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Mathematical modeling of metabolism and hemodynamics

R. Costalat · J.-P. Francoise · C. Menuel · M. Lahutte · J.-N. Vallée · G. de Marco · J. Chiras · R. Guillevin

Abstract We provide a mathematical study of a model of energy metabolism and hemodynamics of glioma allowing a better understanding of metabolic modifications leading to anaplastic transformation from low grade glioma.

Keywords Magnetic resonance spectroscopy \cdot multinuclear spectroscopy \cdot regional cerebral blood flow \cdot differential equation \cdot viability \cdot slow-fast dynamics

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1 Introduction

Because anaplastic transformation of low grade glioma is inescapable and occurs in variable delay, their therapeutic management appears to be a major issue. Until now, clinical and imaging data failed to promote predictive arguments. Only serial volumetric determination may give informations on further evolution, but with a timescale of one year or more. On the other hand, metabolic changes linked to anaplastic transformation occur with smaller timescale. Thus, metabolic glioma investigation may provide informations on further histological and morphological modifications of glioma. Finally, therapeutic management may be influenced by these issues. Multinuclear magnetic resonance spectroscopy ${}^{1}H/{}^{31}P$ allows non invasive follow-up of brain tumors metabolism. Metabolites concentrations measurements as creatine and phosphocreatine, lactate, ATP, intracellular pH can be determined by this technique. In addition, regional cerebral blood flow (rCBF) can be obtained during the same time of examination. Robustness and relevance of these tools have been previously established [4], [7]. However, metabolic and hemodynamic modifications underlying glioma evolution did not receive attention before. With the aim of better understanding pathophysiological mechanisms in this field, we built a mathematical model. In a previous paper [5], we derived a system of ordinary differential equations. This model was based on previously published physiological models [1], [2]. On the basis of this model, we suggested that specific profiles of metabolic changes may constitute an early indicator of further anaplastic transformation. This prompted us to further study the mathematical properties of the model, and this is the aim of this paper. We intent to determine whether non trivial mathematical properties of the model can give useful pathophysiological indications for better management of glioma.

2 Derivation of the Model

The model is inspired by the analysis of brain lactate metabolism developed in [1], [2]. For sake of simplicity, the molar concentration of species X, namely [X], is noted X; units are mM. In this model the two state variables are the intracapillary lactate concentration LAC_c and the intracellular lactate concentration LAC_i . We assume that the total volume of neurons and normal glial cells is smaller than glioma volume within the volume of interest, so that the subscript "*i*" refers to the intracellular milieu of glioma cells; similarly, "cells" designate glioma cells. The model includes the following elements (cf Fig. 1):

(i) intracellular lactate concentration LAC_i and $pH(pH_i = -\log H_i^+)$, cell volume being V_i ;

(ii) cell lactate production J_1 ;

(iii) flux of lactate diffusion from cells to capillaries J_2 ;

(iv) flux J_3 , which is the sum of :

- lactate consumption by the metabolism, taking into account both the conversion lactate-pyruvate catalysed by lactate dehydrogenase and subsequent consumption of pyruvate by mitochondria, and a possible consumption of lactate by remaining neurons (astrocyte-neuron lactate shuttle, as proposed by [9],

- lactate diffusion towards neighbouring regions;

(v) capillary lactate concentration (LAC_c) and $pH(pH_c = -\log H_c^+)$, capillary volume being V_c ;

(vi) arterial lactate concentration LAC_a ;

(vii) cerebral blood flow (CBF);

(viii) flux J_{cap} , which is the difference between lactate input to capillaries and output from capillaries, namely $J_{cap} = CBF.LAC_a - CBF.LAC_v$, where LAC_v is the venous lactate concentration.

Furthermore, volumes and blood flow values are expressed per unit tissue volume. As a consequence, V_c and V_i are dimensionless parameters, and the capillary blood flow *CBF* is expressed in s^{-1} .

Thus the following mass balance equations can readily be obtained:

$$V_i \frac{dLAC_i}{dt} = J_1 - J_2 - J_3, \quad V_c \frac{dLAC_c}{dt} = J_{cap} + J_2.$$

Constitutive equations were derived in the following way. First, transport of lactate is always coupled to H^+ transport via monocarboxylate transporters (*MCTs*). One can simply take into account this passive co-transport by setting (Aubert et al. 2005):

$$J_{2} = T\left(\frac{LAC_{i}H_{i}^{+}}{K_{H} + LAC_{i}H_{i}^{+}} - \frac{LAC_{c}H_{c}^{+}}{K_{H} + LAC_{c}H_{c}^{+}}\right),$$

where K_H is a constant expressed in mM.M, T the maximum transport rate. This formula is a simplified version of a more general equation for carrier-mediated symport [6].

