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Mathematical Cancer Therapy Planning in Deep Regional Hyperthermia

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Abstract

This paper surveys the required mathematics for a typical challenging problem from computational medicine, the cancer therapy planning in deep regional hyperthermia. In the course of many years of close cooperation with clinics, the medical problem gave rise to quite a number of subtle mathematical problems, part of which had been unsolved when the common project started. Efficiency of numerical algorithms, i.e. computational speed and monitored reliability, play a decisive role for the medical treatment. Off-the-shelf software had turned out to be not sufficient to meet the requirements of medicine. Rather, new mathematical theory as well as new numerical algorithms had to be developed. In order to make our algorithms useful in the clinical environment, new visualization software, a virtual lab, including 3D geometry processing of individual virtual patients had to be designed and implemented. Moreover, before the problems could be attacked by numerical algorithms, careful mathematical modelling had to be done. Finally, parameter identification and constrained optimization for the PDEs had to be newly analyzed and realized over the individual patient’s geometry. Our new techniques had an impact on the specificity of the individual patients’ treatment and on the construction of an improved hyperthermia applicator.

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Introduction

The term *hyperthermia* in cancer medicine means a heat treatment of tumour tissue. Already in 1961, P. Scheid had detected “functional specialties in the microcirculation of carcinomas” under the influence of heating, see [81]. In an early attempt, Manfred von Ardenne (1907–1997) had experimented with what is today called full-body hyperthermia: he immersed cancer patients in a hot bath or exposed them to infrared radiation. However, his medical results were unclear: in rare cases, tumour growth was reduced, whereas in most cases there was no effect at all. Today such a procedure is only considered for cancer patients with widespread micro-metastases that cannot be detected by medical imaging. Rather, *regional* hyperthermia evolved as an option. Herein, tumours are only locally heated, while healthy tissue is generally saved. With an increasing understanding about the generation and function of heat shock proteins a lower temperature bound of roughly 42.5°C appeared to be advisable. An upper temperature bound in the tumour should be “not too high” to assure that healthy tissue would be kept close enough to the body temperature 37°C and the systemic body stress remains tolerable. At present, hyperthermia is applied in two typical modes:

- **Downstaging:** Hyperthermia treatment alone can produce sharper boundaries of the tumour, thus facilitating operative tumour extinction by the oncosurgeon – an effect that helps to reduce the possible later occurrence of a relapse of the operated tumour.

- **Combination therapy:** In principle, one might think of a strategy where heat alone just “burns” the tumour which then would be secreted by the kidneys as necrotic tissue. However, as long as healthy tissue cannot be guaranteed to be preserved unburnt in such a process, the therapeutic temperature level must remain moderate. In a temperature window between 41°C and 45°C, so called deep regional hyperthermia (as opposed to surface hyperthermia, e.g., for extremities) is usually combined with chemo- and radiotherapy, see, e.g., R. D. Issels [55]. In fact, testing hyperthermia alone in a research situation, where other therapies like chemo- or radiotherapy might help to cure the patient, is ethically not justified. That is why the actual cases open to our simulation studies were only deep-seated non-operable tumours.

Already in 1984, it became clear that a careful hyperthermia treatment planning required computer simulations, see, e.g., R. B. Roemer [78]. In the late 1980’s, the physician (and, at the same time, physicist) P. Wust came to ZIB seeking advice about the fast numerical solution of a PDE called the bio-heat transfer equation that turned out to be of Helmholtz type in the uncritical case. His interest was in 2D computations, since the input from individual patients were 2D cross sections out of a pile of images. In the beginning, compare [108, 109], he got the recommendation to use the 1985 version of the 2D elliptic solver PLTMG by R. Bank [6]. However, the 2D calculations appeared to be
in strong contradiction with what medical radiologists encountered: various patients complained about “hot spots” quite distant from the tumour to be heated, whereas the simulations did not show any such phenomena. It was the mathematician who suggested that patients should be regarded as 3D! As a consequence, the new version of the 3D elliptic solver KASKADE \[29,13,42\] came into play. This started a close collaboration between the numerical analysis group at ZIB and the radiology department at Charité, a collaboration that continued over more than a decade, see, e.g., \[110,90\]. In the course of time, nearly all mathematical and computer science tools necessary to tackle the challenging hyperthermia problems were put on a test bench. As a result, all pieces were redesigned until finally the whole computational line appeared to be fast, robust, and reliable enough to be used in a clinical environment. For this purpose, an integrated software, the virtual lab HyperPlan, was elaborated (with eventually about half a million lines of code). It included a selection of both visualization and numerical algorithms for the construction of individual 3D virtual patients.

From the numerical side, it contained adaptive multilevel finite element methods for the parabolic bio-heat transfer (BHT) equation coupled with the time-harmonic heterogeneous Maxwell’s equation, the latter in the then still underrated intermediate regime where the wavelength is of the size of the obstacles. For the BHT equation, the hierarchical basis preconditioner of H. Yserentant \[114,29\] for 2D as well as the BPX preconditioner of J. Xu \[111,17\] (in the variant due to \[115,13\]) for 3D appeared just at the right time. For the time-harmonic Maxwell’s equation in the radiowave regime, the key turned out to be the use of (a) Whitney forms (see \[106\]), the lowest order 3D finite edge elements (see J. C. Nédélec \[70\], A. Bossavit \[14\], R. Beck \[7\]) to avoid spurious discrete solutions and (b) the discrete Helmholtz decomposition due to R. Hiptmair \[49,10,9,52\] within multigrid methods. In 2000, a first survey on this kind of work was given in \[26\]. These PDE solvers had been coupled to compute an optimal tuning of the antenna parameters. As a first step, this has been modelled by an unconstrained minimization problem where the medically necessary heating constraints were taken into account via some penalty term.

Unfortunately, the DFG funding of the ZIB–Charité cooperation (within the collaborative research unit SFB 273) terminated at the end of 2002. Fortunately, the topic of regional hyperthermia could be further pursued within the DFG Research Center MATHEON, now on a more mathematical basis. The above unconstrained minimization problem with penalty term was substituted by the medically more reasonable constrained minimization problem where the heating constraints were modelled as strict inequalities. At the time when this substitution was envisaged, the associated mathematical topic (adaptive multilevel methods for PDE constrained optimization) was far from mature enough to be used with medical reliability. It is the purpose of the present survey to report about progress made since then in the context of what has been achieved before.
Outline. The paper is organized as follows. In Section 1 we give the basics of the virtual lab HyperPlan. This software environment fixes all patient-specific details necessary to set the hyperthermia problem on stage: individual geometry, material coefficients, electromagnetic antennas at the applicator, water bolus. After that, in Section 2 the medical problem is transferred into a mathematical model in terms of partial differential equations (PDEs), a Helmholtz equation in the uncritical case for the distribution of heat in the body and the time-harmonic Maxwell equation, which is structurally similar to a Helmholtz equation in the critical case. The therapeutic goal is transformed into a variety of mathematical optimization problems of increasing difficulty, starting from a traditional functional with penalized temperature constraints and continuing to a medically preferable functional with PDE equality constraints and strict temperature inequality constraints. On this basis, Section 3 discusses efficient numerical techniques to solve the PDEs. Section 4 presents the recent theoretical analysis and algorithmic progress for PDE constrained optimization. An interior point method is derived and worked out as a numerical pathfollowing in terms of a barrier parameter. As it turns out, the log-barrier function, well-known from finite dimensional optimization, needs to be replaced by a hierarchy of rational barrier functions in function space optimization. In the last Section 5 the impact of the mathematical developments over years with respect to clinical application is surveyed.

1 Patient-Specific Virtual Lab

The many years of common work between ZIB and Charité in the hyperthermia project caused a learning effect that paid off in all later collaborations between mathematics and medicine. That is why we start with a description of the basic pattern (paradigm) and the patient-specific 3D grid generation.

1.1 Paradigm of Virtual Medicine

For the cooperation between medicine and mathematics the following three-step procedure has evolved, compare Fig. 1.

(I) Construct an individual virtual patient: The construction of a patient-specific geometrical 3D model starts from a stack of 2D images originating from, e.g., CT, MRT, or ultrasound. In a first computational step, 3D details of the patient’s body need to be identified, a procedure called segmentation. In order to obtain reasonable starting grids for a multigrid PDE solution, surface triangulation of interfaces, coarsening of surface grids, and finally tetrahedral grid generation are successively performed.

(II) Model and simulate the treatment in a virtual lab: The modelling of the therapy in question consists of a description of the essential physiological

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1Patient had agreed to be shown recognizably within scientific context.
and physical processes – as accurate as needed for the medical problems to be tackled. Usually, the description consists of a set of partial differential equations (PDEs) and, in the best case, a functional to be minimized under constraints. In order to suggest an “optimal” treatment planning, these PDEs need to be solved fast and reliably. Moreover, in order to be of use as a basis for medical decision, any suggested therapy requires careful visualization. All this must be realized on a hardware and software platform that can be made available in a clinical environment.

(III) Transfer the simulation results back to the clinical environment: In some prephase, the model and its simulation will be validated against the clinical situation. Any discrepancies between model and reality need to be discussed and, if necessary, taken into account by remodelling. Once this adaptation has been done, the computationally suggested “optimal” therapy is transferred back to patient-specific treatment in the clinical environment.

Figure 1: Hyperthermia treatment planning. Left: real patient in hospital. Right: virtual patient in the virtual lab.\(^1\)

The above step (I) is essentially independent of the special medical problem, but only dependent on the body region treated, see Section 1.2 below. Step (II) depends on the medical problem in question, which here is deep regional hyperthermia, see subsequent Sections 2, 3.1 and 4. Step (III) will be illustrated by many examples of virtual patients inserted in the text throughout the paper.

1.2 Construction of Geometrical 3D Model

The above paradigm has been implemented in the virtual lab HyperPlan \([92, 26]\), an integrated software system that has been especially developed to support hyperthermia treatment Planning in a clinical environment. The implementation of roughly half a million lines of code made heavy use of C++-software tools like OpenGL (graphic library) and OpenInventor (object oriented class library) which are of high algorithmic complexity themselves.
HyperPlan combines

- efficient 3D grid generation techniques, which we will describe now,
- fast and reliable simulation techniques, e.g., adaptive multilevel finite element methods, see Section 3.1 below, and
- efficient visualization techniques, which are beyond the scope of the present article (for reference see, e.g. [46, 91]).

Figure 2: Medical imaging input. Left: Stack of 2D computer tomograms (CTs). Right: Segmented raw image data.

Medical imaging input. Patient-specific geometrical input usually enters from medical imaging techniques like CT, MRT, ultrasound or mixtures thereof. We skip the case where different sources of input have to be matched (so-called registration problem, see, e.g., [113]). As shown for CT in Fig. 2, this input comes as a stack of 2D images that merely give mass density information. In view of the solution of PDEs on the specific patient data, we need more than just density information, see below under segmentation. Moreover, for the purpose of presenting numerical results to medical doctors, a 3D grid generation with clear material identification needs to be realized. This is done in several steps which are described next.

Segmentation. As can be seen from Fig. 2 the original CT input contains only information about the mass density distribution in planar cross sections of the body. What we need, however, to solve PDEs on an individual patient, is a

\footnote{More precisely, it gives material property information that is mapped to intensity values.}
clear correspondence of the various body parts with material properties such as electromagnetic or thermal constants. This task is called segmentation. From computer science, several segmentation techniques such as region growing [1] or watershed methods [99] are quite popular. They typically work on the stack of 2D cross sections and, in the best case, exploit some connection between neighbouring 2D images. Such techniques usually require a significant amount of user interaction which is not desirable. Therefore, at ZIB, we followed a rather different approach called statistical shape analysis [22], which transverses through 3D. In principle, this method performs the following steps:

1. Start from a sample set of carefully segmented patient data, a time-consuming interactive job done by experts. For illustration, see the eight pelvic regions ordered along a circle in Fig. 3, left.

2. Compute an “average” shape. See the centre shape in Fig. 3, left. In order to compute a theoretically satisfactory average, one must solve the associated correspondence problem between two surfaces, see, e.g., [61] wherein the way is paved towards a “variational shape analysis”.

3. Compute the differences “individual - average” as fluctuations.

4. Select “essential coordinates” by selecting the fluctuations with the help of PCA (principal component analysis).

5. Whenever new patient data are to be segmented, try to just compute the essential coordinates for the new data, see Fig. 3, right. If this works to sufficient accuracy, then exit. Else enrich the patient data basis by these new data and go to step 1.

In the beginning of the above procedure, the number of essential coordinates will grow with every new patient data, but after a while it will get saturated, thus...
defining the “essential dimension” of the patient data set at hand. Of course, in order to keep this dimension low, patient data should be clustered according to criteria like sex, age, sports activities, biological group etc. Fig. 3 illustrates such a 3D segmentation for some pelvic region. All in all, the techniques summarized here have reached a state of maturity that makes them equivalent in quality to a “second human radiologist, see the award-winning paper [56] by Kainmueller, Lange, and Lamecker - from an international contest on liver segmentation.

Surface triangulation coarsening. Typically, meshes generated after segmentation are much too fine, for an illustration see Fig.1 left. For the purpose of visualization, this large number can be handled, but in view of the numerical solution of PDEs on the virtual patients (e.g., by multigrid methods), this number is far too large. For this reason, some precoarsening of the grids is needed. The associated iterative process is based on minimizing local approximation errors of coarser triangulated surfaces, which is closely related to local curvature: Wherever the local curvature is “low”, the number of local nodes is reduced, whereas it is left unaltered where the local curvature is “high”, see, e.g., [45]. In Fig. 4 the coarsening procedure (including mesh quality control) due to Zilske, Lamecker and Zachow [117] is illustrated for some pelvic bone region. For multigrid methods to be discussed below in Section 5.1 this kind of grid then serves as the initial grid.

Figure 4: Grid coarsening in the pelvic region. From 478.000 nodes via 98.000 nodes to 11.000 nodes.

Tetrahedral grid generation. As a final step, the interface triangular grids are used as starting point to construct spatial tetrahedral grids in the full volume. This is done by a special adaptation of the “advancing front” algorithm of Löhner/Parikh [64]. A more recent improvement is the remeshing technique due to Zilske, Lamecker and Zachow [116]. This final step is illustrated in Fig. 5 right.
Remark 1. HyperPlan was the predecessor of the popular system Amira [93] which applies to general systems in science and medicine.

2 Mathematical Modelling

The purpose of this section is to describe the mathematical modelling necessary to translate the medical treatment planning problem into a mathematical formulation accessible to numerical methods. It covers the physics of heat generation by radio-wave emission in the antennas of the applicator, which is modelled by the time-harmonic Maxwell equation, and the transport of heat inside the human body, which is modelled by some bio-heat transfer equation. Moreover, it includes the identification of unknown parameters in the mathematical models and several choices of quantitative formulations of the therapeutic goal, which leads to optimal control problems that are dealt with in the subsequent Section 4.

Hyperthermia treatment of a real hospitalized patient is depicted in the left hand side of Fig. 1. As the patient has a quite large rectum carcinoma, his abdomen is lying inside a so-called applicator which contains eight antennas (pairwise coupled). The antennas emit radiowaves of frequency 100 MHz which are absorbed in the human body thus generating heat. The associated mathematical model, the time-harmonic Maxwell equation, is discussed in Section 2.1. Between the applicator and the body is a so-called water bolus to ease the entry of the radiowaves into the body and, at the same time, to cool the skin – thus setting a temperature value at the body surface. The absorption of the electromagnetic radiation is different for different body compartments (essentially
depending on the portion of water in the tissue). By virtue of blood flow as well as heat conduction, the heat is distributed in the body. This process is modelled mathematically by a parabolic PDE, the so-called bio-heat transfer equation, see Section 2.2. A solution of the coupled system of PDEs is a necessary prerequisite to compute an optimal tuning of the antenna parameters such that the tumour is heated (within a temperature window), but not healthy tissue. This leads to a formulation in terms of mathematical optimization problems to be elaborated in Section 2.4.

2.1 Emission of Radiowaves

From first principles, the radiowaves emitted by the applicator antennas are known to be governed by Maxwell’s equations. Simplified models are available for the case when the wavelength is much larger or much smaller than the geometry of the objects. In the hyperthermia case, however, the wavelength in water (which is a good approximation for the material of the body) is about 30 cm, which is the order of magnitude of the human body (the “obstacle”), i.e. just in the intermediate regime. As a consequence, there is no way round solving the full Maxwell equations for the heterogeneous medium “water bolus - human body”.

