

### **Linearized General Relativity in Hyperboloidal Coordinates**

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### **Physics Engineering**

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#### Declaration

I declare that this document is an original work of my own authorship and that it fulfills all the requirements of the Code of Conduct and Good Practices of the Universidade de Lisboa.



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A few words about the university, financial support, research advisor, dissertation readers, faculty or other professors, lab mates, other friends and family...

#### Resumo



Palavras-chave: palavra-chave1, palavra-chave2, palavra-chave3,...



# Abstract

Insert your abstract here with a maximum of 250 words, followed by 4 to 6 keywords.

**Keywords:** keyword1, keyword2, keyword3,...



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## **Chapter 1**

### Introduction

#### 1.1 Motivation

Gravitational waves have been in the minds of physicists since Albert Einstein first predicted their existence in 1916 as perturbations in the curvature of spacetime that propagate at the speed of light. However, it was not until nearly a century later, in 2015, that the Laser Interferometer Gravitational-Wave Observatory (LIGO) made the first direct detection of this fascinating phenomenon [1]. This ground-breaking observation not only confirmed a key prediction of Einstein's general theory of relativity but also inaugurated the field of gravitational-wave astronomy, opening an entirely new window through which to observe and understand the universe.

Gravitational waves immediately caught the attention of the scientific community due to their unique interaction with matter. Unlike electromagnetic waves, which can be absorbed, scattered, or reprocessed by intervening media, gravitational waves propagate essentially unimpeded from their astrophysical sources to Earth. This property allows them to carry direct information about highly energetic and compact systems, including binary black hole mergers, neutron star collisions, and potentially processes in the early universe that are otherwise inaccessible to conventional astronomical methods.

Even though this new method of observation opens the door to a wealth of scientific opportunities, it also presents significant challenges. The signals detected by LIGO and other gravitational wave observatories are incredibly weak by the time they reach Earth, necessitating extremely sensitive detectors and sophisticated data analysis techniques to extract meaningful information from the noise. In order to interpret these signals accurately, it is crucial to have precise theoretical models of the gravitational waveforms produced by various astrophysical events. This is where numerical relativity comes into play, providing the tools to simulate the complex dynamics of spacetime and matter under extreme conditions. However, given the distance of Earth from the sources of gravitational waves, to predict the signals that detectors will observe, we must be able to extract the gravitational waves at null infinity, which is a significant challenge in itself.

State-of-the-art numerical relativity codes, such as the Einstein Toolkit [2], do not have this capability built in, working around this issue by calculating the wave front at different radii and then extrapolating

the results until future null infinity, which naturally comes at the cost of some amount of systematic error on the results. In this work, we present an alternative to the current methodology by evolving toy models for gravitational waves in a compactified spacetime, thereby bypassing the need for extrapolation.

#### 1.2 Topic Overview

- Anil proposed hyp layers
- Some work was done by Christian / David / Alex / Anil
- I continue to try and bridge the gap by implementing a toy model for gravitational waves in a compactified spacetime in full 3D
- It may also be important toward developing PDE theory since it is one of the few PDE solver that is spectral full 3d and with AMR for the cubic wave equation
  - mention objectives and their importance

#### 1.3 Objectives and Deliverables

Throughout this work, we aim to:

- Solve the wave equation in hyperboloidal slices using spectral methods in full 3D with Adaptive Mesh Refinement (AMR);
- · Solve the wave equation in hyperboloidal layers using spectral methods in full 3D with AMR;
- Solve the wave cubic equation in hyperboloidal layers using spectral methods in full 3D with AMR;
- Study the blowup rate of the cubic wave equation and compare with theoretical predictions given by modern Partial Differential Equations (PDE) theory;
- Study the late time decay rates of the cubic wave equation and compare with theoretical predictions given by modern PDE theory;
- Study the convergence of the numerical solutions to the cubic wave equation towards an attractor solution, as predicted by modern PDE theory.

#### 1.4 Thesis Outline

Briefly explain the contents of each chapter.

