SocultPaperV6

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Introduction

Adapative Therapy

Worldwide, cancer is the cause of 1 out of 6 deaths. An estimated 90% of these cancer deaths are due to the development of drug resistance (Bukowski, Kciuk, and Kontek 2020). While initial cancer treatment usually shows a positive response in tumor burden, drug resistance develops over time. To highlight the inefficiency of traditional approaches, (Staňková et al. 2019) models cancer treatment as a game-theoretic contest between a physician and a tumor. In this model, the physician's move in each round is to apply a certain treatment, and to which the tumor adapts. While it is a "Stackelberg game"—a game where one player is the leader (the physician) and another player is a follower (the tumor)—this asymmetry is rarely exploited in oncology wards. Instead of using their advantage to steer the evolutionary pressures placed on tumors, physicians let the tumor adapt not only to the current round of the game but also to future rounds. Thus, the advantage of leading the game is lost. The authors analogize the current practice to a game of rock-paper-scisccors:

"in which almost all cells within the cancer play, for example, "paper." It is clearly advantageous for the treating physician to play "scissors." Yet, if the physician only plays "scissors," the cancer cells can evolve to the unbeatable resistance strategy of "rock." (Staňková et al. 2019)

To exploit this assymetry Adaptive Therapy (AT) controls the intra-tumoral evolutionary dynamics by leveraging a likely fitness costs that cancer cells incur when evolving mechanisms to resist drugs. For example, it has been shown that tumor cells can mutate to increase the expression of the PGP membrane pump, which uses ATP to expel drugs from the cell. While this makes the cell more resistant to treatment, it also comes at a metabolic cost. Gatenby et al. (2009) found that PGP activity was responsible for approximately 50% of cell metabolism. If resistant and non-resistant cells are competing for space and resources, drug-sensitive cells should, over time, outcompete resistant cells since they can devote more metabolism to cell proliferation. Thus, AT utilizes Darwinian competition to make cancer suppress itself, a strategy that potentially could make cancer a controllable chronic disease.

Initial results from an on-going pilot clinical trial applying AT in a group of metastatic castrate-resistant prostate cancer (mCRPC) patients are promising, showing both lower cumulative dosages and longer survival times compared to a similar group of patients receiving standard care. The trial utilizes a range-bounded heuristic to make treatment decisions: if the blood marker Prostate-specific Antigen (PSA), a proxy for tumor burden, increases back to pre-trial levels, Abiraterone treatment is applied until PSA drops to 50% of pre-trial levels (Zhang et al. 2017). The trial is expected to run until December 2024 (Center and Research Institute 2024).

Needed Developments in Mathematical Modeling

Key AT researchers West et al. (2023) have recently surveyed the use of mathematical modeling in the discipline and identified needed developments to realize the potential of AT. For example, they argue that modelling approaches will need the capacity to simulate multidrug therapies. Additionally, they argue that it is unlikely that any single treatment approach will be able to delay the emergence of resistant cells, while

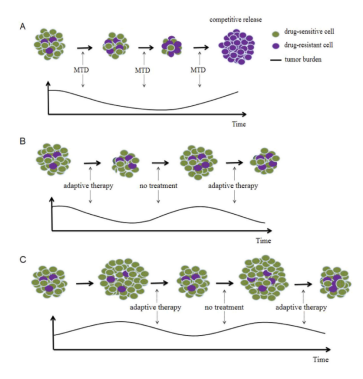


Figure 1: From Zhang et al. 2023. Panel A shows a typical "Maximal Tolerable Dose" treatment protocol. While the initial response in tumor burden is promising, a compettive release of resistant cells ensures. Panel B shows AT approach with a small tumor burden. Panel C shows AT with a high tumor burden, which has been theorized to further increase the suppression of resistance cells through darwinian competition.

lowering tumor burden and minimizing toxicity. Given that patients likely differ in their ability to tolerate tumor burden, drug toxicity, and the evolutionary dynamics of their cancers, therapies will have make informed trade-off between these objectives for each individual patient. The authors also report several hirtherho poorly understood biological mechanisms, such as how competition dynamics change depending on carrying capacities of tumors, or how high tumor burdens might be detrimental due to increased risks of new metastasis. More complicated modelling of tumor dynamics is therefore required, and future modelling approaches will require flexibility in their ability specify complex causal systems.

