A quantitative numerical model for TNF-α mediated cellular Apoptosis

Abstract:
The paper is 3 years old because of dr Liberati's serious sickness, but still worth to be considered in our mind:

Experimental evidence indicates that cells under irradiation induce in the neighbour non-irradiated cells the same biological effects affecting the irradiated ones. This is the so-called bystander effect. Up to now in the scientific literature this kind of effect does not appear to be fully understood, even if several experiments show evidence of its existence. It would be reasonable that bystander effect takes place by means of paracrine chemical transmission mediators that would be broadcasted by the damaged cells to the surrounding cells. Furthermore, a subset of a special class of signaling proteins, namely the cytokines, are probably the very ones involved in such signaling phenomenon. Among them, Tumor Necrosis Factor (TNF) is a particularly relevant protein belonging to the class of cytokines, because it is known to contribute to mediate various relevant cell functions, like apoptosis, the programmed cell death. As a molecule, TNF is quite interesting, because it can issue two opposite signals through different intracellular molecular signaling chains. One signal induces apoptosis, while the other is opposite, inducing the cell resistance to apoptic signals. The crucial point is thus to understand what makes each of such two signals masking the other. Thus a mathematical model related to the TNF signaling pathway is of interest, paying special attention to the study of the TNF reception mechanisms by cells that are not passed through by the radiation beam. In this work we present a new mathematical model of cellular apoptosis - mediated by TNF - and its validation based on data existent in literature. The model that we present will result to be a stable model with respect to large variation of the parameters and simplified with respect to other models already existent.

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A quantitative numerical model for TNF-α mediated cellular *Apoptosis*

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**Abstract**

Experimental evidence indicates that cells under irradiation induce in the neighbour non-irradiated cells the same biological effects affecting the irradiated ones. This is the so called *bystander* effect. Up to now in the scientific literature this kind of effect does not appear to be fully understood, even if several experiments show evidence of its existence. It would be reasonable that bystander effect takes place by means of paracrine chemical transmission mediators that would be broadcasted by the damaged cells to the surrounding cells. Furthermore a subset of a special class of signaling proteins, namely the cytokines, are probably the very ones involved in such signaling phenomenon. Among them, *Tumor Necrosis Factor* (TNF) is a particularly relevant protein belonging to the class of cytokines, because it is known to contribute to mediate various relevant cell functions, like apoptosis, the programmed cell death. As a molecule, TNF is quite interesting, because it can issue two opposite signals through different intracellular molecular signaling chains. One signal induces apoptosis, while the other is opposite, inducing the cell resistance to apoptic signals. The crucial point is thus to understand what makes each of such two signals masking the other. Thus a mathematical model related to the TNF signaling pathway is of interest, paying special attention to the study of the TNF reception mechanisms by cells that are not passed through by the radiation beam. In this work we present a new mathematical model of cellular apoptosis - mediated by TNF - and its validation based on data existent in literature. The model that we present will result to be a stable model with respect to large variation of the parameters and simplified with respect to other models already existent.

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I. INTRODUCTION

In Radiobiology (field of science that studies the biological effect of ionizing and non-ionizing radiation on cells) the bystander effect is the phenomenon inducing unirradiated cells to exhibit irradiated effects as a result of signals received from nearby irradiated cells. Radiation-induced bystander effect is defined as induction of biological effects in cell that are not directly traversed by a charged particle but are receiving signals from the irradiated cells that are in close proximity to them [1]. Bystander effect is thus defined as non-irradiated cells response to signals produced by neighbouring irradiated cells [2]. The bystander effect have many consequences such as genomic instability pertaining the nucleus (that will not be discussed in this paper), and Apoptosis pertaining the cytoplasm, that is the object of this paper.

