A general framework for modeling tumor-immune system competition and immunotherapy: Mathematical analysis and biomedical inferences

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Abstract

In this work we propose and investigate a family of models, which admits as particular cases some well known mathematical models of tumor-immune system interaction, with the additional assumption that the influx of immune system cells may be a function of the number of cancer cells. Constant, periodic and impulsive therapies (as well as the non-perturbed system) are investigated both analytically for the general family and, by using the model by Kuznetsov et al. [V.A. Kuznetsov, I.A. Makalkin, M.A. Taylor, A.S. Perelson, Nonlinear dynamics of immunogenic tumors: parameter estimation and global bifurcation analysis, Bull. Math. Biol. (1994) 56(2) 295–321], via numerical simulations. Simulations seem to show that the shape of the function modeling the therapy is a crucial factor only for very high values of the therapy period \( T \), whereas for realistic values of \( T \), the eradication of the cancer cells depends on the mean values of the therapy term. Finally, some medical inferences are proposed.

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1. Introduction

Millions of people die from cancer every year [1]. And worldwide trends indicate that millions more will die from this disease in the future [2]. Great progress has been achieved in fields of cancer prevention and surgery and many novel drugs are available for medical therapies [3–5]. Biophysical models may prove to be useful in oncology not only in explaining basic phenomena [6,7], but also in helping clinicians to better and more scientifically plan the schedules of the therapies [7,8]. An interesting therapeutic approach is immunotherapy [4,5], consisting in stimulating the immune system in order to better fight and hopefully eradicate, a cancer. In particular, in this paper I will be referring to generic immunostimulations, for example, via cytokines, but for the sake of simplicity I will use the term "immunotherapy". The basic idea of immunotherapy is simple and promising,
but the results obtained in medical investigations are globally controversial [9–12], even if in recent years there has been evident progress. From a theoretical point of view, a large body of research has been devoted to mathematical models of cancer-immune system interactions and to possible applications to cure the disease [13,14,16–24] (and references therein). Analyzing the best known finite dimensional models [13,14,16,20,23], we note that their main features are the following:

• existence of a tumor free equilibrium;
• depending on the values of parameters, there is the possibility that the tumor size may tend to +∞ or to a macroscopic value;
• possible existence of a "small tumor size" equilibrium, which coexists with the tumor free equilibrium.

An "accessory" feature is the existence of limit cycles [16]. From this rough summary, one may understand that the puzzling results obtained up to now by immunotherapy [9] may be strictly linked to the complex dynamical properties of the immune system-tumor competition. In general, it happens that the cancer-free equilibrium coexists with other stable equilibria or with unbounded growth, so that the success of the cure depends on the initial conditions, and – even theoretically – it is not always granted.

2. A general family of models and its properties

In [22], Sotolongo-Costa et al. proposed the following very interesting Volterra-like model (similar to the one in [20]) for the interaction between a population of tumor cells (whose number is denoted by \( X \)) and a population of lymphocyte cells (\( Y \)):

\[
\begin{align*}
X' &= aX - bXY \\
Y' &= dXY - fY - kX + u + p(t),
\end{align*}
\]

where the tumor cells are supposed to be in exponential growth (which is, however, a good approximation only for the initial phases of the growth) and the presence of tumor cells implies a decrease of the "input rate" of lymphocytes. Systems (1) and (2) may be rewritten in non-dimensional form [22]:

\[
\begin{align*}
x' &= ax - xy \\
y' &= xy - \frac{1}{\sigma}y - kx + p(t)
\end{align*}
\]

(in short notation \((x', y') = C(x, y))\). The function \(p(t) \geq 0\) is assumed periodic with period \(T\) and it models the effect of immunotherapy. The model has been studied in depth both in the case of absence of therapy and in the case of therapy by using the test function \(p(t) = 0.5F(1 + \cos(4\pi t))\).

The model shows two equilibria (one of which is tumor-free) and also unbounded growth. However, the systems (3) and (4) allows negative solutions for non-small \( x \), which is not physically acceptable. In fact:

\[
C(x, 0) = ax + p(t) - kx
\]

implies that for \( x > (\sigma + p_{\text{max}})/k \) it is \( C(x, 0) > 0 \), \( -1 > 0 \), and \( y(t) \) becomes negative in finite times. Furthermore, the second equilibrium point is a consequence of the negativity of \( \sigma - kx \).

The model in [22], though it has this problem of lack of physical consistency, is, however, of great interest because the killing of lymphocytes is seen as a function of the \( x \) variable. Alternatively, the influx of lymphocytes may be thought of as a function of the entity of the disease, which we will denote as \( Q(s) \). Indeed, it has been observed that in some cases cancer progression may cause generalized immunosuppression (see [25], and references therein). Thus, in [22] it is \( Q(s) = \sigma(1 - (1/\sigma)s) \), which may be read as a first-order Taylor approximation of a more general non-increasing function.

However, a general influx function is only one of the possible modifications of models (3) and (4): there may be others, which are also biologically reasonable. One might take into the account many factors: different functional forms for the interaction term, saturation in the predation term and, mainly, non-exponential growth of the cancer: logistic, gompertzian, generalized logistic, etc. . . All these modifications are reasonable and useful. Thus, I think that it might be useful to define and study the following general family of models:

\[
x' = x(a f(x) - \phi(x))
\]
\[ y' = f(x)y - \mu(s)y + \sigma q(x) + \theta(t) \]  

(7)

where:

- \( x \) and \( y \) are the non-dimensionalized numbers of, respectively, tumor cells and of effectors cells of immune system;
- \( 0 < f(0) \leq +\infty \), \( f'(x) \leq 0 \) and in some relevant cases we shall suppose that it exists an \( 0 < \lambda \leq +\infty \) such that \( f(\lambda) = 0 \), \( \lim_{x \to +\infty} x f(x) = 0 \). Thus, \( f(x) \) summarizes many widely used models of tumor growth rates, such as the Exponential model: \( f(x) = 1 \) [7], the Gompertz: \( f(x) = \log(A/x) \) [7,50] and its generalizations [7,50], the Logistic model: \( f(x) = 1 - x/A \) [50], the Hart–Schochat–Agur: \( f(x) = x^{-\gamma} \), \( 0 < \gamma < 1 \) [26], the von Bertalanffy: \( f(x) = x^{1/2} - a \) [50,53], the Guo’s et al model: \( f(x) = x^{1/2} - b \) [27], the linear growth model by Bru et al [28] which may be written as follows: \( f(x) = x^{-1/2} \) (note that it may be considered a particular case of the von Bertalanffy model and of the Hart–Schochat–Agur model), etc. . . ;

- \( \phi(x) > 0 \), \( \bar{\phi}(0) = 1 \), \( \tilde{\phi}(x) \geq 0 \) and \( \tilde{\phi}(x) \to 1 \leq +\infty \);
- \( q(x) \) is such that \( q(0) = 1 \) (as a consequence \( \sigma = Q(0) \)) and it may be non-increasing or also initially increasing and then decreasing, i.e. we may assume that either the growth of tumor decreases the influx of immune cells or that, on the contrary, it initially stimulates the influx;
- \( \beta(x) \geq 0 \), \( \beta(0) = 0 \) and \( \beta'(x) \geq 0 \);
- \( \mu(x) > 0 \) and \( \mu'(x) > 0 \).

For the sake of simplicity we define the following function \( \Psi(x) = \mu(x) - \beta(x) \) and write:

\[ x' = x(\sigma f(x) - \phi(x)) \]  

(8)

\[ y' = -\Psi(x)y + \sigma q(x) + \theta(t). \]  

(9)

\( \Psi(x) \) is assumed to be positive, otherwise it may be positive in \( [0, x_1) \cup (x_2, +\infty) \) with \( \Psi(x_1) = \Psi(x_2) = 0 \). We may assume that it has an absolute minimum in \( [0, +\infty) \). We may use \( \Psi(x) \) to classify the tumors depending on their degree of aggressiveness against the immune system:

- \( \Psi(x) > 0 \); in such a case the ability of destroying immune cells is never won by the stimulatory effect on the immune system, therefore the tumor may be indicated as “highly aggressive”/“lowly immunogenic”;
- Variable sign \( \Psi(x) \); since in such a case the destruction of cells may be compensated by the stimulatory effect, we will refer to such a tumor as “lowly aggressive”/“highly immunogenic”.

The above model includes as particular cases the models [13,14,20,23]. For instance, the Stepanova model [13] is such that \( f(x) = 1 \), \( \phi(x) = 1 \), \( \tilde{\phi}(x) = 1 \) and \( \mu(x) = \mu_0 + \mu_1 x^2 \), the de Vladar–Gonzalez model [23] is similar, but: \( f(x) = \log(K/x) \).

Note that Nani and Freedman proposed an interesting model of adoptive cellular immunotherapy in which generic functions are used [19]. However, their approach differs from ours since in their model the proliferation of cells of the immune systems is not stimulated by cancer cells. In other words in the Nani and Freedman model the interaction tumor cells – immune system is only destructive for immune cells. Furthermore, in their model the “loss rates” are proportional (in our notation we might write \( \mu(x) = \mu(0) + \text{const}(\phi(x)) \)).

In the absence of treatment, systems (8) and (9) admit the existence of a cancer-free equilibrium \( CF = (0, \sigma(\Psi(0)) \).

If \( f(0) < +\infty \), we have that if \( \sigma > \sigma_a = \alpha f(0)/\phi(0) \) \( CF \) is locally asymptotically stable (LAS), unstable if \( \sigma < \sigma_a \). Biologically, \( \sigma > \sigma_a \) means that the immune system works very well and that it is able to destroy small tumors. On the contrary \( \sigma \approx 0 \) means that there is immunodepression.

Furthermore, when \( \phi(x) = \text{constant} = \phi_0 \) and \( \Psi(x) \leq \Psi^* \leq +\infty \). If \( \sigma > \sigma^* = \alpha f(0)/\phi(0)/\Psi^* \) it follows that \( CF \) is globally asymptotically stable (GAS). In fact, from \( y' = -\Psi(x)y + \sigma q(x) \geq -\Psi(x)y + \text{constant} \) if follows that asymptotically \( y(t) \geq \text{constant}/\Psi(x) \). As a consequence, asymptotically \( x' \leq (\alpha f(0) - \phi(0)/\Psi(x)) x, \) i.e. if \( \sigma > \sigma^* \) it is \( x(t) \to 0 \Rightarrow y(t) \to \text{constant}/\Psi(x) \).

A relevant problem, up to now, is that the immunotherapeutic agents are characterized by strong toxicity, thus \( \sigma > \sigma^* \) might be too biologically high, even in cases in which it is mathematically small. If \( f(0) = +\infty \), as in the Compartitizan case (used, for example, in [23]) and in other tumor growth models, then \( CF \) is unstable anyway (as previously stressed for the particular model [23]) because in such a case the derivative of \( x f(x) \) at \( x = 0 \) is \( +\infty \). In the light of
and by Konarski and Waliszewski. Gompertz model have been elucidated by Konarski and thermore, some interesting physical properties of the mathematical justification of the Gompertz model. Fur-thermore, some interesting physical properties of the physiological ground in. Using data from multicellular physically the Gomp-Ex model proposed on biological-exponential growth. In other words, they derived bio-mors. From a theoretical point of view, model fits data well from experimental and in vivo tumors[36,35,37–41]. From a theoretical point of view, Gyllenberg and Webb, Calderon and Kwembe [42,43], Calderon and Afenya [44,45] proposed physico-mathematical justification of the Gompertz model. Fur-thermore, some interesting physical properties of the Gompertz model have been elucidated by Konarski and Molski [46] and by Konarski and Waliszewski [47].