Predicting real-time treatment responses of individual patients would likely greatly enhance AT efficacy, since dose modulation could be individualized further to control intratumoral evolutionary dynamics with greater precision. A key issue going forward is rethinking how real-time patient data is collected. West et al. (2023) stress the importance of collecting data not only to detect progression but also to inform future treatment decisions. For example, a risk of using a set testing frequency, as done in the pilot clinical trial, is treatment overshooting. In trial, drops in PSA weren't always detected early enough to minimize induced resistance. This could have limited future treatment opportunities. Additionally, whatever the modelling approach of choice, the authors underline that it allows for careful analysis of the uncertainty in everything from estimated tumor dynamics to treatment outcomes.

Partially Observable Markov Decision Process

Partially Observable Markov Decision Processes (POMDPs) are a class of controller models that model an Markov process. An environment moving through discrete times and states, for which next step of the system only depends on the current step. Partial observability refers to the fact that these models don't directly observe the actual environment but instead only noisy signals emitted its changing states (Åström 1969). This allows POMPDs to differentiate an observed signal from what it "believes" about the actual configuration of states and additionally to use a single reward function to trade-off uncertainty for reaching a certain goal state (Kaelbling, Littman, and Cassandra 1998).

Solving POMPDs with Active Inference

While POMDPs are typically difficult to solve analytically, various approximate approaches exist. One approximation scheme is born out of the neuroscience paradigm *Active Inference* (AI). Typically used to model an agents decision making. The approximation scheme finds a solution by minimizing two objective functions:

- Variational Free Energy (VFE). A measure of fit between a generative model and sensory input.
- Expected Free Energy (EFE). A score of how well a course of action is expected to bring about a set of preferences.

AI has been used to model psychopathology (Da Costa et al. 2020) but also applied to control scenarios such as the mountain-car problem (Friston, Daunizeau, and Kiebel 2009) and, albeit augmented with deep-learning, robotics control (Çatal et al. 2020). AI implementations of the POMDP scheme have been implemented in MATLAB and recently in Python with the python package *pymdp* Heins et al. (2022a).

Typically, AI POMDPs are technically a joint probability:

$$p(o, s, u; \phi)$$

where o are observations, s are hidden states, u are "control states" (actions that an agent can take to influence the enviorment), and ϕ are the hyperparameters of the model, such as α typically used as 'inverse temperature' i.e. how deterministically the model selects actions. By conditioning on certain observations, the POMDP is solvable by approximation for bayes-optimal posterior beliefs at a given time point. This yields a

set of posterior beliefs not only about the configuration of each modeled type of state in the environment, but also posterior probability distribution for the optimal course of action to achieve a set of preferences given the expected uncertainty. As an example, a POMDP could model the joint probability of observing a noisy signal of particular tumor burden in a patient, an actual underlying tumor burden, and the course of action that is most likely to bring the tumor burden down. By modelling modelling, actions, states and observations as discrete events, the joint probability can be made tractable by factorized for tractability into the a number categorical probability distributions describing likelihood mappings between signals and actual states of the environment, transition probabilities of the environment and a desired probability of making certain observations (see appendix A for in depth description of how the joint probability is factorized). Minimizing VFE and EFE to solve a given POMPD at a given time point itself requires approximating the Free Energies. This is done by a neuronally inspired method known as "message-passing", for which multiple implementations exists (Smith, Friston, and Whyte 2022). ¹

The aim of this paper

The treatment decisions used by in the pilot trial reported by Zhang et al. (2017) were based on a well-informed heuristic about how resistance dynamics would develop as consequence of careful timing of applying and withdrawing treatment. However, the actual resistance dynamics of each patient wasn't explicitedly modeled real-time, and thus not used in treatment decisions.