In the scientific literature this kind of effect does not appear to have been fully understood, even if several experiments show evidence of its existence. In particular, one of the most representative experiment sounds as follows: cell population becomes radiated, then its culture medium is conveyed to another environment where non radiated cells live; afterwards the non radiated cells show to suffer the same types of effect as the radiated cells. Such effect is detected even if the power of the ionizing radiation is small. The action mechanism of the bystander effect is intended to be the diffusion of one or more factors from the irradiated cells to the surrounding non-irradiated ones. Such factors bound (at the membrane level) with the conjugate receptors of the target cells. Such a signaling triggers a series of different phenomena among whom genomic instability, and especially Apoptosis are of our main interest. One of the key point is thus to understand what kind of mediator, mediators is, are responsible for the bystander effect. One of the most accredited hypothesis is that the bystander effect takes place by means of a special class of signaling proteins, namely the cytokines: among them, Tumor Necrosis Factor (TNF) seems to be the most relevant protein because it is known to contribute to mediate various cell functions, like cell survival and proliferation, or apoptosis (the programmed cell death) [3][4][5][6].

Thus a mathematical model related to the TNF signaling pathway is helpful to increase our understanding of the underlying biological processes, allowing us to organize existing information from experimental studies and to identify the gaps in our understanding of TNF pathway.

In Sec. II, a short review of TNF biology leading to our mathematical model, will be given. In Sec. III, a mathematical model of TNF citotoxicity will be presented, including: the building of a model; its review and simplification; and the final model. In Sec. IV the simulation results (using initial conditions and parameters taken from literature and reported in the same section) will be presented and then deeply discussed. In this section two aspects will be presented:

- the evolution of the death and survival complexes (compared with respect to other models);
- simulation investigating model robustness in order to better understand both strengths and weaknesses of the proposed model.

The conclusions, summarizing the main results of the simulations, as well as the comparison with respect to other models, will be discussed in sec. V.

II. AN ESSENTIAL REVIEW OF TNF BIOLOGY

For the sake of clarity, here we briefly review the basic facts known about TNF, in order to understand the ground of the model that will be presented in this paper. What follows can be found in more detailed reviews such as [7]. Tumor necrosis factor (TNF) was isolated on the basis of its capacity to kill tumor cells in vitro and to produce hemorrhagic necrosis of transplantable tumors in mice [8] [9].

TNF is a molecule that binds to two different receptors: TNFR1 and TNFR2 [7]. Among them TNFR1 appears to be the key mediator in signaling in both normal and tumor cells; for this reason the large amount of the models are focused on this receptor. TNFR1 has three subunits, whose cytoplasmic tails need to be juxtaposed to trigger intracellular signaling. The signaling pathway has a role of a double-edge sword [10]: it is implicated in tumorigenesis, transplant rejection, septic shock, and in others activities. More commonly, the binding of TNF-α to its receptors can activate two major transcription factors, AP-1 and NF-kB, inducing inflammatory responses. Furthermore, TNF-α can induce an apoptotic response, usually dependent on inhibition of RNA or protein synthesis. Binding of TNF to TNFR1 initiates a series of biochemical events in the cell that take place at the cytoplasmic tails of the receptor subunits and, in particular, at their specialized domains called Death Domains (DD). DD recruit the adaptor protein TRADD that acts as an assembly platform for at least two other proteins, RIP-1 and TRAF-2 initiating the survival arm (on the left in Fig. 1).

Such multiproteic complex initiates the signaling cascade resulting in NF-kB activation and hence gene activation
and cell survival [4][5][7]. Among the genes that are expressed after the NF-kB activation, there are those that code for the two proteins FLIP and IAP that inhibit the TNF apoptotic pathway.

Regarding the TNF apoptotic pathway, it has been demonstrated that this is initiated by TNF-TNFR1 complexes internalized into endocytic vesicles [11]. At this deeper intracellular level, the multiproteic complexes associated to the receptors’ tails modify, and form the Death Inducing Signaling Complex (DISC), whereby TRADD recruits FADD (Fas-Associated protein with Death Domain) and pro-caspase-8. Such caspase then triggers the irreversible pathway leading to apoptosis and cell death.

Both the NF-kB and the apoptotic pathways comprise a series of complex intracellular reactions involving a number of enzymes and substrates [4][5][7]. Current biological data indicate that activation of NF-kB and caspases occurs at different sites in the cells (at the cell membrane and upon internalization in endosome, respectively). The transcription factor NF-kB has been shown to be predominantly localized at cytoplasmic level in its inactive form, i.e. in the absence of an inductive signal. Cell stimulation with TNF-α can induce the degradation of IkB, the inhibitor of NF-kB, allowing nuclear accumulation of NF-kB and regulation of specific gene expression. Although Caspase-8 has the capacity to localize to a number of different cellular locations, activated Caspase-8 have predominantly a cytosolic localization.