Furthermore, following a suggestion by [3] for oxygen balance, we set:

$$J_{cap} = CBF.LAC_a - CBF.LAC_v = 2CBF(LAC_a - LAC_c).$$

In previous papers, we showed that this simple formulation is nearly equivalent to more complex ones, based on partial differential equations [11], and can be applied to lactate, glucose, and oxygen in brain [1], [2]. Finally, since J_1 and J_3 respective contributions are difficult to distinguish on the basis of clinical magnetic resonance studies, we simply write $J = J_1 - J_3$.

Finally, we obtain the following system of two differential equations:

$$V_{i}\frac{dLAC_{i}}{dt} = J - T\left(\frac{LAC_{i}H_{i}^{+}}{K_{H} + LAC_{i}H_{i}^{+}} - \frac{LAC_{c}H_{c}^{+}}{K_{H} + LAC_{c}H_{c}^{+}}\right),$$

$$V_{c}\frac{dLAC_{c}}{dt} = 2CBF.(LAC_{a} - LAC_{c}) + T\left(\frac{LAC_{i}H_{i}^{+}}{K_{H} + LAC_{i}H_{i}^{+}} - \frac{LAC_{c}H_{c}^{+}}{K_{H} + LAC_{c}H_{c}^{+}}\right)$$

It must be noted that the capillary volume V_c is much smaller than the cell volume V_i ; typically, V_c/V_i is about 0.01. Setting $\varepsilon = V_c/V_i$ and $\tau = t/V_i$, we can write:



Fig 1

$$\begin{aligned} \frac{dLAC_i}{d\tau} &= J - T\left(\frac{LAC_iH_i^+}{K_H + LAC_iH_i^+} - \frac{LAC_cH_c^+}{K_H + LAC_cH_c^+}\right),\\ \varepsilon \frac{dLAC_c}{d\tau} &= 2CBF.(LAC_a - LAC_c) + T\left(\frac{LAC_iH_i^+}{K_H + LAC_iH_i^+} - \frac{LAC_cH_c^+}{K_H + LAC_cH_c^+}\right).\end{aligned}$$

In the following mathematical study (paragraphs 3-5), we write t instead of τ , bearing in mind that a moderate change in time scale has been achieved by introducing τ since V_i value is about 0.5.

3 Phase-plane analysis of the system with J and T constant

We change to notations better adapted to the mathematical analysis. We set:

$$x = LAC_i, y = LAC_c, k = K_H/H_i^+, k' = K_H/H_c^+, L = LAC_a, F = 2CBF.$$

We thus obtain the fast-slow system:

$$\frac{dx}{dt} = J - T\left(\frac{x}{k+x} - \frac{y}{k'+y}\right)$$
$$\varepsilon \frac{dy}{dt} = F(L-y) + T\left(\frac{x}{k+x} - \frac{y}{k'+y}\right)$$

In this first paragraph, we consider F, L, k, k' as fixed and J, T as parameters. We discuss the existence and nature of the stationnary point as well as eventual existence of periodic orbits. But we also decide that the system makes only sense in a fixed rectangle that we call the **viable phase space**:

$$V = \{(x, y), 0 \le x \le M, 0 \le y \le N\}$$

This is motivated by the fact that the variables x and y must be positive as they represent concentrations and cannot assume very large values. We say that if an orbit leaves the domain V it is not viable. Our interpretation is that the biological viability of the system is no longer ensured, e.g. cell necrosis occurs.

3.1 Stationary points

Solving the system:

$$0 = J - T\left(\frac{x}{k+x} - \frac{y}{k'+y}\right)$$
$$0 = F(L-y) + T\left(\frac{x}{k+x} - \frac{y}{k'+y}\right),$$

yields

$$y = L + \frac{J}{F} = y_0,$$

which is always positive. This displays:

$$x = \frac{k(\frac{J}{T} + \frac{y_0}{k' + y_0})}{1 - (\frac{J}{T} + \frac{y_0}{k' + y_0})} = x_0.$$

There is, thus, a unique stationary point (x_0, y_0) .

3.2 Nature of the stationary point

The nature of the stationary point (x_0, y_0) can be discussed on the Jacobian of the system at this point. The eigenvalues λ_{\pm} solve the equation:

$$(A+\lambda)(\frac{B+F}{\varepsilon}+\lambda)-\frac{AB}{\varepsilon}=0,$$

with

$$A = \frac{kT}{(k+x)^2}, B = \frac{k'T}{(k'+y)^2}.$$

The eigenvalues are so that:

$$egin{aligned} \lambda_+ + \lambda_- &= -(A + rac{B+F}{arepsilon}) < 0, \ \lambda_+ \lambda_- &= AF/arepsilon > 0, \end{aligned}$$

hence the stationary point is stable. Furthermore,

$$\begin{split} &\Delta = (A + \frac{B+F}{\varepsilon})^2 - 4\frac{AF}{\varepsilon} = \\ &A^2 + 2A(\frac{B+F}{\varepsilon}) + (\frac{B+F}{\varepsilon})^2 - 4\frac{AF}{\varepsilon} > \\ &A^2 - 2A(\frac{B+F}{\varepsilon}) + (\frac{B+F}{\varepsilon})^2 \geq 0, \end{split}$$

hence this unique stationary point is a node.

