Operator splitting for adaptive radiation therapy with nonlinear health dynamics

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ABSTRACT

We present an optimization-based approach to radiation treatment planning over time. Our approach formulates treatment planning as an optimal control problem with nonlinear patient health dynamics derived from the standard linear-quadratic cell survival model. As the formulation is nonconvex, we propose a method for obtaining an approximate solution by solving a sequence of convex optimization problems. This method is fast, efficient, and robust to model error, adapting readily to changes in the patient’s health between treatment sessions. Moreover, we show that it can be combined with the operator splitting method ADMM to produce an algorithm that is highly scalable and can handle large clinical cases. We introduce an open-source Python implementation of our algorithm, AdaRad, and demonstrate its performance on several examples.

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1. Introduction

In radiation therapy, beams of ionizing radiation are transmitted into a patient, damaging both tumor cells and normal tissue. The goal of radiation treatment planning is to deliver enough dose to the tumor so that diseased cells are killed, while avoiding excessive injury to the normal tissue and organs-at-risk (OARs). This is achieved by optimizing the beam intensity profile, or fluence map, subject to constraints on the dose to certain parts of the patient’s anatomy. The fluence map optimization problem is well-studied\cite{1,15,29,30}, and technology like intensity-modulated radiation therapy (IMRT) is now widespread in the clinic\cite{18,41,42,48}.

Treatment in practice usually takes place over multiple sessions. A clinician will divide up the total prescribed dose into smaller dose fractions, which are delivered over the course of several weeks or months. This permits normal tissue time to recover and repair cell damage, but also gives tumors an opportunity to proliferate, especially when the treatment course is long. A study of 4338 prostate cancer patients showed that biochemical failure increases by 6\% for every 1 week increase in treatment time, with a dose equivalent of proliferation of 0.24 Gy/day\cite{38}. Thus, an important question in treatment planning is...
how to choose the sequence of deliverable doses such that they balance these temporal
effects on a patient’s health.

1.1. Related work

Early clinical practitioners split the prescribed dose equally over a fixed number of ses-
sions. While convenient, this method does not account for errors or uncertainty in the
treatment process. For example, due to movements of the patient’s anatomy, the expected
dose may differ from the actual dose to an anatomical structure. If the actual dose is observ-
able, a common way to compensate for this is to divide the residual dose (i.e. the difference
between the prescribed and cumulative actual dose) across the remaining sessions. This
then becomes the new per-session dose goal. In [9], the authors solve for the beam intensi-
ties by minimizing the sum-of-squared difference between this goal dose and the expected
dose. They compare the results when errors are perfectly known, so the expected dose is
equal to the actual dose, with the results when errors are assumed to be zero. A similar
approach is taken in [12], except the errors are modeled explicitly as a random shift in
the surrounding voxels. Instead of the dose to each voxel, [34] work with the equivalent
uniform dose (EUD), a value that captures the biological effect of a dose distribution over
a region. Their objective is to minimize the sum of the EUD over all treatment criteria
subject to bounds on the EUD of the tumor and normal tissues. To solve this problem,
they employ methods from approximate dynamic programming coupled with a discrete
probabilistic model of the dose error.

The papers we have discussed so far only focus on the dose to the patient. By contrast,
Kim et al. [23] introduces a Markov decision process model that includes both the dose
(action) and the patient’s health state. Each choice of dose induces a transition to a partic-
ular health state with some probability. Making this idea concrete, Mizuta et al. [28] define
the health of a tumor (resp. OAR) to be the radiation (resp. damage) effect of the delivered
dose, as calculated from the linear-quadratic (LQ) model of cell survival [13]. They ana-
lyze a simple example with one tumor and one OAR and find that the optimal fractionation
scheme is either a single session delivery of the full dose or equal dose fractions, depending
on the relationship between the LQ parameters. The authors of [3] extend this analysis to
incorporate accelerated tumor repopulation and show that the dose per session increases
over the treatment course. Using simulated annealing, Yang and Xing [45] solve a simi-
lar treatment planning problem based on the LQR model, which captures all 4 Rs (repair
of sublethal damage, repopulation, redistribution, and reoxygenation) of cellular radiation
response [7].

In [8], the authors take a probabilistic approach to patient dynamics. They model the
patient’s breathing motion using a probability mass function (PMF) over a finite set of
states. They then solve a robust optimization problem that enforces dose bounds over a set
of PMFs, which represent uncertainty during treatment. This uncertainty set is updated
after the dose fraction is delivered, and the problem is re-solved for the next session. A
follow-up paper [27] numerically studies the effects of adjusting the target dose based on
the dose delivered to date. It shows that this type of adjustment can lead to a high degree
of heterogeneity in the per-fraction dose distribution, which is undesirable from a medical
standpoint. The paper recommends an alternative method of uncertainty set adaptation to
mitigate these effects.
The above analyzes provide insight into the tradeoffs between different fractionation schemes in a simple setting. However, most clinical cases are more complex, involving multiple tumors, OARs, and nonlinear constraints. For instance, dose-volume (i.e. percentile) constraints are widely used to limit the radiation exposure of a percentage of an anatomical structure, such as the spine. These constraints are nonconvex, but can be approximated by a convex restriction \[14,19,47\]. In \[31\], the authors consider a dynamic setting with multiple OARs and dose-volume constraints. Starting from a given set of beam intensities, they solve for the optimal number of sessions and OAR sparing factors. They also derive sufficient conditions under which the optimal treatment consists of equal dose fractions. In a follow-up paper \[32\], the authors integrate the spatial and temporal aspects of the problem, treating both beam intensities and number of sessions as variables. Restricting their attention to equal fractions, they propose a two-stage solution algorithm: in the first stage, they solve for the optimal beams given each potential fixed number of sessions, and in the second stage, they select the number of sessions based on the optimal objectives from the first stage. They show that their method achieves better tumor ablation than conventional IMRT or the spatiotemporally separated method.

Perhaps the paper most similar to ours is \[24\]. In it, the authors propose a stochastic control formulation of the adaptive treatment planning problem with multiple tumors and OARs. They estimate the radiation response of the tumors with a log-linear cell kill model and the response of the OARs with the standard LQ model. Their goal is to minimize the expected number of tumor cells at the end of treatment subject to bounds on the radiobiological impact on the OARs. Uncertainty arises in the cell model parameters, which may fluctuate randomly between sessions, representing unpredictable changes in the patient’s health status. The authors fix the number of sessions and focus on optimizing with respect to the beam intensities. They show that their problem is convex, so can be solved using a combination of standard stochastic control methods and off-the-shelf convex solvers, and provide several examples demonstrating the effectiveness of their approach.