Fig. 1 summarizes the above reported basics of TNF biology grounding our model. All quantities between square brackets are molar quantities (Tab. I) and k’s are kinetic constants with the meanings shown in Tab. II. To have a better understanding of Fig. 1 we want to underline two general concepts:

- all k’s constants (on\off) are rate of association\dissociation constants giving the information on the speed of association or dissociation reaction;
| Vr | $v_r$ zero-order rate of insertion of receptors R into the membrane |
| k_{on}LR | association rate constant of L and R giving TTR |
| k_{on}TTRC | association rate constant of TTR and C giving TTRC |
| k_{on}TTRF | association rate constant of TTR and F giving TTRF |
| k_{on}CoTTRFC | association rate constant of TTRC and F giving COUPLING |
| k_{on}NFkB | association rate constant of TTRF and F activating NFkB |
| k_{on}ATTRC | association rate constant of TTRC and C giving Apoptosis |
| k_{deg} | rate of degradation of NFkB |

- all $k_{#}$'s constants are the degradation rate of the element #.

In the upper part of Fig. 1 the interaction is schematized between TNF (L) and TNFR1 (R) giving the composite TTR. Starting from this point one can clearly see the branching of two ways:

- the "survival way" represented by composite TTRF (interaction of FLIP with TTR, makes the equilibrium unbalanced toward survival against death);
- the "death way" represented by composite TTRC (interaction of Caspase-8 with TTR, makes the equilibrium unbalanced toward death against survival).

$k_{on}TTRF$ and $k_{on}TTRC$ are thus two key parameters, because they regulate not only the access to the two ways - an higher value of $k_{on}TTRF$ with respect to $k_{on}TTRC$ leads to an increment in cell survival (and viceversa) - but also regulate the sensibility of the cell survival fraction with respect to a variation of every parameter affecting the lower part of each of the two ways. To better understand this point one can think that, for example, for very low value of $k_{off}NFkB$ (0 in the extreme case) with respect to $k_{on}NFkB$, the major part of the composite TTR tends to bound to Caspase-8: thus the survival way is pratically "close", and as a result, we expect that a variation of some parameters in such a way (e.g. $k_{off}NFkB$, $k_{on}NFkB$, ecc..) will not significantly affect the cell survival fraction $f(t)$.

Another important parameter is $k_{deg}TTR$ that models the lysosomal degradation of TTR. An high value for this parameter will result in a increased cell resistance to apoptic signal. In the lower part of Fig. 1 one can see the end of the two ways: on the left the survival way ends with the activation of NFkB (by interaction of FLIP with TTR), while on the right side the death way ends with the activation (by interaction of Caspase-8 with TTRC) of a composite here schematized as Apoptosis. In such death leading pathway $k_{on}ATTRC$, as well as $k_{deg}$, plays a crucial role in cell fate: in particular a small value of $k_{on}ATTRC$ indicates a low affinity of the composite TTRC in bonding to Caspase-8, thus resulting in an increasing cell resistance. The same effect is expected for an high value of $k_{deg}$, that is the degradation term of apoptotic complex, and plays an opposite role in cell survival/death with respect to $k_{on}ATTRC$.

We want to underline that, as it can be seen in Fig. 1, the survival way is bidirectional while death way is not. This will not imply that at the end all the ways will lead to death, because of the terms $k_{deg}$ and $k_{deg}NFkB$.

To conclude it is also important to notice that the possible coupling between the two ways is taken into account by means of the composite named COUPLING. It allows to take into account a new detail in the TNF pathway, not present in our preliminary suggestion [12], nor in [13], but it also significantly increases the number of parameters...
present in the model. Thus, at this stage, we decided to postpone the simulation of the complete model depicted in Fig. 1 to a future paper, then limiting here ourselves to a simplified model like the one published in [13], with respect to whom we will be able to obtain comparable results with the need of much less parameters. Because of the extreme complexity of TNF pathway (yet not completely understood) and of a lot of cross talk between TNF-mediated apoptosis, NF-kB, JNK and other pathways on apoptosis such as AKT pathway, we reported a simple and minimal but efficient model.