However, the doubling time of a population of cells cannot be lower than the minimal time needed by a cell to divide, which is obviously non-null. This biological constraint is in contrast with the unboundedness of \( f(x) \) in the Gompertz and other models, as stressed by Wheldon [7]. More recently, inconsistency at low number of cells have been recognized by Cast-orina and Zappala in their derivation of the Gom-pertzian model based on methods of statistical mechanics [48,49]. They showed that the validity of the Gom-pertz model starts above a minimum threshold for the number of cells, whereas under the threshold there is exponential growth. In other words, they derived bio-physically the Gomp-Ex model proposed on biological ground in [54,7]. Using data from multicellular tumor spheroids, Marusic et al. performed a systematic comparison of many models [50], which showed that Gompertz’s model fitted their data very well, but slightly less well than the Piantadosi model [55], which has finite \( f(0) \). Furthermore, in their fittings, it was not possible to discriminate between the pure Gompertz model and the Gomp-Ex model. Demicheli et al. used Gomp-Ex model on in vitro and in vivo data obtaining results strongly supporting this model [52]. Other comparisons may be found in [44,53]. Moreover, in general, van Leeuwen and Zonneveld [51] claims that it may be not possible to discriminate between exponential, logistic and gompertzian models in the early phases of growth. Recent experimental studies conducted by Bru and coworkers support an initial phase of exponential growth [28]. Summarizing, I consider the results by de Vladar and Gonzalez (and our extensions) to be very valuable, but they may be read in a dichotomic way:

- A tumor is permanent: the innate immune surveil-lance is never able to completely eradicate even the smallest tumor.
- Since there is relevant evidence that the immune sys-tem is able in some cases to eliminate small tumors [57,58] (as we will see in following sections, the ability of eradicate the disease or not depends on initial conditions), the properties of the de Vladar–Gonzalez model (and of our extension) may be seen as an evidence that Gompertzian and other models characterized by \( f(0) = +\infty \) are not appropriate for very small tumors, in coherence with [7,48,49,28].

In case of the absence of influx of immune cells \( q(x) = 0 \) and for laws of growth in which \( x \) exists, there is a different particular equilibrium point, which we shall call “immune free”: \( \Phi = (\bar{x}, 0) \), which is LAS.

Other multiple non-null equilibria may be found by finding the positive intersection of the two nullclines:

\[
\begin{align*}
\gamma_y(x) &= \frac{f(x)}{\Phi(x)} \\
\gamma_x(x) &= \sigma(x) \Phi(x)
\end{align*}
\]

The functions \( \gamma_y(x) \) and \( \gamma_x(x) \) are useful in the deter-mination of the LAS of the equilibria, since the char-acteristic polynomial of the Jacobian, calculated at a given equilibrium point \( (x_e, y_e) \), is:
\[ \dot{\lambda}^2 + \Psi(x) - \lambda^3 \dot{\phi}(x) \ddot{\lambda} = 0. \]

So the LAS condition is:
\[ y'_c(x) < \frac{\Psi(x)}{3\Phi(x)} \quad \text{AND} \quad y'_c(x) > y'_c(\bar{x}). \]

Note that the first part of the AND condition is automatically fulfilled when \( y'_c(\bar{x}) \leq 0 \) (because \( x_c \) cannot lie in an interval where \( \Psi(x) < 0 \), whereas the second part has a straightforward geometrical interpretation.

Finally, it is interesting to note that the above family of model may admit limit cycles if \( f(x) = 1 \) (exponential growth) and \( \phi(x) \) is identically null for \( x > x_0 \) with \( x_0 < x_c \). In fact, in such a case there is the equilibrium point \((x_c, a)\) whose characteristic polynomial is:
\[ \lambda^2 + \dot{h}^2 = -x_1 \Psi(x_1) > 0. \]

In effect, some cases of sustained oscillations (or slow oscillations with very small damping) have been reported in the medical literature [29–31]. Periodic solutions in absence of influx of immunocompetent cells are predicted also in [16].

On the contrary, if \( y'_c(\bar{x}) \leq 0 \) (for example when \( \phi(x) \) is constant), by applying the Dulac–Bendixon theorem with multiplicative factor \( 1/(x\Psi(x)) \) (as in the specific models [14,20]) one obtains that the presence of limit cycles is not possible. In fact:
\[ \text{Div} \left( \frac{1}{x\Psi(x)}(x', y), y'(x, y)) \right) = \dot{y}'_c(x) - \frac{\phi(x)}{x\Psi(x)} < 0. \]

2.1. The global behavior

In some important cases, it is possible to study the global behavior of the family, by means of differential inequalities and of the Poincare–Bendixon trichotomy [56]. We may state the following simple propositions:

**Proposition 1.** When \( \Psi(x) > 0 \) and \( f(x) = 1 \) and \( y'_c(\bar{x}) \geq 0 \), if it is \( y(x) < y_c(\bar{x}) \) then \( x(t) \rightarrow +\infty. \)

**Proof.** Let us define \( y^{MAX}(x) = \max_{x \in [0,1]} y(x) \) and \( y_M \) such that \( y(x_M) = y^{MAX} \). If it is \( y(x) < y_c(\bar{x}) \) it is easy to show that the set \( H = \{(x, y) | x > x_M \quad \text{AND} \quad 0 \leq y \leq y^{MAX} \} \) is positively invariant and absorbing. Thus, since in \( H : x' \geq y(x,y) \quad \text{AND} \quad y' \geq 0 \), it follows readily that \( x(t) \rightarrow +\infty. \)

**Proposition 2.** If \( \Psi(x) > 0 \), it exists \( \lambda \) such that \( f(x) = 0 \), \( y'_c(x) < 0 \) and there is a unique LAS equilibrium point \( S = (x_c, y_c) \), then \( S \) is GAS.

