A_s - \lambda I = \begin{pmatrix} -\lambda & 0 & -\beta U_s \\ 0 & -\delta & \beta U_s \\ 0 & p & (-c - \lambda) \end{pmatrix} \]

Then, considering that \( \text{Det}(A - \lambda I) = \lambda[-\lambda^2 - (c + \delta)\lambda + (\beta U_s p - c\delta)] \), condition \( \text{Det}(A - \lambda I) = 0 \) is given by

\[ \lambda[-\lambda^2 - (c + \delta)\lambda + (\beta U_s p - c\delta)] = 0. \]

The first eigenvalue is trivially given by \( \lambda_1 = 0 \). The other two, are given by:

\[ \lambda_{2,3} = \frac{(c + \delta) \pm \sqrt{(c + \delta)^2 + 4(\beta U_s p - c\delta)}}{-2} \]

To analyze the eigenvalues qualitatively, note that for \( U_s = U_c \) it is

\[ \lambda_{2,3} = \frac{(c + \delta) \pm \sqrt{(c + \delta)^2 + 4(\beta U_s p - c\delta)}}{-2} \]

which means that \( \lambda_2 = 0 \) and \( \lambda_3 = -(c + \delta) < 0 \) (given that \( c, \delta > 0 \)). Furthermore, \( \lambda_2 < 0 \) and \( \lambda_3 < 0 \) for \( U_s < U_c \); and \( \lambda_2 > 0 \) and \( \lambda_3 < 0 \) for \( U_s > U_c \). Given that the maximum eigenvalue is the one dominating the stability behavior of the equilibrium under consideration, it is possible to infer how the system behaves near some segments of \( X_s \). The first intuition is that the equilibrium set

\[ X_s^1 := \{(U, I, V) \in \mathbb{R}^3 : U \in [0, U_c), I = 0, V = 0\} \] (2.8)

is stable, and that the equilibrium set

\[ X_s^2 := \{(U, I, V) \in \mathbb{R}^3 : U \in [U_c, \infty), I = 0, V = 0\} \] (2.9)

is unstable. These are just intuitions, given that one of the eigenvalues of the linearization system is null and so the linear approximation cannot be used to fully determine the stability of the nonlinear system (Theorem of Hartman-Grobman \[ 34, 35 \]). To formally prove the asymptotic stability of \( X_s^1 \) in a given domain, it is necessary to prove its global attractivity (in such domain) and local \( \epsilon-\delta \) stability.

3. Asymptotic stability of the equilibrium sets

A key point to analyze the general asymptotic stability (AS) of system \( (2.1) \) is to consider stability of the complete equilibrium sets \( X_s^1 \) and \( X_s^2 \), and not of the single points inside them (as defined in Definitions \[ 5, 6 \] and \[ 7 \] in Appendix \[ 7 \]. As it is shown in the next subsections, there is no single AS equilibrium points in this system, although there is an AS equilibrium set (i.e., \( X_s^1 \)).

As stated in Definition \[ 7 \] in Appendix \[ 7 \] the AS of \( X_s^1 \) requires both, attractivity and \( \epsilon - \delta \) stability, which are stated in the next two subsections, respectively. Then, in Subsection \[ 5.3 \] the AS theorem is formally stated.
3.1. Attractivity of set \( \mathcal{X}_1 \) in \( \mathcal{X} \)

Before proceeding with the formal theorems of the attractivity of \( \mathcal{X}_1 \), let us consider the following key property of system (2.1) concerning the attractivity of \( \mathcal{X} \).

**Property 1 (Attractivity of \( \mathcal{X} \)).** Consider system (2.1) constrained by the positive set \( \mathcal{X} \), at some arbitrary time \( t_0 \), with \( U(t_0) > 0 \), \( I(t_0) \geq 0 \) and \( V(t_0) > 0 \) (i.e., \( x(t_0) = (U(t_0), I(t_0), V(t_0)) \) \( \in \mathcal{X} \)). Then, \( U_{\infty} := \lim_{t \to \infty} U(t) \) is a constant value smaller than \( U(t_0) \). \( I_{\infty} := \lim_{t \to \infty} I(t) = 0 \) and \( V_{\infty} := \lim_{t \to \infty} V(t) = 0 \), which means that \( x(t) = (U(t), I(t), V(t)) \) tends to some state in \( \mathcal{X} \).

**Proof:** Since \( \dot{U}(t) \leq 0 \) for all \( t \geq 0 \) and all \( (U(t_0), I(t_0), V(t_0)) \) \( \in \mathcal{X} \), by (2.1a) \( U(t) \) is a decreasing function (no oscillation can occur). Since \( U(t_0) > 0 \) and \( V(t_0) > 0 \), then \( U_{\infty} = \lim_{t \to \infty} U(t) \) is a constant value in \( [0, U(t_0)) \). Given that \( U(t) \) converges to a finite fixed value, then \( \dot{U}(t) = 0 \) as \( t \to \infty \), by (2.1a). This implies, by the same equation (2.1a), that \( \dot{U}(t)V(t) = 0 \) as \( t \to \infty \), and so, from equation (2.1b), that \( \dot{I}(t) = -\delta I(t) \) as \( t \to \infty \), whose solution asymptotically goes to zero. Then, \( I_{\infty} = \lim_{t \to \infty} I(t) = 0 \).

Finally, by equation (2.1c), \( \dot{V}(t) = -\delta V(t) \) as \( t \to \infty \), whose solution asymptotically goes to zero. Then \( V_{\infty} = \lim_{t \to \infty} V(t) = 0 \), which completes the proof. \( \square \)

Property 1 states that \( \mathcal{X} \) is an attractive set for system (2.1), in \( \mathcal{X} \), but not the smallest attractive set. Now, conditions are given to show that the smallest attractive set is given by \( \mathcal{X}_1 \).

**Theorem 3.1 (Attractivity of \( \mathcal{X}_1 \)).** Consider system (2.1) constrained by the positive set \( \mathcal{X} \). Then, the set \( \mathcal{X}_1 \) defined in (2.8) is the smallest attractive set in \( \mathcal{X} \). Furthermore, \( \mathcal{X}_2 \), defined in (2.9), is not attractive.

**Proof:** The proof is divided into two parts. First it is proved that \( \mathcal{X}_1 \) is an attractive set, and then, that it is the smallest one.