III. A MATHEMATICAL MODEL OF TNF CYTOTOXICITY

Starting from Fig. 1 we can thus build a mathematical model that is simpler than [13] without losing the main biological details like in [12] (then presented without substantial variations in [14]). This is a two step process:

• to build a general model of TNF citotoxicity;

• to simplify the model, making some biological assumptions, in order to make the model less computationally burdening, in order to be able to simulate it without the help of supercomputing.

A. Building of the model

In this section we present a general model of TNF mediated cellular apoptosis. Such a model is built translating into equations the scheme in Fig. 1:

$$\frac{d[R]}{dt} = V_r - k_d[R] + k_{offTTR}[TTR] - k_{onLR}[L][R]$$

$$\frac{d[L]}{dt} = L_0 + k_{offTTR}[TTR] - k_{onLR}[L][R]$$

$$\frac{d[TTR]}{dt} = -k_{offTTR}[TTR] + k_{onLR}[L][R] - k_{onTTRC}[TTR][C] - k_{onTTRF}[TTR][F] + k_{offTTRC}[TTRC] + k_{offTTRF}[TTRF] - k_{degTTR}[TTR]$$

$$\frac{d[F]}{dt} = V_{inF} - k_{onTTRF}[TTR][F] - k_{onCoTTRCF}[TTRC][F] - k_{onNFkB}[TTRF][F] + k_{offNFkB}[NFkB] + k_{offFCa}[COUPLING] + k_{offTTRF}[TTRF]$$

$$\frac{d[COUPLING]}{dt} = k_{onCoTTRFC}[TTRF][C] + k_{onCoTTRCF}[TTRC][F] - k_{offFCa}[COUPLING] - k_{offCCa}[COUPLING]$$

$$\frac{d[C]}{dt} = V_{inC} + k_{offCCa}[COUPLING] + k_{offTTRC}[TTRC] - k_{onCoTTRFC}[TTRF][C] - k_{onATTRC}[TTRC][C] - k_{onTTT}[C] - k_{onTTRC}[TTR][C]$$

$$\frac{d[NFkB]}{dt} = -k_{offNFkB}[NFkB] + k_{onNFkB}[TTRF][F] - k_{degNFkB}[NFkB]$$
\[
\frac{d[TTRC]}{dt} = k_{offFCo}[COUPLING] + k_{onTTRC}[TTR][C] - k_{offTTRC}[TTRC] - k_{onATTRC}[TTRC][C] - k_{onCoTTRCF}[TTRC][F]
\]

\[
\frac{d[TTRF]}{dt} = k_{offCCo}[COUPLING] + k_{onTTRF}[TTR][F] + k_{offNFkB}[NFkB] - k_{offTTRF}[TTRF] - k_{onCoTTRFC}[TTRF][C] - k_{onNFkB}[TTRF][F]
\]

\[
\frac{d[Apoptosis]}{dt} = k_{onATTRC}[TTRC][C] - k_{deg}[Apoptosis]
\]

\[
\frac{df(t)}{dt} = -k[Apoptosis]f(t)
\]

Eq.(1) describes the dynamics of TNFR1. The two parameters \(V_r\) and \(k_d\) - introduced in [15] and already used in [12] - describe "the zero-order rate of insertion of receptors into the membrane and the turnover rate constant of ligand-free receptors" respectively. In case of absence of TNF the concentration of TNFR1 at the cell surface (in steady state) is given by:

\[
[R]_{(L)=0} = \frac{V_r}{k_d}
\]

The term \(-k_{onLR}[L][R]\) describes the association between TNF and its free receptor. The minus sign is because the association between these two components decreases the concentration of free TNF and TNFR1. The same term is also present in eq.(2). The \(+k_{offTTR}[TTR]\) describes the dissociation rate of TTR and is present obviously with "+" sign also in eq.(2) and with "-" sign in eq.(3) for the reasons just explained above.

All the other equations are built following the same philosophy.

Eq.(2) describes the dynamics of TNF. The term \(I_0\) is the constant rate of arrival of free TNF.

Eq.(3) describes the binding kinetics of TNF\(\backslash TNFR1\). The term \(-k_{degTTR}[TTR]\) takes into account the possible degradation of this compound: thus an high value of \(k_{degTTR}\) results in an increasing resistance of the cell to Apoptosis.