**Proof.** Let us define \( y^{MAX} \) and \( y^{MIN} \) such that \( y(x_M) \) and \( y(x_m) \) are respectively maximum and minimum. Furthermore, if \( f(0) > y^{MAX}(x) \) let it be \( \lambda = y'(y_{y^{MAX}}(x)) \), if \( f(0) \leq y^{MAX}(x) \) let it be \( \lambda = 0 \). Since \( \Psi(x) y^{MIN} \leq y' \leq \Psi(x) y^{MAX} \) it is easy to see that the set \( R = \{ (x, y) | y' \leq \lambda \} \) and \( y^{MIN} \leq y \leq y^{MAX} \) is positively invariant and absorbing and contains \( S \). Since we have ruled out the possibility that there may be limit cycles, as a consequence \( S \) is GAS.

**Proposition 3.** When \( \Psi(x) > 0 \) and \( y'_c(\bar{x}) \) is non-constant and there is a unique LAS equilibrium point \( S = (x_c, y_c) \), if it holds also that \( y'_c(x_c) > y'_c(\bar{x}) \) then \( S \) is GAS.

**Proof.** When \( f(x) \) is unbounded, one may see that there may be a relative minimum followed by a relative maximum in \((0, \lambda)\). On the contrary, when \( f(x) \) is bounded, there is an absolute maximum. Calling now \( x^* \) the point in which \( y_c(x) \) is absolutely or relatively maximum, one has that \( x^* \) and \( y_{y^{MAX}} \leq y \leq y^{MAX} \) is positively invariant and absorbing, contains \( S \). Since in \( R^* \) it is \( y'_c(x) \geq 0 \) (which implies that closed orbits are ruled out), as a consequence, \( S \) must be GAS.

**Proposition 4.** When \( \Psi(x) > 0 \) and \( y(x) > y_c(x) \) for \( x \in [0, 1] \) then \( CF \) is GAS.

**Proof.** It is a particular case of Proposition 2.

**Proposition 5.** If \( \Psi(x) > 0 \), there does not exist a \( \lambda \) such that \( f(x) = 0 \), \( y'_c(\bar{x}) < 0 \) and there is a unique LAS equilibrium point \( S = (x_c, y_c) \), then \( S \) is GAS.

**Proof.** Let us define \( y^{MAX} := \max_{x \in [0,1)} y(x) \) and consider a point \( P_0 = (x_0, 0) \) with \( x_0 > x_c \), and the
orbit starting from it, which intersects the curve $y_C(x)$ in the point $P_o = (x_o, y_C(x_o))$. Let us consider the following points $P_o = (x_o, y_{MAX})$, $P_0 = (0, 0)$ and $P_8 = (0, 0)$. The arc of orbit $P_8, P_0$ and the straight segments $P_0, P_8$, $P_8, P_0$ and $P_0, P_8$ bounds an invariant set for our system. As a consequence of the Bendixson–Poincare’ trichotomy we have that $S$ is GAS.

Proposition 6. When $\Psi(x)$ has variable sign, and $f(x)$ is bounded and $y(x) = y_C(x)$ then CF is GAS.

Proof. The set $X = \{(x, y)|0 < x \leq x AND y \geq 0\}$ is positively invariant and adsorbing and in it closed orbits are impossible, as we have seen. However, it is not a bounded set, so we have to show that all the orbits starting in $X$ are bounded. Firstly, we notice that it cannot be $y(t) \to +\infty$, since in such a case, being $x' = x\varphi (x,y(t))$, it would be $x(t) \to 0 \Rightarrow y(t) \to \sigma/\Psi(0)$. Furthermore hypothetical solutions such that $\min lim_{t \to \infty} y(t) = 0$ and $\max lim_{t \to \infty} y(t) = +\infty$ are not possible since the set $A = \{x, y)|0 < x \leq x_1 AND y \geq y(x)\}$ is positively invariant. As a consequence of these properties, thanks to the Bendixson–Poincare’ trichotomy, CF is GAS.

Proposition 7. When $\Psi(x)$ has variable sign, there is $x$ such that $f(x) = 0$, $y(x) < 0$ and there is a unique LAS equilibrium point $S$ then $S$ is GAS.

Proof. The set $X = \{(x, y)|0 < x \leq x AND y \geq 0\}$ is positively invariant and adsorbing and in it closed orbits are impossible, as we have seen. However, it is not a bounded set. Let us consider $y_C(x)$: it is such that it is split in two branches: $y_{MIN}(x)$ for $x_2 \leq x \leq +\infty$ (which has no intersections with $y_C(x)$) and $y_{MAX}$ for $0 \leq x < x_1$ (on which $S$ lies). Let us consider a point $P_1 = (x_1, y_1)$ lying on the curve $(x, y_{MIN}(x))$ and having $y_1 > y_C(x_1) > y_C(x_2)$. Let the orbit starting from $P_1$ intersect the graph $(x, y_{MIN}(x))$ in a point $P_1 = (x_1, y_1) = (x_1, y_C(x_1))$ (note that it is $y_1 > y_C(x_1)$). Let us define the following points: $P_6 = (0, 0)$, $P_7 = (0, 0)$ and $P_8 = (0, 0)$. It is easy to see that segment of orbit $P_7 P_6$ and the straight segments $P_6 P_8, P_8 P_0, P_0 P_8$ and $P_8 P_0$ bound an invariant set $\mathcal{I}$ for our dynamical system. As a consequence, thanks to the Bendixson–Poincare’ trichotomy, $\mathcal{I}$ is GAS.

Proposition 8. When the sign of $\Psi(x)$ is variable, there is no $x$ such that $f(x) = 0$, $y(x) < 0$ and there is a unique LAS equilibrium point $S$ then $S$ is GAS.

Proof. The proof is easily obtained by applying methods of Propositions 7 and 5 to find a bounded positively invariant set surrounding $S$. □

Proposition 9. When $\Psi(x) > 0$ and $q(x) = 0$ then $\Psi(x(0), y(0))$ it is $y(t) \to 0^+$. Furthermore, in accordance with the growth law $f(x)$, either the tumor tends to an equilibrium value or it grows unbounded.