Attractivity of \( \mathcal{X}_1 \): The attractivity of \( \mathcal{X} \) in \( \mathcal{X} \) is already proved in Property 1. So, to prove the attractivity of \( \mathcal{X}_1 \) in \( \mathcal{X} \) (and to show that \( \mathcal{X}_2 \) is not attractive) it remains to demonstrate that \( U_{\infty} \in [0, U_\infty) \). From system (2.1), by replacing (2.1a) in (2.1b), it follows that

\[
\dot{I}(t) = -\frac{1}{\beta}(\dot{I}(t) + \dot{U}(t)).
\]

From (2.1c) it follows that

\[
V(t) = \frac{1}{c}(pI(t) - \dot{V}(t)).
\]

Then, replacing (3.1) in (3.2), we have

\[
\dot{V}(t) = \frac{1}{c}[p(-\frac{1}{\beta}(\dot{I}(t) + \dot{U}(t)) - \dot{V}(t)] - \frac{1}{c}.\]

Finally, by substituting (3.3) in (2.1a), and multiplying by \( 1/U(t) \) both sides of the equation (without loss of generality we assume that \( U(t) \neq 0 \)), it follows that

\[
\frac{1}{U(t)}\dot{U}(t) = \frac{\beta p}{c \delta}U(t) + \frac{\beta p}{c \delta}I(t) + \frac{\beta}{c}V(t).
\]

This latter equation can be integrated, for general initial conditions \( U_0, I_0 \) and \( V_0 \), as follows:

\[
\ln\left(\frac{U(t)}{U_0}\right) = \frac{\beta p}{c \delta}(U(t) - U_0) + \frac{\beta p}{c \delta}(I(t) - I_0) + \frac{\beta}{c}(V(t) - V_0).
\]

Now, by defining \( U_{\infty} := \lim_{t \to \infty} U(t) \), \( I_{\infty} := \lim_{t \to \infty} I(t) \), \( V_{\infty} := \lim_{t \to \infty} V(t) \), and recalling from Property 1 that \( I_{\infty} = V_{\infty} = 0 \), the latter equation for \( t \to \infty \), reads

\[
\ln\left(\frac{U_{\infty}}{U_0}\right) = \frac{\beta p}{c \delta}(U_{\infty} - U_0) + \frac{\beta p}{c \delta}(I_{\infty} - I_0) + \frac{\beta}{c}(V_{\infty} - V_0)
\]

\[
= \frac{\beta p}{c \delta}U_{\infty} - R_0 - \frac{\beta p}{c \delta}I_0 - \frac{\beta}{c}V_0
\]
\[ \frac{\beta p}{c^2} U_\infty - R_0 + K_0, \]  

(3.6)

where \( R_0 := \frac{\beta p}{c^2} U_0 \) (as it was defined in (2.5)) and

\[ K_0 := -\frac{\beta}{c^2} (\frac{p}{\delta} I_0 + V_0). \]  

(3.7)

Note that \( R_0 \) is a function of \( U_0 \) while \( K_0 \) is a function of \( I_0 \) and \( V_0 \), and, furthermore, \( R_0 > 0 \) and \( K_0 < 0 \) for every \( x_0 = (U_0, I_0, V_0) \in \mathcal{X} \). Then, after some manipulation, (3.6) reads

\[ -\frac{\beta p}{c^2} U_\infty e^{-\frac{\beta}{c^2} U_\infty} = -\frac{\beta p}{c^2} U_0 e^{-R_0} e^{K_0} = -R_0 e^{-R_0} e^{K_0}. \]  

(3.8)

Now, by denoting \( z = z(R_0, K_0) := -R_0 e^{-R_0} e^{K_0} \) and \( y := -\frac{\beta p}{c^2} U_\infty \), the latter equation can be written as

\[ W(z) = y, \]  

(3.9)

or, the same,

\[ W(-R_0 e^{-R_0} e^{K_0}) = -\frac{\beta p}{c^2} U_\infty, \]  

(3.10)

where \( W(\cdot) \) is a Lambert function. Figure 1 shows the graph of such a function, where it can be seen that it has two branches, denoted as \( W_p \) and \( W_m \). However, \( W(\cdot) = W_p(\cdot) \) in this case, since \( W_m \to -\infty \) for \( z \to 0^- \), which is not biological sense (note that \( U_\infty \) is a finite value in \([0, U_0)\)). Besides, function \( -1/e < z(R_0, K_0) \leq 0 \) for \( R_0 > 0 \) and \( K_0 < 0 \) (Figure 2 shows a plot of function \( z(R_0, K_0) \) for negative values of \( K_0 \) and positive values of \( R_0 \)), and function \( W_p \) maps \((-1/e, 0]\) into \((-1, 0]\), which implies that

\[ 1 > -W(z(R_0, K_0)) \geq 0, \]  

(3.11)

for \( R_0 > 0 \) and \( K_0 < 0 \). This way, by (3.10), it follows that

\[ U_\infty = -\frac{c^2}{\beta p} W(-R_0 e^{-R_0} e^{K_0}) \]

\[ = -U_0 W(-R_0 e^{-R_0} e^{K_0}) \]

\[ \in [0, U_\epsilon), \]  

(3.12)

which completes the proof.

\( \mathcal{X}_1^1 \) is the smallest attractive set: It is clear, from the previous analysis, that any initial state \( x_0 = (U_0, I_0, V_0) \) in \( \mathcal{X} \) converges to a state \( x_\infty = (U_\infty, 0, 0) \) with \( U_\infty \in [0, U_\epsilon) \). This means that every state \( x_s \in \mathcal{X}_2^1 \) is not attractive in \( \mathcal{X} \) and so, neither the whole set \( \mathcal{X}_2^1 \). Let us consider now a state \( x_s \in \mathcal{X}_2^1 \), an arbitrary ball of radius \( \epsilon > 0 \), w.r.t. \( \mathcal{X} \), around it, \( \mathcal{B}_\epsilon(x_s) \in \mathcal{X} \). Take two arbitrary initial states \( x_{0,1} = (U_{0,1}, I_{0,1}, V_{0,1}) \) and \( x_{0,2} = (U_{0,2}, I_{0,2}, V_{0,2}) \) in \( \mathcal{B}_\epsilon(x_s) \), such that \( U_{0,1} \neq U_{0,2} \) and \( V_{0,1} \neq V_{0,2} \). These two states converge, according to equation (3.11), to \( x_{\infty,1} = (U_{\infty,1}, 0, 0) \) and \( x_{\infty,2} = (U_{\infty,2}, 0, 0) \), respectively. Given that function \( z(R, K) \) is monotone (injective) in \( R_0 \) (and so in \( U_0 \)) and \( W(z) \) is monotone (injective) in \( z \), then \( U_{\infty,1} \neq U_{\infty,2} \). This means that, although both initial states converge to some state in \( \mathcal{X}_2^1 \), they necessarily converge to different points. Therefore neither single states \( x_s \in \mathcal{X}_2^1 \), nor subsets of \( \mathcal{X}_2^1 \) are attractive in \( \mathcal{X} \). So, \( \mathcal{X}_2^1 \) is the smallest attractive set and the proof is concluded. \( \square \)