# **Chapter 2**

# **Theoretical Background**

In this chapter, we cover the theoretical background necessary for understanding the subsequent material of this work. We start by discussing the well-posedness of Cauchy problems in systems of partial differential equations in section 2.1, which is a necessary condition for our models to have predictive power. We then introduce the 3+1 formalism in section 2.2, which allows us to separate spacetime into three-dimensional space and time, making it easier to study the dynamical evolution of gravitational fields. Next, we discuss hyperboloidal compactification in section 2.3, a technique that enables us to study the asymptotic behavior of solutions at future null infinity. Finally, we present the wave equation and its cubic non linearity in the sections 2.4 and 2.5 respectively, which are the central objects of our study.

#### 2.1 Well-Posedness of Cauchy Problems in Systems of PDEs

When dealing with systems of partial differential equations, one generally works with initial value problems (also sometimes called *Cauchy problems*), which consist of, given initial data  $u(0,x^i)=f(x^i)$  at a time t=0, calculating the solution  $u(t,x^i)$  for a later time. These problems are very relevant in physics due to the predictive character of our models. However, for our systems to possess this predictive power, they must be *well-posed*. *Well-posedness* in a partial differential equation problem is the requirement that there be a unique solution that depends continuously, in some norm, on the given data for the problem [3]. More rigorously, we say an initial value problem is well-posed if there exist constants K and  $\alpha$  such that

$$||u(t,\cdot)||_{L^2} \le Ke^{\alpha t}||f||_{L^2},$$
 (2.1)

where the initial data of the *Cauchy problem* is given by  $u(0, x^i) = f(x^i)$  and the  $L^2$  norm is given by

$$||g||_{L^2}^2 = \int_{\mathbb{D}^3} g^{\dagger} g \, dV. \tag{2.2}$$

Without this property, small changes in the given data may result in arbitrarily large changes in the

solution or render it impossible to find a solution at all. Thus, we must make sure to formulate our field equations in such a way that they form a well-posed system of differential equations.

Let us consider a system of partial differential equations that can be written as

$$\partial_t u = A^p \partial_p u + B u, \tag{2.3}$$

where u is a state vector. We call  $A^p$  the *principal matrix* of the system despite it being an abbreviation for three matrices (one for each spatial dimension). The remaining terms on the right-hand side of our system of differential equations non-principal [3]. Given an arbitrary unit spatial vector  $s^i$ , we define the *principal symbol* of the system as

$$P^s \equiv A^s = A^p s_n. \tag{2.4}$$

If, for every unit spatial vector  $s^i$ , the *principal symbol* of our system has real eigenvalues, our system is considered *weakly hyperbolic*. If, in addition to our system being *weakly hyperbolic*, we have that for every unit spatial vector  $s^i$ , the principal symbol has a complete set of eigenvectors, and there exists a constant K, independent of  $s^i$ , such that

$$||T_s|| + ||T_s^{-1}|| \le K, (2.5)$$

where  $T_s$  is a matrix that has the eigenvectors of  $P^s$  as columns and we have the usual definition of the matrix norm  $||\cdot||$ ; we call our system *strongly hyperbolic*. The components of the vector  $v=T_s^{-1}u$  are called *characteristic variables* in the  $s^i$  direction, which satisfy advection equations with speeds equal to the eigenvectors of the principal symbol up to non-principal terms and derivatives transverse to the  $s^i$  direction. In other words, they satisfy equation (2.3) along the  $s^i$  direction. These variables will be relevant when we discuss how bamps handles patch boundaries in section 3.3.

Let us now consider a system of partial differential equations written as

$$\partial_t u = A^p \partial_p u + F(t, x^i), \tag{2.6}$$

where we only consider solutions on the half-space  $x^1=x\geq 0$ , making it so that we have a boundary. We call F the forcing term, which could be, for example, F=Bu as in equation (2.3). In this initial value problem, we provide initial conditions for the domain  $u(0,x^i)=f(x^i)$ , and boundary conditions  $Lu(t,x^i) \stackrel{\triangle}{=} g(t,x^A)$ , where the index A denotes that the data depends only on  $x^2=y$  and  $x^3=z$ , with L a matrix that will be described later, and  $\stackrel{\triangle}{=}$  denotes equality on the boundary.