This paper will investigate whether the AI implementation of POMDPs could viably model the degree of treatment resistance of individual cancer patients to improve long-term control of the disease. Beyond fitting certain desiderata reported by (West et al. 2023), the use of the AI paradigm for the present project was further motivated by the following considerations:

- Real-time data is likely sparse in clinical settings. Extracting the maximum amount of information from each data point will therefore be essential. Using models that perform optimal Bayesian inference would therefore be preferable.
- Short-term exploitation of tumor vulnerability is essential for keeping the patient alive, but for long-term success, controlling resistance dynamics will be crucial. A therapy plan will have to balance keeping the patient alive now against limiting its future options for treatment. Using a measure such as Expected Free Energy could be instrumental in striking this balance.
- Resistance dynamics are not directly measurable, but should be inferable through observing how the tumor burden responds to treatment. Choosing whether to treat would therefore not only be a consideration of the tumor level but also about how much it informs us about the resistance dynamics.
- The information gain from testing can be quantified and used to prioritize testing for periods of greater uncertainty.

Analysis

Simulated environment

To investigate the appropriateness of the AI implementation of the POMDP scheme, multiple simulations of a clinical setting inspired by the pilot clinical trial were run. To comply with the canonical AI POMPD scheme discrete state and time, discrete states and time were used. In each simulation run, a simulated a maximum of 200 time-steps was created. At each time-step, an agent has to keep a virtual cancer patient alive by deciding when to apply a treatment based on a signal of the tumor burden. The signal could be

¹The Python package pymdp (Heins et al. 2022b) has numerous tutorial notebooks for implementing POMPDs in python. Smith, Friston, and Whyte (2022) describes the process of the constructing POMPDs for different scenearios in detail, albeit in MATLAB and Da Costa et al. (2020) provides in-depth mathematical account of POMPDs in Active inference.

observed at each time point, and treatment decisions were made by solving a slightly modified AI POMPD. The features of the eviroment was inspired by the use of PSA testing and Abiraterone treatment in the pilot trial (Zhang et al. 2017). In the simulations, the patient's tumor state determined whether they survived to the next time-step. If the tumor state every reached maximal state of six possible tumor states, the virtual patient "died" and simulation ended. The tumor was always set to 0 at the beginning of each simulation run, and could increase at a fixed risk at each time-step.

Whether a round of treatment successfully reduced the tumor state depended on an underlying resistance state. This also began at state 0 out of 5. For each round of treatment application, the resistance state had a fixed risk of increasing too. This rendered treatment increasingly ineffective as time progressed. The only remedy, beyond favorable bouts of stochasticity, was to bring down the resistance state by withdrawing treatment. Inspired by the work of (Hansen and Read 2020), higher tumor levels would increase the chance of bringing down the resistance level at each step.

In summary, this means that the simulated agents had balance the tumor state not growing out of control, while not painting itself into a corner by applying treatment too frequently. Striking this balance was further complicated by the stochasticity of the evolving tumor-burdens, changes to resistance levels and realized treatment-responses.

Modifications to POMDP scheme

Typically for AI POMDPs, different types of hidden states such as resistance and tumor states are not modeled to affect each other directly. Instead, the interactions of combinations of different states are modeled as leading to different observations. This means that interactions between types of states would be coded in the likelihood model A.

This setup was deemed incompatible with the clinical setting attempted emulated, since it wouldn't let let a model infer how changes to resistance state should influence the expected transition probabilities of the tumor state. Therefore, custom changes to pymdp were made to allow POMPDs to infer the state of "occluded" nodes – causally upstream nodes that don't themselves produce signals (e.g. the resistance state). Instead the state of the occluded node would have to be inferred, through its effects on downstream nodes that generate signals (e.g. the tumor state).