Proof. Let us define $\Psi_{MIN} = \min_{x \in [x]} \Psi(x)$. If $q(x) = 0$ it is $y' = -\varphi(x,y) \leq -\Psi_{MIN} y \Rightarrow y(t) \to 0^+$. Thus, the equation for $x(t)$ becomes asymptotically autonomous, so that, depending on $f(x)$, either $x(t) \to +\infty$ or $x(t) \to 0$ (i.e. in this case the equilibrium IF $(0, 0)$ is GAS). □

Proposition 10. When $\Psi(x) > 0$ and $f(x) = 1$ and $\phi(x) = const = \gamma$, and there are two equilibria $S = (x_1, y_1)$ (LAS) and $U = (x_2, y_2)$ (unstable) and there is a separatrix curve $y = \Sigma(x)$ which does not join $S$ to $U$, then there are two sets $A$ and $B$ such that if $(x(0), y(0)) \in A$ then $(x(t), y(t)) \to S$, whereas if $(x(0), y(0)) \in B$ then $(x(t), y(t)) \to +\infty$.

Proof. Let us define $\Sigma_{MAX} = \max_{x \in \mathcal{S}} y(x)$ and $\Sigma = \Sigma_{MIN} = \min_{x \in \mathcal{S}} y(x)$ and $\Sigma = \Sigma_{MIN} = \min_{x \in \mathcal{S}} y(x)$ and $\Sigma = \Sigma_{MAX} = \max_{x \in \mathcal{S}} y(x)$. As a consequence, the set $A = \{(x, y)|0 < x < x_1 AND \min_{y} (y(x)) \leq y \leq y_{MAX}\}$ is positively invariant and in it there are no closed orbits, so if $(x(0), y(0)) \in A$ then $(x(t), y(t)) \to S$. It is easy to show that given a $y(x_2)/\sigma < \varphi < a/\sigma$ also the set $B = \{(x, y)|x > x_2 AND 0 \leq y \leq \gamma\}$ is positively invariant. Thus, in $B$ $x' \geq x(a - \varphi)$, it easily follows that $y(t) \to +\infty$. □

Proposition 11. Let it be $\Psi(x) > 0$, $y_C(x) \leq 0$ and it exists $x$ such that $y_C(x) = 0$ Let there be four equilibria $CF$ (unstable), $S_1 = (x_1, y_1)$ (LAS), $U = (x_2, y_2)$ (unstable) and $S_2 = (x_3, y_3)$ (LAS), and let there be a separatrix curve $y = \Sigma(x)$ which does not join $S_2$ to $S_2$ to $U$, then there are two sets $A$ and $B$ such that if $(x(0), y(0)) \in A$ then $(x(t), y(t)) \to S_1$, whereas if $(x(0), y(0)) \in B$ then $(x(t), y(t)) \to S_2$. 

Proof. As in the previous proposition \( A = \{(x, y) \in \mathbb{R}^2 \mid x < 0 \} \) AND \( \text{Min}(0, \Sigma(x) \leq y \leq \Sigma_{\text{MAX}}) \) is positively invariant and in it there are no closed orbits, so if \((x(0), y(0)) \in A\) then \((x(t), y(t)) \to \delta_0\).

In this case \( B = (U(x), 0) \), \( x < 0 \) AND \( \delta_0 \leq y \leq \text{Min}(\Sigma(x), \Sigma_{\text{MAX}}) \), and it is positively invariant as well, and with no closed orbits in it. As a consequence: if \((x(0), y(0)) \in B\) then \((x(t), y(t)) \to \delta_0\). \(\Box\)

Remark. A consequence of the fourth proposition is that if \( y'(0) > 0 \) (or \( y'(0) = 0 \) AND \( y'(0) > 0 \)) then \( \sigma > \sigma_{\text{cr}} \) is a sufficient condition for the GAS of the CF equilibrium.

In case of multiple equilibria with \( \phi(x) = \text{const} \) it may be useful to transform (8) and (9) to a nonlinear oscillator. In fact by setting \( \bar{z} = \log(x) \) it is easy to see that the original family becomes:
\[
z'' + (\bar{\psi}(z) - \bar{\phi}'(z))z' + \omega \bar{\phi}'(z) - \bar{\phi}'(z)\bar{\phi}(z) = 0 \quad (17)
\]
where \( \bar{\psi}(z) = \psi(z) \), etc. . . . . By defining the damping coefficient:
\[
2
\psi(z) = (\bar{\psi}(z) - \bar{\phi}'(z))
\]
and the pseudo-potential:
\[
U(z) = \int_{0}^{\bar{z}} (\omega \bar{\phi}'(z) - \bar{\phi}'(z)\bar{\phi}(z))dz
\]
and the total pseudo-energy:
\[
E_{\text{tot}} = \frac{\zeta^2}{2} + U(z)
\]
it follows immediately that when \( \psi(z) > 0 \):

- Let it be \( \xi < +\infty \) and let there be three equilibria \( \zeta < \zeta_1 < \zeta_2 \) which are, respectively LAS, unstable and again LAS. Let it be \( E_\text{tot}(0) < U(\zeta_1) \), then \( \zeta(0) < \zeta_1 \Rightarrow \zeta(t) \to \zeta_1 \), whereas \( \zeta(0) > \zeta_1 \Rightarrow \zeta(t) \to +\infty \).

- Let it be \( \xi = +\infty \) and let there be two equilibria \( \zeta < \zeta_1 \) which are, respectively LAS and unstable. Let it be \( E_\text{tot}(0) < U(\zeta_1) \), then \( \zeta(0) < \zeta_1 \Rightarrow \zeta(t) \to \zeta_1 \), whereas \( \zeta(0) > \zeta_1 \Rightarrow \zeta(t) \to +\infty \).