**Time-harmonic Maxwell equation: strong formulation.** Fortunately, only a single angular frequency \( \omega \) is emitted from the antennas, which means that we can make the time-harmonic ansatz

\[
E(x,t) = \Re(E(x)e^{-i\omega t}), H(x,t) = \Re(H(x)e^{-i\omega t}),
\]

for the electromagnetic fields \( E, H \in \mathbb{C}^3 \). Insertion into the time dependent Maxwell equations for linear isotropic dielectric media,

\[
\begin{align*}
\text{div}(\mu H) &= 0 \quad \text{(1a)} \\
\text{div}(\epsilon E) &= 0 \quad \text{(1b)} \\
\text{curl} \, H &= \sigma E + \epsilon \frac{\partial E}{\partial t} \quad \text{(1c)} \\
\text{curl} \, E &= -\mu \frac{\partial H}{\partial t}, \quad \text{(1d)}
\end{align*}
\]

applying the curl operator to (1d), and inserting (1c) yields the so-called *double-curl equation* or *time-harmonic Maxwell equation*

\[
\text{curl} \, \frac{1}{\mu} \text{curl} \, E = \epsilon' \omega^2 E \quad \text{(2)}
\]

for the complex electrical field \( E \). Herein \( \mu \) is the magnetic permeability and \( \epsilon' = \epsilon + i\sigma/\omega \) the complex dielectric constant related to Ohm’s law \( j = \sigma E \) with electric conductivity \( \sigma \geq 0 \) for any current density \( j \) and the generic dielectric constant \( \epsilon > 0 \). The material constants \( \mu, \epsilon, \sigma \) depend, as the name indicates,
Figure 6: Antenna modelling with Dirichlet boundary conditions.

on the local materials (e.g. muscle, fat, bone etc.) and should therefore, more precisely, be written as $\mu(x), \epsilon(x), \sigma(x)$. They are allocated on the basis of the geometrical patient model, see the above Section 1.2.

For spatially constant material parameters (homogeneous material), the term on the left side may be reformulated to lead to

$$- \Delta E + \nabla \text{div} E = \mu \epsilon' \omega^2 E.$$  \hspace{1cm} (3)

Herein the Laplace operator $\Delta$ is applied componentwise, such that (3) represents three decoupled Helmholtz-type equations in the critical case. Otherwise, with material jumps, the three PDEs are coupled, see Section 3.1.

**Boundary conditions.** The computational domain of interest, see Fig. 1 right, consists of an artificially chosen sphere coating the water bolus, the antenna array, and the virtual patient model. Hence we have three types of boundary conditions to be imposed. First, the metallic antennas on the physical boundary of the applicator are regarded as perfect conductors, which implies homogeneous Dirichlet boundary conditions for the tangential components $E_t = (n \times E) \times n$ of the electric field $E$, i.e.

$$E_t = 0 \quad \Leftrightarrow \quad n \times E = 0 \quad \text{on} \quad \Gamma_{\text{ant}}. $$

Herein $\Gamma_{\text{ant}}$ denotes the thin metallic antennas shown in Fig. 1 and $n$ the outer unit normal of the surface $\Gamma_{\text{ant}}$. Second, the small gap between the trapezoidal antenna sheets is modelled by an artificial boundary $\Gamma_{\text{gap}}$ on which the tangential component of $E$ is prescribed such as to yield the voltage of the power generator when integrated across the gap:

$$E_t = E_t^0 \quad \text{on} \quad \Gamma_{\text{gap}}. $$

Third, on the external sphere $\Gamma_{\text{ext}}$, the Sommerfeld radiation condition (also...
named Silver-Müller condition in the electrical engineering community)

\[
\mathbf{n} \times \left( \frac{1}{\mu} \text{curl } E \right) - \mathbf{n} \times \left( \frac{i\omega}{\mu c} \mathbf{n} \times E \right) = 0 \quad \text{on } \Gamma_{\text{ext}}
\]  

(4)
is prescribed as an approximate transparent boundary condition in order to avoid reflection at the artificial boundary \(\Gamma_{\text{ext}}\). Herein \(c = 1/\sqrt{\varepsilon \mu}\), the electromagnetic wave velocity in the dielectric material, has been inserted.

**Time-harmonic Maxwell equation: weak formulation.** The weak formulation starts from the double-curl equation (2) directly. For lossy materials (i.e. for \(\sigma > 0\)), the material constant \(\epsilon'\) is complex. As a consequence, the above PDE with boundary conditions represents a non-selfadjoint operator. Then, following [10], we introduce the appropriate Hilbert space for the operator equation as

\[
H_{\Gamma}(\text{curl}; \Omega) := \{ v \in L_2(\Omega)^3; \text{curl } v \in L_2(\Omega)^3, \mathbf{n} \times v = 0 \text{ on } \Gamma_{\text{ant}} \cup \Gamma_{\text{gap}}, \mathbf{n} \times v \in L_2(\Gamma_{\text{ext}})^3 \text{ on } \Gamma_{\text{ext}} \},
\]
equipped with the scalar product

\[
\langle v, w \rangle_{H(\text{curl})} := \langle \text{curl } v, \text{curl } w \rangle_{L_2(\Omega)} + \langle v, w \rangle_{L_2(\Omega, \Gamma_{\text{ext}})}
\]
and a sesquilinear form \(a : H_{\Gamma}(\text{curl}; \Omega) \times H_{\Gamma}(\text{curl}; \Omega) \rightarrow \mathbb{C}\) (conjugate linear in the first component):

\[
a(E, v) := \int_{\Omega} \left( \frac{1}{\mu} (\text{curl } E)^* \text{curl } v - \omega^2 (\epsilon' E)^* v \right) dx
\]

\[
- \int_{\Gamma_{\text{ext}}} \frac{i\omega}{\mu c} (\mathbf{n} \times E)^* (\mathbf{n} \times v) ds.
\]

(5)
We end up with the following variational problem: find \(E \in H_{\Gamma}(\text{curl}; \Omega)\) such that

\[
a(E, v) = f(v) \quad \forall \ v \in H_{\Gamma}(\text{curl}; \Omega),
\]

(6)
where \(f : H_{\Gamma}(\text{curl}; \Omega) \rightarrow \mathbb{C}\) is a linear form containing the inhomogeneity of the Dirichlet conditions.

**Properties of (6).** In order to study the question of uniqueness of a solution, we follow the usual procedure, see [69, Theorem 4.12]. Suppose there exists some \(\bar{E}\) such that \(a(E, \bar{E}) = 0\), which implies that both its real and its imaginary part vanish. We assume homogeneous, possibly lossy material, i.e. \(\sigma \geq 0\). For the imaginary part we get

\[
-\Im a(\bar{E}, \bar{E}) = \omega \left(\sigma \int_{\Omega} |\bar{E}|^2 dx + \frac{1}{\mu c} \int_{\Gamma_{\text{ext}}} |\mathbf{n} \times \bar{E}|^2 ds \right).
\]

Both integrals are positive. For lossless material \((\sigma = 0)\), the first term vanishes. The second one is positive, unless the external boundary integral vanishes, which
would be the case for homogeneous Dirichlet or Neumann conditions. For the radiation condition (4), however, this is not the case. For lossy material ($\sigma > 0$), both terms are positive, which leads to a contradiction to the assumption that they should vanish. If a solution of (5) exists at all, it is unique whenever $\sigma \geq 0$.

Two further features of (5) deserve special attention: (a) the first term in the volume integral, representing the principal part of the PDE, has an “ample” nullspace

$$H^0_T(\text{curl}; \Omega) = \ker \text{curl}$$

containing all gradient-functions, in particular also functions of high frequency, (b) in this nullspace, the second term in the volume integral, representing the zero order term in the PDE, is negative definite, since the factor $-\omega^2\Re \epsilon' = -\omega^2\epsilon$ is negative. Thus, the functional $a$ is highly indefinite. The indefiniteness is the more marked, the higher the angular frequency $\omega$, a fact clearly given in hyperthermia (with 100 MHz).

On simply connected Lipschitz domains $H^0_T(\text{curl}; \Omega)$ gives rise to a regular decomposition of $H^1_T(\text{curl}; \Omega)$ (cf. [51]):

$$v = \phi + \text{grad} \, p, \quad \phi \in H^1(\Omega)^3, \quad p \in H^1(\Omega), \quad \text{grad} \, p \in H^0_T(\text{curl}; \Omega) \quad (7)$$

such that

$$\|\phi\|_{H^1} + \|p\|_{H^1} \leq C\|v\|_{H(\text{curl})}. \quad (8)$$

In this way, our problem is split into a vectorial $H^1$-part and a negative definite $H^1$-component for the potentials. This decomposition is a modification of the classical Helmholtz decomposition.

**Interface conditions at material jumps.** Both quantities $\mu$ and $\epsilon$ have jumps at material interfaces that have been determined by segmentation, see Section 1.2 above. At these interfaces special conditions must be observed, which in weak formulation are automatically satisfied (via the therein incorporated vanishing of the remainder terms stemming from an integration by parts). These conditions imply a kink in the electric field vector $E$ at the boundaries, for an illustration in FastLIC representation [46] see Fig. 7.

**Superposition.** The radiowaves (100 MHz) are emitted from $k_E$ channels of pairwise coupled $2k_E$ antennas. Since the Maxwell equations are linear, superposition can be applied. Let $E_k(x)$ denote the electric field (the antenna profile) originating from channel $k$ for $k = 1, \ldots, k_E$. Then the total electrical field can be written as a linear combination

$$E(x) = \sum_{k=1}^{k_E} \alpha_k E_k(x), \quad \alpha_k \in \mathbb{C}, \quad (9)$$

in terms of complex amplitudes $\alpha_1, \ldots, \alpha_{k_E}$. They may be parametrized via

$$\alpha_k = a_k \exp(-i\theta_k) \quad k = 1, \ldots, k_E, \quad (10)$$

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with real amplitudes $a_k$ and phase delays $\theta_k$ that can be controlled independently. Where appropriate, the $k_E$ complex parameters $\alpha_k$ will be interpreted as $2k_E$ real parameters $\{\Re\alpha_k, \Im\alpha_k\}$.

**Remark 2.** In [58], Kremer and Louis treated the problem of approximating a given electrical field (obtained, e.g., from optimization) via excitation through $n$ antennas. For their theoretical analysis, they define a hyperthermia operator that maps arbitrary antenna excitations at a boundary circle of a cylinder to any electrical fields in that cylinder. This operator is shown to be compact in $L_2$. By means of singular value decomposition they can conclude that hot spots outside the tumour volume can only be suppressed for sufficiently large $n$.

### 2.2 Heat Distribution inside the Human Body

Compared to the generation of heat as discussed in the preceding section, the distribution of heat inside a specific patient’s body is much harder in terms of modelling.

**Heat absorption.** Inside the body, the electromagnetic waves generate heat by absorption to be characterized via the absorption rate density

$$S = \frac{1}{2} \sigma |E|^2_{C_3},$$

(11)

wherein $\sigma$ is the electric conductivity of the specific tissue. If one exploits the superposition (9), one ends up with the quadratic term

$$S(x) = \frac{1}{2} \sigma(x) |E(x)|^2_{C_3} = \frac{1}{2} \sigma(x) \sum_{j,l=1}^{k_E} \alpha_j^* \alpha_l E_j^*(x) E_l(x),$$

(12)
wherein $\sigma$ varies among different tissues. From the definition (12) it is apparent that simultaneous phase shifts of the controls $\alpha_k$ lead to a corresponding phase shift of the electrical field $E$ and thus have no thermal impact at all. Therefore, one of the phases can be fixed a priori, e.g. by setting $\Im(\alpha_k) = 0$. We will therefore think of $\alpha$ containing only $2k_E - 1$ real parameters, which will be important in Section 4.1 to get rid of a trivial local non-uniqueness of optimal solutions.

**Bio-heat-transfer (BHT) equation: strong formulation.** The standard heat transfer model used in our simulations has been the so-called bio-heat-transfer (BHT) equation over the domain $\Omega \subset \mathbb{R}^3$ covered by the patient’s body:

$$\rho c \frac{\partial T}{\partial t} = \text{div}(\kappa \text{grad} T) - c_b w (T - T_{\text{blood}}) + S \quad \text{in } \Omega, \quad (13)$$

where $\rho$ is the density, $c$ the specific heat capacity of tissue, $\kappa$ the thermal conductivity, $c_b$ the specific heat capacity of blood, $w$ the mass flow rate of blood per unit volume of tissue, the so-called perfusion, $T_{\text{blood}}$ the blood temperature in the unheated body (e.g. 37°C), and $S$ the absorption rate density as defined in (12). All material constants are understood to be local constants, different in different tissues. Already in 1948, this Helmholtz-type equation in the uncritical case was proposed by Pennes [74] on the basis of theoretical consideration and experimental measurements – a rare scientific combination!

The boundary conditions are of Robin type:

$$- \kappa \frac{\partial T}{\partial n} = h(T - T_{\text{bolus}}) \quad \text{on } \partial \Omega. \quad (14)$$

For the time dependent case initial conditions

$$T(0, x) = T_a$$

are prescribed. In hyperthermia, the steady state is typically reached after about 15 – 20 minutes and maintained for 40 – 60 minutes. For this reason, we may well be content with solving the stationary elliptic problem rather than the time dependent parabolic problem.

As with the other material coefficients, the discontinuity of the thermal conductivity here also yields interface conditions across the tissue interfaces. Again, these conditions are automatically satisfied, if the PDE is solved in its weak form.

**BHT equation: weak formulation.** Here we interpret the stationary BHT equation (with notation as in (13)) as an operator equation:

$$\langle A(T) - B(\alpha), \varphi \rangle = 0 \quad \forall \varphi \in C^\infty(\Omega), \quad (15)$$
where \( A \) and \( B \) are defined as
\[
\langle A(T), \varphi \rangle := \int_{\Omega} \kappa (\text{grad } T)^T \text{grad } \varphi + c_b w (T - T_{\text{blood}}) \varphi \, dx \\
+ \int_{\partial \Omega} h(T - T_{\text{bolus}}) \varphi \, dS,
\]
\[
\langle B(\alpha), \varphi \rangle := \int_{\Omega} \sigma_2 |E(\alpha)|^2 C_3 \varphi \, dx.
\]

The domain \( \Omega \) consists of a number of subdomains \( \Omega_i \), corresponding to various types of tissue. All coefficients depend on the tissue type and are chosen piecewise constant on each subdomain associated with a tissue type. They may vary significantly from tissue to tissue. For a more detailed description of the parameters we refer to [33].

**Multiscale heat transport.** Inside the patient’s body, heat is distributed only by diffusion in tissue and by direct transport in blood vessels (with velocity \( v(x) \), say), but not by any process described by the zero order term in the BHT equation (13) above. Therefore, from first principles, a PDE of diffusion-advection type including the source term seems to be more appropriate:
\[
\rho c \frac{\partial T}{\partial t} = \text{div } (\kappa \text{grad } T) - \rho c v^T \text{grad } T + S,
\]
(17)

However, due to the multiscale structure of the velocity field \( v \), both from the modelling aspect and from the point of view of computational complexity, such a PDE cannot be used to successfully describe the situation within an individual patient.

The multiscale vascular situation is schematically represented in Fig. 8. Three scales can be distinguished:

1. On the macroscale, large blood vessels should be treated as full 3D objects (e.g., as objects in a finite element setting, compare also Lagendijk et al. [60]).
2. On some mesoscale, smaller vessels could be modelled as 1D objects (but with some finite width, compare \([23, 75, 66]\)).
3. On the microscale, detailed phenomena within the capillaries cannot be resolved, both from a modelling and a computational perspective. That is why some kind of averaging has to take place which results in a macroscale model for some porous medium region.

In [108], the above BHT equation had been suggested as the averaged microscale model. It had been derived by some potential flow assumption in view of Darcy’s law – known to arise from homogenization techniques in reservoir simulation. That assumption was certainly not self-explaining in the present
context. Therefore we directly turn to the mathematical homogenization approach (see, e.g., Bensoussan et al. [12]) that has been worked out by Hochmuth and Deuflhard [28, 54] for the case of hyperthermia.

