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Article

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Keywords: keyword 1; keyword 2; keyword 3 (list three to ten pertinent keywords specific to the article, yet reasonably common within the subject discipline.).

1. Introduction

1.1. POMDP

1.2. Simulation details

The simulated environment features discretized of planning cancer treatments. At each timestep a "cancer state" has a certain risk of increasing. If the tumor ever reaches the state 5, the simulation ends. To avoid this, a model has decide when to apply treatment. Applying treatment can reduce the tumor level - whether the treatment successfully reduces the "cancer state" depends on a underlying *resistance state*. When the resistance is low,

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Table 1. MDPI article types.

abstract	entry	retraction
addendum	expressionofconcern	review
article	extendedabstract	perspective
book	datadescriptor	protocol
bookreview	editorial	shortnote
briefreport	essay	studyprotocol
casereport	erratum	systematicreview
comment	hypothesis	supfile
commentary	interestingimage	technicalnote
communication	obituary	viewpoint
conferenceproceedings	opinion	guidelines
correction	projectreport	registeredreport
conferencereport	reply	tutorial

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the chance of treatment succeeding is high, but probability declines as the resistance state increase. The resistance state has fixed probability of increasing when treatment is applied, and it can decrease when treatment is withdrawn. Whether the resistance state decreases depends on the tumor level. At high tumor levels, the resistance state is more likely to decrease, but this carries the risk of tumor growing out of control and "killing the patient". In order to successfully manage the disease, a model will therefore also have to manage the resistance state. However, the resistance state is directly orbserable in the given simulation. It must therefore be inferred from how the tumor state responds to treatment.

Both the resistance state, and tumor states start at state 0 out of 5, meaning that a total is six states are possible in each. Two slightly different simulations are used:

- Performance simulation, where a range-bounded approach to managing the tumor is used, where the treatment is begun each time the tumor state increases above level 3, and is withdrawn when the tumor state drops below 2. 100 runs at 4 different probabilities of increasing the tumor state is run for maximally 200 timesteps. For each run, vectors of outcomes are pregenerated, meaning that vector of whether the tumor increases of length 200 is predetermined where each entry has the probability specified for the entire simulation. Likewise vectors are generated for treatment outcomes at each resistance level, and outcomes for resistance drops. This is done to ensure comparibility between the range-bounded approach and POMDP models that consider different future outcomes at different lengths. In these runs the tumor state is perfectly observable, meaning a tumor state always generates tumor observation that corresponds to hidden factor. The POMPD can only observe the tumor level, and its prior preferences are uniform over all tumor states except for the highest. These POMPDS only consider a combination of policies at each timestep. They can only consider treating or not treating for three timesteps in a row. The shortest horizon model only considers one of these blocks, while the medium considers two blocks of treating or not treating, for total horizon of 6 steps into the future. The longest horizon model considers three blocks for a total of 9 steps into the future. This is done to ease the computational burden, since it greatly reduces the total number of policies to be evaluated.
- In the capacities simulation, a single run of a POMDP with a slightly different model structure is used. This model can observes a perfect signal of whether the patients is alive, whether the model is testing, treating and a noised signal of the tumor state if the model has decided to test the patient on the given run. Its prior preferences are heavily skewed against observing a dead patient, somewhat against treating and little against treating. This means that it will have to balance the "cost" of all these actions". It is also handicapped further by the fact that the tumor signal is noised, and the resistance state, which must be inferred through the tumor signal is therefore doubly obfuscated.

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2. Results2.1. Performance of POMDPs vs ADT heuristic

POMDP performance against ADT 0.3 0.075 0.050 0.025 Density 00000 0.5 0.7 0.075 0.050 0.025 0.000 150 200 0 50 50 100 100 150 200

Four different simulation runs each consisting of 100 runs, of maximally 200 timesteps for different rates of tumor growth (respectively 1. .3, .5, and .7 probability of increasing the tumor state) . On each run the the outcomes on each of the potential 200 random variables are generated and each strategy is therefore tested on a the same environment. On the lowest tumor growth rate almost all the runs reach maximal length.