1.2. Contribution

In this paper, we integrate the stochastic control approach with a distributed optimization algorithm to produce a method for efficient large-scale adaptive treatment planning. As clinical cases are quite complex, with tens of thousands of beams and treatment that takes place over months, such methods are necessary to construct plans in a timely fashion. (See \[21\] for a review of previous work on high-performance computing in radiation therapy, particularly treatment optimization). We formulate the adaptive treatment planning problem as a finite-horizon nonconvex optimal control problem. To solve it, we introduce an operator splitting algorithm, which is based on solving a sequence of convex approximations. Our algorithm is naturally parallelizable and can handle a large number of beams, sessions, and anatomical targets or OARs. Moreover, it can be combined with model predictive control to produce treatment plans that are robust to errors and uncertainty about the patient’s health status. We illustrate our algorithm’s performance on a synthetic case, as well as a large prostate cancer case, and provide an implementation in the Python package AdaRad: https://github.com/anqif/adarad.
2. Problem formulation

In radiation treatment, beams of ionizing radiation are delivered to a patient from an external source. The goal is to damage or kill diseased tissue, while minimizing harm to surrounding healthy organs. A course of treatment is generally divided into $T$ sessions. At the start of session $t$, the clinician chooses the intensity levels of the $n$ beams, denoted by $b_t \in \mathbb{R}_+^n$. Typically, $T \approx 20$ to 40 and $n$ is on the order of $10^3$ to $10^4$. We are interested in determining the best sequence of beam intensities $b = (b_1, \ldots, b_T)$, otherwise known as a treatment plan, subject to upper bounds $B_t \in \bar{\mathbb{R}}_+^n$ on $b_t$ for $t = 1, \ldots, T$.

2.1. Anatomy and doses

The beams irradiate an area containing $K$ anatomical structures, labeled $i \in \{1, \ldots, K\}$, where usually $K < 10$. A subset $\mathcal{T} \subset \{1, \ldots, K\}$ are targets/tumors and the rest are OARs. The dose delivered to each structure is linear in the beam intensities. We write the dose vector $d_t = A_t b_t$ with $A_t \in \mathbb{R}^{K \times n}_+$ a known matrix that characterizes the physical effects and define $d = (d_1, \ldots, d_T)$. Notice that since $b_t$ and $A_t$ are nonnegative, $d_t \geq 0$.

In every session, we impose a penalty on $d_t$ via a dose penalty function $\phi_t : \mathbb{R}^K \to \mathbb{R} \cup \{\infty\}$. A common choice is

$$\phi_t(d_t) = \theta^T_t d_t + \xi^T_t d_t^{\circ 2},$$

where $\theta_t \in \mathbb{R}^K$ and $\xi_t \in \mathbb{R}_+^K$ are constants. Here $d_t^{\circ 2} = d_t \odot d_t$ denotes the vector $d_t$ with each element squared. The total dose penalty over all sessions is

$$\phi(d) = \sum_{t=1}^T \phi_t(d_t).$$

Additionally, we enforce upper bound constraints $d_t \leq D_t$, where $D_t \in \bar{\mathbb{R}}_+^K$ is the maximum dose in session $t$.

2.2. Health dynamics

To assess treatment progress, we examine the health status of each anatomical structure and encode these statuses in a vector $h_t \in \mathbb{R}^K$. For now, the details of this encoding do not matter. Typically, $h_{t,i}$ represents an estimate of the total surviving cells in structure $i$. Hence if $i \in \mathcal{T}$, a smaller $h_{t,i}$ is desirable (since the tumor is shrinking), while if $i \notin \mathcal{T}$, a larger $h_{t,i}$ is desirable.

From an initial $h_0$, the health status evolves in response to the radiation dose and various other biophysical factors that depend on the patient’s anatomy, generating a health trajectory $h = (h_1, \ldots, h_T)$. Here we represent its dynamics as

$$h_t = f_t(h_{t-1}, d_t), \quad t = 1, \ldots, T,$$

where $f_t : \mathbb{R}^K \times \mathbb{R}^K \to \mathbb{R}^K$ is a known mapping function. In this paper, we focus on the linear-quadratic (LQ) model in which

$$f_{t,i}(h_{t-1}, d_t) = h_{t-1,i} - \alpha_{t,i} d_{t,i} - \beta_{t,i} d_{t,i}^2 + \gamma_{t,i}, \quad i = 1, \ldots, K, \quad t = 1, \ldots, T$$

(2)
with constants $\alpha_t \in \mathbb{R}^K$, $\beta_t \in \mathbb{R}^K_+$, and $\gamma_t \in \mathbb{R}^K$. This model is commonly used to approximate cellular response to radiation [6,13,37]. Specifically, in the LQ + time framework [39], $h_{t,i}$ is the log of the fraction of surviving cells in structure $i$ after a dose $d_{t,i}$, while $\alpha_{t,i}/\beta_{t,i}$ and $\gamma_{t,i}$ are constants related to the structure’s survival curve and repair/repopulation rate, respectively. Notice that Equation (2) implies that the health status of each structure evolves independently of the others.

### 2.3. Health penalty and constraints

In order to control the patient’s health, we introduce a health penalty function $\psi_t : \mathbb{R}^K \to \mathbb{R} \cup \{\infty\}$ that imposes a penalty on $h_t$. Moreover, we assume that

$$\psi_t(h_t) = \psi_t(h_{t,1}, \ldots, h_{t,K}) \text{ is monotonically increasing in } h_{t,i} \quad i \in T$$

$$\psi_t(h_t) = \psi_t(h_{t,1}, \ldots, h_{t,K}) \text{ is monotonically decreasing in } h_{t,i} \quad i \notin T$$

for $t = 1, \ldots, T$. This means that for a target, the health penalty increases as the health status increases, while for an organ-at-risk, the health penalty decreases as the health status increases. The assumption is reasonable if, for instance, $h_t$ is a measure of cell survival in session $t$, so a lower (higher) status is desirable for a target (organ-at-risk). An example of a penalty function that satisfies (3) is

$$\psi_t(h_t) = \bar{w}^T(h_t - h_t^{\text{goal}})_+ + \bar{w}^T(h_t - h_t^{\text{goal}})_-,$$

where $h_t^{\text{goal}} \in \mathbb{R}^K$ is the desired health status and $\bar{w} \in \mathbb{R}^K_+$ and $\bar{w} \in \mathbb{R}^K_+$ are parameters with $\bar{w}_i = 0$ for $i \in T$ and $\bar{w}_i = 0$ for $i \notin T$. Here we define $(x)_+ = \max(x, 0)$ and $(x)_- = -\min(x, 0)$ applied elementwise to $x$. The total health penalty is

$$\psi(h) = \sum_{t=1}^T \psi_t(h_t).$$

In addition, we enforce bounds $H_t \in \mathbb{R}^K$ on the health status such that $h_{t,i} \leq H_{t,i}$ for $i \in T$ and $h_{t,i} \geq H_{t,i}$ for $i \notin T$.