We say that our system is  $symmetric\ hyperbolic$  if there exists a Hermitian positive definite symmetrizer H such that  $HA^ps_p$  is Hermitian for every spatial vector  $s_p$ . This property of a system is very relevant as it is related to  $strong\ well-posedness$  when the boundaries of our system are  $maximally\ dissipative$ , as we will see briefly. It is important to note that every  $symmetric\ hyperbolic$  system is  $strongly\ hyperbolic$ , even though the opposite is not true.

Saying that an initial value problem is *strongly well-posed* means that we can bind the solution in the bulk and restrict it to the initial boundary data with the addition of growth caused by non-principal terms or the boundaries [3]. That is, an initial value problem is *strongly well-posed* if, for every time T, there exists a constant  $K_T$  independent of the initial data and the forcing terms such that, for  $0 \le t \le T$ , we have

$$||u(t,\cdot)||_{\Sigma}^{2} + \int_{0}^{t} ||u(t',\cdot)||_{\partial\Sigma}^{2} dt' \leq K_{T}^{2} \left( ||f|_{\Sigma}^{2} + \int_{0}^{t} \left( ||F(t',\cdot)||_{\Sigma}^{2} + ||g(t',\cdot)||_{\partial\Sigma}^{2} \right) dt' \right), \tag{2.7}$$

where  $||\cdot||_{\Sigma}$  denotes the  $L^2$  norm on the half-space, and  $||\cdot||_{\partial\Sigma}$  denotes the  $L^2$  norm on the boundary plane.

Since every symmetric hyperbolic system is strongly hyperbolic, there exists a matrix  $T_x$  such that

$$T_x^{-1} P^x T_x = \Lambda_x = \begin{pmatrix} \Lambda_x^I & 0\\ 0 & \Lambda_x^{II} \end{pmatrix}. \tag{2.8}$$

If we have  $\Lambda_I>0$  and  $\Lambda_{II}<0$ , our system has *maximally dissipative* boundaries. If those conditions are not met, our system has characteristic boundaries in the variables where the characteristic speed vanishes at the boundaries.

We can prove the well-posedness of our initial value problem by defining the energy

$$E = \int_{\Sigma} \epsilon \ dV = \int_{\Sigma} u^{\dagger} H u \ dV \ . \tag{2.9}$$

By taking the time derivative of this energy and performing an integration by parts, we get

$$\partial_t E + c_1 \int_{\partial \Sigma} \left( u^{\dagger} H u \right) \ dS \le \int_{\Sigma} \left( u^{\dagger} H F + F^{\dagger} H u \right) \ dV + c_2 \int_{\partial \Sigma} \left( g^{\dagger} H g \right) \ dS, \tag{2.10}$$

for some  $c_1, c_2 > 0$ , from which we get *strong well-posedness* [3]. It is important to note that *maximally dissipative* boundary conditions only guarantee *well-posedness* for *symmetric hyperbolic* systems, not for *strongly hyperbolic* ones.

#### 2.2 The 3+1 Formalism

One usually finds the Einstein field equations written in a fully covariant form, where there isn't a clear distinction between space and time. However, in numerical relativity, we are interested in studying the dynamical evolution of a gravitational field in "time". It is thus convenient to split spacetime into three-dimensional space and time. That way, we can provide spacelike initial conditions and obtain the subsequent evolution along our time coordinate. To perform that separation, we use the so-called 3+1 Formalism [4–6].