Remark 3. Note that \( \mathcal{X}_2^1 \) and \( \mathcal{X}_2^2 \) are in the closure of the open set \( \mathcal{X} \), which is not in \( \mathcal{X} \). In other words, what Theorem 3.7 shows is that any initial state in \( \mathcal{X} \) converges to a point onto the boundary of \( \mathcal{X} \) that does not belong to \( \mathcal{X} \). Furthermore note that, an initial state of the form \((U_0, 0, 0), U_0 > U_\epsilon \), (i.e., a state in \( \mathcal{X}_2^2 \) cannot be attracted by any set since it is - by definition - an equilibrium state (every state in \( \mathcal{X}_2^2 \) will remain unmodified). This is the reason why it is not possible to consider the attractiveness of \( \mathcal{X}_2^2 \) in \( \mathcal{X} \).
Figure 1: Lambert function $W(z)$ has two branches, denoted as $W_p$ (in blue) and $W_m$ (in red). Both branches are defined for $z \in [-1/e, 0]$; however, $\lim_{z \to 0^-} W_p = 0$ while $\lim_{z \to 0^-} W_m = -\infty$, which means that only the branch $W_p$ will be used in our analysis, as it is shown in the proof.

Figure 2: Function $z(R_0, K_0)$, for $R_0 \geq 0$ and $K_0 \leq 0$. 
Proof: Let us consider a particular equilibrium point $\mathbf{x}_e$. Then a Lyapunov function candidate is given by (similar to one used in [36] for chronic infections)

$$J(x) := U - U_s - U_x \ln\left(\frac{U}{U_s}\right) + I + \frac{\delta}{p} V. \quad (3.13)$$

This function is continuous in $\mathfrak{X}$, is positive for all nonegative $x \neq x_e$ and $J(x_e) = 0$. Function $J$ evaluated at the solutions of system (2.1) reads:

$$\dot{J}(x(t)) = \frac{\partial J}{\partial x} \dot{x}(t) = \begin{bmatrix} \frac{dJ}{dU} \frac{dJ}{dI} \frac{dJ}{dV} \end{bmatrix} \begin{bmatrix} -\beta U(t)V(t) \\ \beta U(t)V(t) - \delta I(t) \\ \delta I(t) \end{bmatrix} = \begin{bmatrix} \frac{1}{U(t)} - \frac{U_s}{U(t)} \frac{\delta}{p} \end{bmatrix} \begin{bmatrix} -\beta U(t)V(t) \\ \beta U(t)V(t) - \delta I(t) \\ \delta I(t) \end{bmatrix} = (-\beta U(t)V(t) + U_s \beta V(t)) + (\beta U(t)V(t) - \delta I(t)) + (\delta I(t) - \frac{\delta c}{p} V(t)) = U_s \beta V(t) - \frac{\delta c}{p} V(t) = V(t)(U_s \beta - \frac{\delta c}{p}). \quad (3.14)$$

Now, given $U_s \in [0, U_c]$, with $U_c = \frac{\delta c}{p}$, it follows that $\dot{J}(x(t)) \leq 0$ for every $x \in \mathfrak{X}$ (note that it is not true that $\dot{J}(x(t)) < 0$ for $x \neq x_e$, as shown next, in Remark 4). Then, $J$ is a Lyapunov function for system (2.1), which means that each $x_e \in \mathcal{X}_1$ is $\epsilon - \delta$ stable. Therefore, it is easy to see that the equilibrium set $\mathcal{X}_1$ as a whole is also $\epsilon - \delta$ stable, which completes the proof. □

Remark 4. Note that, in the latter proof, it is not true that $\dot{J}(x(t)) < 0$ for every nonegative $x \neq x_e$. If for instance, the function $J(x(t))$ is evaluated at $\dot{x}_e = (U, 0, 0)$, with $U \not\in U_s$, we have that $\dot{J}(\dot{x}_e(t)) = 0$. In fact, $\dot{J}(x(t))$ is null along the whole $U$ axis, given that this axis is an equilibrium set. This means that the (individual) states in $\mathcal{X}_1$ are $\epsilon - \delta$ stable, but not attractive.

A schematic plot of such a behavior can be seen in Figure 3.
Consider system \( U \) \( V \). Property 2 states some characteristics of the general relationship between the peak times of the MPC \([39, 40]\). \( U \), \( V \) will be fully successful in controlling system \((2.1)\).

\[ t \in X \in \epsilon \] states the local infectivity (by reducing the infection rate \( \beta \)) or the production of infectious virions (by reducing the replication rate \( p \)) \([20]\). So, the previous stability analysis is still valid for controlled system, since only a modification of some of the parameters defining \( U_c \) is done. In such a context, only a controller able to consider the whole set \( \mathcal{X}_0^1 \) as a target (a set-based control strategy, as zone MPC \([39, 40]\)) will be fully successful in controlling system \((2.1)\).

3.3. Asymptotic stability of \( \mathcal{X}_0^1 \)

In the next Theorem, based on the previous results concerning the attractivity and \( \epsilon - \delta \) stability of \( \mathcal{X}_0^1 \), the asymptotic stability is formally stated.

**Theorem 3.3.** Consider system \((2.1)\) constrained by the positive set \( X \). Then, the set \( \mathcal{X}_0^1 \) defined in \((2.8)\) is smallest asymptotically stable (AS) equilibrium set, with a domain of attraction given by \( X \).

**Proof:** The proof follows from Theorems \(3.1\) which states that \( \mathcal{X}_0^1 \) is the smallest attractive in \( \mathcal{X} \), and \(3.2\) which states the local \( \epsilon - \delta \) stability of \( \mathcal{X}_0^1 \). \( \Box \)

A critical consequence of the latter Theorem is that no equilibrium point in \( \mathcal{X}_0 \) (neither in \( \mathcal{X}_0^1 \), nor in \( \mathcal{X}_0^2 \)) can be used as setpoint in a control strategy design. The effect of antivirals (pharmacodynamic), for instance, is just to reduce the virus infectivity (by reducing the infection rate \( \beta \)) or the production of infectious virions (by reducing the replication rate \( p \)) \([20]\). So, the previous stability analysis is still valid for controlled system, since only a modification of some of the parameters defining \( U_c \) is done. In such a context, only a controller able to consider the whole set \( \mathcal{X}_0^1 \) as a target (a set-based control strategy, as zone MPC \([39, 40]\)) will be fully successful in controlling system \((2.1)\).