Specifically, modifications were made to the structure of B, the generative model's transition beliefs. Usually, three-dimensional B-tensors describe expected transition probabilities within type of state state. This means a B-tensor for type of state typically holds one dimension for the current state, one dimension for the next state, and a third dimension for every allowed action. Each cell of the tensor then contains the transition probability from one state to another conditional on a certain action. Another dimension was added to relevant b-tensors which corresponded to another type of state. Concretely, this meant that the tumor b-tensor had a fourth dimension corresponding to the possible resistance states. The expected transition probabilities could then be estimated by matrix multipliying the tumor state B- with a vector the posterior of expected resistance at a given time point. One can imagine that instead of having only one B-tensor for the transition probabilities of the tumor state, multiple B-tensors of tumor transition probabilities were specified, were each tensor corresponded to a specific resistance level. This modification to the generative model necessitates taking an average of all these tensors weighted by the posterior probabilities for each resistance state at given time point. This amounts to calculating the expected tumor state conditional on a set of beliefs about the resistance state. This modification strategy was repeated for the resistance level, since decreases in the resistance state depended on the tumor state. It should be noted that this modification means that the order in which beliefs about states are evaluated must be specified. For example, for the present simulations, the effect of resistance level on treatment efficacy was evaluated first since this was evaluated first for each step of the simulation, while the transition probabilities between resistance states were evaluated after an action had been taken.

While these changes rendered much of the functionality of pymdp immediately unusable, the modification strategy should be able to model arbitrarily complex dependencies between states. This could potentially

allow POMPDs to model more complex biological mechanism as desired by West et al. (2023). Hopefully, a robust implementation can be designed without overhauling pymdp.

Exploratory Simulation

An exploratory simulation was conducted, where for each run an agent instituted as a modified POMPD controlled whether to treat and test. Its modalities were:

- A noiseless signal of whether the patient was alive.
- A noiseless signal of wheter it was applying tests.
- A noiseless signal whether it was applying treatment
- A noised signal of the tumor state, but only if tests were being applied.

The agent's prior preferences were skewed heavily against observing a dead patient, somewhat against treating, and slightly against testing. This was done to simulate a cost to both treating and testing. This means that the agent had to balance the cost of its actions while considering the potential information gain of each choice. The costs to treating and testing were introduced to investigate whether the POMPD scheme potentially could be used to strike a balance between drug toxicity, tumor burden while optimizing real-time data collection as desired by West et al. (2020). The model was further handicapped by noise in the tumor signal. This means that the resistance state, which had to be inferred through the tumor signal, was doubly obfuscated.

However, the generative model used did perfectly map the expected noise in the tumor signal, and modeled transition probabilities identical to the actual transition probabilities of the environment. Additionally, the agent was given uniform priors over initial states, meaning it had no knowledge of the configuration of states at the beginning of each run. The same set of predetermined policies that considered the next 6 timesteps was evaulated at each step. This limited set of policies was done to ease computation by limiting the search space of possible actions. The set of policies consisted of two blocks of either testing or treating for three timesteps in a row for permuted by each possible sequence of applying testing or not for the time horizon of the policies.

Learning the Resistance State

On a the following example of a explotaroy simulation run, the agent managed to keep the patient alive for 68 timesteps and chose to test on 55 of these (see Fig. 2).

The agent appears to be somewhat sensibly applying treatment, but it doesn't apply treament at pivotal period approximately between steps 30 and 39. Generally, it refrains from applying treatment when the tumor state is low, and applies treatment when the tumor state is high. Interestingly, it does applies treatment at the first timestep, despite the tumor state and resistance state being at their lowest possible values. Considering that the generative model had uniform priors over both resistance state and tumor states, this behavior is more appropriate from the perspective of the agent, since treating and testing would immediately provide information about both states. Qualitatively, the model also seems to prefer testing when treatment is being applied. This suggests that it finds testing to be more worthwhile while also treating. A plot of the model's beliefs about the resistance state at each time-point (see Fig. 3) shows that its beliefs about the resistance state generally follow the development of the actual resistance state despite the lack of direct signal.

In summary, the modification to POMPD structure appears to have allowed the agent to otherwise occluded resistance state by selectively applying treatment and collecting noised signals of the tumor burden. While the results are intially promising, the particular agent does make questionable choices regarding the timing of treatment. The decision-making behind these choices is dissected in-depth the appendix.