In the homogenization framework, we consider a periodic arrangement of microscopic cubic cells $\varepsilon Y$, periodic along the three spatial coordinate axes $y_1, y_2, y_3$, with insertions of blood $\varepsilon Q$, the capillaries, assumed to have constant temperature $T_{\text{blood}}$, see Fig. 9. Let $\Omega_\varepsilon$ denote the solid tissue volume without the capillaries. Therein heat transfer takes place only by diffusion. For simplicity, the authors in [28] scaled the thermal conductivity to $\kappa = 1$. As a consequence, the temperature $T_\varepsilon$ is defined by the simple model equation

$$-\Delta T_\varepsilon = S_\varepsilon \quad \text{in } \Omega_\varepsilon$$
with thermal source term \( S_\varepsilon = \frac{1}{2}\sigma_\varepsilon |E_\varepsilon|^2 \), exterior Dirichlet boundary conditions
\[
T_\varepsilon = T_{\text{bas}} \quad \text{on } \partial \Omega_\varepsilon ,
\]
in terms of some basic temperature \( T_{\text{bas}} \) and interior Robin boundary conditions
\[
\frac{\partial T_\varepsilon}{\partial n} = \varepsilon \alpha (T_{\varepsilon \text{blood}} - T_\varepsilon) \quad \text{on } \partial (\varepsilon Q) .
\]
Due to [28], the following asymptotic result can be shown to hold.

**Theorem 2.1.** Notation as just introduced. Let \( |\cdot| \) denote the Lebesgue measure and define relative surfaces and volumes with and without blood by
\[
\theta_{\text{surface}} := \frac{|\partial Q \setminus \partial Y|}{|Y \setminus Q|} , \quad \theta_{\text{volume}} := \frac{|Y \setminus Q|}{|Y|} < 1 .
\]
Assume that, in weak* -convergence, we have
\[
S_\varepsilon \rightharpoonup \theta_{\text{volume}} S , \quad T_{\varepsilon \text{blood}} \rightharpoonup \theta_{\text{volume}} T_{\text{blood}} .
\]
Then, for \( \varepsilon \to 0 \), the temperature converges weakly as
\[
T_\varepsilon - T_{\varepsilon \text{blood}} \to \theta_{\text{volume}} (T_0 - T_{\text{blood}}) \quad \text{in } L_2 ,
\]
where the limit temperature \( T_0 \) satisfies the PDE
\[
- \text{div} (A \text{grad } T_0) + \alpha \theta_{\text{surface}} (T_0 - T_{\text{blood}}) = S \quad \text{in } \Omega
\]
with Dirichlet boundary conditions
\[
T_0 = T_{\text{bas}} \quad \text{on } \partial \Omega
\]
and perturbed conductivity
\[
A = a_{ij} , \quad a_{ij} = \delta_{ij} - \frac{1}{|Y \setminus Q|} \int_{Y \setminus Q} \frac{\partial \chi^j}{\partial y_i} dy ,
\]
where the functions \( \chi^j \), \( j = 1, 2, 3 \) are understood to be solutions of
\[
- \Delta \chi^j = 0 \quad \text{in } Y \setminus Q , \quad \frac{\partial (\chi^j - y_j)}{\partial n} , \quad \chi^j \quad Y - \text{periodic} . \tag{18}
\]
The proof is a bit technical, mainly because of the occurrence of Robin boundary conditions (for \( T_{0 \text{blood}} = 0 \) a proof has been given earlier, see, e.g., the book [2] and references therein). It starts from the above diffusion-advection equation \( (17) \) assumed locally in a microscale periodic cell structure which is spatially averaged. Details are left to [28]. Summarizing, the above theorem confirms the validity of the BHT equation in the microvascular regime – apart from some perturbation factors \( \theta_{\text{surface}}, \theta_{\text{volume}} \approx 1 \) that take care of the relative blood volume versus the full volume.

**Remark 3.** An alternative approach aiming at directed advective heat transport in tumours has been developed by Gerber [40], but not validated.
Temperature dependent perfusion. In a more detailed investigation, temperature induced vasodilation leads to an increased perfusion of heated tissue and thus gives rise to a more realistic, nonlinear version of the BHT equation. Experiments [89] have shown that the blood flow in normal tissues, e.g., skin and muscle, increases significantly when heated up to 41–43°C, whereas in the tumour zone the blood flow may increase or decrease with temperature depending on the type of tumour. On this experimental basis, the perfusion \( w = w(x, T) \) can be chosen monotonically increasing with respect to \( T \) in the muscle and fatty tissue, but monotonically decreasing in tumour tissue.

Simulations show significant qualitative differences between the temperature distributions predicted by the linear and the nonlinear heat transfer model, as illustrated in Fig. 10. Generally speaking, the self-regulation of healthy tissue reflected by the nonlinear model reduces “hot spots” caused by local maxima of the absorbed electromagnetic fields and hence allows to rise the total energy deposition. This is one reason for a slightly better tumour heating (ca. 0.5°C) predicted by the nonlinear model. An analogous result is reported in [97] for ferromagnetic thermoseed hyperthermia. An important finding is that the nonlinear model has an impact on the optimal treatment parameters as well. Maximal discrepancies turned out to be 22° for the phases \( \theta_k \) and 0.22 for the relative amplitudes \( a_k \) — for the notation see (10). For a more detailed discussion see [33, 63].

Figure 10: Optimized temperature distributions in a frontal section of the pelvic region. Black lines: body outline and tumour contour. Light grey to dark grey shading: regions heated from 39°C to 43°C. Left: linear model. Right: nonlinear model.

Besides local vasodilation, systemic thermoregulation can play an important role, e.g., via centrally controlled vasodilation of the skin and a corresponding shift of perfusion from the body core to the extremities, or an increased heart
rate. A feedback mechanism observed rather recently is the so-called **steal effect** where different regions compete for a joint blood supply. An increased perfusion in one region, e.g., a heated muscle that actively reduces its peripheral resistance by vasodilation, will lead to a larger blood flow and hence a greater decrease of effective blood pressure along the vessels feeding both regions. In this case, another region with less ability of thermoregulation, e.g. a tumour, will experience reduced perfusion and a smaller cooling effect. Such a nonlocal interaction may open up new possibilities for therapy optimization. It might lead to higher tumour temperatures not to focus the power on the tumour, but to heat up muscles supplied by the same vessels in order to cut off the tumour from cooling perfusion.

### 2.3 Parameter Identification

All of the above thermoregulation models contain a number of parameters, most of which are unknown. Such parameters must be determined either from direct measurements or from computations comparing indirect measurements with the applied model. The most relevant parameters are:

- **Material coefficients:** Thermal, electrical, and physiological properties of tissue depend on the specific patient under consideration. Several techniques are available to get hold of reasonable values. Parts of these data are obtained from measurements, see, e.g. [109, Table 1] for electromagnetic and bioheat data, or to [88] with values from [73], where data for vessels, lungs, spleen were taken from [39]. In [110, Fig. 1], only slight differences between different patients for the absorption rate density are reported. One of the most important parameters is the perfusion $w$, both due to its dominant effect on the temperature and its high inter-individual variability. It can be measured by MRI using a contrast agent, but this is not generally indicated.

- **Geometry:** Since therapy planning and in particular segmentation takes a significant amount of time, the patient’s geometry is acquired some time before the therapy starts. Moreover, several therapy sessions are performed based on this single geometry. Positioning the patient in exactly the same location every time is a major practical difficulty. Even with utmost care, spatial deviations of roughly 15mm are unavoidable. Movements of the patient during therapy lead to additional offsets. Changes in body mass and intestine or bladder contents also affect the actual geometry.

- **Technical parameters:** Clinically available HF power generators are less stable in maintaining requested amplitudes and phase shifts than would be desirable. Moreover, part of the HF energy is reflected at contact resistances in coaxial cable connectors between the power generator and the applicator. These resistances depend on the mechanical force with which the connectors have been put together. While the transmission properties
of the cable network can, in principle, be measured, this is not always possible in clinical practice.

Figure 11: Hyperthermia applicator system in open MRT.

**Computational identification.** A validation of the models describing the electrical field and the heat transport requires in-vivo measurements of relevant quantities, in particular the temperature. If sufficient data is acquired, the parameters entering the models can be identified, which allows to improve the therapy. 3D distributed data sets can be obtained from magnetic resonance tomography (MR) measurements \[77\] in a hybrid MR-hyperthermia applicator, see Fig. [11]. At a rather limited number of discrete times, temperature probes can be placed inside the patient’s body. By MR thermometry based on positron resonance frequency shift (PRFS), a combination of tissue temperature increase \(T - T_0\) and perfusion \(w\) can be measured. Due to lack of better understanding of the relevant processes, a simple measurement model of the form

\[m \approx a(T - T_0) + bw\]

has been used, where the coefficients \(a\) and \(b\) depend on the tissue type. In particular in fatty tissue, the perfusion influence vanishes almost completely \((b \approx 0)\). Resolution, signal-to-noise ratio, and measurement time are mutually dependent. For useful thermometry data, a measurement takes at least 30s with current MR scanners.

**Adaptation of electric fields.** As mentioned above, several model parameters are not exactly known in clinical practice. These modelling errors lead to predictions significantly deviating from measurements (see Fig. [12]).

A radical remedy would be to omit the numerical solution of Maxwell’s equation completely and to identify the selfadjoint heating effect \(M(x) \in C^{k_E, k_E}\) of applicator control settings using only the \(k_m\) MR thermometry measurements \[57\]. Assuming the correctness of the computed absorption rate density
and the validity of the BHT equation. Tikhonov regularization leads to the quadratic optimization problem

\[
\min_{M \in L^2(\Omega; C^{k_E \times k_E})} \sum_{i=1}^{k_m} \| m_i - aT(t_i; M) - bw \|^2_{L^2(\Omega)} + \gamma \| M \|^2
\]

subject to the equality constraints and source term \(S(x) = \frac{1}{2} \alpha^* M(x) \alpha\). An implementation of this concept in clinical practice is prevented by the number and type of required measurements. Since for each point or voxel \(x \in \Omega\) the number of real parameters in \(M(x)\) for an applicator with \(k_E\) channels is \(k_E^2\), many measurements have to be taken for a complete identification (the Sigma-Eye applicator with \(k_E = 12\) would require more than one hour just to calibrate the model). Moreover, in order to allow a reasonable identification, sufficiently different controls \(\alpha \in C^{k_E}\) have to be used, most of which are of questionable benefit for the patient. Motion artefacts at tissue boundaries prevent the identification of heating effect in a significant part of the domain.

The number of parameters to be identified can be reduced by considering the antenna profiles \(\tilde{E} = (E_j)_{j=1,...,k_E}\) instead of the heating effect \(M\). Since \(M(x) = \tilde{E}(x)^* \tilde{E}(x)\), this leads to a nonlinear identification problem with \(6k_E\) parameters per voxel. Because simultaneous phase shifts and simultaneous spatial rotation of field vectors have no thermal effect, a rank defect of at least three reduces the number of parameters further. Still, relying on models identified from scratch is impractical.

A promising approach is to start with antenna profiles obtained as the solutions \(\tilde{E}_0\) of Maxwell’s equation and adapt them by least-change updates such as to reproduce measurements. This allows to start the therapy with a reasonably good model to be improved further whenever new measurements are available. Without the need to sample the whole space of applicator controls, it is moreover possible to deliver the best therapy that is known at any time during the whole hyperthermia session.

Solving identification problems of the type subject to the time-dependent BHT equation giving \(T(t; M)\) in real time is a challenging task. Spatial decoupling by neglecting heat conduction in the identification leads to a set of smaller parameters.
problems to be solved independently for every voxel, e.g., by a Gauß-Newton method:
\[
\min_{\tilde{E} \in \mathbb{C}^{3 \times kE}} \sum_{i=1}^{k_m} |m_i - aT(t_i; M) - bw|^2 + \gamma \|\tilde{E} - \tilde{E}_0\|^2 \tag{20}
\]

Heat conduction can nevertheless be taken into account without solving a spatially coupled identification problem. Splitting the antenna profiles \(\tilde{E}\) into a prediction part \(\tilde{E}_0\) based on Maxwell’s equations and a correction part \(\delta\tilde{E}\) to be identified, and the temperature \(T = T_0 + \delta T\) into corresponding contributions, the time-dependent BHT equation (13) reads
\[
\rho c \frac{\partial (T_0 + \delta T)}{\partial t} = \text{div}(\kappa \text{grad}(T_0 + \delta T)) - c_b w(T_0 + \delta T - T_{\text{blood}}) + \frac{\sigma}{2} \alpha^* \tilde{E}_0 \tilde{E}_0 + \frac{\sigma}{2} \alpha^* (2 \tilde{E}_0^* + \delta \tilde{E}^*) \delta\tilde{E} \alpha.
\]

Assuming \(\delta\tilde{E}\) and \(\delta T\) to be comparatively small, we may neglect heat conduction in \(\delta T\) only and obtain
\[
\rho c \frac{d\delta T}{dt} = -c_b w \delta T + \frac{\sigma}{2} \alpha^* (2 \tilde{E}_0^* + \delta \tilde{E}^*) \delta\tilde{E} \alpha
\]
as the constraint to (20). The temperature \(T_0\) satisfying (13) does not depend on \(\delta\tilde{E}\) to be identified and can thus be computed before or during the MR measurement. This approach yields much better results than neglecting heat conduction completely, see Fig. 13. For details we refer to [76].

One important point to notice is that, despite being underdetermined, the identification problem is severely ill-conditioned and hence leads to significant noise amplification whenever similar applicator controls \(\alpha_i\) are used – a fact that had been long overlooked in literature. Applied in a feed-back loop to improve the applicator controls during therapy, the process will converge towards some optimum. Hence, very similar controls are bound to occur, and sufficient regularization is necessary, see Fig 13.

**Identification of perfusion.** Among the heat transfer model parameters the perfusion is the most influential one. Hence, measurement or identification of *individual* perfusion values would be very valuable. Quite accurate values can be obtained from MR data using contrast agents [65]. The application of contrast agents may incur an undesirable risk for the patient, which is why they are only used if absolutely necessary. Moreover, due to the required injection of the contrast agent, only very few of these measurements can be obtained, such that a continuous monitoring of thermoregulatory feedback is impossible. Consequently, such data are not generally available.

As a noninvasive alternative, the MR thermometry data \(m\) mentioned above can be considered. This leads to an identification problem for temperature and perfusion [41]:
\[
\min_{T, w} \|m - aT - bw\|_{L_2(\Omega)}^2 + \gamma \|w - w_0(T)\|_{H}^2 + b(w, w_0(T))
\]

22
Figure 13: $L_2$ error of the predicted temperatures (in K) over the number of adaptation steps on a simulated therapy. The curves correspond to antenna profile adaptation with and without regularization (reg) as well as with and without taking heat conduction (con) into account. The straight dotted line at 0.1 is the $L_2$ norm of a normally distributed noise applied to generate artificial measurements from exact temperatures (from [76]).

subject to the equality constraints (13) and (14). Herein, $w_0(T)$ is the a-priori perfusion model. The norm

$$
\|w\|^2_H := \|w\|^2_{L_2(\Omega)} + \sum_{i=1}^{\# \text{tissues}} \|\nabla w\|^2_{L_2(\Omega_i)}
$$

for the regularization term is constructed based on the assumption that the perfusion depends smoothly on the temperature in every tissue region, but may jump across tissue boundaries. The barrier function $b$ enforces nonnegativity of the perfusion, but has to be skewed such that its minimizer is the reference perfusion $w_0(T)$ in order not to introduce a systematic bias.

Clinical application of that concept is prevented by the fact that there can be two different value pairs of temperature and perfusion which are compatible with both the BHT equation and the measurements, which is a consequence of the bilinearity of the BHT equation. Hence, while local consistency is fostered by regularization, global consistency cannot be guaranteed. In fact, numerical experiments indicate that the occurrence of both compatible values must be expected [103].

The consequence is that more data is required to provide the clinical staff with reliable identified temperature and perfusion values. Such data might come from additional measurements taken with MR sequences which exhibit impact coefficients $a$ and $b$ different from PRFS.
2.4 Therapy Planning Problem

In a first step, we formulate the medical problem of therapy planning which then has to be translated into mathematical terms.