4. Characterization for different initial conditions

In this section some further properties of system \((2.1)\) concerning its dynamic are stated, based on the initial conditions at the infection time \( t = 0 \). The objective is to fully characterize the states behavior in a qualitative way, including the times at which the virus and the infected cells reach their peaks. First, Property \(2\) states some characteristics of \( U^* \) for different initial conditions. Then, Theorem \(4.1\) states a general relationship between the peak times of \( V \) and \( I \) and the time at which \( U \) reaches its critical value \( U_c \).

**Property 2.** Consider system \((2.1)\), constrained by the positive set \( X \), at the beginning of the infection, i.e., \( U(0) = U_0 > 0 \), \( I(0) = 0 \) and \( V(0) = V_0 > 0 \) (i.e., \( x(0) = (U(0), I(0), V(0)) \in X \)). Consider also that \( V_0 \) is small enough. Then,

i. \( U_\infty \rightarrow 0 \) when \( U_0 \rightarrow \infty \) or \( U_0 \rightarrow 0 \).

ii. \( U_\infty \rightarrow U_c \) when \( U_0 \rightarrow U_c \).

iii. \( 0 < U_\infty(U_{0,1}, I_0, V_0) < U_\infty(U_{0,2}, I_0, V_0) < U_c \), for initial conditions \( U_{0,1} < U_{0,2} < U_c \).

iv. \( 0 < U_\infty(U_{0,2}, I_0, V_0) < U_\infty(U_{0,1}, I_0, V_0) < U_c \), for initial conditions \( U_{0,2} < U_{0,1} < U_c \).

**Proof:** If \( I_0 = 0 \) and \( V_0 \approx 0 \) then \( K_0 \approx 0 \). Therefore \( W(-R_0 e^{K_0 - R_0}) \approx W(-R_0 e^{-R_0}) \), and \( U_\infty \approx -U_c W(-R_0 e^{-R_0}) \) by \((3.10)\).

i. \( W(-R_0 e^{-R_0}) \rightarrow 0 \) when \( -R_0 e^{-R_0} \rightarrow 0 \), which means that either \( R_0 \rightarrow 0 \) or \( R_0 \rightarrow \infty \). This implies that \( U_0 \rightarrow 0 \) or \( U_0 \rightarrow \infty \), respectively.

ii. \( W(-R_0 e^{-R_0}) \approx -1 \) when \( -R_0 e^{-R_0} \approx -1/e \), which is true if \( R_0 \approx 1 \) or, the same, when \( U_0 \rightarrow U_c \).

iii. Function \( z(R_0) = R_0 e^{-R_0} \) is strictly decreasing for \( R_0 \in (0, 1) \) (note that \( R_{0,1} := 10^{-0.1} p \) and \( R_{0,2} := 10^{-0.2} p \) are in \((0, 1)\), since they are smaller than \( U_c \), while \( W_p(.) \) is strictly decreasing in \((-1/e, 0)\). So, \( 0 < -W_p(-R_0 e^{R_0}) < -W_p(-R_0 e^{-R_0}) < 1 \), which implies that \( 0 < U_\infty(U_{0,1}, I_0, V_0) < U_\infty(U_{0,2}, I_0, V_0) < U_c \).
iv. Function\( z(R_0) = R_0 e^{-R_0} \) is strictly increasing for \( R_0 \in (1, \infty) \), while \( -W_p(\cdot) \) is strictly decreasing in \((1/e, 0)\). So, \( 0 < -W_p(-R_0 e^{-R_0}) < -W_p(-R_0 e^{-R_0}) < 1 \), which implies that \( 0 < U_\infty(U_0, 1, I_0, V_0) < U_\infty(U_0, 2, I_0, V_0) < U_c \). Figure 4 shows \( U_\infty \) as a function of \( U_0 \), taking \( V_0 \) as a parameter. □

Figure 5 shows a phase portrait of system (2.1), where all parameters are equal to 1 for simplicity, which means that \( U_c = 1 \).

**Theorem 4.1** (Virus behavior from the infection time). Consider system (2.1), constrained by the positive set \( \mathbb{X} \), at the beginning of the infection, i.e., \( U(0) = U_0 > 0 \), \( I(0) = 0 \) and \( V(0) = V_0 > 0 \) (i.e., \( x(0) = (U(0), I(0), V(0)) \in \mathbb{X} \)). If the virus spreads (according to Definition 1), then there exist positive times \( t_V, t_I, t_c \), and \( t_V \), such that \( t_V < t_I < t_c < t_V \), where \( t_V \) and \( t_V \) are the times at which \( V(t) \) reaches a local minimum and a local maximum, respectively, \( t_I \) is the time at which \( I(t) \) reaches a local maximum, and \( t_c \) is the time at which \( U(t) \) reaches \( U_c \). Furthermore, \( V(t) < 0 \) for all \( t > t_V \).

**Proof:** If \( I(0) = 0 \) then \( R_V(0) < 1 \), and Theorem 4.1 is a particular case of Theorem 9.1(ii).

Next, a Remark concerning a particularity of Theorem 4.1 is introduced, that may help to approximately determine the global maximum of the virus load.

**Remark 6.** Consider the hypothesis of Theorem 4.1. Then, it can be shown that for \( V_0 \rightarrow 0, \dot{V} \rightarrow t_c \) from the right, and \( t_I \rightarrow t_c \) from the left, meaning that the peaks of \( V \) and \( I \) tends to occurs simultaneously at time \( t_c \). This fact can be seen in Table 2 when data coming from real patients are used to identify the model.

A main consequence of Theorem 4.1 and Remark 6 is that it guaranties that the virus will monotonically go to zero only after \( U \) is below \( U_c \). That is, any action devoted to steers \( V \) to zero before \( t_c \) may be counterproductive, since slowing down \( V \) implies to soften the decreasing behavior of \( U \), delaying time \( t_c \), and maintaining \( V \) large for a longer time. This critical fact has a direct effect in a potential controlled system, when parameters \( p \) or \( \beta \) are scaled down by antiviral treatments. Assuming that \( R_0 \) remains greater than 1, any attempt to steer \( V \) to zero before \( U \) has taken values below \( U_c \) may be unsuccessful, if \( U \) is not controlled first.
5. Particularization of the model with patient data

In this section, the parameters of model (2.1) will be associated to data from 9 patients with COVID-19 - labeled as A, B, C, D, E, F, G, H and I - reported in [41]. Here, we consider work in [17], where different models for in-host SARS-CoV-2 were proposed and identified to fit the virus load data collected in [41].