3. On immunotherapies

3.1. Therapy schedulings

A realistic anticancer therapy may be modeled with sufficient approximation as constant (e.g. via a constant intravenous infusion) or periodic (e.g. the agent is delivered each day as a bolus):
\[
\theta(t) = \theta_0 + \sum_{n=0}^{\infty} \delta(t-nT) \geq 0, \quad \theta(t+T) = \theta(t),
\]
\[
\theta_0 = \frac{1}{T} \int_{0}^{T} \delta(t)dt
\]
(21)

For humans, typical periods ranges between 8 h and 7 days \([9,5]\). A particular case of periodic therapy is pulsed therapy, i.e. a therapy which induces an instantaneous increase of the number of lymphocytes:
\[
\theta(t) = \theta_0 + \sum_{n=0}^{\infty} \delta(t-nT)
\]
(22)

In the case of constant infusion therapy (CIT) \( \theta(t) = \theta_0 \) by defining:
\[
\hat{\sigma} := \sigma + \theta_0, \quad \hat{\phi}(t) := \frac{\sigma + \theta_0}{\hat{\sigma}}
\]
(23)

Remark. In the next subsections some asymptotic analyses of therapies shall be conducted. The meaning of the underlying \( t \to +\infty \) limits is the following: the therapies are administered for a time interval \([0, T]\) which is finite but sufficiently high to guarantee that the number of cancer cells is zero or that other targets have been reached.

3.2. Continuous infusion therapy

All the considerations we have done the absence of therapy hold also in case of CIT. In particular, for \( f(0) = +\infty \), the condition for the LAS of the cancer-free equilibrium is:
\[
\sigma + \theta_0 > \sigma_c
\]
(24)

Because of the co-presence of other equilibria, the above criterion is not global, i.e. the immunotherapy is not able to guarantee the disease eradication from whatever initial values \((x(0), y(0))\). However, observ-
ing that in models in which $\Psi(x) > 0$:

$$y_I(x) = \frac{c_0(x) + \theta m}{\Psi(x)} > y_U(x)$$

(e.g. in Stepanova’s model with low $\mu_1$) it happens that, roughly speaking, the stable equilibrium size of the cancer becomes smaller and the unstable equilibria greater, so that the basin of attraction of the unbounded solution is reduced.

Let us consider now some typical situations in case of $\Psi'(x) < 0$:

- Non-aggressive tumor (i.e. $\Psi(x) \leq 0$ in $[x_1, x_2]$). In such a case, in absence of therapy there may be in the most complex case four equilibria: CF (unstable), a small tumor equilibrium $E_{micro}$ (LAS), a macroscopic equilibrium $E_{macro}$ (LAS) and an intermediate unstable equilibrium $E_U$, as in Fig. 1, subplot 1. $E_{macro}$ is determined by the intersection between $\Psi(x)$ and the branch $y'_I(x)$, $E_{macro}$ by the intersection between $\Psi(x)$ and $y'_U(x)$. Increasing $\theta$ there are new equilibria. For $\theta > \theta_{cf} = y_C(0) - y_I(0)$ CF becomes at least LAS and $E_{macro}$ disappear. On the right, as a consequence of the elementary properties of continuous decreasing functions, increasing $\theta$ the equilibria move and it is $x_{EU}(\theta) > x_{EU}(0)$, $x_{macro}(\theta) < x_{macro}(0)$, and there exists $\theta_r > 0$ such that for $\theta > \theta_r$ all equilibria disappear. Summarizing, when $\theta > \theta_{cf}$ then CF is GAS (Fig. 1, subplot 3), because of Proposition 4 of Section 2.1.

- Aggressive tumors with variable sign $\Psi'(x)$. In such a case, in the absence of therapy there may in the most complex case be one macroscopic equilibrium

![Illustration of the effect of a CIT on a typical configuration in a lowly aggressive tumor. The case is shown in which $\theta_r < \theta_{cf}$. $y_I(x)$ is plotted as a solid line, whereas $y_C(x)$ is dashed. The equilibria are plotted as black points and they are labeled $U$ when unstable, otherwise $S$. First subfigure: in the absence of therapy there are four equilibria among which CF. Second subfigure: with a therapy with $\theta_r < \theta < \theta_{cf}$ CF is unstable and coexists with a microscopic tumor equilibrium which is GAS. Fourth subfigure: for a high dose therapy $\theta > \theta_{cf}$ CF becomes GAS.](image-url)
equilibrium: $E_{\text{Macro}}$ (GAS) and, of course, CF (unstable). Increasing $\theta$ two further equilibria may appear. The analysis is similar to the previous one (cf. Figs. 3 and 4) and we may find a $\bar{\theta}$ such that for $\theta > \bar{\theta}$ CF is GAS. Note that when the tumor is aggressive it is very likely that $\bar{\theta}$ is "extremely high"; $\bar{\theta} \gg \sigma$.

- Aggressive tumors with $\Psi(x) < 0$ [17]. In such a case, in the absence of therapy there may in the worst case be one macroscopic equilibrium equilibrium: $E_{\text{Macro}}$ (GAS) and, of course, CF (unstable). Increasing $\theta$, if when $y_I(0) = y_C(0)$ it is $y_I'(0) < y_C'(0)$ then we may find two values $\theta_1$ and $\bar{\theta}$ such that for $\theta_1 < \theta < \bar{\theta}$ CF is LAS and there is the birth of
a third unstable equilibrium $E_3$. Finally for $\theta > \hat{\theta}$
CF is GAS. Note that if when $y(t) = z(t) = y_C(0)$ it is
$y_C(0) > y_C(0)$ then $\tilde{\theta}_i = \hat{\theta}$.