Eq.(4) describes the dynamics of FLIP. The term \(V_{inF}\) is a constant rate of arrival of free FLIP that once bound to TTRF activates NFkB and starts the survival pathway of the cell.

Eq.(5) describes the dynamics of COUPLING that is a compound that models the possible coupling between the two ways (survival and death).

Eq.(6) describes the dynamics of Caspase-8, that once bound to TTRC starts the irreversible death pathway of the cell.

Eq.(7) describes the dynamics of the survival complex NFkB. The term \(-k_{degNFkB}[NFkB]\) takes into account the possible degradation of this complex.

Eq.(8) and (9) describe the dynamics of the intermediate complexes TTRC and TTRF.

Eq.(10) describes the temporal evolution of the death complex. The term \(-k_{deg}\) takes into account the possible lysosomal degradation of this complex: thus an high value of \(k_{deg}\) results in an increasing resistance of the cell to Apoptosis.

The last equation, (11) already introduced in [12] describes the temporal evolution of the surviving fraction \(f(t)\).

B. Review and simplification of the model

From a mathematical point of view the model consists of 11 non-linear first order differential equations with 23 parameters, most of whom never experimentally measured. Because of the huge number of free parameters, before
proceeding with the simulations we try to simplify the model on a biological basis and existent literature (see for example [13]).

As a first assumption we consider an "isolated sistem" from the TNF, FLIP and Caspase-8 point of view: thus we can put $L_0 = V_{inF} = V_{inC} = 0$. In this case the crucial role in cell survival\death is played by initial conditions and parameters, since no TNF, FLIP or Caspase-8 are entered into the system after $t=0$.

As a second assumption we consider the death way and survival way completely independent (as considered in [13]). Thus we can completely remove eq. (5), and all the kinetic parameters connected to it ($k_{onCoTTRCF}$, $k_{offFCo}$, $k_{onCoTRFC}$, $k_{offCCo}$ and $k_{offFCo}$) can be set to 0.

The resulting model can be schematized as in Fig. 2 and resumed in the following equations:

$$\frac{d[R]}{dt} = V_r - k_d[R] + k_{offTTR}[TTR] - k_{onLR}[L][R]$$

$$\frac{d[L]}{dt} = L_0 + k_{offTTR}[TTR] - k_{onLR}[L][R]$$

$$\frac{d[TTR]}{dt} = -k_{offTTR}[TTR] + k_{onLR}[L][R] - k_{onTTRC}[TTR][C] - k_{onTTRF}[TTR][F]$$

$$+ k_{offTTRC}[TTRC] + k_{offTTRF}[TTRF] - k_{degTTR}[TTR]$$

$$\frac{d[F]}{dt} = -k_{onTTRF}[TTR][F] - k_{onNFkB}[TTRF][F] + k_{offNFkB}[NFkB] + k_{offTTRF}[TTRF]$$

$$\frac{d[C]}{dt} = k_{offTTRC}[TTRC] - k_{onATRC}[TTRC][C] - k_{onTTRC}[TTR][C]$$

$$\frac{d[NFkB]}{dt} = -k_{offNFkB}[NFkB] + k_{onNFkB}[TTRF][F] - k_{degNFkB}[NFkB]$$

$$\frac{d[TTRC]}{dt} = k_{onTTRC}[TTR][C] - k_{offTTRC}[TTRC] - k_{onATRC}[TTRC][C]$$

$$\frac{d[TTRF]}{dt} = k_{onTTRF}[TTR][F] + k_{offNFkB}[NFkB] - k_{offTTRF}[TTRF] - k_{onNFkB}[TTRF][F]$$