### 2.4. Optimal control problem

Given an initial health status $h_0$, we wish to select a treatment plan that minimizes the total penalty across all sessions. Thus, our problem is

$$\begin{align*}
\text{minimize} & \quad \sum_{t=1}^T \phi_t(d_t) + \sum_{t=1}^T \psi_t(h_t) \\
\text{subject to} & \quad h_t = f_t(h_{t-1}, d_t), \quad t = 1, \ldots, T, \\
& \quad h_{t,i} \leq H_{t,i}, \quad i \in T, \quad h_{t,i} \geq H_{t,i}, \quad i \notin T, \quad t = 1, \ldots, T, \\
& \quad d_t = A_t b_t, \quad 0 \leq d_t \leq D_t, \quad 0 \leq b_t \leq B_t, \quad t = 1, \ldots, T
\end{align*}$$

(4)

with variables $(b_1, \ldots, b_T), (d_1, \ldots, d_T)$, and $(h_1, \ldots, h_T)$. This is a discrete-time optimal control problem. If $\phi_t$ and $\psi_t$ are convex and $f_t$ is affine, e.g. $f_t$ is given by (2) with quadratic dose effect $\beta_t = 0$, it is also convex and can be solved directly using standard convex solvers [5].
The equality constraint has been replaced with two inequality constraints: a lower bound for targets and an upper bound for OARs. Notice that the first inequality is the only nonconvex constraint in (5). Our relaxed problem has the same solution set as (4) because these two inequalities are tight at the optimum.

**Proposition 3.1:** Let \((b^*, d^*, h^*)\) be a solution to problem (5). If conditions (2) and (3) hold,

\[ h_t^* = f_t(h_{t-1}^*, d_t^*), \quad t = 1, \ldots, T. \]

**Proof:** Suppose there exist some \(t \in \{1, \ldots, T\}\) and \(i \in \mathcal{T}\) such that \(h_{t,i}^* > f_{t,i}(h_{t-1}^*, d_t^*)\). Then, we can choose an \(\epsilon > 0\) such that \(h_{t,i}^* > h_{t,i}^* - \epsilon > f_{t,i}(h_{t-1}^*, d_t^*)\). Since \(f_{s,i}(h_{s-1}^*, d_s)\) is nondecreasing in \(h_{s-1}^*\) for all \(s \in \{1, \ldots, T\}\), the point \((b^*, d^*, h^*)\) with

\[ \hat{h}_{s,j} = \begin{cases} h_{s,j}^* - \epsilon & s = t, j = i \\ h_{s,j}^* & \text{otherwise} \end{cases} \]

is feasible for problem (5) because

\[ \hat{h}_{t,i} > f_{t,i}(h_{t-1}^*, d_t^*) \geq f_{t,i}((h_{t-1}^*, d_t^*)), \quad h_{t+1,i}^* \geq f_{t+1,i}(h_{t,i}^*, d_t^*) \geq f_{t+1,i}(h_{t,i}^*, d_t^*), \]

and \(\hat{h}_{t,i} < h_{t,i}^* \leq H_{t,i}\). Moreover, by condition (3), \(\psi_t(h_t^*) < \psi_t(h_t^*)\) so \((b^*, d^*, h^*)\) achieves a lower objective value than \((b^*, d^*, h^*)\), contradicting our original assumption. An analogous argument holds for \(t \in \{1, \ldots, T\}\) and \(i \notin \mathcal{T}\) such that \(h_{t,i}^* < f_{t,i}(h_{t-1}^*, d_t^*)\) with \(\hat{h}_{t,i} = h_{t,i}^* + \epsilon\)

**4. Sequential convex optimization**

**4.1. Algorithm description**

Problem (5) is in general nonconvex because the target’s health dynamics constraint

\[ h_{t,i} \geq f_{t,i}(h_{t-1}, d_t), \quad i \in \mathcal{T}, \quad t = 1, \ldots, T \]

is nonconvex when any \(\beta_t \neq 0\). However, we can derive an estimate of its optimum by solving a sequence of convex approximations. Each approximation is formed by linearizing...
the health dynamics function (2) around a fixed dose point and replacing the right-hand side of (6) with this linearization minus a slack variable. The slack allows for a degree of error in the approximation and is penalized in the objective.

More precisely, let \( d^t_i \in \mathbb{R}^K \) for \( t = 1, \ldots, T \). Define the linearized dynamics function

\[
\hat{f}_{t,i}(h_{t-1}, d^t_i; d^t_i) = h_{t-1,i} - \alpha_{t,i} d^t_i - \beta_{t,i} d^t_i (2d^t_i - d^t_i) + \gamma_{t,i}, \quad i = 1, \ldots, K. \tag{7}
\]