The initial number of target cells \( U_0 \) is estimated approximately \( 10^7 \) cells [17]. \( I_0 \) is assumed to be 0 while \( V_0 \) is determined by interpolation considering an incubation period of 7 days (note, that \( V_0 \) ranges from 0.02 to 5.01 copies/mL which is below the detectable level of about 100 copies/mL). Moreover, the onset of the symptoms is assumed to occur 4 to 7 days after the infection time (day 0, Figure 6 and 7). Since the viral load is measured in logarithmic scale, the parameter fitting was performed minimizing the root mean square (RMS) difference on logarithmic scale between the model predictive output (\( \hat{V}_i \)) and the experimental measurements (\( V_i \)), employing the Differential Evolution (DE) algorithm proposed in [17, 20]. The parameters and the initial conditions (\( U_0, I_0 \) and \( V_0 \), with \( t_0 = 0 \) the infection time) of each patient are collected in Table 1.

Table 1: Target limited cell model parameter values for different patients with COVID-19 [17].

| Patient | \( \beta \)       | \( \delta \)  | \( p \) | \( c \)  |
|---------|------------------|--------------|--------|--------|
| A       | \( 9.98 \times 10^{-8} \) | 0.61         | 9.3    | 2.3    |
| B       | \( 1.77 \times 10^{-7} \) | 14.11        | 20.2   | 0.8    |
| C       | \( 8.80 \times 10^{-7} \) | 79.51        | 134.4  | 0.4    |
| D       | \( 3.15 \times 10^{-8} \) | 45.51        | 620.2  | 2.0    |
| E       | \( 5.61 \times 10^{-8} \) | 7.51         | 96.4   | 5.0    |
| F       | \( 1.41 \times 10^{-8} \) | 37.61        | 995.0  | 0.6    |
| G       | \( 1.77 \times 10^{-8} \) | 8.21         | 338.4  | 5.0    |
| H       | \( 1.58 \times 10^{-8} \) | 21.11        | 927.8  | 1.8    |
| I       | \( 4.46 \times 10^{-9} \) | 4.21         | 994.6  | 4.3    |

According to the system analysis developed in previous sections, some relevant dynamical values are shown in Table 2. Constant \( \alpha(0) \) (defined in Theorem 9.1) is small than \( 10 \times 10^{-4} \) for all the patients, so it is not taken into account for the study.
Table 2: Characterization Parameters of patients with COVID-19.

| Patient | $U_c$ | $U_\infty$ | $R_0$ | $K_0$ | $t_f$ | $t_e$ | $t_V$ | $V_{max}$ |
|---------|-------|-------------|-------|-------|-------|-------|-------|-----------|
| A       | $1.51 \times 10^6$ | $1.36 \times 10^4$ | 6.61 | $-2.17 \times 10^{-7}$ | 10.16 | 10.24 | 10.58 | $1.73 \times 10^4$ |
| B       | $3.15 \times 10^6$ | $4.88 \times 10^5$ | 3.18 | $-6.87 \times 10^{-8}$ | 11.54 | 12.26 | 12.32 | $4.35 \times 10^6$ |
| C       | $2.66 \times 10^5$ | $4.81 \times 10^{-10}$ | 37.57 | $-6.89 \times 10^{-7}$ | 1.43 | 1.67 | 1.69 | $1.47 \times 10^7$ |
| D       | $4.65 \times 10^6$ | $1.67 \times 10^6$ | 2.15 | $-4.89 \times 10^{-9}$ | 9.04 | 9.42 | 9.44 | $2.33 \times 10^7$ |
| E       | $6.94 \times 10^6$ | $4.58 \times 10^6$ | 1.44 | $-3.48 \times 10^{-9}$ | 15.02 | 15.16 | 15.24 | $4.03 \times 10^6$ |
| F       | $1.61 \times 10^6$ | $2.03 \times 10^4$ | 6.21 | $-7.28 \times 10^{-9}$ | 7.12 | 7.76 | 7.78 | $1.42 \times 10^8$ |
| G       | $6.84 \times 10^6$ | $4.43 \times 10^6$ | 1.46 | $-1.1 \times 10^{-9}$ | 14.80 | 14.92 | 15.00 | $1.44 \times 10^7$ |
| H       | $2.59 \times 10^6$ | $2.3 \times 10^5$ | 3.86 | $-2.72 \times 10^{-9}$ | 5.16 | 5.44 | 5.48 | $1.577 \times 10^8$ |
| I       | $4.08 \times 10^6$ | $1.14 \times 10^6$ | 2.45 | $-3.21 \times 10^{-10}$ | 9.28 | 9.38 | 9.50 | $2.60 \times 10^8$ |

Figures 6 and 7 shows the evolution of $V$ and $U$ for all patients. As expected, the states converges to $\lambda_1^*$, although significantly different behavior can be observed for the different patients. From Figure 7 it can be seen that the healthy cells final value $U_\infty$ is reduced in cases of patients with large values of $R_0$, in spite all patients have the same initial $U_0$. This can be explained from the fact that $W(R_0e^{-R_0e^{K_0}})$ is monotonically decreasing for $R_0 > 1$ (see Figures 1 and 2), and therefore, $0 < U_\infty(R_{01}) < U_\infty(R_{02})$ for $R_{01} > R_{02} > 1$. Note that the healthy cells of patient C converges to $U_\infty$ equals to $4.810 \times 10^{-10}$ [cell], which can be explained by the fact that this patient has a reproduction number ($R_0$) of 37.57, which is 5.2 times above the cohort mean value of 7.21. Figure 6 and Table 2 show that the viral load of patient C reaches the peak at 1.69 days post infection (dpi) (40.56 hours post infection, hpi).

Furthermore, from Figure 6 it can be seen that for all the cases the viral load spreads (i.e.: the virus presents a peak) although $R_V(0) < 0$ for all patients (i.e., $I_0 = 0$). This can be justified since $U_0 \gg U_c$ and, therefore, $R_0$ will be greater than $1 + \alpha(0)$ for all patients (note that, $\alpha(0) < 10 \times 10^{-4}$). Moreover, from Table 2 we can corroborate that $t_f > t_e > t_V$ which is in accordance to what is stated in Theorem 4.1.

Concerning the immune response, this model makes the assumption that it is constant and independent on viral load as well as infected cells. Furthermore, neither innate or adaptive response are modeled, being the viral load dynamic mainly limited by target cells availability. Since recent studies have shown a dysfunctional immune response (i.e.: lymphogenia, desegulated secretion of pro-inflammatory cytokines, excessive infiltration of monocytes, macrophages and T cells, among others) [33, 42], this effect should be added in the proposed model, in order to have a more reliable representation (and, eventually, a more realistic control objective). In addition, a more reliable standard to measure the severity of disease could be related with the viral spreadability as well as the deregulated inflammatory response.