We start by considering a spacetime with the metric  $g_{ab}$ . To maintain hyperbolicity in our evolution equations, we must restrict our spacetimes to be globally hyperbolic, meaning they possess a Cauchy

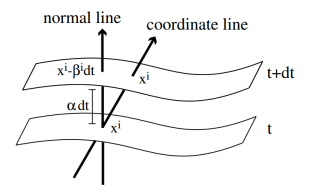


Figure 2.1: Geometric definition of the lapse function  $\alpha$  and the shift vector  $\beta^i$ . The lapse function  $\alpha$  is the proper time between two adjacent hypersurfaces  $\Sigma_t$  and  $\Sigma_{t+dt}$  measured by *Eulerian observers*. The shift vector  $\beta^i$  is the relative velocity between the *Eulerian observers* and the lines of constant spatial coordinates. [5]

surface. These globally hyperbolic spacetimes can be completely foliated in a way such that each three-dimensional slice is spacelike. Additionally, we can identify each three-dimensional hypersurface with the level set of a parameter t, which we consider to be a universal time function.

Given two adjacent hypersurfaces  $\Sigma_t$  and  $\Sigma_{t+dt}$  in a specific foliation, we can determine the geometry of the region of spacetime between them by studying the movement of observers moving along the direction normal to the hypersurfaces. We call those observers *Eulerian*. Considering those observers, we define three quantities that can describe our region of interest: the three-dimensional metric  $\gamma_{ij}$  (which measures proper distances within the hypersurfaces), the lapse function  $\alpha$  (which is the lapse of proper time between both hypersurfaces measured by *Eulerian observers*), and the shift vector  $\beta^i$  (which is the relative velocity between the *Eulerian observers* and the lines of constant spatial coordinates). It is also useful to define the extrinsic curvature  $K_{\mu\nu}$  (which measures the change of the normal vector to the hypersurface when parallel transported from one point in the hypersurface to another) and the acceleration of the *Eulerian observers*  $a^i$  [4–6].

In this work, we are interested in specifying initial conditions starting from a known spacetime metric  $g_{ab}$ . Thus, we need to calculate the previously mentioned 3+1 quantities from  $g_{ab}$ , which can be done by performing a 3+1 split of  $g_{ab}$ . This split is done by writing  $g_{ab}$  as a function of  $\alpha$ ,  $\beta^i$  and  $\gamma_{ij}$ :

$$ds^{2} = (-\alpha^{2} + \beta_{i}\beta^{i}) dt^{2} + 2\beta_{i} dt dx^{i} + \gamma_{ij} dx^{i} dx^{j},$$
(2.11)

or, more explicitly,

$$g_{\mu\nu} = \begin{pmatrix} -\alpha^2 + \beta_k \beta^k & \beta_i \\ \beta_j & \gamma_{ij} \end{pmatrix}, \qquad g^{\mu\nu} = \begin{pmatrix} -1/\alpha^2 & \beta^i/\alpha^2 \\ \beta^j/\alpha^2 & \gamma^{ij} - \beta^i \beta^j/\alpha^2 \end{pmatrix}. \tag{2.12}$$

Using this split, it becomes easy to obtain lpha ,  $eta^i$  and  $\gamma_{ij}$  from  $g_{ab}$  , from which we can obtain the

extrinsic curvature  $K_{\mu\nu}$  using the fact that

$$K_{ij} = \frac{D_i \beta_j + D_j \beta_i - \partial_t \gamma_{ij}}{2\alpha},\tag{2.13}$$

where  $D_{\mu} := (\delta_{\mu}^{\alpha} + n^{\alpha}n_{\mu})\nabla_{\alpha}$  is the projection of the covariant derivative onto the hypersurface and n is the normal vector to the hypersurface, with components given by

$$n^{\mu} = (1/\alpha, -\beta^{i}/\alpha), \qquad n_{\mu} = (-\alpha, 0).$$
 (2.14)

We can also obtain the acceleration of the Eulerian observers  $a^i$  as

$$a^i = \gamma^{ij} \partial_i(\ln \alpha).$$
 (2.15)

#### 2.3 Hyperboloidal Compactification

In this work, we are interested in studying how waves behave at future null infinity, making it useful to compactify our spacetime in such a way that allows us to study the asymptotic behavior of our solutions numerically. To accomplish this, we will employ a specific set of coordinates called *hyperboloidal* coordinates [7, 8].