When $f(t) = +\infty$ the total elimination cannot be
achieved by immunotherapy alone. Furthermore, even the
suboptimal target of reducing the cancer to a micro-
scopic size in many relevant cases cannot be achieved for
therapies of finite duration, however they may be
long. In fact, let it be $\Psi(t) > 0$ (aggressive tumor) and
let there be a unique GAS microscopic equilibrium
$E_{MACRO}$. By applying a CIT with $\theta$ sufficiently high
there is a unique GAS microscopic equilibrium. How-
ever, when the therapy ceases $\theta$ falls to zero and the can-
cer restarts growing macroscopically, since $E_{MACRO}$ is
again GAS. We note in brief that if the original equilib-
rium is microscopic (e.g. micrometastasis) the effect of
the therapy is simply to create another and temporary
microscopic equilibrium.

We note that $\theta$ acts a global bifurcation parameter,
and we point out that these behavior may be observed
in case of bounded $f(t)$ when therapy is applied for an
insufficient time.

Finally, this simple analytical analysis may explain
theoretically some numerical results of [15] on the re-
relationships between the efficacy of the cure and the
proliferation rate of cancer, and on the correlation be-
tween the burden of initial size and the probability of
effectiveness of a therapy.

3.3. Periodic scheduling

In the case of periodic drug schedulings, there is a periodically varying cancer-free solution $C^p = (0, z(t))$, where $z(t)$ is the asymptotic periodic solution of:

\[ y' = -\Psi(0)y + \sigma + \theta_0 + \Omega(t) \]  

that, assuming $\Omega(t) = \sum_{k=1}^{\infty} C_k \cos(2\pi/k \cdot \tau - \zeta_k)$, can be rewritten as:

\[ z(t) = \frac{\sigma + \theta_0}{\Psi(0)} + \sum_{k=1}^{\infty} C_k \frac{1}{\Psi(0) + k^2(2\pi/k)^2} \times \cos \left( k \frac{2\pi}{\tau} t - \zeta_k - \text{Arg} \left( \frac{\Psi(0) + ik \cdot 2\pi}{\tau} \right) \right) \]  

Note that if $T \ll 1/\Psi(0)$ there is a filtering effect and
$z(t) \sim (\sigma + \theta_0)/\Psi(0)$.

Two basic models of therapy may be:

\begin{itemize}
  \item $\theta_0(t) = A(1 + b \cos(at))$ (28)
    which is rather unrealistic, but whose functional
    form is commonly used to assess the effect of peri-
   odic forcing on nonlinear systems. The asymptotic
    solution of (26) corresponding to (28) is given by:
    \[ z_a(t) = \frac{\sigma + A}{\Psi(0)} + \frac{Ab}{\sqrt{\Psi(0) + \omega^2}} \times \cos(at - \text{Arg}(\Psi(0) + i\omega)) \]  
    \item the more realistic function:
    \[ \theta_0(t) = \frac{G}{1 - \exp(-\tau t)} \exp(-c \text{Mod}(t, T)), \]
    \[ \theta_0 = \frac{G}{cT}, \]  
    \end{itemize}

which represent a bolus-based delivery. The “shape”
of $\theta_0(t)$ depends on $c$ and the corresponding asymp-
totic periodic solution of (26) is given by:

\[ z_a(t) = \frac{\sigma}{\Psi(0)} + \frac{G}{\Psi(0) - \epsilon} \times \left( \frac{E^{-\text{Mod}(T)}}{1 - E^{-\tau}} - \frac{E^{-\Psi(0)\text{Mod}(T)}}{1 - E^{-\Psi(0)}} \right) \]  

In case of impulsive therapy, by solving the impulsive
differential equation

\[ y' = -\Psi(0)y + \sigma, \quad y(T^-) = y(T^+) \]  

one obtains that:

\[ z(t) = \frac{\sigma}{\Psi(0)} + \frac{\gamma}{1 - \exp(-\Psi(0)T)} \times \exp(-\Psi(0)\text{Mod}(t, T)). \]  

Furthermore, it is easy to show that the condition $\sigma + \theta_m > \sigma_\ast$ guarantees the LAS of CF. In fact, since the variational equations around $(0, z(0))$ are: $U' = (aX(0) - \psi X(0))U$, $W' = (aq(0) - \psi q(0))U - \psi'(0)z(t)U$, we obtain that $aX(0) - \psi(0) < z(0) < 0 \Rightarrow U(t) \rightarrow 0 \Rightarrow W(t) \rightarrow 0$, and since $< z(t) \geq (\sigma + \theta_m)/\psi(0)$ we recover the LAS condition $\sigma + \theta_m > \sigma_\ast$. Similarly, one may demonstrate the GAS condition: $\sigma + \theta_m > \sigma_\ast$.

3.4. Numerical simulations

We performed a set of simulations of immunotherapy on the basis of the model proposed by Kuznetsov et al. [14], in which:

$$a(\mathbf{x}) = 1.636(1 - 0.002x), \quad \phi(x) = 1,$$

$$\beta(x) = \frac{1.1311}{20.19 + x}, \quad \eta(x) = 0.1181,$$

$$\mu(x) = 0.00311x + 0.3743,$$

and

$$t_{\text{ onc}} = 9.915 \text{ days}, \quad (X, Y) = 10^6 (x, y) \text{ cells}$$

We chose this model since its parameter values were fitted from real data of chimeric mice [14]. Note that the dynamic of tumors in mice is faster than that of human tumors, and that for periods of about 1 day or less (i.e. $T < 0.101$) it results that $(1/\mu(0)) \gg T$. Moreover, $\mu(x) = 0.00311 \ll 1$ and the tumor is not aggressive. We also performed simulations in a case of a more aggressive tumor, for which we set $\mu(x) = 1000.00311x + 0.3743$. For the non-aggressive tumor $\sigma_\ast \approx 0.612$ and $\sigma_\ast^* \approx 1.44 \gg \sigma$.

It is worth noticing that in other kinds of anticancer therapies the shape of the therapy may be critical in determining whether or not the cancer will be eradicated [8].