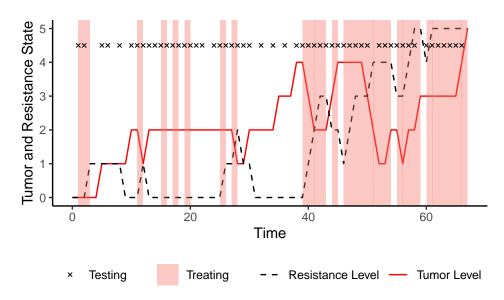


Figure 2: A simulation run for exploratory purposes.

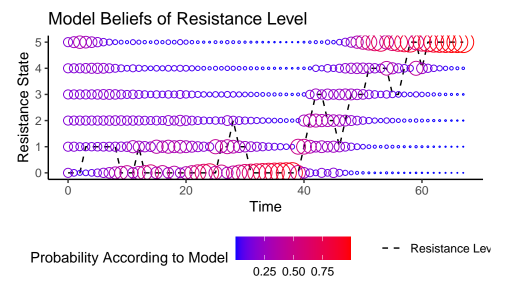


Figure 3: Plot of the models belief about the reistance level at every timepoint

Performance of POMPDs against a Range-Bounded Strategy

Several agents instiantied as modified POMPDs were tested against a range-bounded strategy. This was done to benchmark the performance of POMPDs against a strategy inspired by the treatment protocol used in the pilot clinical trial (Zhang et al. 2017). The generative models used were simplified for ease of computation POMPDs and as result they observed a noiseless signals of the tumor state and whether treatment was being applied at every timestep. They were also only given prior preferences for observations of tumor states. The preferences were uniform over all survivable tumor states, and the probability of the highest tumor state was set to 0. The following three agents were tested:

- A short policy horizon POMPD that could consider a block of treating or not treating for the next three steps.
- A medium policy horizon POMPD that could consider two blocks of treating or not treating for three steps, yielding a total horizon of 6 timesteps.
- A long policy horizon model that could consider four blocks of treating or not treating for three steps, resulting in a total horizon of 12 steps.

The agents were benchmarked on four different "tumor aggressivity settings". The tested tumor growth rates were 0.1, 0.3, 0.5, and 0.7 probability of increasing the tumor state at each timestep. For each setting, 100 simulation runs was done. For each run, the environment was predetermined by generating 200 outcome variables for each combination of tumor state, resistance state, and treatment state to ensure comparability between the tested agents and the RB-strategy at the level of each run.

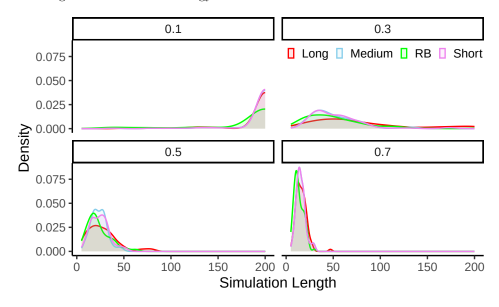


Figure 4: Simulated patient survival times for each agent and baseline strategy.

At the lowest tumor growth rate, nearly all the runs reached the maximal length (see Fig. 7). On the other runs, the long horizon model also appears to be slighly outperforming the others. Contrasts as the percentage difference in survival from swithching from the baseline RB-strategy to the long horizon POMPD are computed for each run (see Fig 5.).coc

In the 100 runs with a 0.1 risk of tumor growth, the long horizon POMPD seems to perform similarly to the RB strategy. However, as the tumor growth risk increases, the POMPD agent increasingly shows large percentage improvements compared to the RB strategy. Potentially a consequence of a greater the need for proactive action when tumor dynamics accelerate, which in turn would favor models that explicitly model how treatment will effect future resistance dynamics.

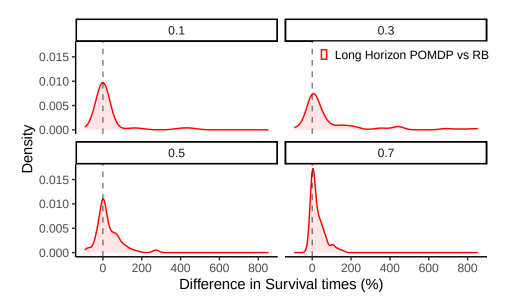


Figure 5: Change in percent of between the range-bound approach and the longest horizon POMPD for different tumor growth rates.