Medical problem setting. Due to the production of so-called heat shock proteins, the temperature should best be above 42.5°C. Up to now the heating cannot safely be restricted to tumour cells only, not the least, since it is not possible to distinguish tumour cells reliably from healthy ones. Therefore, in order to be on the safe side for healthy tissue, local temperatures in tumours cannot be allowed to be too high. Thus the therapeutic aim is in general not to burn tumour cells, but to make them more sensitive to other treatments like radiotherapy or chemotherapy (cytostatica). In vitro, experiments have confirmed an increased sensitivity of tumour cells by a factor of 7 over healthy ones, an effect also be seen in a milder form in vivo, see, e.g., [19]. The physiological mechanism behind this effect is captured in the so-called oxygen enhancement ratio (OER). Consequently, the general aim can be specified as:

\textit{Control the antenna parameters such that tumour tissue is heated, while healthy and uncertain tissue is not heated above a prescribed threshold temperature.}

After these preparations, we are now ready to translate the above medical problem into a mathematical one. Let $T$ denote the temperature inside the body. Following the medical intention, an optimal temperature distribution should satisfy the following items:

- In the region $V_{\text{tumour}}$ populated by tumour cells, the temperature should be sufficiently high to incur a therapeutic benefit. In view of the metabolism of heat shock proteins, a temperature level $T_t \geq 42.5^\circ\text{C}$ should be maintained.

- In the region $V_{\text{healthy}}$ populated by a significant portion of healthy tissue, the temperature should not exceed a tissue-type dependent threshold $T_h \approx 42 - 44^\circ\text{C}$ in order to avoid subcutaneous burns.

In addition there is a region where both tumour and healthy cells are present, which can be treated in a special way (cf. Section 5.2).

Apart from the medical intention, technical restrictions are imposed by the applicator. The amplitude of each antenna is limited, as is the total power emitted. Moreover, the antenna phases are only reliable as long as the amplitudes are not too different.

These aims and restrictions, partially conflicting, have been modelled and formulated as mathematical optimization problems in a variety of ways, which we will present in the following.

Cost functionals. First approaches to the quantification of the therapeutical benefit to be maximized have been formulated in terms of the absorption rate
density alone, based on the idea that high power deposition leads to high temperature. The associated cost functional to be minimized is

$$f(E) = -\frac{1}{2} \int_{V_{\text{tumour}}} \sigma |E(\alpha)|^2 \, dx.$$  \hspace{1cm} (21)

An advantage is that this functional can be easily minimized by solving a low-dimensional eigenvalue problem, as long as no temperature constraints are prescribed.

For medical reasons, a temperature-based objective functional is preferable. Therefore, in [71, 73], such a functional has been suggested. The idea is to penalize “cold spots” in the tumour below a therapeutical temperature $T_t \approx 43^\circ C$, since the impact of hyperthermia on cancer cells drops quickly below this threshold. This leads to the functional

$$f(T) = \int_{V_{\text{tumour}}} (T_t - T)^2_+ \, dx$$  \hspace{1cm} (22)

with $(\cdot)_+ = \max(\cdot, 0)$. This approach can be extended to the concept of thermal dose, with the aim of minimizing the fraction of surviving tumour cells. The thermal damage rate inflicted on cancer cells follows the Arrhenius law [34, 95], with a rate constant

$$k(T) = A \exp \left( -\frac{\Delta E}{RT} \right).$$

Herein, the frequency factor $A$ and the activation energy $\Delta E$ depend on the tissue type, $R$ is the universal gas constant. An estimate $s$ of the fraction of surviving cells is given by the simple differential equation

$$\dot{s}(t, T) = -k(T)s(t, T), \quad s(0, T) = 1,$$

which gives rise to

$$s(t, T) = \exp(-tk(T)).$$  \hspace{1cm} (23)

Thus the objective functional to be minimized is the fraction of cancer cells surviving after a prescribed duration $t$ of a therapy session:

$$f(T) = \int_{V_{\text{tumour}}} \exp \left( -tA \exp \left( -\frac{\Delta E}{RT} \right) \right) \, dx$$  \hspace{1cm} (24)

A schematic representation of such a function is given in Fig. 14. The thermal isoeffect dose is an established quantity for assessing the therapeutic benefit of a treatment [34, 79]. Treatment planning based on the tumour cell survival has been proposed for thermoseed placement [98], but up to now rather ad hoc cost functionals like (22) have been used in regional hyperthermia planning.

**Temperature constraints.** At the time when mathematical therapy planning was started, a sufficient mathematical understanding of state constraints as well as an efficient algorithmic treatment were lacking. A common approach
was therefore an external penalization of “hot spots” arising in healthy tissue with temperatures above a tissue-dependent threshold $T_{\text{lim}}$, occasionally augmented by an additional, weaker penalization of heating healthy tissue above a comfortable temperature $T_h < T_{\text{lim}}$. This leads to extended cost functionals

$$J(T) = f(T) + \int_{V_{\text{healthy}}} \left( \gamma_1 (T - T_h)^2 + \gamma_2 (T - T_{\text{lim}})^2 \right) \, dx$$

with fixed penalty parameters $0 \leq \gamma_1 \ll \gamma_2$. Due to external penalization, however, “hot spots” cannot be completely prevented. In fact, they are bound to arise unless $\gamma_2 \to \infty$. An analysis of the asymptotic behaviour performed in [85] suggests that even under favourable circumstances the maximum constraint violation only decreases by a rate of $O(\gamma_2^{-2/5})$. We would like to point out that in hot spots with less regularity of the temperature distribution, in particular due to discontinuous temperature bound or singularities of the absorption rate density, the maximum violation decreases even more slowly.

A strict prevention of hot spots in healthy tissue will require the explicit respection of state constraints of the kind

$$T(x, \alpha) \leq T_{\text{lim}}, \quad x \in V_{\text{healthy}}. \quad (25)$$

One way to guarantee feasibility of the temperature distribution is to use a barrier method. In Section 4 below, we will work out results in terms of both theory and algorithms.

**Control constraints.** Additional constraints directly on the control $\alpha$ are imposed to model the limited power of the microwave applicator:

$$|\alpha_k| \leq \alpha_{\text{max}}, \quad k = 1, \ldots, k_E$$

$$\sum_{k=1}^{k_E} |\alpha_k|^2 \leq \alpha_{\text{total}}^2. \quad (26)$$
If desired, stability of antenna phases can be ensured by confining the amplitude ratios to
\[
\alpha_{ratio}^{-1} \leq \frac{\alpha_k|_C}{\alpha_j|_C} \leq \alpha_{ratio}, \quad j, k = 1, \ldots, k_E, \quad \text{with } \alpha_{ratio} > 1,
\]
or by requesting all antennas to have the same amplitude:
\[
|\alpha_j|_C = |\alpha_k|_C, \quad j, k = 1, \ldots, k_E.
\]

**Optimal control problems.** Putting all pieces together, different optimization problems can be formulated. Omitting the control constraints, we state two variants that have been used for therapy planning.

**Penalized problem:**

\[
\min_{\alpha \in \mathbb{C}^k, T \in H^1(\Omega)} \int_{V_{tumour}} (T - T)^2 \, dx + \int_{V_{healthy}} \left( \gamma_1 (T - T_h)^2 + \gamma_2 (T - T_{lim})^2 \right) \, dx \tag{27a}
\]

subject to the BHT equation (15)
\[
A(T) - B(\alpha) = 0. \tag{27b}
\]

**Constrained problem:**

\[
\min_{\alpha \in \mathbb{C}^k, T \in H^1(\Omega)} \int_{V_{tumour}} \exp \left( -tA \exp \left( -\frac{\Delta E}{RT} \right) \right) \, dx \tag{28a}
\]

subject to the BHT equation (15) and temperature constraints
\[
A(T) - B(\alpha) = 0, \quad T \leq T_{lim}. \tag{28b}
\]

This above set of goal functionals and constraints supplies the starting point for the actual computation of optimal therapies for individual patients.

### 3 Efficient Simulation

In the numerical simulation of the whole hyperthermia procedure computational efficiency is of utmost importance, since clinical decision making requires numerical algorithms that run on a local PC both fast and reliably. The simulation involves solving both the time-harmonic Maxwell equation and the BHT equation. The latter, a Helmholtz equation in the non-critical case, can be readily solved by algorithms which nowadays are standard textbook material, see, e.g., [32]. That is why we skip this part here. Rather, we concentrate on the two main sources of computational cost, the adaptive multilevel finite element methods for the time-harmonic Maxwell equation in Section 3.1 and their unidirectional coupling with the BHT equation in the subsequent Section 3.2. A first survey on the decisive role of these methods for hyperthermia treatment planning has been given in [31].
3.1 Adaptive FEM for Time-Harmonic Maxwell Equation

The first algorithmic attempts to solve the time-harmonic Maxwell equation were based on the rather popular FDTD method, see [94]. These methods started directly from the voxel discretization, i.e. the raw data, as input for numerical integration, certainly an intriguing feature. In addition, the methods appeared to be very fast; but, after testing them in a hyperthermia environment, it turned out that they were unable to treat the interface conditions correctly, which in an FEM would be automatically satisfied (see Section 3.1). A comparison of results for a special virtual patient is given in Fig. 15.

![Comparison of electrical field computations. Left: Popular FDTD method [94] based on the strong formulation (without correct interface conditions). Right: Adaptive finite element method based on the weak formulation (with automatically correct interface conditions), see Section 3.1.](image)

Figure 15: Comparison of electrical field computations. Left: Popular FDTD method [94] based on the strong formulation (without correct interface conditions). Right: Adaptive finite element method based on the weak formulation (with automatically correct interface conditions), see Section 3.1.

The typical ingredients of an adaptive multilevel finite element method (FEM) are (a) an a-posteriori local error estimator, (b) an adaptive mesh refinement strategy, and (c) a fast solver for the arising large linear systems. For Poisson-type PDEs, such techniques are well-developed. For the time-harmonic Maxwell equation, however, a deeper understanding of the indefinite structure is necessary to reach optimal multigrid complexity.

**Edge Elements.** We return to the weak formulation (5) in Section 2.1 above. In order to obtain a finite element discretization of the weak equation (6), we generate a tetrahedral triangulation $T_h$ of the domain $\Omega$ and employ Nédelec’s curl-conforming finite elements of first order [70], also called edge elements [14] or Whitney 1-forms. Let us denote the space of linear Lagrange elements by $S_h$ and the nodal basis functions associated to vertex $i$ by $\lambda_i$. Then the edge element basis function associated with the edge $e = \{ij\}$ is defined as

$$w_{\{ij\}} = \lambda_i \text{grad} \lambda_j - \lambda_j \text{grad} \lambda_i,$$

(29)
Figure 16: The tetrahedral edge element shape function $w_{ij}$ associated to edge $\{ij\}$ is a circular field rotating around the opposite edge.

see Fig. 16. The corresponding FE space is

$$ND_h := \text{span}\{w_{ij}\}.$$  

Edge elements possess several distinct advantages compared to vectorial Lagrange elements: They guarantee the desired continuity of the tangential components of the electric field while preserving jumps of the normal components. In particular, concave metallic edges and internal boundaries between materials with jumps in the coefficients can be naturally incorporated in agreement with physical continuity relations. Moreover, they avoid the occurrence of spurious modes, whereas nodal basis functions may give rise to wrong solutions (see [15]). The convergence behaviour of edge element discretizations of [68] is studied in [68]. For the actual realization we note that for a vector field $E$ the degree of freedom $E_e \in \mathbb{C}$ associated with each edge $e$ in $T_h$ is given by the path integral

$$E_e = \int_e E \cdot t_e \, ds$$  \hspace{1cm} (30)

along the edge $e$ with direction vector $t_e$.

**Spectral properties.** Before we dive into algorithmic details, let us have a closer look at the spectral structure of the time-harmonic Maxwell operator from (5) that occurs in the weak equation (6). As illustrated in Fig. 17, its spectrum divides into three distinct parts: (i) eigenvalues with negative real part originating from the nullspace $H^0_t(\text{curl}, \Omega)$ of the curl-operator, (ii) eigenvalues with negative real part to be associated with some space $\mathcal{M}^-$ spanned by the corresponding eigenfunctions, and (iii) eigenvalues with positive real part, which go with some space $\mathcal{M}^+$.  

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Coarse mesh. Suppose we formally expand both sides of the weak equation \([6]\) in terms of eigenfunctions of the Maxwell operator. Then the system would decouple in terms of these eigenfunctions. Then those terms on the right side with smallest coefficients will dominate the solution structure in the generic case – assuming, of course, that all eigenfunctions are present in the actual solution. Recall that, in the Laplace operator, the smooth eigenfunctions have the smallest eigenvalues and thus dominate the solution structure. The situation is very different for the indefinite time-harmonic Maxwell operator. To see that, a simple plane wave analysis has been given in [9], which we repeat here.

For simplicity, let the material be homogeneous, so that the coefficients \(\mu, \epsilon\) are constants. Let \(E(x) := \exp(ikx)e\) denote a plane spatial wave in a polarization direction \(e\). Upon ignoring the boundary terms in \(A\), eigenvalues \(|k|^2 - \omega^2 \mu \epsilon\) will come up. Obviously, wavenumbers \(k\) close to the critical wavenumber \(k_C = \frac{\omega}{\sqrt{\mu \epsilon}}\) give rise to the smallest eigenvalues of the time-harmonic operator \(A\). The associated eigenfunctions are bound to dominate the solution generically.

Let the critical wavelength be defined by

\[
\lambda_C = \frac{2\pi}{k_C} = \frac{2\pi}{\omega \sqrt{\mu \epsilon}}.
\]

Then an appropriate coarse mesh able to resolve solutions in the space \(\mathcal{M}^-\), compare Fig. [17] should obey a mesh size restriction of the kind

\[
h \leq \frac{\lambda_C}{2},
\]

(31)
which is often called Nyquist condition. It will again show up below in the context of multigrid methods. In passing we note that the above restriction depends, of course, on the material. In the hyperthermia setting, it is only restrictive in the water bolus, producing a nearly uniform mesh therein.

Large linear system solvers. Replacing $H^r_\Gamma(\text{curl}; \Omega)$ by $ND_h$ in the variational formulation (6) leads to a large, very sparse linear equation system

$$Au = b$$

with complex symmetric and indefinite matrix $A$. The solution of (32) can be obtained by either direct or iterative solvers. Direct solvers have experienced a significant progress in recent years [24]. Despite a still considerable memory requirement due to fill-in, they are very effective on coarse up to moderately fine meshes. Only for very fine grids or if low memory requirement is important, iterative solvers have to be used. Iterative solvers face two difficulties, as can be seen from the eigenvalue distribution of the operator $A$ in Fig. [17] which carries over to the matrix $A$. First, the large eigenvalues grow with decreasing mesh size $h$. The growing condition number of the matrix $A$ renders both matrix decomposition methods and Krylov methods inefficient on fine meshes. Second, due to the eigenvalues with negative real part, usual positive smoothers such as Jacobi or Gauß-Seidel methods do not converge: the spectral radius of their iteration matrix $I - B^{-1}A$ exceeds 1, so that the corresponding modes in $H^0_\Gamma \cup M^-$ are amplified rather than reduced.

Adaptive mesh refinement. Due to the complex geometrical structure and strongly varying material properties of virtual patients, corner singularities and other local solution features are bound to arise. Hence we consider adaptive mesh refinement to be of crucial importance to ensure both efficiency of the solver and reliability of its computational results. The foundation of adaptive mesh refinement are a-posteriori error estimators, which are most often based on localized operations. As we are dealing with equations of partly hyperbolic nature in this context, it is clear that any local estimator will be unable to capture far-field contributions. Accordingly we prefer to use the name error indicator in the sense of Babuška [4].

Among the different flavours of error indicators we consider hierarchical error indicators in the spirit of [29], which have been observed to yield very favourable results with efficiency quite close to 1 on simple test problems [9, Section 8.2]. Quantitative reliability is, of course, of utmost importance in medical applications.