This is the tangent line to the LQ function (2) at \( d^t_i \) and, as such, provides a uniform upper bound because \( \beta_t \geq 0 \). We replace the nonconvex constraint (6) in problem (5) with the affine constraint

\[
h_{t,i} = \hat{f}_{t,i}(h_{t-1}, d^t_i; d^t_i) - \delta_{t,i}, \quad i \in T, \quad t = 1, \ldots, T, \tag{8}
\]

where \( \delta_t \in \mathbb{R}^K_+ \) is a slack variable. (The inequality can be tightened into an equality due to Proposition 3.1). Convex approximation \( s \) is then

\[
\begin{aligned}
\text{minimize} & \quad \sum_{t=1}^T \phi_t(d^t_i) + \sum_{t=1}^T \psi_t(h_t) + \lambda \sum_{t=1}^T 1^T \delta_t \\
\text{subject to} & \quad h_{t,i} = \hat{f}_{t,i}(h_{t-1}, d^t_i; d^t_i) - \delta_{t,i}, \quad i \in T, \quad \delta_t \geq 0 \quad t = 1, \ldots, T, \\
& \quad h_{t,i} \leq H_{t,i}, \quad i \in T, \quad h_{t,i} \geq H_{t,i}, \quad i \not\in T, \quad t = 1, \ldots, T, \\
& \quad d_t = A_t b_t, \quad 0 \leq d_t \leq D_t, \quad 0 \leq b_t \leq B_t, \quad t = 1, \ldots, T.
\end{aligned} \tag{9}
\]

with variables \((b_1, \ldots, b_T), (d_1, \ldots, d_T), (h_1, \ldots, h_T), \) and \((\delta_1, \ldots, \delta_T)\) and slack penalty parameter \( \lambda > 0 \). The parameter \( \lambda \) is typically determined empirically or by trial-and-error. This problem is convex and can be solved using standard convex solvers. Given a solution to (9), we set the next linearization point \( d^{s+1} = (d^{s+1}_1, \ldots, d^{s+1}_T) \) equal to the optimal dose.

**Algorithm 4.1** Sequential convex optimization.

**input:** initial point \( d^0 \), parameter \( \lambda > 0 \).

**for** \( s = 0, 1, \ldots \) **do**

1. **Linearize.** For \( t = 1, \ldots, T \), form the linearization (7) around \( d^t_i \).
2. **Solve.** Set \( d^{s+1}_t \) equal to an optimal dose of problem (9).

**until** stopping criterion (10) is satisfied.

Algorithm 4.1 is a special case of the convex-concave procedure (CCP) [25,33,46], which is itself a form of majorization-minimization [20,36]. CCP is a heuristic for finding a local optimum of a nonconvex optimization problem. It is guaranteed to converge; indeed, when certain differentiability conditions are met, it converges to a stationary point [35]. As a descent algorithm, CCP is usually terminated when the change in the objective falls below
some user-specified threshold $\epsilon > 0$, i.e.

$$p_{\text{opt}}^i - p_{\text{opt}}^{i+1} < \epsilon,$$

where $p_{\text{opt}}^i$ is the optimal objective of problem (9). In our simple experiments, we have found that an initial linearization point of $d^0 = 0$ and threshold of $\epsilon = 10^{-3}$ produce good results.

### 4.2. Illustrative example

#### 4.2.1. Problem instance

We consider an example with $n = 1000$ beams divided into 50 bundles of 20 parallel beams each, positioned evenly around a half-circle. There are $K = 5$ structures, a single target $T = \{1\}$ and four OARs (including generic body voxels) depicted in Figure 1. Treatment takes place over $T = 20$ sessions, so the basic problem has $nT + 2KT = 20200$ variables.

The patient’s initial health status is $h_0 = (1, 0, 0, 0, 0)$. His status evolves according to Equation (2) with

$$\alpha_t = (0.01, 0.50, 0.25, 0.15, 0.005),$$

$$\beta_t = (0.001, 0.05, 0.025, 0.015, 0.0005),$$

$$\gamma_t = (0.05, 0, 0, 0, 0)$$

over all sessions $t = 1, \ldots, T$.

We set the health penalty function to

$$\psi_t(h_t) = (h_{t,1})_+ + \sum_{i=2}^{5} (h_{t,i})_-, \quad t = 1, \ldots, T.$$

![Figure 1. Anatomical structures for Example 4.2. Red is the target ($i = 1$), while green ($i = 2$), blue ($i = 3$), and orange ($i = 4$) are specific OARs. White denotes the non-target body voxels ($i = 5$).](image)
This function penalizes positive statuses of the target and negative statuses of the OARs. Moreover, we constrain the target’s health status to be $h_{t,1} \leq 2.0$ for $t = 1, \ldots, 15$ and $h_{t,1} \leq 0.05$ for the remaining sessions, and we enforce a bound on the other structures’ health statuses of $(h_{t,2}, h_{t,3}, h_{t,4}, h_{t,5}) \succeq (-1.0, -2.0, -2.0, -3.0)$. Thus,

$$H_t = \begin{cases} 
(2.0, -1.0, -2.0, -2.0, -3.0) & t = 1, \ldots, 15 \\
(0.05, -1.0, -2.0, -2.0, -3.0) & t = 16, \ldots, T.
\end{cases}$$

For the dose penalty function, we choose

$$\phi_t(d_t) = \sum_{i=1}^{4} d_{t,i}^2 + 0.25d_{t,5}^2, \quad t = 1, \ldots, T.$$ 

In addition, we restrict the dose and beam intensity to be no more than $D_t = 20$ and $B_t = 1.0$, respectively, over all sessions $t$.

### 4.2.2. Computational details

We implemented Algorithm 4.1 in Python using CVXPY [10] and solved problem (9) with MOSEK [2]. From an initial $d^0 = 0$ and $\lambda = 10^4$, the algorithm converged in 11 iterations to a threshold of $\epsilon = 10^{-3}$. Total runtime was approximately 17 seconds on a 64-bit Ubuntu OS desktop with 8 4-core Intel i7-4790k / 4.00 GHz CPUs and 16 GB of RAM.

### 4.2.3. Results and analysis

The optimal treatment plan is depicted in Figure 2. Beams are densely clustered diagonal from the vertical, striking the target while largely sparing the OARs. As the sessions continue, the number of beams slowly increases, damaging some of the less sensitive organs ($i = 3$ and $4$). Then at $t = 16$, when the target’s health bound becomes more stringent, the beam density drops precipitously so that only a narrow bundle remains focused on the target, keeping its health status at the desired level.

![Figure 2. Optimal beam intensities for Example 4.2.](image)
Figure 3. Optimal (a) radiation dose and (b) health status trajectories for Example 4.2. Dashed lines represent the bounds on the dose and health status ($D_{t,i}$ and $H_{t,i}$ respectively).

Figure 3 shows the radiation dose and health status resulting from this plan. The latter was computed by plugging the optimal dose into Equation (2). Total dose to the target ($i = 1$) and body voxels ($i = 5$) far exceed the dose to any other structures. By the end of treatment, the target’s health status has fallen to a steady 0.05, while the health statuses of the OARs remain within their respective lower limits.