6. Conclusions

In this work a full dynamical characterization of a COVID-19 in-the-host target-cell model is performed. Opposite to what happens in other similar models, it is shown that there exists a minimal non-punctual stable equilibrium set depending only on the system parameters. Furthermore, it is shown that there exists a parameter-depending threshold for the susceptible cells that fully characterizes the virus and infected cells qualitative behavior. Simulations performed with real-patient data demonstrate the potential utility of such system dynamic characterization to tailor the the most valuable pipeline drugs against SARS-CoV-2.

7. Appendix 1. Stability theory

In this section some basic definitions and results are given concerning the asymptotic stability of sets and Lyapunov theory, in the context of non linear continuous-time systems. All the following definitions are referred to system

$$\frac{dx(t)}{dt} = f(x(t)), \quad x(0) = x_0,$$

where $x$ is the system state constrained to be in $\mathbb{R}^n$, $f$ is a Lipschitz continuous nonlinear function, and $\phi(t; x)$ is the solution for time $t$ and initial condition $x$. 

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Definition 4 (Equilibrium set). Consider system (7.1) constrained by $X$. The set $X_s \subset X$ is an equilibrium set if each point $x \in X_s$ is such that $f(x) = 0$ (this implying that $\phi(t; x) = x$ for all $t \geq 0$).

Definition 5 (Attractivity of an equilibrium set). Consider system (7.1) constrained by $X$. A closed equilibrium set $X_s \subset X$ is attractive in $X \subset X$ if $\lim_{t \to \infty} \|\phi(t; x)\|_{X_s} = 0$ for all $x \in X$.

Any set containing an attractive set is attractive, so the significant attractive of a constrained system set is given by the smallest one.

Definition 6 ($\epsilon - \delta$ local stability of an equilibrium set). Consider system (7.1) constrained by $X$. A closed equilibrium set $X_s \subset X$ is $\epsilon - \delta$ locally stable if for all $\epsilon > 0$ it there exists $\delta > 0$ such that in a given boundary of $X_s$, $\|x\|_{X_s} < \delta$, it follows that $\|\phi(t; x)\|_{X_s} < \epsilon$, for all $t \geq 0$.

Definition 7 (Asymptotic stability (AS) of an equilibrium set). Consider system (7.1) constrained by $X$. A closed equilibrium set $X_s \subset X$ is asymptotically stable (AS) in $X \subset X$ if it is $\epsilon - \delta$ locally stable and attractive in $X$.

Theorem 7.1 (Lyapunov theorem). Consider system (7.1) constrained by $X$ and an equilibrium state $x_s \in X_s \subset X$. Let consider a function $V(x) : \mathbb{R}^n \to \mathbb{R}$ such that $V(x) > 0$ for $x \neq x_s$, $V(x_s) = 0$ and $\dot{V}(x(t)) \leq 0$, denoted as Lyapunov function. Then, the existence of such a function implies that $x_s \in X_s$ is $\epsilon - \delta$ locally stable. If in addition $\dot{V}(x(t)) < 0$ for all $x \neq x_s$ and $V(x_s) = 0$, then $x_s \in X_s$ is asymptotically stable.

8. Appendix 2. Derivation of the basic reproduction number $R_0$

The derivation of the basic reproduction number $R_0$ will be given by means of the concept of next-generation matrix [43]. Consider system (2.1) and assume that a healthy equilibrium exists, of the form $x_0 = (U_0, 0, 0)$, and it is stable in absence of disease. Of the complete state of system (2.1), $x = (U, I, V)$,
only two states depend on infected cells, that is $I$ and $V$. Let us rewrite the ODEs for this two states in the form

$$
\dot{I}(t) = F_I(x) - G_I(x)
$$

$$
\dot{V}(t) = F_V(x) - G_V(x)
$$

where $F_i(x)$, $i = \{I, V\}$, is the rate of appearance of new infections in compartment $i$, while $G_i(x)$, $i = \{I, V\}$, is the rate of other transitions between compartment $i$ and the other infected compartments, that is

$$
F_I(x) = \beta U(t)V(t) \quad \text{and} \quad G_I(x) = \delta I(t)
$$

$$
F_V(x) = 0 \quad \text{and} \quad G_V(x) = -p I(t) + e V(t)
$$

If we now define

$$
F = \begin{bmatrix}
\frac{\partial F_I(x)}{\partial I} & \frac{\partial F_I(x)}{\partial V} \\
\frac{\partial F_V(x)}{\partial I} & \frac{\partial F_V(x)}{\partial V}
\end{bmatrix}_{x=x_0} = \begin{bmatrix}
0 & \beta U_0 \\
0 & 0
\end{bmatrix}
$$

and

$$
G = \begin{bmatrix}
\frac{\partial G_I(x)}{\partial I} & \frac{\partial G_I(x)}{\partial V} \\
\frac{\partial G_V(x)}{\partial I} & \frac{\partial G_V(x)}{\partial V}
\end{bmatrix}_{x=x_0} = \begin{bmatrix}
\delta & 0 \\
-p & e
\end{bmatrix}
$$

then matrix $FG^{-1}$, represents the so-called next-generation matrix. Each $(i, j)$ entry of such a matrix represents the expected number of secondary infections in compartment $i$ produced by an infected cell introduced in compartment $j$. The spectral radius of this matrix, that is, the maximum absolute value of its eigenvalues, defines the basic reproduction number $R_0$. 

Figure 7: Healthy cells time evolution for all patients. Pat C shows a very slow value of $U_\infty$ (practically zero), which suggest that the selected value of $U_0 = 1.0e^7$ is too large.
For the specific case of system (2.1), the next-generation matrix is given by

\[
FG^{-1} = \begin{bmatrix}
\frac{\beta p U_0}{c d} & \frac{\beta U_0}{c} \\
0 & 0
\end{bmatrix}
\]

Therefore, the basic reproduction number \( R_0 \) is given by

\[
R_0 = \frac{\beta p U_0}{c d}
\]

Notice that \( R_0 \) coincides with the entry \((1,1)\) of matrix \( FG^{-1} \), thus meaning that \( R_0 \) represents the expected number of secondary infections produced in compartment \( I \) by an infected cell originally in \( I \).

9. Appendix 3. General virus characterization for the dynamic (2.1)

The next theorem characterizes all possible virus behavior, depending on the arbitrary initial conditions and the parameters.