Conceptually, the ansatz space $V^L = ND_h$ is extended by a hierarchical surplus $V^H$ to give a higher order ansatz space $V^+ = V^L \oplus V^H$, in which (32) assumes the block structure

$$
\begin{pmatrix}
A^{LL} & A^{LH} \\
A^{HL} & A^{HH}
\end{pmatrix}
\begin{pmatrix}
\begin{bmatrix}
u^L \\
w^H
\end{bmatrix}
\end{pmatrix}
= 
\begin{pmatrix}
\begin{bmatrix}b^L \\
b^H
\end{bmatrix}
\end{pmatrix}.
$$

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For the mere purpose of error indication, (33) needs only be solved approximately via a defect equation for the hierarchical components:

\[ A^{HH} u^H = b^H - A^{HL} u^L . \]  

Hierarchical extensions of the lowest-order edge elements have been described in [80], where the basis functions are given in terms of Lagrangian nodal functions \( \lambda_i \) associated to the vertices \( i \) of the tetrahedral mesh. With each edge \( \{ij\} \), one basis function

\[ w^H_{\{ij\}} = \lambda_i \text{grad} \lambda_j + \lambda_j \text{grad} \lambda_i \]

and with each face \( \{ijk\} \), two basis functions

\[ w^{H,1}_{\{ijk\}} = \lambda_i \lambda_j \text{grad} \lambda_k - \lambda_i \lambda_k \text{grad} \lambda_j, \quad w^{H,2}_{\{ijk\}} = \lambda_i \lambda_j \text{grad} \lambda_k - \lambda_j \lambda_k \text{grad} \lambda_i \]

are associated. An approximate solution of (34) may be obtained via one block-Jacobi step only (!), where it is essential to keep the \( 2 \times 2 \)-block-entries of both functions \( w^{H,l}_{\{ijk\}} \), \( l = 1, 2 \), attached to each face of the triangulation (see [9] for details). At first glance, this block-Jacobi sweep appears critical, as the matrix contains negative eigenvalues and the related eigenmodes may be amplified. However, the modulus of all negative eigenvalues being comparatively small, these modes can be expected to give only minor contributions. Let \( \tilde{u}^H \) denote the first Jacobi iterate of the exact \( u^H \). For the definition of a relative discretization error estimator we set

\[ \eta^2 = \frac{\| \tilde{u}^H \|^2_E}{\| u^L \|^2_E} , \]

wherein, in the absence of a better idea, we employ the norm

\[ \| v \|^2_E := \int_{\Omega} \left( \frac{1}{\mu} \text{curl} v^* \text{curl} v + \omega^2 \epsilon v^* v \right) dx + \int_{\Gamma_{\text{ext}}} \beta (n \times v^*) (n \times v) ds . \]  

This error estimator is the key ingredient for an adaptive refinement strategy in the spirit of a technique suggested by I. Babuška and W.C. Rheinboldt [5] for purely elliptic problems. As an illustration of an adaptive grid constructed along these lines, results for a virtual patient mesh are shown in Fig. 18.

Table 1 gives a list of error indicators as a function of the refinement levels starting from the mesh of Fig. 18.

**Multilevel preconditioning.** In the hyperthermia problem, a multigrid V-cycle preconditioner is used within a conjugate residual iterative method (CR), see [38]. On the subspace \( \mathcal{M}^+ \), standard multigrid methods are effective in reducing the error – however, at the expense of amplifying error components in \( H^0_0 \cup \mathcal{M}^- \). This leads to a deterioration of the whole multigrid procedure in the present case, see Table 2 under columns Std. Therefore, the standard multigrid approach needs to be modified.
Figure 18: Adaptive mesh of a virtual patient consisting of about 128.000 nodes. Note that a uniform grid would have required estimated 16.000.000 nodes. *Left:* frontal cut. *Right:* sagittal cut.

Table 1: Estimated discretization errors versus refinement level $j$ with node numbers $N_j$. The “coarse” mesh ($j = 0$) is taken from Fig. 18.

<table>
<thead>
<tr>
<th>$j$</th>
<th>$N_j$</th>
<th>$\eta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>128 365</td>
<td>10.1 %</td>
</tr>
<tr>
<td>1</td>
<td>373 084</td>
<td>4.96 %</td>
</tr>
<tr>
<td>2</td>
<td>1 085 269</td>
<td>2.54 %</td>
</tr>
</tbody>
</table>

The general applicability of multigrid methods to indefinite problems has been investigated in [16]. In order to capture the error part in $\mathcal{M}^-$, certain restrictions on the mesh size of the coarse grid have to be observed – unfortunately, given there in terms of unknown theoretical quantities. Fortunately, both local mode analysis and numerical experience confirmed that it is enough to observe the above Nyquist condition (31). The coarse grid correction is then computed by a direct solver. Treating the part originating from $H^0_\Gamma(\text{curl}, \Omega)$ is more delicate, since it contains functions with all spatial frequencies and therefore cannot be captured on the coarse mesh.

For Nédélec finite elements there exists a discrete counterpart of the representation of $H^0_\Gamma(\text{curl}, \Omega)$ as space of gradients of potentials. Thus, on simply connected domains each curl-free element of $\mathcal{ND}_h$ can be represented as the gradient of a scalar potential in the space $\mathcal{S}_h$ of nodal finite elements:

$$\mathcal{ND}_h \cap H^0_\Gamma(\text{curl}, \Omega) = \text{grad} \mathcal{S}_h,$$

(36)
Table 2: Convergence history for multilevel solvers with standard Gauss-Seidel smoothing (Std) compared to multiplicative (M-H) and additive (A-H) versions using hybrid Gauss-Seidel smoothing on each level. The iteration is terminated if the ratio of the Euclidean norms of residuals and right hand sides is below $10^{-5}$. From [31].

<table>
<thead>
<tr>
<th>Ref.</th>
<th>Nodes</th>
<th>#Iter</th>
<th>CPU [min]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Std</td>
<td>M-H</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Std</td>
<td>M-H</td>
</tr>
<tr>
<td>0</td>
<td>128 365</td>
<td>4250</td>
<td>354</td>
</tr>
<tr>
<td></td>
<td></td>
<td>150</td>
<td>24</td>
</tr>
<tr>
<td>1</td>
<td>373 084</td>
<td>4832</td>
<td>265</td>
</tr>
<tr>
<td></td>
<td></td>
<td>800</td>
<td>76</td>
</tr>
<tr>
<td>2</td>
<td>1 085 269</td>
<td>&gt; 10000</td>
<td>186</td>
</tr>
<tr>
<td></td>
<td></td>
<td>&gt; 2000</td>
<td>215</td>
</tr>
</tbody>
</table>

Moreover, the stable decomposition (7) has a (perturbed) discrete counterpart, too. To this end, let $I^{\text{curl}} : (S_h)^3 \rightarrow ND_h$ denote the edge interpolation operator, defined via (30). Then

$$v = \tilde{v} + I^{\text{curl}} \phi + \text{grad} p, \quad \phi \in (S_h)^3, \ p \in S_h, \ \tilde{v} \in ND_h$$

(37)

such that

$$\|h^{-1} \tilde{v}\|_{L^2} + \|\phi\|_{H^1} + \|p\|_{H^1} \leq C\|v\|_{H(\text{curl})}.$$  

(38)

This outstanding feature of Nédelec elements (see e.g. [10, 51]) is of crucial importance for the construction of efficient multigrid solvers for the arising linear systems, as we will illustrate at the simpler example of positive definite problems.

A multigrid method, applied to a linear system $Au = b$ on a space $V$ consists of a (non-direct) splitting

$$V = \sum V_i$$

and preconditioners $B_i$, defined on each $V_i$. Here we will choose $V_i$ as one dimensional spaces that come from a multilevel hierarchy and $B_i = A$. The convergence theory then depends on the stability condition

$$\inf_{v_0 = \sum v_i} \sum \|v_i\|^2_A \leq K_1 \|v\|^2_A,$$

(39)

and the strengthened Cauchy-Schwarz inequality

$$(v_k, v_l)_A \leq \gamma_k \|v_k\|_A \|v_l\|_A \quad K_2 := \|\gamma\|_2,$$

(40)

which have to be established for the given splitting [112]. In the additive case (e.g., Jacobi type smoothing) the product $K_1 K_2$ gives an estimate for the condition number of the preconditioned system. If $K_1 K_2$ admits an upper bound that is independent of $h$ we obtain an optimal preconditioner, i.e., a bounded
condition number. Similar results hold in the multiplicative case, e.g., for Gauss-Seidel type smoothing.

To understand, why standard multigrid smoothers are inefficient in an $H(\text{curl})$-setting, consider $\| \cdot \| = \| \cdot \|_{H(\text{curl})}$ and a multilevel splitting

$$ND_h = \sum \Phi_i$$

where $\Phi_i$ are one dimensional spaces, spanned by the edge element basis functions on different levels. Now let $v \in ND_h \cap H^1_0(\text{curl}, \Omega)$ (recall $\text{curl} v = 0$), such that

$$v = \sum_{\phi_i \in \Phi_i} \phi_i.$$

The basis functions satisfy $\| \phi_i \|_{L^2} \leq c h_i \| \phi_i \|_{H(\text{curl})}$. Thus we obtain

$$\| v \|_{H(\text{curl})}^2 \leq C \sum_{v = \sum \phi_i} \| \phi_i \|_{L^2}^2 \leq C \sum_{v = \sum \phi_i} h_i^2 \| \phi_i \|_{H(\text{curl})}^2. \quad (41)$$

Whenever $v$ bears high spatial frequencies, this implies

$$K_1 \geq C h^{-2},$$

and thus leads to mesh-dependent condition numbers and contraction rates.

Returning again to (36), we may introduce a potential $p \in S_h$, such that $v = \text{grad} \ p$. Then, for a multilevel decomposition

$$S_h = \sum P_i,$$

where $P_i$ are one dimensional subspaces spanned by the nodal basis functions of $S_h$ on different levels, standard $H^1$-multigrid theory yields:

$$\inf_{p = \sum P_i} \sum \| p_i \|_{H^1}^2 \leq K_{1,H^1} \| p \|_{H^1}^2 = K_{1,H^1} \| \text{grad} \ p \|_{L^2}^2 = K_{1,H^1} \| v \|_{H(\text{curl})}^2. \quad (42)$$

For general $v \in ND_h$ we can apply the stable discrete decomposition (37) which finally yields a stable multigrid decomposition of the form

$$V = \sum \Phi_i + \sum \text{grad} \ P_i, \quad (43)$$

where $\Phi_i$ and $P_i$ are taken from a multilevel hierarchy of $ND_h$ and $S_h$, respectively. Then a similar multigrid estimate as (42) for the rotational part of $v$, together with (42) yields an $h$-independent constant $K_1$. Note that the mere existence of such a splitting is sufficient, it need not be computed explicitly. More details can be found in [10, 9, 8, 50, 53, 113].

On the basis of the splitting (43) a hybrid smoothing procedure was constructed in [10, 9]. There, as a basic solver the conjugate residual (CR) method was used, which is similar to the well-known conjugate gradient algorithm, but adjusted to symmetric indefinite systems. Its basic operations are Gauss-Seidel sweeps both in the Nédélec space $ND_h$, coping with the elliptic part of $A$, and in
the space $S_h$ of potentials. Within this framework, efficient transfer operators between field representations in $S_h$ and $ND_h$ are essential.

If we represent a vector field $v \in ND_h \cap H^1_0(\text{curl}, \Omega)$ by $v = \text{grad} \, p$, then the representation in $ND_h$ can be obtained easily from (30):

$$v_e = \int_{P_1}^{P_2} \text{grad} \, p \cdot t \, ds = p(P_2) - p(P_1).$$

Here $P_1$ and $P_2$ denote the positions of the endpoints of the edge $e$.

In the following $P_{S_h}$ denotes the transfer operator from the potential space $S_h$ into the Nédélec space $ND_h$ on $T_h$. $P^*_{S_h}$ will denote the adjoint operator, defining the canonical restriction. Then the smoother $A_S$ for the nullspace is constructed by the Galerkin product $A_S = P^*_{S_h} AP_{S_h}$. The following hybrid smoothing algorithm for a given right hand side vector $b \in ND_h$ will provide an updated solution vector $u \in ND_h$ via the following steps:

One **Gauss-Seidel** step for $Au = b$

- $\tilde{b} \leftarrow b - Au$
- $b_S \leftarrow P^*_{S_h} \tilde{b}$
- $u_S \leftarrow 0$

One **Gauss-Seidel** step for $A_S u_S = b_S$

- $u \leftarrow u + P_{S_h} u_S$

In order to be on safer theoretical grounds, we realize symmetric preconditioning, which means an immediately following step of the adjoint procedure. Note that in this approach both spaces are treated in a **multiplicative** fashion in the sense of domain decomposition methods.

As an alternative, we propose a symmetric **additive** version:

One **symmetric Gauss-Seidel** step for $Au = b$

- $b_S \leftarrow P^*_{S_h} b$
- $u_S \leftarrow 0$

One **symmetric Gauss-Seidel** step for $A_S u_S = b_S$

- $u \leftarrow u + P_{S_h} u_S$

If we employ adaptive mesh refinement, thus creating a sequence of nested triangulations $T_0 \subset T_1 \subset \ldots \subset T_h$, then the extension to a multilevel solver is quite straightforward. Taking into consideration that the associated sequence of Nédélec spaces is nested, i.e. $ND_0 \subset ND_1 \subset \ldots \subset ND_h$, we may adopt the classical multigrid idea by using canonical grid transfer operations between these spaces, but employing a hybrid smoother on each level.

Finally, in order to demonstrate the efficiency of this hybrid algorithm, we present comparative results for both the multiplicative and the additive hybrid algorithm using the discrete Helmholtz decomposition in Table 2 above. As can be seen, the hybrid versions exhibit a superior performance and do not deteriorate with increasing refinement level. Note that we have no direct factorization available on the coarse grid, thus facing comparatively large iteration counts. A proof of optimal multigrid complexity of this type of algorithm can be found in [8] [43].
3.2 Coupling of BHT and Maxwell Equation

In the covered temperature range of 30–50°C, the electromagnetic material parameters are essentially constant, such that the coupling of the time-harmonic Maxwell equation and the BHT equation is unidirectional. Thus, we can first solve the time-harmonic Maxwell equation and insert the results into the BHT equation. Since the Maxwell equation is linear, we can compute the antenna profiles \( E_k \) for each antenna channel \( k = 1, \ldots, k_E \) and use superposition afterwards. For the insertion into the BHT equation, however, a distinction between the linear and the nonlinear case is necessary.

**Linear BHT equation.** As can be seen from equation (12), the superposition of the electric field \( E \) from \( k_E \) antenna profiles gives rise to a superposition of \( k_E^2 \) source modes to the source term \( S \):

\[
\sum_{j,l=1}^{k_E} \alpha_j^* \alpha_l E_j^* E_l = \sum_{j=1}^{k_E} |\alpha_j|^2 |E_j|^2 + 2 \sum_{j=2}^{k_E} \sum_{l=1}^{j-1} (\Re(\alpha_j^* \alpha_l) \Re(E_j^* E_l) - \Im(\alpha_j^* \alpha_l) \Im(E_j^* E_l))
\]

The source modes

\[
S_{jl} = \begin{cases} 
\Re(E_j^* E_l), & j \leq l \\
\Im(E_j^* E_l), & j > l 
\end{cases}
\]

correspond to temperature elevations \( T_{jl} \) above the basic tissue temperature \( T_{bas} \) for \( \alpha = 0 \) and can be computed in advance by solving the homogeneous BHT equation

\[
-\text{div}(\kappa \text{grad} T_{jl}) = -c_b w T_{jl} + \frac{1}{2} \sigma S_{jl} \quad \text{in } \Omega \\
-\kappa \frac{\partial T_{jl}}{\partial n} = h T_{jl} \quad \text{on } \partial \Omega.
\]

With coefficients \( \gamma_{jl} = 2 \Re(\alpha_j^* \alpha_l) \) for \( j < l \), \( \gamma_{jj} = |\alpha_j|^2 \), and \( \gamma_{jl} = -2 \Im(\alpha_j^* \alpha_l) \) for \( j > l \), the temperature is obtained as

\[
T(\alpha) = T_{bas} + \sum_{j,l=1}^{k_E} \gamma_{jl} T_{jl}.
\]

Let \( \text{cost}_{\text{linBHT}} \) denote the computational cost for one solution of the BHT equation. Then the total computational cost sums up as

\[
\text{cost}_{\text{total}} = k_E \text{cost}_{\text{Maxwell}} + (k_E^2 + 1) \text{cost}_{\text{linBHT}} + \text{cost}_{\text{Opt}}.
\]

(44)

As an illustration, see Table 3 below in Section 5. The cost for the optimization as a whole depends, of course, on the number of iterations.
**Nonlinear BHT equation.** In the case of a temperature dependent perfusion \( w(T) \), some kind of iteration is needed to take care of this nonlinearity. In [31], the following fixed point iteration has been suggested:

- **\( k_E \) Maxwell solves:** \( E_k \)
- let \( \alpha^0 := 0 \)
- for \( m = 0, \ldots, n \):
  - one nonlinear BHT equation solve for \( S(\alpha^m) \) supplies temperature \( T^m \) and perfusion \( w^m := w(T^m) \)
  - \( k^2 + 1 \) linear BHT equation solves for \( S_{jl} \) using \( w^m \) supplies \( T_{bas}^m \) and temperature modes \( T_{jl}^m \)
  - optimization \( \min_\alpha f(T_{bas}^m + \sum_{j,l=1}^{k_E} \gamma_{jl}(\alpha)T_{jl}^m) \) supplies \( \alpha^{m+1} \)

The idea behind this algorithm has been that the Maxwell solves are considerably more expensive than the BHT solves. The iteration converges linearly and quite fast at an observed contraction rate \( \theta \approx 0.3 \). So the total computational cost sums up as

\[
  \text{cost}_{\text{total}} = k_E \text{cost}_{\text{Maxwell}} + n_{\text{iter}} (\text{cost}_{\text{nonlinBHT}} + (k_E^2 + 1) \text{cost}_{\text{linBHT}} + \text{cost}_{\text{Opt}}) \quad (45)
\]

In typical computations, \( n_{\text{iter}} \approx 6 \) optimization iterations are required. The nonlinear BHT equation has been solved by a Newton iteration in function space [33], typically requiring no more than 2 steps due to the fast local convergence.