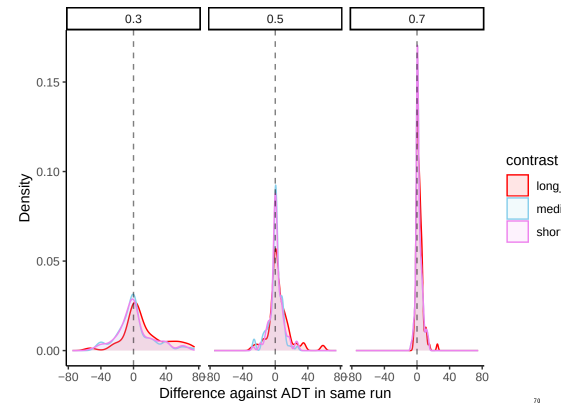
Simulation Length

Contrasts against ADT-performance for growth rate .3, .5 and .7 are plotted. .1 is omitted since it appears that maximal perforamnce was achieved for each policy horizon of the POMPD strategies.

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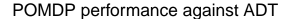
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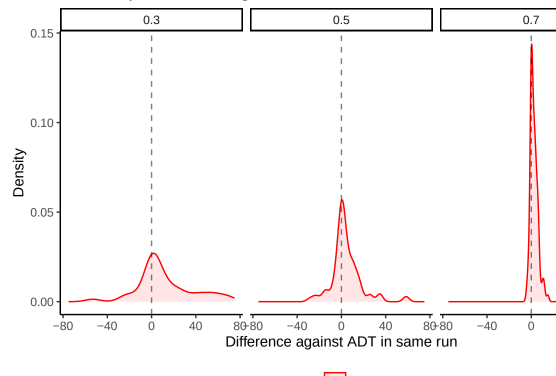




Two interpretations emerge from the contrasted simulation lengths. As the the growth rate increases, the POMDP seems to run for longer than the specific instance of the range bounded approach. It also seems that longer policy horizon seems beneficial. When the tumor state had .7 probability of increasing each timestep, the longest POMPD with the longest policy horizon performed as follows.

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At more aggressive cancer rates, the simulation consistently to outperform the range bounded approach. The absoulute increase in timesteps decreases however.

contrast

long_vs_adt

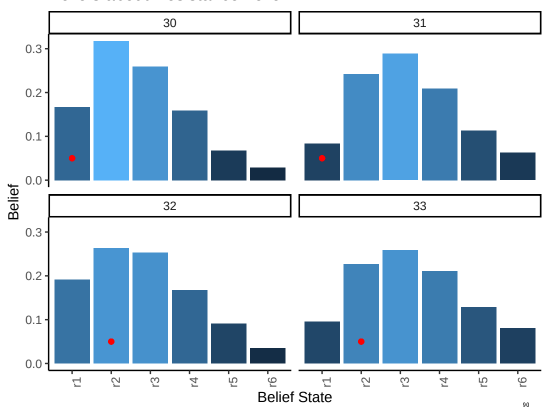
2.2. Capabilities

2.2.1. Learning underlying hidden state

The POMDP structure is capable of inferring an underlying hidden state: the resistance state. Even though this not directly observable, this can be inferred to how the tumor state responds to treatment. Applying treatment for longer time without any beneficial effect would suggest that the resistance level is high, while immediately observing that the tumor level decreases would suggest that the resistance level is low. At each timestep the POMDPs perform infer the most likely state of every state factor, given their current observation and prior beliefs. Through custom changes to the 'get_expected_states' function in PYMDP that allowed the models to consider how one hidden state factor (resistance factor) would influence another (the tumor factor).

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Beliefs about Resistance Level

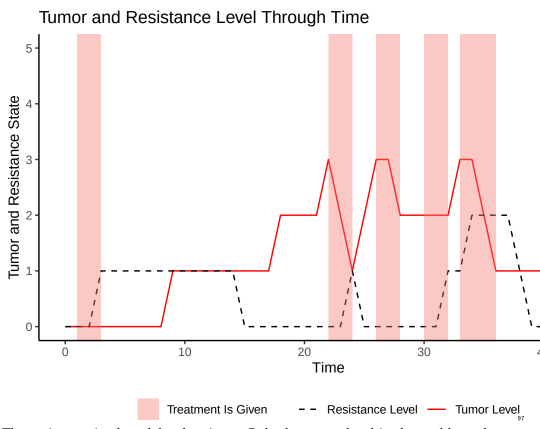


The above plot shows the strength of beliefs in t probabilities for a subset of timesteps [30 - 34], and the red dot show the actual the resistance levels. A time progresses the resistance level increases, and the model adjusts its beliefs. While the tumor level doesn't increase durings this time_period, this would also signal the model that the resistance level is low. This is the case since the environment always has change of increasing the tumor state, but such an increase would be negated by successful round of treatment.