In our simulations we assumed $\sigma + \theta_m > \sigma_\ast$, which means that the mean value of the therapy, if given as CIT, would ensure the LAS of the disease free equilibrium. Since for each $T$ the mean value is constant, this means that in the limit $t \rightarrow +\infty$ the therapy $\theta(t)$ tends to become impulsive.
points characterized by low values of the number of immune system cells are characterized by an initial rapid growth of the tumor size, followed by a regression to 0. Biologically, the therapy might seem to help the tumor growth, instead of fighting it. For the highly aggressive tumor, the cancer free equilibrium is LAS, but there is also a high size LAS equilibrium (Fig. 7).

• In the presence of periodic therapy with \( \theta_r(t) \), for both types of tumors the phase portrait is roughly similar to that of the constant therapy: the cancer-free periodic solution remains GAS for the non-aggressive tumor (Fig. 8). For the aggressive tumor there is the coexistence of the cancer free solution with a solution fluctuating around high values of the cancer size (near the equilibrium of the constant therapy). The two basins of attraction for the aggressive tumor remain unvaried with respect to those of the constant therapy (Fig. 9).

• For \( \theta_r(t) \) the dependence of the qualitative properties of the system on the parameter \( c \) is not critical.

• For aggressive tumor and \( \theta_u(t) \), it may occur that, given an initial point, the eradication is also a function of parameters \( b \) and \( \omega \), but this happens only for unrealistically high values of the therapy period (Fig. 10), e.g. \( T > 100 \) days. These results may be roughly explained considering that for \( T \gg \text{Max}(1/\Psi(0), 1/\alpha) \), one may approximately consider \( \theta_u(t) \) as constant.

• Both with CIT and with periodic therapy \( \gamma(t) \) may reach values considerably higher than the physiological value \( \sigma/\mu(0) \), which might model some serious side effects of immunotherapies due to the excess of immunocompetent cells [4,5].

For the sake of completeness, we also performed some simulations in which \( 0 < A < \sigma \alpha - \sigma \) and for which there were high oscillations (\( b = 1 \)). We obtained the result that for low frequencies, there may
be points in the \((\omega, A)\) plane for which eradication is possible (see Fig. 11).

Finally, we performed simulations for a hybrid model similar to that by Kuznetsov et al. \[14\], but in which we assumed:

\[
\alpha = 0.626, \quad f(x) = \log \left( \frac{500}{x} \right).
\]

The other parameters being as before. We choose the value \(\alpha = 0.626\) in order to minimize the difference with \(f(x)\) in \[14\]. The results of the simulations are very close to those relative to the logistic case: Figs. 12 and 13. In order to obtain via CIT the reduction to the microscopic state \(\theta > 8.4\sigma\) about is required.

The analytical and numerical results obtained in this section may be usefully compared with two similar works of the recent literature which focus on Adoptive Cellular Immunotherapy. An excellent analytical work is \[19\], who, however, cannot be fully compared with our results because it refers to tumors which have no action in stimulating immune cells. Furthermore, its formulae for the global stability of the cancer free equilibrium are not expressed as a function of the parameters of the therapy. In a very interesting paper \[16\] some
results similar to ours are obtained through numerical bifurcations on a three dimensional model in which the direct immunogenicity of tumors is expressed as an additive term $c_\sigma$. As previously stressed, in the absence of therapy and of influx of immunocompetent cells both our model and the model in [16] show the possibility of having periodic solution, which in [16] are shown to be present also in some cases in which there is therapy. We notice in brief that a term $c_\sigma$ may be formally embedded in our generic function $\sigma(x)$.

4. Concluding remarks

It is interesting to use well established conceptual frameworks of ecological models to model competition phenomena in human biology, but it is important to pay attention to the whole ecological modeling aspect, such as the basic requirement of the positivity of the solutions. Even if model [22] violates the positivity rule, it is valuable because it may be read as a model which takes into account a disease-induced depression in the influx of lymphocytes. Then, instead of proposing another specific model, we preferred to add this new feature to a family of equations, and to analyze its properties. We stressed also that models which do not allow the possibility to have LAS tumor-free solutions should be cautiously considered. The general family (8) and (9) may be, of course, further generalized following Volterra’s ecological theory, i.e. by considering that there may be a delay between the consumption of a prey and the birth of a predator (see also [15,20,21]), i.e. by allowing a delay $\tau$ with probability density $q(\tau)$.

This delayed model and stochastic models will be the subject of further investigations. Finally, we would like to illustrate some qualitative medical inferences from the investigations that we have here proposed. The main problem of immunotherapy is that, as it is clear from our analysis and simulations, in general, eradication may be possible but is dependent on the initial conditions $(x(0), y(0))$. However, the ICs are in medical practice unknown or known with very large confidence intervals (cfr. [59] for the cancer cells at the start of a radiotherapy and). This makes it impossible to plan an anticancer therapy based solely on this therapy. This is a peculiarity of immunotherapy, since there are other kinds of anticancer cures for which a globally stable eradication is possible [8]. However, in our simulations we have seen that in some particular cases the model [14] predicts that globally stable eradication is possible also in case of immunotherapy, but that it depends on the “degree of aggressiveness” of the cancer, i.e., on the framework of the model [14], on the parameter $\mu_1$. However, $\mu_1$ is difficult to be estimated (as a range) and, in particular, on single patients. If in the future it might be possible, the option to use immunotherapy as main strategy, for relatively small “non-aggressive” tumors, could be seriously considered. Furthermore, we showed that the behavior of the system does not depend on the amplitude of fluctuations of $\sigma(t)$, so that the option of continuous intravenous infusion is not, dynamically, better than the bolus based therapy. This result may be of interest, since continuous intravenous infusion may cause major practical problems to the patients. Finally, in case of disease aggressive towards the immune system, since our simulations indicated that all the positive quadrant is GAS towards a macroscopic disease in absence of therapy and low $\sigma$, whereas in the presence of therapy the eradication is possible in an adequate basin (see Fig. 7), we may infer that a conventional therapy should be followed by immunotherapy to increase the probability of total remission.

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