Discussion

Findings

This project tested a simple modification to how POMDPs are typically implemented in AI to investigate whether POMDPs could learn and control the transition of state that could whose configuration inferred based on its down stream consequences on other states. In a simulated environment designed to mimic the dynamics of AT, the modification successful allowed agents instantiated as POMDPs to model an underlying resistance state that controlled the efficacy of cancer treatment, despite the underlying state didn't any produce observable signals itself. Additionally, costs to testing and treating implemented and crucially these could be holistically integrated using free energy minimization to trade-off utility for reductions in uncertainty. Modified POMPDs were further tested against a range-bounded treatment strategy inspired by a pilot clinical trial using AT, and were found to outperform the range-bounded strategy substatantially. The gap in performance was larger in simulations of aggressive of tumor growth rates, and for agents with an ability to consider longer future courses of actions. These findings underscore the potential of using the AI paradigm in real-time planning of treatment and testing decisions in AT, and, albeit under simplified circumstances, the modified AI POMDP is promising model choice for improving real-time patient specific treatment decisions in AT.

Additionally, this paper demonstrates that the AI implementation of POMPDs shows several desirable qualities for AT:

• The modified AI POMPDs show great flexibility in their ability to specify the causal structure of an environement. Computation and our ability to inform likelihood and transition dynamics are the only limits to complexity that could be specified. This paper implemented a POMPD capable of combining multiple decisions, such as treating and testing, but more complex generative models could include multiple testing and treatment actions allowing the planning multidrug therapies, integrating multiple types of testing and modelling more complicated modelling of the biology underlying tumor development. But to avoid having to figure out how quarks and leptons might interact to produce a cancer cell, we will have to limit our desire for higher causal resolution. While simulating finer grained mechanism of tumor biology likely will benefit clinical adaptions of AT, abstracting away physical mechanisms as random fluctuations inherent to the modelled environment will at some point be necessary. Models

for real-time treatment decisions will therefore have to need the capacity model the uncertainty of the environment, and account for it in decision-making.

- Another quality is the capacity to balance expected information gain against expected utility. The certainty of the model is incorporated directly in selecting when to apply treatments and when to test. This suggests that there is more information to be gained from testing when treatment is being applied, as it provides a view into the underlying resistance dynamics. Immediate testing seems to suggest that the models are more likely to apply testing when also applying treatment. Another interesting corollary to basing therapy plans in AI is that POMPD solutions could believe that two patients have exactly the same tumor burden but suggest treatment for one patient and not the other. This is a feature, not a bug. If certaintuy about an underlying state (e.g., resistance in the models in this paper), there will expected information gain from treating. Yet if the model is uncertain about the resistance state of the other patient, it might be worthwhile to treat simply to gain information. This perspective slurs the line between testing and treating and drastically alters the decisions on when to collect real-time data patient data by trading uncertainty and utility in a quantitative fashion.
- Another key quality of models desired by West et al. (2023) is the ability to carefully monitor the uncertainty of outcomes. Given a pre-specified generative model, the uncertainty of both current states, and expected outcomes can easily be extracted and interogated (See appendix C for examples).

Future Research

While the results are initially promising, the simple nature of the simulated clinical setting prohibits drawing any strong conclusions about whether the promise of using AI POMPDs in a clinical setting. For example, they might simply be too computationally expensive to be practically viable for more complex generative models.