5. Model predictive control

5.1. Algorithm description

So far, we have assumed that at the time of planning, $f_t$ perfectly captures the health dynamics from $t = 1, \ldots, T$. This is rarely true in practice. A patient’s anatomy changes
unpredictably between sessions, affecting the dispersion of radiation beams and the course of their health status. We can incorporate these changes into problem (4) using model predictive control (MPC).

MPC is a powerful technique for automatic control of complex, nonlinear, stochastic systems. It performs extremely well even when the dynamics are approximated by a simple model, since the system’s state is updated regularly and new information is incorporated into the solution. This is particularly fitting for radiation treatment planning.

As is customary in MPC, we first convert the state variable constraints in the original problem into soft constraints, i.e. we remove the inequality constraints on \( h \) in (4) and add a penalty for violating them to the objective. Let \( c_\tau : \mathbb{R}^K \to \mathbb{R} \) be the corresponding health violation penalty function, defined as

\[
c_\tau(h_\tau) = \sum_{i \in T} (h_{\tau,i} - H_{\tau,i})_+ + \sum_{i \notin T} (H_{\tau,i} - h_{\tau,i})_+, \quad \tau = 1, \ldots, T.
\]

This penalty function allows us to accommodate new and unexpected changes in the patient’s health, such as the metastasis of a tumor that renders it impossible to control without exceeding the health damage limit of an OAR.

We are now ready to describe MPC for our model. At the beginning of each session \( t \), we observe \( A_t, f_t \), and the patient’s true health status, \( h_{t-1} \), then form the problem

\[
\begin{align*}
\text{minimize} & \quad \sum_{\tau=t}^{T} \phi_\tau(d_\tau) + \sum_{\tau=t}^{T} \psi_\tau(h_\tau) + \eta \sum_{\tau=t}^{T} c_\tau(h_\tau) \\
\text{subject to} & \quad h_\tau = f_t(h_{\tau-1}, d_\tau), \quad \tau = t, \ldots, T, \\
& \quad d_\tau = A_t b_\tau, \quad 0 \leq d_\tau \leq D_\tau, \quad 0 \leq b_\tau \leq B_\tau, \quad \tau = t, \ldots, T
\end{align*}
\]

with variables \((b_t, \ldots, b_T), (d_t, \ldots, d_T)\), and \((h_t, \ldots, h_T)\) and violation penalty parameter \( \eta > 0 \). Since \( c_\tau \) is convex, problem (11) is convex and can be solved using a slight variation on Algorithm 4.1. Let \( \bar{b} = (\bar{b}_t, \ldots, \bar{b}_T) \) be the optimal treatment plan. We carry out only the first treatment, \( \bar{b}_t \), and update our observations \( A_{t+1}, f_{t+1} \), and \( h_t \) based on the patient’s response. This process repeats until all \( T \) sessions have been completed.

5.2. Illustrative example

5.2.1. Problem instance

We return to the setting of Example 4.2, except now, the health dynamics are modeled with some error. Specifically, let \( h_{t-1} \) be the patient’s health status at the beginning of session \( t \) and \( d_t \) the dose delivered during session \( t \). Our model predicts the status will become \( \hat{h}_t = f_t(h_{t-1}, d_t) \). In fact, at the beginning of the next session, we observe the true health status to be

\[
h_{t,i} = \begin{cases} 
\max(\hat{h}_{t,i} + \omega_{t,i}, 0) & i \in T \\
\min(\hat{h}_{t,i} + \omega_{t,i}, 0) & i \notin T
\end{cases}
\]

where \( \omega_t \in \mathbb{R}^K \) is drawn from \( N(\mu, \sigma^2 I) \). This random process continues for \( t = 1, \ldots, T \).

For this example, we choose \( \mu = 0 \) and \( \sigma = 0.1 \). The rest of the functions and parameter values are identical to 4.2. In particular, we still employ the LQ model (2) with constant
\(\alpha_t, \beta_t, \text{ and } \gamma_t\) even though the health status is now stochastic. We plan the treatment using MPC with \(\eta = 10^4\) and compare the results to those generated by the naive approach, which simply solves problem (4) once prior to session 1.

### 5.2.2. Computational details

We solved problem (11) using Algorithm 4.1 with \(\lambda = 10^4\) and \(\epsilon = 10^{-3}\). For the initial dose in session 1, we chose \(d^0 = 0\). In each subsequent session \(t\), we set \(d^0\) to be the (truncated) optimal dose point from the previous session, \((d^*_1, \ldots, d^*_T)\). With these parameters, the algorithm took an average of 7 iterations per session to achieve convergence; most runs completed in only 3–4 iterations. The total runtime was 116 seconds.

### 5.2.3. Results and analysis

Figure 4 depicts the treatment plan output by MPC. Most beams are aimed slightly diagonal from the vertical, similar to the naive plan (Figure 2) up to session 14. Then, the bundles of beams start to grow sparser and fan out, hitting more areas of the OARs. This sparse irradiation pattern continues until the final session, when there is a brief spike in intensity to bring the target’s health status into the desired range.

In Figure 5(a), we plot the dose trajectories of the MPC plan (green) and the naive plan (blue). The MPC curves are more jagged with a large spike at the end of treatment. However, in each structure, the area under the MPC and naive dose curves remains on par. Thus, we conclude that the MPC plan delivers about the same amount of radiation as the naive plan, only spread across a wider range of beam angles/intensities so as to compensate for uncertainty in the health dynamics model.

This strategy results in better patient health as shown in Figure 5(b). The MPC plan reduces the target’s health status to 0.05, while maintaining the health status of the OARs at a high level. Indeed, the health of these organs under the MPC plan exceeds their health under the naive plan by a significant margin in all but structure 4, where the two are relatively equal up until the last session.

![Figure 4](image-url)  
**Figure 4.** Optimal beam intensities for Example 5.2 using MPC.
Figure 5. Optimal (a) radiation dose and (b) health status trajectories for Example 5.2 using MPC (green) and a naive planning approach (blue). Dashed lines represent the bounds on the dose and health status ($D_t$ and $H_t$, respectively). The MPC plan’s health trajectories all remain within the desired bounds, despite the error in the health dynamics model.

6. Operator splitting

MPC enables us to robustly handle uncertainty over time. However, another challenge in radiation treatment planning is the sheer size of problems, which makes them computationally difficult to solve in practice. A typical case with $K = 15$ and $n = 10^4$ requires approximately $10^5$ floating-point operations for the beam-to-dose calculation alone. Over a month of sessions, that comes out to 4.5 million operations on a single machine.