### 4 PDE Constrained Optimization

In Section 2.4, we had already worked out the medically preferable optimization problem with strict temperature limits in healthy tissue. At the time, when this aspect had come into the focus of interest, there were no efficient algorithms to tackle such problems. That is why thorough theoretical investigations had started, which we present in Section 4.1, while algorithmic details are postponed to Section 4.2. In order to derive efficient *adaptive* algorithms, we follow a multistage computational paradigm. In a first stage, we develop the necessary numerical analysis background in a function space setting, which covers both optimality conditions and formal algorithms for the infinite dimensional problem. In a second stage, we construct an iterative method, again in the infinite dimensional setting, for the arising nonlinear problems. Only in the last stage, the innermost loop, where linear operator equations are to be solved, we discretize the infinite dimensional problem up to a prescribed error tolerance. This kind of scheme has emerged as the general basis for any adaptive PDE algorithm.
4.1 Theoretical Background

The hyperthermia treatment planning problem can be formulated as an optimization problem in function space subject to equality constraints and inequality constraints in the following form:

\[
\min f(T, \alpha) \quad (46a)
\]

subject to

\[
A(T) - B(\alpha) = 0 \quad (46b)
\]

\[
\|\alpha_k\|_{\mathbb{R}^2} \leq \alpha_{\text{max}} \quad T \leq T_{\text{lim}} \quad (46c)
\]

Here \(A\) and \(B\) represent the BHT equation in weak form \(\text{(15)}\). From Section 2.2 we have \(\alpha\) as a set of \(2k_E - 1\) real parameters.

Various cost functionals \(f\) and inequality constraints have been discussed in Section 2.4. The most challenging part are the constraints on the temperature \(T \leq T_{\text{lim}}\). They can be considered as an infinite number of inequality constraints in function space. In PDE constrained optimization and optimal control they are classified as pointwise state constraints and have been subject to intense research in recent years (cf. e.g. [20, 67, 47, 25]).

The following derivations are based on [86, 83]. More details and proofs can be found therein.

**Barrier regularization and strict feasibility.** In finite dimensions, barrier regularizations give rise to the highly popular class of interior point methods [96, 107]. With \(y = (y_1, \ldots, y_n)\), the principal idea is to replace an inequality constrained problem of the kind

\[
\min_{y \in \mathbb{R}^n} f(y) \text{ subject to } y_i \geq 0 \quad i = 1 \ldots n
\]

by an unconstrained problem, depending on a parameter \(\mu > 0\):

\[
\min_{y \in \mathbb{R}^n} f(y) - \mu \sum_{i=1}^{n} \log(y_i) ,
\]

wherein the functional \(f\) has been extended by a sum of logarithmic barrier functions. This new minimization problem with minimizer \(y^\mu\) can then be solved approximately. By driving the barrier parameter to its limit \(\mu = 0\) we can approximate the solution of the original problem. The logarithmic terms assure strict feasibility \(y_i^\mu > 0\) for all \(\mu > 0\), so that Newton’s method can be applied to the corresponding first order optimality conditions in a feasible neighbourhood of \(y^\mu\).

If we want to extend this idea to infinite dimensional function spaces, e.g., for the problem

\[
\min_{y \in Y} f(y) \text{ subject to } y(x) \geq 0 \quad \forall x \in K ,
\]
then a straightforward idea might be to substitute the above finite sum by an integral. Along this line, the associated barrier regularization then would read
\[ \min_{y \in Y} f(y) - \mu \int_K \log(y(x)) \, dx. \]

However, we have to take into account two additional issues

(i) We have to achieve strict feasibility of \( y^\mu \) in the sense that \( y^\mu(x) \geq \varepsilon(\mu) > 0 \) for \( x \in K \) and for all \( \mu > 0 \).

(ii) The neighbourhood of \( y^\mu \) that contains the Newton iterates has to be taken with respect to the \( \| \cdot \|_\infty \) norm (or stronger). Any norm that is finite for a pointwise unbounded function yields neighbourhoods that are not contained in the feasible set.

Unfortunately, the logarithmic barrier functional guarantees strict feasibility of \( y^\mu \) only \textit{almost everywhere} in \( K \), and possibly allows touch points where \( y^\mu(x) = 0 \) for some \( \mu > 0 \), even when \( y^\mu \) is a smooth function. A simple example of such a case has been constructed in [83]. Of course, in such a case no feasible neighbourhood exists, even in the \( \| \cdot \|_\infty \)-norm.

A way out of this difficulty is shown by the following lemma:

\textbf{Lemma 4.1.} Let \( K \subset \mathbb{R}^d \) be a compact set, satisfying a cone property. Let \( y \geq 0 \) be in the space \( C^\beta(K) \) of Hölder continuous functions, \( 1/y^m \in L_1(K) \), \( 0 < \beta \leq 1 \). Assume that \( m \geq d/\beta \). Then \( 1/y \in C(K) \).

\textbf{Proof.} Assume that without loss of generality \( 0 \in K \) and \( y(0) = 0 \). We will show that this contradicts the assumption \( 1/y^m \in L_1(K) \). By the Hölder-continuity of \( y \) we infer that \( y(x) < cr^\beta \) for all \( x \in B(0,r) \cap K \). Using the cone property of \( K \), there is a cone \( C_0 \subset K \), and we can compute for sufficiently small \( R < 1 \)

\[ \| y^{-m} \|_{L_1(K)} \geq \int_{C_0} y(x)^{-m} \, dx \geq c \int_{[0,R]} r^{-\beta m} r^{d-1} \, dr \geq c \int_{[0,R]} r^{-1} \, dr = \infty, \]

which shows \( 1/y^m \notin L_1(K) \). Hence, by contradiction, \( y > 0 \) in \( K \), which implies by compactness of \( K \) that there is \( \psi > 0 \) with \( y \geq \psi \) and thus \( 1/y \in C(K) \). \( \square \)

Hence, in order to obtain strict feasibility of barrier minimizers in Hölder spaces, we may use a hierarchy of higher order barrier functions

\[ l_m(y) := \begin{cases} -\mu \log(y) & : m = 1 \\ \mu^m & : m > 1 \end{cases} \]

and solve

\[ \min_{y \in C^\beta(K)} f(y) + \int_K l_m(y(x)) \, dx. \]
Then, for sufficiently large $m$ and Hölder continuous $y$, existence of a touch point $y(x) = 0$ would imply

$$
\int_K l_m(y(x)) \, dx = \infty
$$

in contradiction to the optimality of $y$.

**BHT equation as operator equation in function space.** In order to study the weak form of the BHT equation \([15]\) in a functional analytic setting, we have to impose some regularity assumptions. Our set of assumptions has to cover the case of discontinuous coefficients, which arise due to the different types of tissues in the human body.

The presence of state constraints forces us to impose an $\| \cdot \|_{\infty}$-topology on the space of temperature distributions. For this reason, we have to depart from the usual $H^1$-framework for elliptic equations, since $H^1(\Omega) \not\subset C(\Omega)$ for spatial dimension greater than one.

To proceed, let $q > 3$, and define its dual exponent $q'$ via $1/q + 1/q' = 1$. Let $D_q \subset H^1(\Omega)$ be the set of all $T$, such that $A(T) \in W^{1,q'}(\Omega)^*$, i.e.

$$
\langle A(T), \varphi \rangle \leq M\|\varphi\|_{W^{1,q'}(\Omega)} \forall \varphi \in C^\infty(\Omega).
$$

We have to achieve that the operator

$$
A(\cdot) - B(\cdot) : D_q \times \mathbb{R}^{2kE-1} \to W^{1,q'}(\Omega)^*
$$

is well defined, and $D_q$ can be embedded into $C(\Omega)$. For a strictly monotone perfusion term $w(T, x)(T - T_{\text{blood}})$, general results from elliptic regularity theory (cf. \([14]\)) show that $A$ has a continuous inverse $A^{-1} : W^{1,q'}(\Omega)^* \to C(\Omega)$, i.e., $D_q \subset C(\Omega)$, and even $\|T\|_{C^\beta(\Omega)} \leq c\|A(T)\|_{W^{1,q'}(\Omega)^*}$ for some $\beta > 0$, where $C^\beta(\Omega)$ is the space of Hölder continuous functions. Moreover, $D_q$ only depends on the principle part of $A$, and is thus independent of $T$.

To assure that $B$ maps into $W^{1,q'}(\Omega)^*$, we have to impose a technical regularity assumption on the antenna profiles, which requires slightly more regularity than the usual one for Maxwell’s equation, namely $E_k \in H(\text{curl}, \Omega)$. We assume that each $E_k$ is contained in $L_q(\Omega, \mathbb{R}^3)$. Then $B(\alpha)$ is contained in $L_{q/2}(\Omega)$, which, in turn, is contained in $W^{1,q'}(\Omega)^*$ by the Sobolev embedding theorem. In this way singularities in the temperature distribution $T$ can be excluded. Such regularity has been shown under certain smoothness assumptions on the tissue interfaces in \([36, 35]\). Practical experience anyway shows that such singular temperature distributions do not occur in the hyperthermia setting. In this framework $A$ and $B$ are then twice continuously Fréchet differentiable as long as the perfusion term is twice differentiable with respect to $T$. The derivative of $A$ with respect to $T$,

$$
A' : C(\Omega) \ni D_q \to W^{1,q'}(\Omega)^*
$$
The function $A$ is not continuous, but it has a continuous inverse. Moreover, since the principal part of $A$ is linear, $A'(T) - A'(	ilde{T})$ does not contain a differential operator. These facts allow us to apply results, such as the inverse function theorem to $A$, and thus to establish continuous differentiability of the control to state mapping

$$\alpha \mapsto T(\alpha) := (A^{-1} \circ B)(\alpha).$$

Application of the barrier regularization to the pointwise state constraints

$$T(x) \leq T_{\lim}(x) \quad \forall x \in \Omega$$

immediately leads us to identify the above variable $y$ with $T_{\lim}(x) - T(x)$. Consequently, we arrive at the following barrier minimization problem:

$$f(T, \alpha) + \int_{\Omega} l_m(T_{\lim} - T) \, dx$$

subject to

$$A(T) - B(\alpha) = 0.$$ (47)

Our regularity considerations yield $T \in C^2(\Omega)$, and since $T_{\lim}$ is constant on each type of tissue, Lemma 4.1 applies separately on each tissue. With these considerations a generalization of the Weierstrass theorem on existence of minimizers yields (for details of the proof see [86]):

**Theorem 4.2.** There is $m < \infty$, such that for every $\mu > 0$ the barrier problem (47) has an optimal solution, which is strictly feasible with respect to the inequality constraints.

**First and second order optimality conditions.** For a strictly feasible barrier minimizer $(T, \alpha)$ first and second order optimality conditions have been derived in [86]. To this end, it is convenient to define the Lagrangian function of the problem:

$$L(T, \alpha, \lambda) := f(T, \alpha) + \int_{\Omega} l_m(T_{\lim} - T) \, dx + \langle \lambda, A(T) - B(\alpha) \rangle.$$

Here, since $A(T) - B(\alpha) \in W^{1,q}(\Omega)^*$, $\lambda$ is a Lagrangian multiplier in $W^{1,q'}(\Omega)$.

**Theorem 4.3 (KKT Conditions).** If $(T, \alpha)$ is a locally optimal strictly feasible solution of (47), then there exists a unique Lagrange multiplier $\lambda \in W^{1,q}(\Omega)$, such that

$$0 = F(T, \alpha, \lambda) := \begin{cases} f_T(T, \alpha) + \int_{\Omega} l'_m(T_{\lim} - T) \, dx + A'(T)^* \lambda, \\ f_\alpha(T, \alpha) - B'(\alpha)^* \lambda, \\ A(T) - B(\alpha). \end{cases}$$ (48)

42
Of course, $F$ also depends on the parameter $\mu$, since $l'_m$ does so. Since $A'(T)^*$ is a differential operator, the first line of this system is another (linear elliptic) partial differential equation, the adjoint equation and $\lambda$ is called adjoint state.

Our algorithm will be based on this version of the KKT conditions and can therefore be considered as a primal interior point method. Alternatively one could also introduce Lagrangian multipliers for the state constraints, which would lead to primal-dual interior point methods. These are known to be more efficient than their primal counterparts for finite dimensional problems. However, the analysis in function space yields that these multipliers can only be represented as Borel measures [20], which are very hard to treat numerically. Below, a specific pointwise damping step is presented that recovers some of the benefits of primal-dual methods.

**Theorem 4.4** (Second Order Optimality Conditions). Let $(T, \alpha, \lambda)$ be a solution of (48). Moreover, denote by

$$\mathcal{K} := \ker (A'(T), -B'(\alpha)) := \{(\delta T, \delta \alpha) : A'(T)\delta T - B'(\alpha)\delta \alpha = 0\}$$

the kernel of the linearised constraints and by

$$H(T, \alpha, \lambda)(\delta T, \delta \alpha)^2 := L_{TT}(T, \alpha, \lambda)\delta T^2 + L_{\alpha\alpha}(T, \alpha, \lambda)\delta \alpha^2$$

(49)

the Hessian of the Lagrangian.

(i) If $(T, \alpha)$ is a minimizer of (47), $H(T, \alpha, \lambda)$ is positive semi-definite on $\mathcal{K}$.

(ii) $H(T, \alpha, \lambda)$ is positive definite on $\mathcal{K}$, if and only if $(T, \alpha)$ is a local minimizer of (47) and $J$ satisfies a local quadratic growth condition.

Then for each $(r_1, r_2, r_3) \in H^1(\Omega)^* \times \mathbb{R}^{2k_E-1} \times W^{1,q'}(\Omega)^*$ the linear system

$$\begin{pmatrix}
L_{TT}(T, \alpha, \lambda) & 0 & A'(T)^* \\
0 & L_{\alpha\alpha}(T, \alpha, \lambda) & -B'(\alpha)^* \\
A'(T) & -B'(\alpha) & 0
\end{pmatrix}
\begin{pmatrix}
\delta T \\
\delta \alpha \\
\delta \lambda
\end{pmatrix}
= \begin{pmatrix}
r_1 \\
r_2 \\
r_3
\end{pmatrix}$$

(50)

has a unique solution $(\delta T, \delta \alpha, \delta \lambda) \in D_q \times \mathbb{R}^{2k_E-1} \times H^1(\Omega)$, depending continuously on $(r_1, r_2, r_3)$.