To further investigate whether POMDPs could viably be applied in a clinical setting, multiple problems will need to be solved. For example, in the present study, the environment featured only discrete values. Since biomarkers and dosage intensities will likely take continuous values, some combination of binning continuous values to discrete values or adapting the model structure to work with continuous data is necessary. Deep learning has been used to construct likelihood mappings and transition probabilities in POMDPs from continuous data (Çatal et al. 2020). Another key issue is determining a way to inform likelihood and transition probabilities in a fashion that would be viable in a clinical setting. The present study used the actual transition probabilities of the simulated environment and, depending on the simulation run, noiseless likelihood mappings. Possibly, simulations of cancer dynamics, such as those by Zhang et al. (2017), could be used to inform transition probabilities, and a clinical model could potentially readjust to patient data. The AI implementation of POMDPs has a developed literature of online learning of the categorical probability distributions of neede to accurately represent the environment. Even the potential information gain form learning transition and likelihood probabilities can be factored in to decision-making ((Smith, Friston, and Whyte 2022) and Da Costa et al. (2020)]).

Another key issue is constructing a more rigorous benchmarking system. In the present paper, the ranges of the baseline range-bounded strategy were selected fairly arbitrarily. It produced convincing results during initial testing and did manage to control tumor levels for a substantial amount of time. However, a systematic method for comparing models is necessary, especially considering the difficulty of real-world tests. No matter the choice of modeling framework, building a benchmarking could help propel modeling work in AT. Given the extreme barriers to real-world testing performance testing of patient-specific models, maximizing the information that can be extracted from simulation work seems crucial. The detailed qualitative account of needed developments on the needed developments mathematical modelling of AT dynamics by West et al. (2023) could be translated to a set of simulation environments used for benchmarking models. Potentially useful simulation scripts already exist. For example, Zhang et al. (2017) released their simulations on mCRCP dynamic. Wrapping existing simulations into environments that can easily faciliates actions and observations with simulated agents could be done by using an API like the one used in Gymnasium (formerly OpenAI Gym) (Towers et al. 2024). This would hopefully minimize friction for non-oncology researchers wishing to

contribute to the field, since it would allow them to prioritize improving model performance instead of having too spend excessive time deciphering how to construct useful metrics themselves.

POMPDs could also be "reverse-engineered" from more complex models, which could more easily be trained if a benchmarking suite existed. For example, if a black-box model, like a neural network, can be shown to successfully control treatments, fitting a POMPD to its behavior would probably be a fairly straight-forward endavour since AI POMPDs often have been used in computational psychiatry to model human decision-making. The same techniques could potentially allow us to translate of otherwise non-transparent models into the structure of a POMPD, thus combining the performance of the black-box model with the transparency of POMPD scheme.

Appendix A - In Depth Description of the AI POMPD Scheme

The canonical implenation of POMDPs in Active Inference requires specificing the following categorical probability distributions.

The likelihood model, A. A is typically a set of arrays where each array corresponds to a modality — a category of observations. The each cell of the array describes the likelihood of making a particular observation in the modality given some configuration of states. For example, PSA readings could be considered a modality, and different test results would be the observations within the modality. The likehoods within the modality could then be modeled as a perfect signal of trumor burden, i.e., the same tumor burden always gives the same test result, or as a noisy signal.

The transition model, B, describes the probability of each possible stransitions of within a type of state between time-steps. B also encodes how actions are expected to influence the transition probabilities between states. For example, it could describe how a tumor is likely to evolve from one time-step to the next depending on whether treatment is being applied. The transition model is usually coded as a collection of three-dimensional arrays, a first dimension for the next state, a dimension for the current state, and a third dimension with the length of each action that would influence transition probabilities of the state.

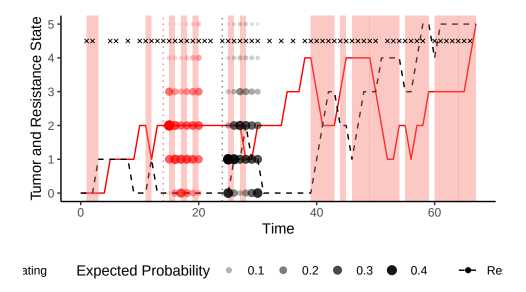
A prior over preferred states C. A particularity of AI POMPDs is that utility is a probabilities for making certain observations. It can be considered the states that the system attempts to "self-organize" around. This method of modelling preferences can superficially seem odd, but its benefits will be clear later.