In this section, we propose a fast, efficient method for solving the radiation treatment planning problem using operator splitting. Our method is distributed and scales readily with the number of beams as well as the length of treatment. It can be applied both to the
original problem (4) and the soft constrained MPC variant (11). Below, we describe the mathematical details for the former; the latter is a straightforward extension.

6.1. Consensus form

We first rewrite problem (4) in an equivalent consensus form:

\[
\begin{align*}
\text{minimize} & \quad \sum_{t=1}^{T} \phi_t(d_t) + \sum_{t=1}^{T} \psi_t(h_t) \\
\text{subject to} & \quad h_t = f_t(h_{t-1}, d_t), \quad 0 \leq \tilde{d}_t \leq D_t, \quad t = 1, \ldots, T, \\
& \quad h_{t,i} \leq H_{t,i}, \quad i \in T, \quad h_{t,i} \geq H_{t,i}, \quad i \notin T, \quad t = 1, \ldots, T, \\
& \quad d_t = A_t b_t, \quad 0 \leq d_t \leq D_t, \quad 0 \leq b_t \leq B_t, \quad t = 1, \ldots, T, \\
& \quad d_t = \tilde{d}_t, \quad t = 1, \ldots, T
\end{align*}
\]

with additional variable \( \tilde{d} = (\tilde{d}_1, \ldots, \tilde{d}_T) \). This splits the problem into two parts, one that encapsulates the radiation physics and the other that contains the health dynamics. The parts share no variables. They are only linked by the consensus constraint, \( d_t = \tilde{d}_t \), which requires their doses be equal.

6.2. ADMM

We solve problem (12) using an iterative algorithm called the alternating direction method of multipliers (ADMM) [4]. In ADMM, the beams and health statuses are optimized separately, taking into account the difference between their resulting dose values. This difference is associated with a dual variable \( u = (u_1, \ldots, u_T) \), where each \( u_t \in \mathbb{R}^K \), which is updated every iteration in order to promote consensus.

\begin{algorithm}
\caption{ADMM algorithm.}
\begin{algorithmic}
\STATE \textbf{input:} initial point \((\tilde{d}^0, u^0)\), parameter \( \rho > 0 \).
\FOR {\( k = 0, 1, \ldots \)}
\STATE 1. \textit{Calculate beams.} For \( t = 1, \ldots, T \), set the value of \((b_t^{k+1}, d_t^{k+1})\) to a solution of the problem
\[ \begin{align*}
\text{minimize} & \quad \phi_t(d_t) + \frac{\rho}{2} \| d_t - \tilde{d}^k_t - u_t^k \|_2^2 \\
\text{subject to} & \quad d_t = A_t b_t, \quad 0 \leq d_t \leq D_t, \quad 0 \leq b_t \leq B_t.
\end{align*} \]
\STATE 2. \textit{Calculate health trajectory.} Set the value of \((h_t^{k+1}, \tilde{d}^{k+1})\) to a solution of the problem
\[ \begin{align*}
\text{minimize} & \quad \sum_{t=1}^{T} \psi_t(h_t) + \frac{\rho}{2} \| \tilde{d} - d^{k+1} + u^k \|_2^2 \\
\text{subject to} & \quad h_t = f_t(h_{t-1}, \tilde{d}_t), \quad 0 \leq \tilde{d}_t \leq D_t, \quad t = 1, \ldots, T, \\
& \quad h_{t,i} \leq H_{t,i}, \quad i \in T, \quad h_{t,i} \geq H_{t,i}, \quad i \notin T, \quad t = 1, \ldots, T.
\end{align*} \]
\STATE 3. \textit{Update dual variables.} \( u_t^{k+1} := u^k + \tilde{d}^{k+1} - d^{k+1} \).
\ENDFOR
\STATE \textbf{until} stopping criterion (17) is satisfied.
\end{algorithmic}
\end{algorithm}
Here $1/\rho > 0$ may be interpreted as the step size. Notice that the first step of Algorithm 6.1 can be parallelized across sessions. We impose the dose bound constraint on both the beam and health subproblems because it produces faster convergence in practice.

### 6.2.1. Initialization

For complex problems, the initial dose point $\tilde{d}^0$ can have a significant impact on the performance of Algorithm 6.1. Below, we describe one heuristic that produces a good starting point by solving a series of simple optimization problems. The idea is to first find the optimal beams in the static setting, where the entire treatment is delivered in a single session, then rescale these static beams each session to form a plan in the dynamic setting. By limiting ourselves to scaling factors, we significantly reduce the size of the dynamic problem.

![Equation](image)

with respect to $b_1 \in \mathbb{R}^n$, $d_1 \in \mathbb{R}^K$, $h_1 \in \mathbb{R}^K$, and $\zeta \in \mathbb{R}^K$, where $\mu > 0$ is a slack penalty parameter. A reasonable choice for $\mu = \frac{1}{K-|T|}$, assuming there is at least one non-target structure. Problem (13) is convex and can be easily handled on a single machine (e.g. via interior-point methods) for up to $10^5$ beams. Denote the optimal beam intensities by $b^{\text{stat}}$.

Next, we consider the dynamic treatment planning problem in which the beams for each session are restricted to be a scalar multiple of $b^{\text{stat}}$,

![Equation](image)

with variables $(\nu_1, \ldots, \nu_T, d_1, \ldots, d_T, (h_1, \ldots, h_T), (\xi_1, \ldots, \xi_T))$, where each $\nu_t \in \mathbb{R}$ and $\zeta_t \in \mathbb{R}^K$. This problem can be solved using a slight variation on Algorithm 4.1. (For the initial CCP point, we may use the optimal time-invariant $\nu_t = v$ when $\beta_t = 0$; finding this value entails solving a small convex problem). Since there are only $O(TK)$ variables, convergence is generally quick, taking less than 5 iterations in our experiments. We use the resulting doses as our initial dose point for ADMM, i.e. $\tilde{d}_t^0 = \nu^* b^{\text{stat}}$ for $t = 1, \ldots, T$.