4 Control of the position of the stationary point

We aim in this section to write explicitly the conditions on the control (J,T) and the parameters so that:

$$0 \le y_0 \le M, 0 \le x_0 \le N.$$

It was observed previously that y_0 is always positive. But there is a condition so that $x_0 > 0$. This yields

$$1 - (\frac{J}{T} + \frac{y_0}{k' + y_0}) > 0,$$

with

$$y_0 = L + \frac{J}{F},$$

this displays the condition

$$T > J[1 + \frac{1}{k'}(L + \frac{J}{F})].$$

This shows, in particular that the condition T > J is necessary.

The condition $y_0 \leq M$ displays:

$$\frac{J}{F} \leq (M-L).$$

Note that the rectangle should be such that $M \ge L$. The last condition

$$x_0 \leq N$$
,

yields:

$$\frac{y_0}{k'+y_0} < \frac{N}{k+N} - \frac{J}{T},$$

and

$$L + \frac{J}{F} < \frac{k'(\frac{N}{k+N} - \frac{J}{T})}{\frac{k}{k+N} + \frac{J}{T}}.$$

5 The slow curve

The geometry of the slow curve is important as it allows to explain how the orbits may eventually leave the viability domain.

The equation of the slow curve is:

$$f(x,y) = F(L-y) + T(\frac{x}{k+x} - \frac{y}{k'+y}) = 0.$$

This slow curve is always attractive because:

$$f_y'(x,y) = -F - \frac{k'T}{(k'+y)^2} < 0.$$

Note as well that it is a graph over the *y*-axis: f(x, y) = 0 if and only if $x = \phi(y)$, with:

$$x = \frac{k(Fy - FL + \frac{Ty}{k'+y})}{T - (Fy - FL + \frac{Ty}{k'+y})}$$

The function ϕ is increasing. Generic orbits are almost parallel to the *y*-axis then reach a neighborhood of the slow curve and follow inside this neighborhood untill they tend to the stationary point.

Numerical computations using MATLAB software clearly confirm this point: orbits in the $(x,y) = (LAC_i, LAC_c)$ phase plane are displayed in Figure 2, where $\varepsilon = V_c/V_i = 0.006875$. Moreover, even values of ε as high as 1 result in a somewhat similar phase portrait (not shown).



Fig. 2 Please write your figure caption here

6 Discussion and Conclusion

Whatever the parameters values, the model has a unique stationary point. Furthermore, the stationary point is asymptotically stable. This finding is consistent with a clinically observed fact that, within a short time scale from minutes to days, metabolite concentrations within the tumor appear nearly constant. Moreover, we derived explicit and sufficient conditions which ensure that a stationary point is in a viability domain in the first quadrant. These conditions can give useful pathophysiological insights on tumor viability. For instance, "x(t) is positive" implies that T, the maximum transport of lactate via MCT, must be sufficiently high. This strongly suggests modifications of density and /or kinetic properties of MCT during glioma evolution. In fact, both MCT in glioma cells and blood-brain barrier should have enhanced density or individual maximal rate. This also suggests that MCT could be potential target for glioma therapeutics [8]. Furthermore, the two variables of the system display distinct time evolutions. Thus, the system could be studied using asymptotic and geometric analysis of slow-fast systems. Quite interestingly, the model has an associated viability domain, and generic orbits are almost parallel to the $Y(LAC_c)$ axis, then remain in neighborhood of the slow curve while tending to the stationary point. As a consequence, generic orbits do not leave the viability domain. Thus, in the framework of the slow-fast dynamics approximation, the problem of the viability of trajectories (solution curves) can fully be solved. This will allow using a larger study frame where some parameters of the model can be replaced by control variables. There is indeed the perspective to build a hierarchy of models for cerebral metabolism and hemodynamics, in analogy with the SAPHIR model built for renal physiology [10] by replacing successively each control variables by a compartmental adding of supplementary models, on the basis of the Aubert-Costalat equations. As a conclusion, the model provides pathophysiological mechanisms of glioma metabolism. Further mathematical and clinical studies can provide a better understanding of the natural history of glioma and then may allow improvement of therapeutic management.

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