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Model

Singel interval SD

# Control and Decision Making in Systems Biology

Northeastern University

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Prof. Mark Niedre Prof. Carey Rappaport

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## Presentation outline

- 1. Background
- 2. Chemotherapy
- 3. Immunotherapy
- 4. Epidemics
- 5. Acknowledgement

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### Outline

Background

Chemotherapy

Model Metronomic

Optimal cont

MDOR

Epidemic
Singel interval SD



### Background

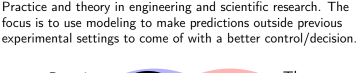
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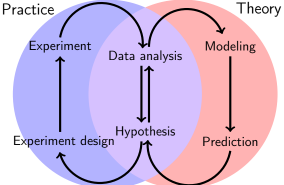
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► Motivation: Metronomic/intermittent experiments¹ in mice. A lower dose with a higher frequency than MTD were shown to requit immune system and reduce the tumor volume.

The current standard of care limits the regiments used primaritly

to daily dose and maximum-tolerated dose (MTD) treatments.

Objective: Use optimal control techniques in order to have a better treatment outcome among all possible dosing strategies.

<sup>&</sup>lt;sup>1</sup> Junjie Wu and David J Waxman. "Metronomic cyclophosphamide schedule-dependence of innate immune cell recruitment and tumor regression in an implanted glioma model". In: *Cancer letters* 353;2 (20±4), pp. 272±289 c

## Chemotherapy

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Now, a new generation of quantitative experiments made it

Early efforts in using optimal control techniques for cancer treatment started in the 1970s for Radiotherapy<sup>2</sup> and

Chemotherapy<sup>3</sup>.

possible to have more realistic models of the system.

The goal is to use optimal control techniques to find a mathematically derived optimal regimen (MDOR) to be tested in a similar experimental settings.

<sup>&</sup>lt;sup>2</sup>K Bahrami and M Kim. "Optimal control of multiplicative control systems arising from cancer therapy". In: IEEE Transactions on Automatic Control 20.4 (1975), pp. 537-542.

<sup>&</sup>lt;sup>3</sup>Thomas L Swan George W Vincent. "Optimal control analysis in the chemotherapy of IgG multiple myeloma". In: Bulletin of mathematical biology 39.3 (1977), pp. 317-337. 4日 → 4周 → 4 目 → 4 目 → 9 Q P

### Model

Singel interval SD

$$\dot{T}(t) = k_a T(t) - \frac{k_b C(t) T(t)}{k_c C(t) + T(t)} - k_d T(t) I(t), \tag{1a}$$

$$\dot{I}(t) = qX(t) - k_e T(t)I(t) - k_f C(t)I(t) - k_g Y(t)I(t) - k_h I,$$
 (1b)

$$\dot{X}(t) = \frac{qC(t)T(t)}{k_iC(t) + T(t)} - k_jX(t) - k_kX(t)Y(t),$$
 (1c)

$$\dot{Y}(t) = \frac{I(t)}{k_I + I(t)} - k_m Y(t) C(t), \tag{1d}$$

$$\dot{C}t = u(t) - \frac{k_1 C(t)}{k_2 + C(t)}.$$
 (1e)

Where Tumor T represents the tumor volume, and the phenamenological variables immune system I, immunostimulatory X, immunossuppressor Y, and drug C represent the dynamics in the tumor microenvinronment.

Model is fitted to the tumor and immune data in mouse experiments.

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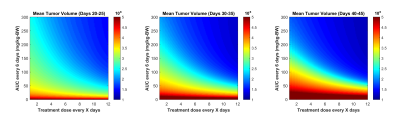
#### Metronomic Optimal contr

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The average tumor volume at different time ranges of 20-25 days (left), 30-35 days (middle), and 40-45 days (right) after starting a metronomic regimens. The horizontal axis represent the number of dose between each dose, y axis is the total amount of drug given to the animal every 6 days.

for the optimal control problem.