A prior over initial states D. These are vectors of probabilities specifying intial beliefs. In continuation with example above, D could specify how probable a model believes different levels of tumor burden are before making any observations. When AI POMPDs are solved on multiple time-steps, the posterior beliefs of the preceding time-step replaces D.

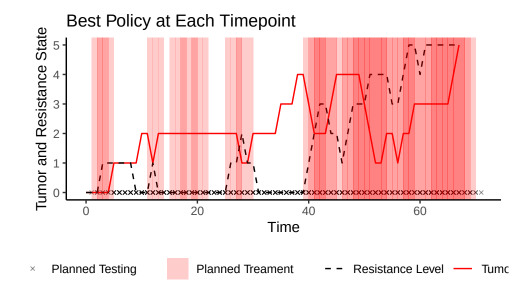
Appendix B - A Dissection POMPD Behavior.

Expected trajectories

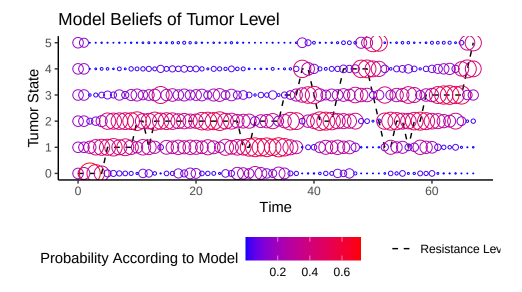
For each the policy, the agent evaluates the expected trajectory of the hidden states (see Fig. x). These evaluations provide insights into the model's uncertainty and what it expects to happen under the best policy at each time point.



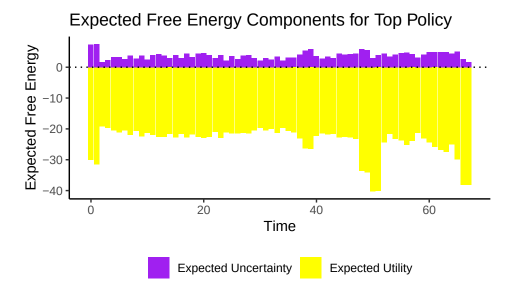
The amount of "flip-flopping" on decisions can also be investigated by examining what policy the model finds the most promising at each time step (see Fig. 4).



Curiously, there is a long stretch, approximately from timesteps 30 to 39, where the model never considers treating to be the optimal course of action, even though the resistance level is low. The model also holds accurate beliefs about the resistance level at this timepoint (see Fig. 2). However, this is likely due to a combination of multiple factors. First, there is a cost to treating, which means that the model generally prefers not to treat. It also underestimates the tumor burden at this timepoint (see Fig. 5) and gets "caught off guard" by a sudden rise in the tumor level. Since there is a cost associated with testing, it is also hesitant to apply tests during this period. Because it only considers six timesteps into the future, it likely doesn't consider the long-term consequences of not reducing the tumor level.



The exact "decision making" for each policy can be evaluated. Each policy is evaluated for its EFE, which is a combination of the expected utility penalized for expected uncertainty (see Fig. 6).



The free energy components reveal several interesting decision-making points. At first, the model, given its uniform priors, is very uncertain about outcomes, and decreasing uncertainty weighs heavily in its decision making. This uncertainty also means that the model believes it is more likely to be in a risky situation, and reducing this risk is a priority. During the earlier identified pivotal period between timesteps 30-39, there is a concerning lack of increase in the weight of the utility component. This could mean that the model is too concerned with avoiding both treating and testing. It is only from timestep 40 onwards that the model realizes the severity of the situation, i.e., the rising tumor level (see Fig. 5). This analysis warrants strategies to increase the model's ability. Changing the prior preferences to make testing and treatment "cheaper" in terms of utility might help. However, simply treating and testing more would translate to actual costs, both financial and to patient well-being. While it is important that the prior preferences are tuned to costs in a clinical setting, it would probably be more fruitful to give the model longer policies to consider. This would hopefully yield models that are better tuned to the risks of not detecting and controlling tumor levels at low values.

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