**Theorem 9.1** (Virus behavior from an arbitrary time). Consider system (2.1), constrained by the positive set \( \mathbb{R} \), at some arbitrary time \( t_0 \), with \( U(t_0) > 0 \), \( I(t_0) \geq 0 \) and \( V(t_0) > 0 \) (i.e., \( x(t_0) = (U(t_0), I(t_0), V(t_0)) \in \mathcal{X} \)). Let define \( R(t) := \frac{p U(t)}{c V(t)} \) and \( R_V(t) := \frac{p U(t)}{c V(t)} \). Then,

i. if \( R_V(t_0) < 1 \) and \( R(t_0) < 1 + \alpha(t_0) \), where \( \alpha(t_0) \) is a positive value, depending on \( I(t_0), V(t_0) \) and the parameters, then, \( V(t) < 0 \) for all \( t > t_0 \) (i.e., \( V(t) \) is strictly decreasing for all \( t > t_0 \) and, so, the virus does not spread in the body host),

ii. if \( R_V(t_0) < 1 \) and the virus spreads in the body host (i.e. \( V(t) \) reaches a local maximum, at some time \( t_V > t_0 \), before it goes to zero), then \( R(t_0) > 1 + \alpha(t_0) \). Even more, \( V(t) \) has a local minimum at some time \( t_V > t_0 \), with \( V_V < t_V \), and after the local maximum it is strictly decreasing, i.e., \( \dot{V}(t) < 0 \) for all \( t > t_V \). On the other hand, \( I(t) \) has only one local maximum at time \( t_1 \), being \( t_0 < t_1 < t_0 < t_V \), where \( t_0 \) is the time when \( R \) reaches 1 from above (or, the same, when \( U(t) \) reaches its critical value \( U_c \)). This means that the virus spreads in the host body.

**Proof:** First note that for positive parameters, equation (2.1.a) implies that \( \dot{U}(t) < 0 \) for all \( t \geq t_0 \) and, so, \( R(t) := U(t) \frac{dU}{c} \) is strictly decreasing for all \( t \geq t_0 \). Given that \( R_\infty := \lim_{t \to \infty} R(t) = U_\infty \frac{dU_c}{c} \) and \( U_\infty < U_c := \left\langle \frac{c}{\beta p} \right\rangle \) (see Theorem 3.1), then \( R_\infty < 1 \). So, for \( R(t_0) > 1 \), there exists only one time \( t_c > t_0 \),

\[
t_c > t_0,
\]

at which \( R(t_c) = 1 \), being \( R(t) > 1 \) for \( t_0 < t < t_c \) and \( R(t) < 1 \) for \( t > t_c \).

i. By hypothesis, \( R(t_0) = \frac{p(t_0)}{c V(t_0)} < 1 \), which implies that \( \dot{V}(t_0) < 0 \) (\( V \) starts decreasing at \( t_0 \)). On the other hand, by Lemma 1 \( V \) reaches a minimum or an inflection point at at time \( t^*_V > t_0 \) if \( R(t^*_V) \geq 1 \). But \( R(t) \) is strictly decreasing and, so, to reach 1 (at least) at \( t^*_V > t_0 \) it must be

\[
R(t_0) = 1 + \alpha(t_0),
\]

for some \( \alpha(t_0) > 0 \). However, by hypothesis it is \( R(t_0) < 1 + \alpha(t_0) \), which implies that \( \dot{V}(t) < 0 \) for all \( t > t_0 \). This concludes the proof.

ii. By hypothesis, \( R_V(t_0) < 1 \), which means that \( \dot{V}(t_0) < 0 \), and \( V(t) \) reaches a local maximum at some time \( t_V > t_0 \). Therefore, \( V(t) \) must reach a local minimum at some time \( t_0 < t_V < t_V \). At both, the local minimum and maximum it is \( \dot{V}(t_V) = 0 \) and \( \dot{V}(t_V) = 0 \). Then, by Lemma 1 it is \( R(t_V) > 1 \) and \( R(t_V) < 1 \), respectively.
Since $R(t)$ is strictly decreasing for $t > t_0$ and $\dot{t}_V > t_0$ (note that $\dot{V}(t_0) < 0$ and $\dot{V}(\dot{t}_V) = 0$, so $\dot{t}_V$ cannot be equal to $t_0$), then $R(t_0) > R(\dot{t}_V) > 1$, which implies that $R(t_0) > 1 + \alpha(t_0)$ for some $\alpha(t_0) > 0$. Furthermore, $t_0 < \dot{t}_V < t_c < t_V$ ($R$ crosses 1 at time $t_c$ between $\dot{t}_V$ and $t_V$) and, given that $R(\dot{t}_V) < 1$, $V$ cannot reach another local minimum after its local maximum. This implies that $\dot{V}(t) < 0$ for $t > \dot{t}_V$.

From the minimum and maximum conditions of $V$, at times $\dot{t}_V$ and $t_V$, they are $\dot{V}(\dot{t}_V) = 0, \ddot{V}(\dot{t}_V) > 0$ and $V(\dot{t}_V) = 0, \ddot{V}(\dot{t}_V) < 0$, respectively. After some algebraic computation, it is easy to see that $\dot{V}(\dot{t}_V) > 0$ and $\ddot{V}(\dot{t}_V) < 0$, which means that $I(t)$ must reach a maximum at some time $\hat{t}_I$, fulfilling $\dot{t}_V < \hat{t}_I < t_V$. Even more, it must be $\dot{I}(\hat{t}_I) = 0$, or

$$\beta U(\hat{t}_I)V(\hat{t}_I) - \delta I(\hat{t}_I) = 0. \quad (9.3)$$

Given that $\dot{V}(t) > 0$ for $\dot{t}_V < t < \hat{t}_V$ (it goes from its minimum to its maximum), then by (2.1a), $I(\hat{t}_I) > \frac{p}{\beta}V(\hat{t}_I)$. Replacing this later condition in (9.3), it follows that

$$\left(\beta U(\hat{t}_I) - \frac{\delta c}{\rho}\right)V(\hat{t}_I) > \beta U(\hat{t}_I)V(\hat{t}_I) - \delta I(\hat{t}_I) = 0, \quad (9.4)$$

which implies that $R(\hat{t}_I) = \frac{\beta U(\hat{t}_I)}{\delta c} > 1$ and, then, $\hat{t}_I < t_c$. Therefore, $t_0 < \dot{t}_V < \hat{t}_I < t_c < \dot{t}_V$, which concludes the proof.

iii. By hypothesis, $R_V(t_0) = \frac{pI(t_0)}{\delta cV(t_0)} > 1$, which implies $\dot{V}(t_0) = pI(t_0) - cV(t_0) > 0$ ($V$ starts increasing at $t_0$). Since $V_\infty = 0$ (Theorem 3.1), then, there exists $\dot{t}_V > t_0$ such that $V(\dot{t}_V)$ is a maximum. According to Lemma 1 if $V$ has a maximum at $\dot{t}_V$, then $R(\dot{t}_V) < 1$.