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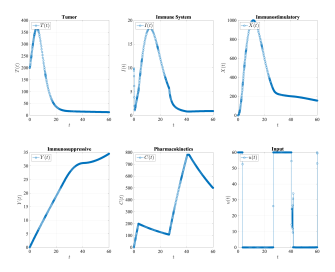
 $\min_{u(t)} T(t_f), \tag{2a}$ 

Numerical software GPOPS II is used to solve the following setup

s.t.  $\begin{bmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{bmatrix} \le \begin{bmatrix} C(t) \\ T(t) \\ I(t) \\ X(t) \\ Y(t) \\ U(t) \end{bmatrix} \le \begin{bmatrix} C_m \\ T_m \\ I_m \\ X_m \\ Y_m \\ U_m \end{bmatrix}, \tag{2b}$ 

$$\int_{0}^{t_f} u(t)dt \le U_m. \tag{2c}$$

# Numerical result for a low input upper bound



Circles show the final collocation points.



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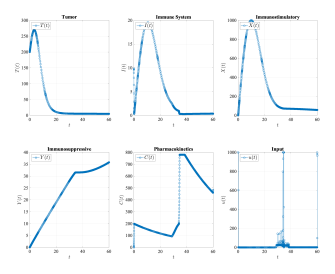
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# Numerical result for a high input upper bound



Circles show the final collocation points.



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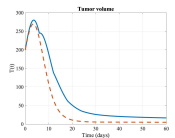
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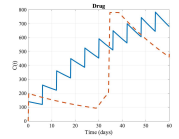
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# Optimal control vs. metronomic regimen





Comparing a standard 140 mg/kg Q6D metronomic chemotherapy plan (solid lines) with the obtained optimal control (dashed lines).

However, The maximum tolerated dose is 300 mg/kg/day for CPA.

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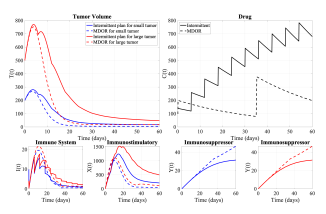
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# Mathematically derived optimal regiment



Comparing a standard 140 mg/kg Q6D metronomic/intermittent plan (solid lines) and the mathematically derived optimal regimen (dashed lines).

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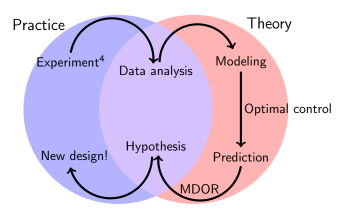
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# A new viable regimen to be tested experimentally



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<sup>&</sup>lt;sup>4</sup> Junjie Wu and David J Waxman. "Metronomic cyclophosphamide schedule-dependence of innate immune cell recruitment and tumor regression in an implanted glioma model". In: *Cancer letters*, 353(2) (2014), pp. 272=280; q.

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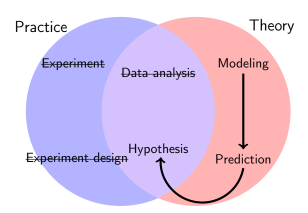
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Acknowledgment

During the COVID-19 epidemic, social distancing as a form of non-pharmaceutical intervention has been enacted in the US and other countries.

- ► Motivation: Shortening the period of time that populations are socially distanced is economically advantageous.
- Objective: To reduce the disease burden (here measured as the peak of the infected population) while simultaneously minimizing the length of time that the population is socially distanced.

# Early days and limited data!



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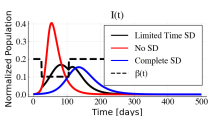
MDOR

## Epidemic

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$$\beta(t) = \begin{cases} \beta_n & 0 \le t_s \\ \beta_d & t_s \le t < t_s + t_d \\ \beta_n & t_s + t_d \le t \end{cases}$$
 (3)

Normalized infected population in SIR model, with no re-infection.



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# This work is a result of teamwork

Advisor: Eduardo Sontag

Lab members: M. Ali Alradhawi, Anh Phong Tran, Zheming An,

William Cho, Shu Wang, Tianchi Chen.

Presented projects

Chemotherapy: Anh Phong Tran, Irina Kareva, M. Ali Alradhawi,

and Waxman Lab.

Epidemics: James Greene, M. Ali Alradhawi.

Other projects

Immunotherapy: Irina Kareva, Kumpal Madrasi, Abed Alnaif, Anup Zutshi, and EMD Serono Inc team.

Parkinson's Disease: AMP-PD research community, and Sanofi team.

Ribosome: M. Ali Alradhawi, Michael Margaliot, Nikolai Slavov,

Edward Emmott.

Open-source community: Julia team, Gleb Pogudin, Esteban Vargas. Bioconductor project.

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