$$\frac{d[Apoptosis]}{dt} = k_{onATRC}[TTRC][C] - k_{deg}[Apoptosis]$$

$$\frac{df(t)}{dt} = -k[Apoptosis]f(t)$$

The resulting model consists in 10 differential equations with 15 kinetics parameters and therefore it is possible to simulate it without supercomputing, differently from the complete model presented in the previous section.
In this section, the models and some of the simulations presented in [12] and [13] will be used as a term of comparison for the simulations results of our model. In order to simulate the model, we need the initial conditions and kinetic parameters. Some of the parameters are taken from [12] and [13], as for example \( \frac{V}{r}, k_d, k_{onLR}, k_{offTTR}, k_{degTTR} \) and \( k \). Since the model is structurally different, and at a different level of abstraction (intermediate with respect to the lower detail in [12] and the higher detail in [13]), we cannot take all the parameters from these models and compare all the results. One way to avoid this problem, and get a reasonable guess for all the parameters, is to look to the equations of our model with respect to the equations of the two cited models, and try to understand the underlying biological meaning, in order to find a sort of “contact points” between the equations. Thanks to the stronger bond to the biological reality of our model with respect to [12], we can find many of these contact points with [13] in terms of common equations/parameters. An example of this kind of parameter is \( k_{deg} \) that can be taken as \( k_{deg} = k_{19} + k_{20} \) (in [13]), since the equation describing \( c_{21} \) (death complex) dynamics in [13] can be loosely identified with our eq. (21) and the term \(-(k_{19} + k_{20})c_{21}\) can be seen as lysosomal degradation of this complex. The same can be done for \( k_{onATTRC}, k_{offNFkB}, k_{onNFkB}, k_{offTTRF} \). The remaining parameters are identified by fitting the temporal evolution of death and survival complexes in [13]. This was done for TNF(0)=1nM as shown in figures 3(a) and 3(b). Initial conditions and kinetic parameters reported in Tab. III and Tab. IV are then used to perform all the simulations.

FIG. 3: Death/survival complex temporal evolution for 1nM of TNF(0)
TABLE III: Initial conditions of the species involved.

<table>
<thead>
<tr>
<th>Species</th>
<th>Initial Value (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R</td>
<td>100</td>
</tr>
<tr>
<td>L</td>
<td>a (this parameter change during simulations)</td>
</tr>
<tr>
<td>TTR</td>
<td>0</td>
</tr>
<tr>
<td>F</td>
<td>100</td>
</tr>
<tr>
<td>C</td>
<td>80</td>
</tr>
<tr>
<td>TTRF</td>
<td>0</td>
</tr>
<tr>
<td>NFkB</td>
<td>0</td>
</tr>
<tr>
<td>TTRC</td>
<td>0</td>
</tr>
<tr>
<td>Apoptosis</td>
<td>0</td>
</tr>
</tbody>
</table>

Survival fraction | Initial Percentage
f(t) | 1

TABLE IV: kinetic parameters values. First order constants are in s\(^{-1}\) while second order constants are in nM\(^{-1}\)s\(^{-1}\)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(V_r)</td>
<td>(1 \cdot 10^{-3})</td>
</tr>
<tr>
<td>(k_d)</td>
<td>(1.67 \cdot 10^{-3})</td>
</tr>
<tr>
<td>(k_{offTTR})</td>
<td>(1.25 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{degTTR})</td>
<td>(2.67 \cdot 10^{-3})</td>
</tr>
<tr>
<td>(k_{onLR})</td>
<td>(185 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{onTTRC})</td>
<td>(750 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{onTTRF})</td>
<td>(580 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{ofjTTRC})</td>
<td>(0.3 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{ofjTTRF})</td>
<td>(1.25 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{onNFkB})</td>
<td>(185 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{offNFkB})</td>
<td>(371.25 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{deg})</td>
<td>(371.25 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k_{deg})</td>
<td>(300 \cdot 10^{-6})</td>
</tr>
<tr>
<td>(k)</td>
<td>(6.7 \cdot 10^{-4})</td>
</tr>
</tbody>
</table>

A. Death\(\backslash\)Survival complex dynamics

Simulations show that even if dramatically simplified, the proposed model inherited the significant properties of the more detailed model presented in [13]. Figures 4(a) and 4(b) show the comparison between the temporal evolution of the death and survival complexes simulated with either our model or model [13], computed for 5 and 10nM of TNF(0). The two models keep agreeing at different TNF(0) values.

Fig. 5 shows the temporal evolution of the surviving fraction computed for all the three values of TNF(0). The surviving fraction decreases as the TNF(0) increases.