Finally, in

$$L_{TT}(T, \alpha, \lambda)\delta T^2 = f_{TT}(T, \alpha)\delta T^2 + \int_\Omega l'_m(T - T)\delta T^2 \, dx + \langle \lambda, A'(T)\delta T \rangle$$

(51)

we encounter the second derivative $l'_m$ of the barrier term. This factor can be very large if $T$ is close to its upper bound, and thus induces a strong scaling to be taken into account in solution algorithms.
Central path. The solvability of (50) in a neighborhood of a locally unique solution according to Theorem 4.4 allows us to compute Newton steps for the solution of the KKT system (48). Here, it is important to obtain $\delta T \in D_q \subset C(\Omega)$, instead of the weaker result $\delta T \in H^1(\Omega)$. This ensures that Newton’s method stays in an $L_\infty$-neighborhood of the minimizer and yields feasible iterates.

Introducing the shorthand notation $z := (T, \alpha, \lambda)$, we can formulate

**Corollary 4.5.** If $H(z)$ is positive definite on $\ker (A', -B')(T, \alpha)$, Newton’s method, applied to $F(z)$ converges locally superlinearly to a zero $z^\mu$ of $F$.

**Proof.** Since $F'$ depends continuously on $z$, we may just use a qualitative local convergence result for Newton’s method (see, e.g., [72, Thm. 10.2.2]). \qed

In particular, the implicit function theorem yields local existence and differentiability of the so called central path of barrier minimizers:

**Corollary 4.6.** If $H(z)$ is positive definite on $\ker (A', -B')(T, \alpha)$, then, locally, there is a differentiable path $\mu \mapsto z^\mu$ of local minimizers of the barrier problems, defined in some open interval $[\mu, \mu]$.

A convergence theory for the path of barrier minimizers has been established for the case of a linear partial differential equation as equality constraint and a convex functional. In this case, estimates of the form

$$f(T^\mu, \alpha^\mu) - f(T_{\text{opt}}, \alpha_{\text{opt}}) \leq C\mu$$

have been established, which implies under a quadratic growth condition on $\alpha$ the estimate

$$\|T^\mu - T_{\text{opt}}\|_{L_\infty(\Omega)} + |\alpha^\mu - \alpha_{\text{opt}}|_{\mathbb{R}^{2k_E - 1}} \leq C\sqrt{\mu}.$$ 

Furthermore, the qualitative result of Corollary 4.5 can be quantified in this case, giving a $\mu$-dependent lower bound on the radius of convergence. Then a global convergence result can be established (cf. [82]) for a barrier path-following method.

In hyperthermia, the equality constraints are nonlinear, and so our problem is generically non-convex. However, in a neighborhood of the central path, under second order sufficient optimality conditions, our barrier problems behave like convex ones, so that we can expect these results to carry over. Numerical results give clear evidence to this conjecture, as can be seen in Figure 19.

From an algorithmic point of view, the second order optimality conditions yield well definedness of a Newton path-following method, as presented in the following section, once an initial guess close to the central path is found. To find such a point, algorithms have to be used that are robust with respect to non-convexity. For our numerical experiments we employ a new algorithm (see the forthcoming report [87]) that is based on the idea of a composite step method (cf. e.g. [21]) with a cubic regularization [104] of the functional and an affine covariant treatment of the constraints.
4.2 Adaptive Central Path Following in Function Space

The previous section laid the ground for a Newton path-following method with respect to a barrier parameter $\mu$. The aim of this section is to describe the development of an efficient algorithm based on this principle. We base our algorithmic approach on an affine covariant adaptive path-following method, as described in [27, Chapter 5]. However, compared to the standard case, additional difficulties appear:

- Our problem is posed in function space, and thus Newton steps cannot be computed exactly, which means that an inexact Newton corrector has to be used.

- The radius of convergence of Newton’s method tends to zero as $\mu \to 0$. Thus, the subproblems become more and more difficult to solve. This has to be taken into account in an adaptive update strategy for $\mu$.

An algorithm that takes these issues into account will consist of three loops:

(i) In the outer path-following loop, the barrier parameter $\mu$ is reduced successively. This requires an adaptive strategy.

(ii) Each step of the outer loop invokes an inexact Newton method in function space. Here we have to design termination criteria, and criteria on the success of the Newton iteration. An estimate of the local contraction quantity $\Theta(z)$ is crucial in this respect.

(iii) In the inner loop each Newton step is computed up to a desired relative accuracy. Typical sources of errors are truncation errors of iterative solvers, but also discretization errors which arise during the approximate computation of Newton steps in function space by finite element discretizations. We have to devise error estimators that are suited to the particular problem structure.
**Local convergence of an inexact Newton method.** A local affine covariant convergence theory of inexact Newton methods lays the groundwork for our algorithmic considerations. A detailed discussion of this topic can be found in [27]. For the convenience of the reader we give a simplified presentation, based on the analysis of [84].

The starting point is the following very simple local convergence result for Newton’s method. It is a generalization of the affine covariant refined Newton-Mysovskikh theorem due to [30] that can be applied to semi-smooth systems $F$ as well. It is formulated exclusively in terms of quantities in the domain space of the problem, i.e. affine covariant, and covers the inexact computation of Newton steps.

**Theorem 4.7.** Let $R$ be a linear space, $Z$ a normed space, $D \subset Z$ and $F : D \to R$. Assume there is $z^* \in D$ with $F(z^*) = 0$. For given $z \in D$ assume that there exists an invertible linear mapping $F'(z)(\cdot) : Z \to R$ such that

$$\Theta(z) := \frac{\|F'(z)^{-1}[F'(z)(z - z^*) - (F(z) - F(z^*))]\|}{\|z - z^*\|}. \quad (53)$$

is well defined. Assume that an inexact Newton step results in

$$z_+ := z - F'(z)^{-1}F(z) + e,$$

where the relative error $\gamma(z) := \|e\| / \|z - z^*\|$ is bounded by

$$\gamma(z) + \Theta(z) \leq \beta < 1. \quad (54)$$

Then

$$\|z_+ - z^*\| \leq \beta \|z - z^*\|. \quad (55)$$

**Proof.** We compute for one inexact Newton step:

$$\|z - z^*\| = \|z - F'(z)^{-1}F(z) + e - z^*\|
\leq \|F'(z)^{-1}[F'(z)(z - z^*) - (F(z) - F(z^*))]\| + \|e\|.$$

Inserting the definition of $\Theta(z)$ and assumption (54) we obtain

$$\|z - z^*\| \leq (\Theta(z) + \gamma(z)) \|z - z^*\| \leq \beta \|z - z^*\|. \quad \Box$$

Equations (54) and (55) show that local convergence of Newton’s method is governed by the local contraction rate $\Theta$ and the relative accuracy $\gamma$. As we will show, in an adaptive Newton path-following algorithm both quantities can be controlled algorithmically.

To be able to react adaptively on the nonlinearity of the problem we have to provide computable estimates for $\Theta$ and $\gamma$. We will construct an estimate for $\gamma$ in the paragraph about adaptive refinement, below.
Next we derive a computable estimate for $\Theta(z)$. From (53) we observe that $\Theta(z)$ cannot be computed exactly, unless $z_*$ is known. However, with the help of a simplified Newton step:

$$\delta z := -F'(z)^{-1}F(z_+)$$

we can replace the solution $z_*$ in (53) by the new iterate $z_+$. Then compute

$$[\Theta(z)] := \frac{\|F'(z)^{-1}[F'(z)(z - z_+) - (F(z) - F(z_+))])\|}{\|z - z_+\|} = \frac{\|\delta z\|}{\|\delta z\|}.$$  (56)

In view of local quadratic convergence of Newton’s method, we may define a parametrized model for $\Theta(z)$ as follows

$$[\tilde{\Theta}(z)] := \left[\frac{\omega}{2}\right] - \|z - z_*\|.$$  (57)

The constant $[\omega]$ can be computed for given $z$ via (56):

$$[\omega] := \frac{2[\Theta(z)]}{\|z - z_+\|} = \frac{2\|\delta z\|}{\|\delta z\|^2}.$$  (58)

Then $[\omega]$ can be interpreted as an estimate from below for an affine covariant Lipschitz constant $\omega$, as used in [27]. This gives an estimate for the radius of contraction, i.e., the radius of the largest ball around $x_*$, where a contraction $\Theta(z) \leq \Theta$ takes place:

$$r_{\Theta} \geq \frac{2\Theta}{[\omega]}.$$  (59)

Replacing $\omega$ by our estimate $[\omega]$ we get a quantitative estimate on the size of the region of convergence. Moreover, the remaining error via of Newton’s method satisfies the triangle inequality:

$$\|z_+ - z_*\| = \Theta(z)\|z - z_*\| \leq \Theta(z)([z - z_+\| + \|z_+ - z_*\|),$$

and thus, solving for $\|z_+ - z_*\|$ and replacing $\Theta(z)$ by $[\Theta(z)]$, we obtain the estimate

$$[\|z_+ - z_*\|] := \left[\frac{\Theta(z)}{1 - [\Theta(z)]}\right] \|z - z_+\|.$$  (58)

for the remaining error.

Adaptive reduction of barrier parameter. The geometric picture of a successful path-following method is sketched in Figure 20. For some prescribed desired contraction $\Theta_d < 1$ (in practice one chooses $\Theta_d \in [0.1, 0.5]$), each iterate $z_j$ should remain in the region of convergence around a target solution $z^{\mu_j}$ which lies on the central path, i.e.,

$$\|z_j - z^{\mu_j}\| \leq r_{\Theta_d}(\mu_j).$$  (59)
Figure 20: Sketch of a Newton path-following method. In each corrector step the iterates approach a target point on the central path and eventually enter the region of convergence of the next target.

We want to achieve this inductively by a suitable adaptive choice of $\mu_j$. First of all, $r_{\Theta_{\mu}}$ can be estimated via (57), if we assume a relation between $\omega$ and $\mu$. Numerical experiments, and estimates for related problem classes [105] suggest the relation $\omega \sim \mu^{-1/2}$.

To obtain a choice of $\mu_j$ that yields (59), we also have to provide an estimate for

$$\|z_j - z^\mu\| \leq \|z_j - z^{\mu_{j-1}}\| + \|z^{\mu_{j-1}} - z^\mu\|.$$  

The first term of the right hand side is the remaining error of the last Newton corrector and can be estimated via (58). The second term depends on the sensitivity of $z^\mu$ with respect to $\mu$, and we have the estimate

$$\|z^{\mu_{j-1}} - z^{\mu_j}\| = \left\| \int_{\mu_j}^{\mu_{j-1}} \frac{d}{d\mu} z^\mu \, d\mu \right\| \leq \int_{\mu_j}^{\mu_{j-1}} \left\| \frac{d}{d\mu} z^\mu \right\| d\mu. \quad (60)$$

For finite dimensional problems or for fixed discretizations, we may estimate this slope simply by finite differences in the iterates. If, as in our approach, adaptive grid refinement is performed during the corrector, finite differences are not reliable due to the presence of discretization errors. Then, at the cost of one system solve, an estimate $[\eta(\mu_{j-1})]$ can be computed via the formula

$$\frac{\partial}{\partial z} F(z^{\mu_{j-1}}) \frac{d}{d\mu} z^\mu - \frac{\partial}{\partial \mu} F(z^{\mu_{j-1}}),$$
where we have to replace $z^{\mu-1}$ by its approximation $z_j$. Then, using the relation $\eta(\mu) \sim \mu^{-1/2}$, we may compute the estimate

$$
\|[z^{\mu_j-1} - z^{\mu_j}]\| := \int_{\mu_j}^{\mu_{j-1}} [\eta(\mu_{j-1})] \sqrt{\mu_{j-1} - \mu} \, d\mu
= 2[\eta(\mu_{j-1})] \sqrt{\mu_{j-1} - \mu_j}.
$$

To summarize, we may write down (59) as an implicit inequality in $\mu_j$: \[ ||z_j - z^{\mu_j-1}|| + 2[\eta(\mu_{j-1})] \sqrt{\mu_{j-1} - \mu_j} \leq 2\Theta \sqrt{\sigma_j}. \tag{61} \]

and compute the smallest $\mu$ that satisfies this inequality as a prediction for $\mu_j$. In the subsequent inexact Newton corrector the estimated error is reduced up to a tolerance $TOL_{corr}$, which is chosen in a way that usually one corrector step is sufficient, so that we terminate, if

$$
\|[z_j - z^{\mu_j-1}]\| \leq TOL_{corr} := \rho \frac{2\Theta_d}{\omega(\mu_{j-1})} \sqrt{\mu_{j-1}}, \quad \rho \in [1, 1.2].
$$

This rule suggests a multiplicative update strategy. Define $\sigma_j := \mu_j / \mu_{j-1}$. Then (61) can be written as an implicit inequality in $\sigma_j$

$$
\|[z_j - z^{\mu_j-1}]\| \omega(\mu_{j-1}) + 2[\eta(\mu_{j-1})] \omega(\mu_{j-1}) (1 - \sqrt{\sigma_j}) \leq 2\Theta_d \sqrt{\sigma_j}. \tag{62} \]

Under our model assumptions $[\omega] \sim \mu^{-1/2}$, $[\eta] \sim \mu^{-1/2}$, which implies $TOL_{corr} \sim \sqrt{\mu}$ we observe that $\mu_{j-1}$ cancels out in (62), and thus $\sigma$ is constant. Of course our model assumptions only hold approximately in computational practice, but their use leads to an unbiased and strictly positive update $\mu_{j-1} \rightarrow \mu_j > 0$.

If it terminates successfully, we accept the result of the corrector as new iterate $z_j$ and compute $\mu_{j+1}$ as new homotopy parameter, just as before. In case of unsuccessful termination (for example, if $[\Theta(z)] > 1$ for some iterate $z$), with the additional information gained on $[\omega]$ in this correction step, a more conservative choice for $\mu_j$ can be computed and the corrector is restarted at $z_{j-1}$.

**Local norms for barrier subproblems.** Up to now our considerations have not included any special structure of our problem. A common possibility to do this is the choice of a problem suited norm. In the context of barrier methods the strong scaling property of the Hessian, as observed in (51) should be taken into account. Thus, (for fixed $\mu$) a suitable local norm can be defined as follows:

$$
\|\delta z\|_2^2 := \|\delta T\|_{L_2(\Omega)}^2 + \int_{\Omega} l_m^\prime (T_{lim} - T) \delta T^2 \, dx + \|\delta \alpha\|_{H_{\infty}}^2 + \|\delta \lambda\|_{L_2(\Omega)}^2. \tag{63}
$$

Qualitatively speaking, the inclusion of the barrier function $l_m$ into the norm has the effect that errors are given more weight in regions of $\Omega$, where $T$ is close...
to its upper bound $T_{\text{lim}}$. Thereby it is taken into account that the problem is more nonlinear and thus more sensitive to errors in these regions.

In practical computations, the multiplier term $\|\delta \lambda\|_{L^2(\Omega)}^2$ can be omitted. The reason is that $\lambda$ can be reconstructed approximately from the primal variables $T, \alpha$ by suitable multiplier update rules (cf. [21]) and plays the role of an auxiliary variable.

In a neighbourhood of the central path, where positive definiteness of the Hessian $H$ of the Lagrangian from (49) can be assumed on $\ker(A', -B')$, the generic $L_2$ norms of $\delta T$ and $\delta \alpha$ in (63) can be substituted by the norm induced by $H$ for the tangential components, which dominate during the homotopy due to the mild nonlinearity of the state equation. This leads to the affine conjugate norm

$$\|\delta z\|_{H(z)}^2 := H(z)(\delta T, \delta \alpha)^2.$$ (64)

The normal components can also be taken into account if desired, see e.g. [100].

As the local norm changes in every Newton step, contraction $\Theta < 1$ for a fixed norm need not translate to contraction between different local norms. Whether this is guaranteed to hold depends on the actual impact of $z$ on $\|\cdot\|_{z}$. For details, we refer to [27, 100].

**Pointwise damping.** Barrier methods rely on iterates that are feasible with respect to the inequality constraints. Since in the barrier context Newton’s method approximates a rational function by a linear one, Newton steps tend to violate the constraints. On the other hand, the nonlinearity, introduced by the barrier functions is given in a pointwise fashion and highly structured. This should be exploited.