Besides providing a good starting point, this initialization heuristic also gives us a way to quickly tune problem parameters. If the health trajectory from $\tilde{d}_t^0$ is poor, it is much faster to modify weights and re-solve problems (13) and (14) than it is to re-run the full ADMM algorithm.
6.2.2. Stopping criterion

If problem (12) is convex, then under mild conditions, ADMM converges to a solution assuming one exists. Moreover, the primal and dual residuals

\[ r^k_{\text{prim}} = d^k - \tilde{d}^k \]  
\[ r^k_{\text{dual}} = \rho(\tilde{d}^k - \tilde{d}^{k-1}) \]

also converge to zero. Thus, a reasonable stopping criterion is

\[ \|r^k_{\text{prim}}\|_2 \leq \epsilon_{\text{prim}} \text{ and } \|r^k_{\text{dual}}\|_2 \leq \epsilon_{\text{dual}}, \]

where \(\epsilon_{\text{prim}} > 0\) and \(\epsilon_{\text{dual}} > 0\) are tolerances for primal and dual feasibility, respectively. Typically, these tolerances are chosen with respect to absolute and relative cutoffs \(\epsilon_{\text{abs}} > 0\) and \(\epsilon_{\text{rel}} > 0\) using the relation

\[ \epsilon_{\text{prim}} = \epsilon_{\text{abs}}\sqrt{TK} + \epsilon_{\text{rel}}\max(\|d^k\|_2, \|\tilde{d}^k\|_2) \]
\[ \epsilon_{\text{dual}} = \epsilon_{\text{abs}}\sqrt{TK} + \epsilon_{\text{rel}}\|u^k\|_2. \]

A common choice for \(\epsilon_{\text{rel}} = 10^{-3}\), while the choice for \(\epsilon_{\text{abs}}\) depends on the scale of the treatment planning problem [4, §3.3.1].

6.2.3. Convergence and choice of \(\rho\)

When the problem is convex, i.e. the health dynamics function is affine, Algorithm 6.1 converges to a solution for any \(\rho > 0\), although the value of \(\rho\) may have an impact on the practical convergence rate. When the problem is nonconvex, ADMM is a heuristic and the final beam/dose plan can depend directly on \(\rho\) [4, §9]. The question of how to choose \(\rho\) is still unsettled; see [17,43,44] for further discussion on the topic. We have found that for data on the order of one, values of \(\rho\) between \(10^{-2}\) and \(10^2\) work reasonably well.

6.3. Clinical example

6.3.1. Problem instance

We test our method on a fluence map optimization of a prostate cancer IMRT case with \(n = 34848\) beams and \(K = 7\) structures consisting of a single PTV \((i = 1)\), five OARs, and generic body voxels \((i = 7)\). Treatment is carried out over \(T = 45\) sessions, so the planning problem has about 1.6 million variables. The matrix \(A_T\) remains constant over time and maps the beam intensities to the average dose per structure, i.e. \(d_{t,i}\) is the total dose to structure \(i\) divided by the number of voxels in \(i\). Each beam’s intensity cannot exceed \(B_t = 0.025\).

The LQ model parameters, initial health status, and dose and health status bounds can be found in Table 1; these have been adapted from prior clinical datasets [16,22,26,40]. We choose the health and dose penalty functions to be

\[ \psi_t(h_t) = (h_{t,1})_+ + \frac{1}{6} \sum_{i=2}^7 (h_{t,i})_-, \quad \phi_t(d_t) = \sum_{i=1}^6 d_{t,i}^2 + 0.25d_{t,7}^2, \quad t = 1, \ldots, T. \]

These penalties place greater importance on reducing the health status of the PTV compared to sparing the OARs or generic body tissue.
Table 1. Prostate IMRT problem parameters.

| i  | Structure          | $\alpha_{ij}$ | $\beta_{ij}$ | $\gamma_{ij}$ | $h_{ij}$ | $H_{ij}$ | $D_{ij}$ |
|----|-------------------|---------------|--------------|--------------|----------|---------|---------|
| 1  | Prostate          | 0.15          | 0.05         | 0            | 0        | 5.8579  | 10      |
|    |                   |               |              | $t \leq 28$  |          |         |         |
|    |                   |               |              | 0.0173       |          |         |         |
|    |                   |               |              | $t > 28$     |          |         |         |
| 2  | Urethra           | 1             | 0.2          | 0            | 0        | -4.8    | 10      |
| 3  | Bladder           | 1             | 0.2          | 0            | 0        | -4.8    | 10      |
| 4  | Rectum            | 1             | 0.2          | 0            | 0        | -4.8    | 10      |
| 5  | L. Femoral Head   | 1             | 0.25         | 0            | 0        | -3.0    | 10      |
| 6  | R. Femoral Head   | 1             | 0.25         | 0            | 0        | -3.0    | 10      |
| 7  | Body              | 1             | 0.3333       | 0            | 0        | -6.0    | 10      |

6.3.2. Computational details

The computational setup is the same as in Example 4.2. To solve the ADMM subproblems, we used MOSEK and ran CCP ($\lambda = 10^4$) on the health trajectory subproblem. With $\rho = 80$, ADMM converged in 82 iterations to cutoffs of $\epsilon_{\text{abs}} = 10^{-2}$ and $\epsilon_{\text{rel}} = 10^{-3}$. The normed residuals, $\|r^k_{\text{prim}}\|_2$ and $\|r^k_{\text{dual}}\|_2$, are shown in Figure 6. Total runtime was about 43 minutes, with the bulk of that time spent on the main ADMM loop (initialization took only 32 seconds). By contrast, a straightforward application of Algorithm 4.1 to this problem required over an hour.

6.3.3. Results and analysis

Figure 7 depicts the dose trajectories resulting from the initial plan (green) and the final plan output by ADMM (blue). The initial plan is essentially a piecewise equal-dose fractionation scheme, reflected by the flat plateaus in the corresponding dose trajectories. This
already gives us a good approximation of the final plan: both plans maintain a relatively high dose to the PTV of about 0.9 Gy until session 31, then drop off sharply to the same constant doses thereafter. However, during the high dose phase, the final plan gradually increases the dosage over time to all structures except the bladder \((i = 3)\). By adapting dynamically to changes in the patient’s anatomy, it is able to deliver more dose per session and thus achieve better tumor control, while still respecting the limits on the OARs’ health statuses.