Fig. 6 shows the TNF cytotoxicity as a function of both TNF concentration and time. There is a saturation effect in the survival fraction in both concentration and time. Results are in good agreement with [12]. The apparent minimum in Fig. 6 around TNF(0)=100nM, for all the observation times, has probably no biological meaning, being more likely a mathematical artifact illustrating one limit of the proposed model at the boundary of its validity: in the following robustness analysis the behavior of \(k_{onTTRF}\) will confirm such boundary limits. There is thus probably no biological explanation for this kind of behaviour, but there is a mathematical one: since to perform these simulations we are using parameters already present in literature, some of them are at the boundary of validity for our model (see e.g. \(k_{onTTRF}\) in sec. IV B 3): thus little unexpected behavior or oscillations can arise. In this case the undershoot is negligible and the results shown in Fig. 6 are still to be considered acceptable.

B. Robustness simulations

As already mentioned, experimental determinations of the parameter values for intermediate binding reactions are not available. Moreover the small number of available experimental data did not allow us to estimate the error bounds for the parameter values. Because of these problems our aim is to test the stability of our model (i.e. if the model
(a) Death complex evolution computed with our model (green line 5nM and red line 10nM of TNF(0)) and the model proposed by [13] (black lines, continuous for 5nM, dotted for 10nM of TNF(0)).

(b) Survival complex evolution computed with our model (green line 5nM and red line 10nM of TNF(0)) and the model proposed by [13] (black lines, continuous for 5nM, dotted for 10nM of TNF(0)).

FIG. 4: Death/survival complex temporal evolution for 5nM and 10nM of TNF(0)

FIG. 5: Surviving fraction temporal evolution computed with our model for 1 (blue), 5 (green), 10 nM (red) of TNF(0).

FIG. 6: Computing TNF cytotoxicity as a function of both TNF concentration and time.
gives reasonable results or fail) over a wide range of parameter values, instead of concentrating ourselves on the accurate determination of the parameters themselves, thus providing a useful tool for the simulation of cellular Apoptosis in many possible different experimental conditions.

1. $\Delta$parameter vs $\Delta$TNF(0)

In this section we report the results of such simulations plotting the survival fraction after 48 hours (transient already ended) and or 1 hour (during transient). Both the initial condition of TNF(0) and some key parameters of the model vary over several decades, with discretization of 40 logarithmic spaced samples over the whole range, around the "nominal value" given in the previous sections. In Tab. V the parameters values are reported, used in all simulations, of whom only the more significant ones are reported in the following.

TABLE V: kinetic parameters values. First order constants are in s$^{-1}$ while second order constants are in nM$^{-1}$s$^{-1}$

<table>
<thead>
<tr>
<th>Initial Condition</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNF(0)</td>
<td>logspace(-2,3.5)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_{onattre}$</td>
<td>logspace(-7,-4)</td>
</tr>
<tr>
<td>$k_{log}$</td>
<td>logspace(-4.5,1)</td>
</tr>
<tr>
<td>$k_{deg}$</td>
<td>logspace(-7,-3)</td>
</tr>
<tr>
<td>$k_{onNFkB}$</td>
<td>logspace(-6,-3)</td>
</tr>
<tr>
<td>$k_{offNFkB}$</td>
<td>logspace(-6,-3)</td>
</tr>
<tr>
<td>$k_{onttrc}$</td>
<td>logspace(-6,-3)</td>
</tr>
<tr>
<td>$k_{offttrc}$</td>
<td>logspace(-8,-4)</td>
</tr>
<tr>
<td>$k_{onttrf}$</td>
<td>logspace(-6,-2)</td>
</tr>
<tr>
<td>$k_{offttrf}$</td>
<td>logspace(-8,-4)</td>
</tr>
<tr>
<td>$k_{degtrc}$</td>
<td>logspace(-5,0)</td>
</tr>
</tbody>
</table>

FIG. 7: Surviving fraction computed varying TNF(0) and $k_{onattre}$

Increasing TNF(0) always leads to an increment of the cell death. Fig. 7 shows that an increasing value of $k_{onattre}$ leads to an increasing value of cell death, as expected from the model, as well as to a difference in the survival fraction.
The clear difference between Fig. 7(a) and 7(b) is due to the different simulation time (1h and 48h). In fact, for 1h of simulation time we can consider that the system is still evolving (simulation time shorter than time constants present in the system, e.g. k’s) thus only higher values of konattrc lead to a significative fraction of cell death. On the contrary, for 48h of simulation time, the system can be considered "in steady state" (simulation time much longer than time constants present in the system): thus only lower values of konattrc lead to an increasing value of cell survival.