In the following we propose a modification of Newton’s method, which addresses these issues and may be considered as a pointwise damping strategy. The idea exploits the pointwise structure of the problem and guarantees strict feasibility of the iterates. In the whole discussion $\mu > 0$ is fixed.

For what follows, define the Lagrangian function of the unconstrained problem:

$$\tilde{L}(z) := f(T, \alpha) + \langle \lambda, A(T) - B(\alpha) \rangle.$$ Further, let $z_+ = (T_+, \alpha_+, \lambda_+)$ be the possibly infeasible next Newton iterate, which satisfies

$$F(z) + F'(z)(z_+ - z) = 0.$$ (65)

Our principle idea is to construct a modified feasible iterate $z_C = (T_C, \alpha_+, \lambda_+)$ that satisfies the first row of (48), i.e.

$$\tilde{L}_T(z_C) + \iota_{\mu}(T_{\text{lim}} - T_C) + A'(T_C)^*\lambda_+ = 0.$$ (66)

This is an equation in $(H^1)^*$, which seems to be difficult to solve for $T_C$. However, if we subtract the first row of (65) from (66) we observe that the temperature independent differential operators in $A'(\cdot)^*$ cancel out, and we obtain a pointwise equation in $T_C$ that depends on $z$ and $z_+$. If we neglect those higher
order terms that are independent of $l_m$ we end up with a pointwise equation of the form
\[ \tilde{L}_{TT}(z)T_C + l'_m(T_{\text{lim}} - T_C) = r(z, z_+). \]
This equation can be solved for $T_C$ at each node of the discretization. For scalar equations there are efficient and robust algorithms available \( \text{e.g.} \ [18] \), and the computational cost of performing this step is marginal compared to the overall computation time.

To obtain a pointwise damping step we use only those $T_C = T_C(x)$, for which $|T_C - T| \leq |T_+ - T|$, these are exactly those, for which $T_+ \geq T_C \geq T$ holds. Hence, we are able to compute a strictly feasible corrected iterate $T_C$ from a possibly infeasible iterate $T_+$ in a natural way. It can be shown that
\[ \lim_{\|T - T^0\|_{\infty} \to 0} \frac{\|T_C - T_+\|_{\infty}}{\|T - T_+\|_{\infty}} = 0. \]
This implies that undamped steps are recovered asymptotically, close to a strictly feasible solution. An estimate for the Newton contraction $\Theta(z)$ can be computed in analogy to the case, described above.

Our pointwise damping strategy has a marked effect on the numerical performance of the path-following method. As Figure 21 indicates, an algorithm without pointwise damping shows relatively slow linear convergence behaviour for an academic test problem. In contrast, with pointwise damping the reduction of $\mu$ often takes place at a superlinear rate. This is particularly beneficial in the context of our hyperthermia problem.

Figure 21: Comparison of path-following algorithm with and without pointwise damping. The horizontal axis: barrier parameter $\mu$. Vertical axis: reduction factor $\sigma := \mu_+ / \mu$. For pointwise damping one can observe $\sigma \to 0$ for $\mu \to 0$. 

10
−8
10
−7
10
−6
10
−5
10
−4
10
−3
10
−2
10
−1
10
0
0.1
0.2
0.3
0.4
0.5
0.6
0.7
0.8
0.9
1
µ
σ
no pw. damping
pw. damping, conservative
pw. damping, aggressive
Discretization and adaptive grid refinement. So far, we assumed that our algorithm can compute Newton steps in function space exactly. However, this is not realistic, because the Newton steps have to be discretized for computation. Discretization of the temperature $T$ and the Lagrange multiplier $\lambda$ is done straightforwardly by linear finite elements, and the precomputed electric fields are discretized using edge elements of first order, as described in Section 3.

As an additional numerical difficulty the barrier integrals and their derivatives cannot be evaluated exactly. However, it can be shown that the use of the trapezoidal rule is sufficient to obtain satisfactory a-priori error estimates (cf. [48]).

For a-posteriori error estimation and adaptive grid refinement we rely once more on hierarchical error estimators as already presented in Section 3.1.

Discretizing (50) with an extended ansatz space, we obtain a block linear system of the form (33):

$$
\begin{pmatrix}
F'(z)^{LL} & F'(z)^{LH} \\
F'(z)^{HL} & F'(z)^{HH}
\end{pmatrix}
\begin{pmatrix}
\delta z^L \\
\delta z^H
\end{pmatrix}
= -
\begin{pmatrix}
F(z)^L \\
F(z)^H
\end{pmatrix}.
$$

(67)

The only difference is that each block of this system is in turn a block matrix of the form (50). Finally, as in Section 3.1 we end up with the problem of solving the block linear system

$$
F'(z)^{HH} \delta z^H = r^H := -(F(z)^H + F'(z)^{HL} \delta z^L).
$$

(68)

Once $\delta z^H$ is computed, one can compute cheaply an estimate in the affine conjugate norm defined in (64). If $\delta z^H = (\delta T^H, \delta \alpha^H, \delta \lambda^H)$, define $\delta z^H := (\delta T^H, \delta \alpha^H, -\delta \lambda^H)$. Then one can compute via (50) that

$$
[\|e\|_{H(z)}]^2 := \langle \delta z^H, r^H \rangle = \langle \delta z^H, F'(z)^{HH} \delta z^H \rangle = \|\delta z^H\|^2_{H(z)}.
$$
because the off-diagonal blocks of $F'(z)H^H$ cancel out. Hence, we can estimate
the error in the same norm that is used to measure the length of the Newton
steps.

Now we can use these estimates to control the relative error

$$\gamma(z) := \frac{\langle \delta z^H, \tilde{r}^H \rangle^{1/2}}{\|\delta z^L\|_{H(z)}}$$

by adaptive mesh refinement. The refinement loop is stopped as soon as the
estimate for $\gamma(z)$ drops below a given relative tolerance $TOL_{\text{disc}} \in [0.1, 0.5]$. A closely related adaptive refinement scheme results from goal-oriented error
estimation for optimal control problems looking at the error in the cost func-
tional [11, 102].

In the context of Theorem 4.7 it can be seen that the corresponding inexact
Newton iteration will converge asymptotically linearly in function space. A
superlinear or quadratic convergence rate is not aimed at, since this usually leads
to needlessly high accuracy and a corresponding dramatic increase of unknowns,
impeding the efficiency of the algorithm.

**Solution of linear systems.** Finally, we have to solve discretized versions of
the system (50) in order to compute the discretized Newton steps, and also to
solve the defect equation (68).

This can be done by reducing it to a linear system in the control $\alpha \in \mathbb{R}^{2k_E-1}$
via a Schur complement approach. To obtain the reduced linear system, each
of the discretized PDEs $A'(T)v = r$ and $A'(T)^*w = s$ has to be solved for
$2k_E - 1$ right hand sides. For a wide range of discretizations a sparse direct
solver (e.g., MUMPS [3]) performs quite efficiently. Alternatively, one can solve
the PDEs iteratively up to a certain accuracy by a preconditioned conjugate
gradient method and perform an outer iteration to obtain a solution of the total
system.

In particular, when solving (68) approximately, the solution of the PDEs
$A'(x)v = r$ and its adjoint can be found iteratively via a Jacobi preconditioned
conjugate gradient method.

Putting all pieces together, which have been described above, we now arrive
at the following pseudo-code.
Algorithm AdaptivePathfollowing($z_0, \mu_0$) (Initial Guess)

for $j = 1, \ldots$ (Path-following Loop)
  ($\tilde{z};$ success) $\leftarrow$ InexactNewtonCorrector($z_{j-1}; \mu_{j-1}$)
  if success
    compute new $\mu_j < \mu_{j-1}$
    $z_j \leftarrow \tilde{z}$
  else
    compute corrected $\mu_j > \mu_{j-1}$
    $z_j \leftarrow z_{j-1}$

Subroutine InexactNewtonCorrector($z; \mu$):

do (Newton Iteration Loop)
  do (Inexact Newton Step)
    RefineMarkedElements
    $z_+ := z - F_h'(z)^{-1} F_h(z)$
    $\|e\|_{H(z)} := $ ErrorEstimator:Discretization($z$)
    MarkElements
  while $\|e\|_{H(z)} > \text{TOL}_{\text{disc}} \|z_+ - z\|
  compute a pointwise damped iterate $z_C$
  compute $[\Theta(z)]$
  compute $\|z_C - z\|$
  success = ($\|z_C - z\| < \text{TOL}_{\text{corr}}$)
  failure = ($\Theta(z) > \text{RequiredContraction}$)
  $z \leftarrow z_C$
  while not(success $\lor$ failure)
return($z; \text{success}$)

5 Impact on Medical Technology

Already the earlier versions of our virtual lab HyperPlan have led to a significant impact in the treatment of deep regional hyperthermia, part of which we will present here.

5.1 Redesign of Applicator

During the past ten years the applicator most frequently used for regional hyperthermia had been the Sigma-60 applicator of BSD Medical Corp., Salt Lake City, Utah \footnote{meanwhile Munich, Germany} - see Fig. 23 left. This applicator consists of 8 antennas (i.e.
$k_E = 4$ channels) arranged on a ring with a diameter of 60 cm, hence its name. The space between the antenna ring and the patient’s body (see Fig. 1 left, in reality and Fig. 23 left, as computer image) is filled with a water bolus containing de-ionized water. In early 1988, based on results of our first simulations, the new Sigma-Eye applicator (see Fig. 23 right), was introduced. It has 24 antennas (i.e. $k_E = 12$ channels) arranged in three parallel rings thus allowing for an additional power steering along the patient’s axis. The much smaller bolus volume has an “eye”-shaped cross-section, hence the name.

Figure 23: Applicator design. Left: Old applicator Sigma-60 with 8 antennas ($k_E = 4$) and circular cross section, compare also Fig. 1 right. Right: New applicator Sigma-Eye with 24 antennas ($k_E = 12$) and eye-shaped cross section, designed on the basis of our simulations.

In order to illustrate the progress, let us document some comparative results. In Table 3, the computation times for the whole simulation process (on a SUN UltraSparc), based on the linear BHT as heat transfer model, are listed.

Table 3: Coarse grid computation times (SUN UltraSparc) for the whole simulation process with linear BHT equation. From [31].

<table>
<thead>
<tr>
<th></th>
<th>Sigma-60 $k_E = 4$</th>
<th>Sigma-Eye $k_E = 12$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grid Generation</td>
<td>15 min</td>
<td></td>
</tr>
<tr>
<td>Field Calculation</td>
<td>80 min</td>
<td>120 min</td>
</tr>
<tr>
<td>Temperature Calculation</td>
<td>2 min</td>
<td>20 min</td>
</tr>
<tr>
<td>Optimization</td>
<td>6 sec</td>
<td>1 min</td>
</tr>
</tbody>
</table>

Note that, essentially due to its smaller bolus volume, the field calculation time per channel of the Sigma-Eye is only about half the one of the Sigma-60. Moreover, recall from the cost model (44) that the temperature calculation times roughly increase with $k_E^2$, whereas the Maxwell solves enter with $O(k_E)$. 

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In Table 4, the progress in hyperthermia is documented at three patients with typical tumour locations. Not only in these cases new applicator has proven to significantly improve tumour heating, keeping “hot spots” in healthy tissue at a constant level or slightly reducing them. By the way, the second location is the most difficult one since the tumour is to a large extent enclosed by bony structures.

Table 4: Comparison of old applicator Sigma-60 and the new applicator Sigma-Eye for three patients with different tumour locations. From [31].

<table>
<thead>
<tr>
<th>part of tumour volume heated to above 43°C</th>
<th>Sigma-60</th>
<th>Sigma-Eye</th>
</tr>
</thead>
<tbody>
<tr>
<td>distal (supraanal) rectal carcinoma</td>
<td>17.5%</td>
<td>62.5%</td>
</tr>
<tr>
<td>highly presacral rectal carcinoma</td>
<td>0.7%</td>
<td>18.4%</td>
</tr>
<tr>
<td>cervical carcinoma at pelvic wall</td>
<td>24.8%</td>
<td>49.1%</td>
</tr>
</tbody>
</table>

Figure 24: Optimal temperature distribution for some rectum carcinoma at a frequency of 100 MHz. Left: Old applicator (Sigma-60). Right: New applicator (Sigma-Eye).

In Fig. 24 we illustrate the progress in the resulting therapy by visualizations of the optimal solution in both cases.

Frequency study. First frequency studies have already been performed in 1999 by Paulsen et al. [73] for 100, 150, and 200 MHz, but only for power functionals like (21); for their applicator test cases, they came up with an “optimal” frequency of about 150 MHz. In Fig. 25 we show our results for the new...
Figure 25: Optimal temperature distribution in new applicator for some rectum carcinoma at frequencies different from the one in Fig. 24 right. Left: 150 MHz. Right: 200 MHz.

Over all our simulations in various patient cases, a frequency around 150 MHz also appeared out to be preferable.

5.2 Case Study: Femoral Tumour

In order to give an impression of the computational impact by the new optimization techniques, we finally present a case study. The anonymous patient data belong to some femoral tumour, see Fig. 5.2 top left. Among the various objectives from Section 2.4 we selected the Arrhenius functional (24) inside the tumour. Outside the tumour, in healthy tissue, we imposed the strict restriction $T \leq 44^\circ C$ and a mild quadratic penalty term, if tissue were heated above $41^\circ C$. To treat the unsure tissue around the tumour, we coated the tumour by some region (of width just one coarse mesh size): for this region we replaced the constraints by a strong quadratic penalty term – thus smoothing the temperature profiles to be computed. The $2k_E$ electrical fields have been computed off-line once; they are a property of the given applicator. In order to satisfy the temperature constraints in healthy tissue, we applied the barrier algorithm from Section 4. The intermediate values of the barrier parameter $\mu$ are automatically determined from some adaptive control as described in Section 4.2. In the present example, the path-following method started at $\mu = 0.5$ and took the number of $\mu$-steps documented in Table 5 down to to $\mu = 10^{-6}$. The state, shown in Figure 5.2 bottom left, was reached already after 7 $\mu$-steps. The remaining 39 steps have merely been computed for reference purposes. In real world applications, one, of course, employs an automatic termination criterion.

Numerical central path following. The computational homotopy chain for $\mu \to 0$ started at $\mu = 0.5$ and a conservative solution (no 43°C isothermal surface active), see again Fig. 5.2 top left. Upon examination of the other figures
in Fig. 5.2, one observes that intermediate steps may well supply interesting planning options, depending on additional medical information to be taken into account, which opens new possibilities to the medical staff in charge of the hyperthermia treatment: For large $\mu$, the thermal dose is lower than for smaller values. For smaller and smaller $\mu$, the appearance of hot spots becomes clearly visible. However, due to the barrier regularization, these regions are still below the prescribed upper bounds of $44^\circ C$. Hence, some individual weighting of the occurrence of hot spots versus a better covering of the tumour by the temperature isosurface may be done.

**Mesh history.** In contrast to the old penalty approach, the new approach via the PDE constrained optimization solves the full nonlinear problem including adaptive mesh refinement within one algorithmic sweep. The geometry of the tissue regions is adjusted during mesh refinement. For practical reasons, the maximal number of degrees of freedom had to be restricted. Table 5 gives details about three refinement levels at different barrier parameters $\mu$.

Table 5: Computational effort spent on the therapy planning problem for the femoral tumour. The first line represents a computation on the coarse grid.

<table>
<thead>
<tr>
<th>tetrahedra</th>
<th>unknowns</th>
<th>$\mu$-steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>74457</td>
<td>28691</td>
<td>39</td>
</tr>
<tr>
<td>617178</td>
<td>222972</td>
<td>46</td>
</tr>
<tr>
<td>3658357</td>
<td>1275057</td>
<td>55</td>
</tr>
</tbody>
</table>

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Figure 26: Femoral tumour: Temperature distributions along the numerical central path. Light grey grid: bone, medium grey grid: tumour, dark grey surface: isothermal surface for 43°C. Top left: Just bone and tumour (identical with \( \mu = 0.5 \)). Top right: \( \mu = 0.44 \). Center left: \( \mu = 0.42 \). Center right: \( \mu = 0.38 \). Bottom left: \( \mu = 0.26 \). Bottom right: \( \mu = 10^{-6} \). Obviously, the difference to the bottom left result is marginal.
References


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