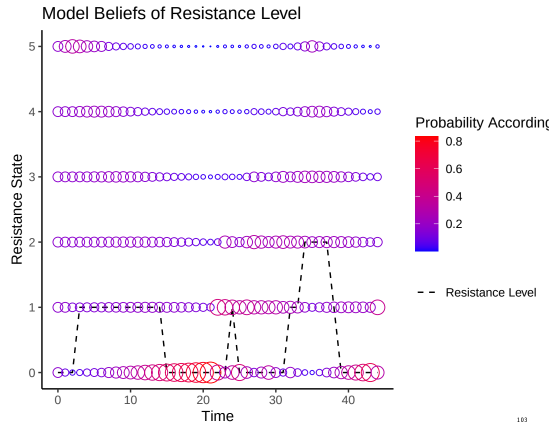
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The entire run is plotted for the given. Only the tumor level is observable to the POMDP. It must combine its knowledge about how resistance level likely increases after applying treatment, and how the tumor level responds to treatment depenping on the the resistance level. While the model far from perfectly knows the resistance level. Model beliefs for the entire run is plotted.



The model begins fairly agnostic. Given that the model was initialized with a uniform prior over resistance states this makes sense. Thorought out the course of the simulation it then finds lower values of resistance state more likely.

2.2.2. Beliefs, current/future and uncertainty is accessible

This current simulated model used a slightly different implementation than those compared to performance of rangebounded therapy planning. While the those models had longer policy horizons, they didn't consider the entire space possible actions. A model with a shorter policy horizon that would instead search every single possible action four steps is plotted to investigate the structure of decision-making by the POMDP model. The environment was also more unceartain. Instead of featuring a 1-to-1 mapping of the observations of the tumor to actual tumor state, it recieved a noised signal. It must therefore. The beliefs about how tumor state will evolve as a consequence of the most promising policy at time step 14 is plotted, and the expectation of resistance states at t=24 is plotted too.

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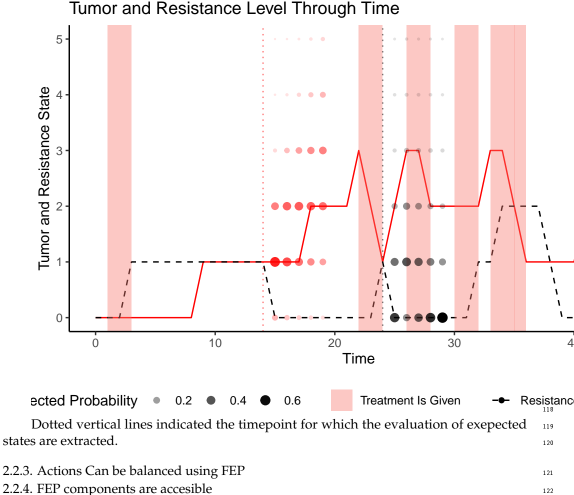
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states are extracted.

3. Discussion 3.1. Does it work for other sorts of medical planning 3.2. Would longer policy search be better 3.3. Only one range bound 3.4. Future Research 3.4.1. Continuous state space or binning illness 3.4.2. Learning the transition parameters

3.4.3. searching policy space better 3.4.4. other fep models

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/1010000/s1, Figure S1: title; Table S1: title; Video S1: title.

Author Contributions: For research articles with several authors, a short paragraph specifying their individual contributions must be provided. The following statements should be used "X.X. and Y.Y. conceive and designed the experiments; X.X. performed the experiments; X.X. and Y.Y. analyzed the data; W.W. contributed reagents/materials/analysis tools; Y.Y. wrote the paper." Authorship must be limited to those who have contributed substantially to the work reported.

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Written informed consent for publication must be obtained from participating patients who can be identified (including by the patients themselves). Please state "Written informed consent has been obtained from the patient(s) to publish this paper'' if applicable.

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Sample Availability: Samples of the compounds are available from the authors.

Abbreviations

The following abbreviations are used in this manuscript:

MDPI Multidisciplinary Digital Publishing Institute

DOAJ Directory of open access journals

TLA Three letter acronym LD linear dichroism

Appendix D

Appendix D.1

The appendix is an optional section that can contain details and data supplemental to the main text. For example, explanations of experimental details that would disrupt the flow of the main text, but nonetheless remain crucial to understanding and reproducing the research shown; figures of replicates for experiments of which representative data is shown in the main text can be added here if brief, or as Supplementary data. Mathematical proofs of results not central to the paper can be added as an appendix.

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All appendix sections must be cited in the main text. In the appendixes, Figures, Tables, etc. should be labeled starting with 'A', e.g., Figure A1, Figure A2, etc.

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