Fig. 8 can be understood if one looks at the value of kontrc, and realizes that this is the parameter that "regulates the access" to the apoptotic way. A high value of kontrc with respect to kontrf means that all the parameters variations in the survival way do not significantly affect the solution.

2. |Δ|parameter#1 vs |Δ|parameter#2 |

In this section the results are shown about the survival fraction, when varying two parameters across several decades (resolution is 40 points) around the "nominal value" given in the previous sections, for the initial condition of TNF(0) = 10nM. In Tab. VI the parameters values are reported that are used to generate all the figures. Fig. 9 refers to a variation of kontrc and kontrf: thus this figure shows clearly the competition between the two ways. As expected an high value of kontrc with respect to kontrf leads to an increasing value of cell death, and vice versa.

<table>
<thead>
<tr>
<th>Initial Condition Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>TNF(0)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Kinetic Parameter</th>
<th>Ranges</th>
</tr>
</thead>
<tbody>
<tr>
<td>kontrf</td>
<td>logspace(-6,-1)</td>
</tr>
<tr>
<td>kofftrf</td>
<td>logspace(-7,-1)</td>
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<tr>
<td>kontrc</td>
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</tr>
<tr>
<td>kofftrc</td>
<td>logspace(-8,-4)</td>
</tr>
<tr>
<td>konNFkB</td>
<td>logspace(-6,0)</td>
</tr>
<tr>
<td>koffNFkB</td>
<td>logspace(-8,-3)</td>
</tr>
<tr>
<td>kdeg</td>
<td>logspace(-8,-3)</td>
</tr>
</tbody>
</table>

(a)Contour of surviving fraction @t=1h. (b)Contour of surviving fraction @t=48h.

FIG. 9: Surviving fraction computed varying kontrf and kontrc

3. Limits of the model: an example

The model shows computational limits for certain values of the parameters. For example Fig. 10 shows an unexpected behaviour for lower values of kontrf; in fact, for these values of kontrf we see a new rise in cell survival even for increasing level of TNF(0). Such a behavior is probably related to the already reported curious non monotonic decay in Fig. 6. This is most likely not correct, and can thus be considered one of the limits of validity of the model for this parameter.
A new general mathematical model of TNF cytotoxicity has been presented and discussed. The model has been simplified and simulated under the main biological assumptions that the survival and death ways are completely independent, and that the system has to be considered completely isolated with respect to Caspase-8, TNF and FLIP. The final model results to be less computational demanding with respect to [13] but not oversimplified as the one presented in [12] and discussed in [14]. Despite to its simplicity, the results of our model are in good agreement with [13]. We want to stress that with ”good agreement” we intend that the functional form of the common outputs of our model (death/survival complex) and of the model [13] are nearly the same, even if the two models are structurally different: thus the meaning of some of the involved parameters are different.

We have extensively explored the parameters space (robustness simulation) of the model: the simulations show that the model has a stable behaviour for a broad range of parameter values and that no unexpected patterns (such as oscillations) emerge, except for the one shown in Fig. 10 (and possibly the one occurring in Fig. 6).

Our model (as the one presented in [12]) is not specific for a given cell type: then it could be used to simulate effects of TNF in most cell types, thus quite independently of the experimental settings.

It is important to underline that in our model there is no switching mechanism that selects cell survival or death signals, but rather a balance between the two pathways that produces partial cell killing, even for long lasting and intense TNF treatments. The equilibrium between the pathways might further be balanced in favour of cell survival or death in real cells by increasing/decreasing the degradation terms and association/dissociation rate constants (as well as by re-integrating the here not yet fully discussed COUPLING pathway).

To conclude we would like to propose new experimental campaigns in order to measure the temporal evolution of some output of the model (such as survival fraction, survival complex, death complex,...), for instance varying the temperature. The aim of such campaigns would be to understand if there is a substantial modification in TNF cytotoxicity in terms of functional form of the outputs, or only a simple change in the ”temporal constants” (k’s factors). In the second case, our model will acquire an even more general validity.

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Marco Pappalettera, MD, has been the Chief Immunologist within the Biochemical Clinical Chemistry of the University Hospital San Gerardo of the University of Milano Bicocca.

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