On the other hand, for $V(t)$ to reach a minimum after time $\dot{t}_V$, it must be $R(\dot{t}_V) > 1$. But $R(t)$ is strictly decreasing for $t > t_0$, which means that no further minimum exists after $\dot{t}_V$. This implies that $\dot{V}(t) < 0$ for all $t > \dot{t}_V$, which concludes the proof. □

**Lemma 1.** Consider system (5.1), constrained by the positive set $\mathcal{X}$, at some arbitrary time $t_0$, with $U(t_0) > 0$, $I(t_0) \geq 0$ and $V(t_0) > 0$ (i.e., $x(t_0) = (U(t_0), I(t_0), V(t_0)) \in \mathcal{X}$). Then, (i) if $V(t)$ reaches a local minimum at time $t_V^* > t_0$, then $R(t_V^*) > 1$, (ii) if $V(t)$ reaches a local maximum at time $t_V^* > t_0$, then $R(t_V^*) < 1$, and (iii) if $V(t)$ reaches an inflection point at time $t_V^* > t_0$ (a point in which $\ddot{V} = 0$), then $t_V^* = t_c$, where $t_c$ is the (unique) time at which $R$ reaches 1 (i.e., $R(t_c) = 1$ or, the same, $U(t_c) = U_c$).

**Proof:** Any of the three hypothesis ($V(t)$ reaches a local minimum, a local maximum or a inflection point) implies that

$$\dot{V}(t_V^*) = pI(t_V^*) - cV(t_V^*) = 0, \quad (9.5)$$

which means that

$$I(t_V^*) = \frac{p}{c}I(t_V^*). \quad (9.6)$$

Consider the critical case of an inflection point, i.e.,

$$\dot{V}(t_V^*) = p\dot{I}(t_V^*) - cV(t_V^*) = p\dot{I}(t_V^*) = 0. \quad (9.7)$$

From (9.7) it is $\dot{I}(t_V^*) = 0$ which, by (2.1b) at $t_V^*$, is equivalent to

$$\dot{I}(t_V^*) = \beta U(t_V^*)V(t_V^*) - \delta I(t_V^*) = 0. \quad (9.8)$$

Now, by (9.6), we have

$$\left(\frac{\beta p}{c} U(t_V^*) - \delta\right)I(t_V^*) = 0. \quad (9.9)$$
Given that \( I(t^*_v) > 0 \) (note that \( I(t) \) is positive for all \( t > 0 \)), then \( \frac{\beta p}{c_0} U(t^*_v) - \delta = 0 \), or

\[
\mathcal{R}(t^*_v) = \frac{\beta p}{c_0} U(t^*_v) = 1. \tag{9.10}
\]

This way if an inflection point does occur at \( t^*_v \), then \( t^*_v = t_c \), where \( t_c \) is the time at which \( \mathcal{R} = 1 \). This proves item (iii).

Furthermore, if \( V \) reaches a local minimum at \( t^*_v \), then \( \ddot{V}(t^*_v) > 0 \) (instead of \( \ddot{V}(t^*_v) = 0 \), as it is in (9.7), which by (9.6) implies that

\[
\mathcal{R}(t^*_v) = \frac{\beta p}{c_0} U(t^*_v) = 1. \tag{9.11}
\]

This proves item (i).

On the other hand, if \( V \) reaches a local maximum at \( t^*_v \), then \( \ddot{V}(t^*_v) < 0 \) (instead of \( \ddot{V}(t^*_v) = 0 \), as it is in (9.7), which by (9.6) implies that

\[
\mathcal{R}(t^*_v) = \frac{\beta p}{c_0} U(t^*_v) < 1. \tag{9.12}
\]

This proves item (ii). □

Figure 8 shows schematic plots of cases i, ii and iii of Theorem 9.1. For the sake of simplicity, \( \beta = \delta = p = c = 1 \), which means that \( t_c = 1 \), while the initial time is selected to be zero, i.e., \( t_0 = 0 \).

Figure 8 first column, illustrates case (i), in which \( \mathcal{R}_V(0) < 1 \) and \( \mathcal{R}(0) < 1 + \alpha(0) \), being \( \alpha(0) = 0.43 \) (numerically computed). This simulation shows that even if \( \mathcal{R}(0) > 1 \), but it is not greater than \( 1 + \alpha(0) \), the virus does not spread in the body host, i.e, it is strictly decreasing. Note that the maximum of \( \mathcal{R}_V \) does not reach 1. Figure 8 second column, illustrates case (ii), in which \( \mathcal{R}_V(0) < 1 \) and \( \mathcal{R}(0) > 1 + \alpha(0) \), being \( \alpha(0) = 0.43 \). This simulation shows that for \( \mathcal{R}(0) > 1 + \alpha(0) \), the maximum of \( \mathcal{R}_V \) is greater than 1, and the period of time in which \( \mathcal{R}_V > 1 \) is precisely the period of time the virus increases and reach a maximum (i.e., it spreads in the body host). As expected, \( V \) reaches a minimum first, then \( I \) reaches a maximum, then \( \mathcal{R} \) reaches 1 (at \( t_c \)) and, finally, \( V \) reaches a maximum, before to strictly decrease to zero. Figure 8 third column, shows case (iii), in which \( \mathcal{R}_V(0) > 1 \) and \( \mathcal{R}(0) > 1 + \alpha(0) \). As expected, the virus spreads on the body host an it has only one global maximum.

Remark 7. The value of \( \alpha \) is necessary to properly understand and characterize the system behavior according to the initial conditions and parameters. Although it cannot be explicitly defined, it can be computed numerically. Furthermore, it should be noted that for real patients data, this values use to be small in comparison with \( \mathcal{R} \), given that \( \beta \) is small (see Table 2).

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Figure 8: Theorem 9.1 schematic plot. $\beta = \delta = p = c = 1$, which means that $U = 1$. The initial time is selected to be $t_0 = 0$. First column: case (i). System time evolution when $U(0) = 1.2, I(0) = 0.25, V(0) = 0.4$, such that $R_V(0) = 0.63 < 1$ and $R(0) = 1.2 < 1 + \alpha(0)$, being $\alpha(0) = 0.43$. Second column: case (ii). System time evolution when $U(0) = 1.8, I(0) = 0.25$ and $V(0) = 0.4$, such that $R_V(0) = 0.63 < 1$ and $R(0) = 1.8 > 1 + \alpha(0)$, being $\alpha(0) = 0.43$. Third column: case (iii). System time evolution when $R_V(0) > 1$ and $R(0) > 1 + \alpha(0)$. 
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