Indeed, we see in Figure 8 that the final plan exactly attains the desired PTV health status of zero for \(t > 31\). It must sacrifice some OARs to do this, reducing the health statuses of the urethra, rectum, and right femoral head \((i = 2, 4, \text{and } 6)\) to their lower bounds, but never violates those bounds. In fact, by shifting radiation to other structures, the final plan actually improves the health of the bladder over that from the initial plan, which results in a \(h_3(t)\) curve far below the limit of \(-4.8\) for \(t \geq 35\). Overall, it is clear that the combination of a solid initialization heuristic and ADMM produces a treatment plan that satisfies or even exceeds all of our clinical goals.

7. Implementation

We provide an implementation of our adaptive radiation treatment planning method in AdaRad, an open-source Python software package based on CVXPY [10]. Our
Figure 8. Optimal health trajectories resulting from the doses in Figure 7. Dashed lines represent the health status bounds $\mathcal{H}_{t,i}$.

The code below imports some patient data and a prescription, solves for the optimal treatment plan, and plots the resulting dose and health trajectories.

```python
import adarad, numpy
from adarad import Case, CasePlotter

# Construct the clinical case.
> case = Case()
> case.import_file("/examples/patient_01-case.yaml")
> case.physics.dose_matrix = numpy.load("/examples/patient_01-dmat.npy")

# Solve using ADMM algorithm.
> status, result = case.plan(slack_weight = 50, max_iter = 100,
    solver = ECOS, use_admm = True)
> print("Solve status: {}".format(status))
> print("Solve time: {}".format(result.solver_stats.solve_time))
> print("Iterations: {}".format(result.solver_stats.num_iters))

# Plot the dose and health trajectories.
> caseviz = CasePlotter(case)
> caseviz.plot_treatment(result, stepsize = 10)
> caseviz.plot_health(result, stepsize = 10)
```

implementation is fully distributed, leveraging Python’s built-in multiprocessing library to execute solves in parallel. Users can quickly import patient data, define clinical goals, construct treatment plans, and visualize the results. They can also rapidly modify and re-plan a case, allowing for comparisons between different prescriptions and treatment lengths. Moreover, since AdaRad is a Python library, it can be easily integrated with other libraries (e.g. for image processing) used in radiation therapy.

The code below imports some patient data and a prescription, solves for the optimal treatment plan, and plots the resulting dose and health trajectories.
In this example, the dose matrix \( A_t \) is the same for all \( t \) and stored in a single \( * .npy \) file. AdaRad also supports other sparse data representations, such as \texttt{scipy.csc_matrix}.

To specify a time-varying dose matrix, the user would input a list of matrices in order \([A_1, \ldots, A_T]\).

We start by constructing a Case, which contains Anatomy, Physics, and Prescription objects. The Anatomy and Physics must be defined prior to planning, either by manually specifying them in the code or importing a case description. A description is a YAML file that contains at minimum the keys treatment_length and structures, where the latter is a list of anatomical structures \( i = 1, \ldots, K \), each of which has a name, \texttt{is_target} boolean indicator, and \texttt{alpha}, \texttt{beta}, and \texttt{gamma} values corresponding to the LQ model parameters. The initial health status and health and dose bounds may also be specified.

Once the Case is defined, we can solve for the optimal treatment plan. The \texttt{plan} function implements Algorithms 4.1 and 6.1 (the latter with use_admm = True). It takes as optional input \texttt{d_init}: the initial dose point, \texttt{use_slack}: a boolean indicating whether to include slack variable \( \delta \), \texttt{slack_weight}: the slack penalty parameter \( \lambda \), \texttt{max_iter}: the maximum number of iterations, and \texttt{solver}: the convex solver to use for the beam and health subproblems. In the above example, we call the solver ECOS [11], one of several free, open-source solvers packaged with CVXPY. If MOSEK is installed, we can call it as well by passing \texttt{solver = MOSEK} into the planning function.

After the algorithm finishes, \texttt{plan} saves the results in \texttt{case.current\_plan} and returns the final solve status along with a \texttt{RunRecord} object that carries solver performance data, such as the total runtime, and the optimal variable values. To visualize the resulting plan, we instantiate a \texttt{CasePlotter} object and call \texttt{plot\_treatment} and \texttt{plot\_health} on the \texttt{RunRecord} to display the dose and health trajectories, respectively. We can also extract the optimal beams, doses, and health statuses with, e.g. \texttt{result.beams} for further processing.

If we wish to explore alternate plans, we can easily modify the dose and health status constraints of any structure and re-plan the case. Re-planning is generally fast, since AdaRad uses the previously stored solution as a warm start point. In a typical workflow, we may import a prescription formed from general clinical guidelines, then repeatedly adjust the dose/health status bounds until we obtain a treatment plan with our desired properties. The \texttt{case.current\_plan} will be updated with the new optimal values after each run. To keep a history of plans for comparison, we can save our results in the \texttt{Case} by calling \texttt{save\_plan} before re-optimizing. The code below provides an example of changing the upper dose bound on the PTV to \( D_{th} = 10 \) Gy for all sessions and plotting the dose and health trajectories under this new constraint alongside the trajectories of the original plan.

```python
# Save previous treatment plan.
> case.save_plan('Original Plan')
# Constraint allows maximum of 10 Gy per session on the PTV.
> case.prescription['PTV'].dose_upper = 10
# Re-plan the case with new dose constraint.
> status2, result2 = case.plan(slack_weight = 50, max_iter = 100,
                                   solver = ECOS, use_admm = True)
> print('Solve status: {}'.format(status2))
# Compare original and new treatment plans.
> caseviz.plot_treatment(result2, stepsize = 10, label = "New Plan",
                          plot_saved = True)
```
> caseviz.plot_health(result2, stepsize = 10, label = "New Plan", plot_saved = True)

For more details on AdaRad’s functions as well as additional examples, see the documentation at https://github.com/anqif/adarad.

8. Conclusion

To achieve the best outcomes, radiation therapy must adapt to new information about the patient’s health and anatomy during treatment. We have described one method for adaptive radiation treatment planning using an operator splitting algorithm. Our method is highly scalable, parallelizable, and can efficiently handle a large number of beams and sessions. Moreover, it is robust to errors in the patient’s health response model, as well as other sources of uncertainty in the clinic. We demonstrated its effectiveness on a large prostate cancer case and showed that the resulting plan improves markedly on a standard equal-dose fractionation scheme.

Future work will focus on expanding our health response model to include sublethal damage repair, redistribution, and reoxygenation effects. We will also incorporate dose-volume constraints into the optimal control problem. Finally, to increase our algorithm’s speed, we intend to release an implementation that takes advantage of the parallel processing capabilities of the GPU.

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