The hyperboloidal coordinates (t,r) are related to the usual spherical coordinates (T,R) by the *height* function H(R) and the *compress function*  $\Omega(r)$  as follows:

$$T = t + H(R),$$
  $R = \Omega^{-1}(r) r.$  (2.16)

The angular coordinates remain unchanged. This transformation gives rise to the Jacobian matrix

$$(J^{Hyp})_{\alpha'}^{\beta} = \begin{pmatrix} 1 & -H'(r) & 0 & 0 \\ 0 & \frac{L(r)}{\Omega^2(r)} & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix} ,$$
 (2.17)

where H'(r) denotes the derivative of the height function with respect to R written as a function of r, and L(r) is defined as

$$L(r) \equiv \Omega(r) - r \,\partial_r \Omega(r) \; . \tag{2.18}$$

It is important to note that, even though there is freedom to choose the height and the compress function, we must require that, asymptotically in R, or equivalently, as  $\Omega$  approaches zero, we have

$$1 - H' \sim O(\Omega^2). \tag{2.19}$$

Since the solutions to our equations decay as we approach null infinity, it is useful to rescale our spacetime in such a way that our solutions are of order one through space. Given a spacetime with metric  $g_{ab}$ , we can rescale it by a strictly positive, smooth scalar function  $\Omega$  to obtain a new metric  $\tilde{g}_{ab}$ :

$$\tilde{g}_{ab} = \Omega^2 g_{ab},\tag{2.20}$$

This rescaling is called a *conformal transformation* and the function  $\Omega$  is called the *conformal factor*. If  $g_{ab}$  is a Lorentzian metric, then a vector  $v^a$  is timelike, null or spacelike with respect to  $g_{ab}$  if and only if it is timelike, null or spacelike with respect to  $\tilde{g}_{ab}$ , meaning that the causal structure of the spacetime is preserved under conformal compactification [9].

After applying both the coordinate transformation to hyperboloidal coordinates and the conformal transformation, we obtain a new metric  $\tilde{g}_{ab}$ , which on Cartesian coordinates is given by

$$\tilde{g}_{ab} = \begin{pmatrix} -\Omega^{2}(r) & -\frac{xH(r)L(r)}{r} & -\frac{yH(r)L(r)}{r} & -\frac{zH(r)L(r)}{r} \\ -\frac{xH(r)L(r)}{r} & \frac{(y^{2}+z^{2})\mathcal{R}^{2}(r)+r^{2}x^{2}L^{2}(r)\Xi(r)}{r^{4}} & \frac{xy(-\mathcal{R}^{2}(r)+r^{2}L^{2}(r)\Xi(r))}{r^{4}} & \frac{xz(-\mathcal{R}^{2}(r)+r^{2}L^{2}(r)\Xi(r))}{r^{4}} \\ -\frac{yH(r)L(r)}{r} & \frac{xy(-\mathcal{R}^{2}(r)+r^{2}L^{2}(r)\Xi(r))}{r^{4}} & \frac{(x^{2}+z^{2})\mathcal{R}^{2}(r)+r^{2}y^{2}L^{2}(r)\Xi(r)}{r^{4}} & \frac{yz(-\mathcal{R}^{2}(r)+r^{2}L^{2}(r)\Xi(r))}{r^{4}} \\ -\frac{zH(r)L(r)}{r} & \frac{xz(-\mathcal{R}^{2}(r)+r^{2}L^{2}(r)\Xi(r))}{r^{4}} & \frac{yz(-\mathcal{R}^{2}(r)+r^{2}L^{2}(r)\Xi(r))}{r^{4}} & \frac{(x^{2}+y^{2})\mathcal{R}^{2}(r)+r^{2}z^{2}L^{2}(r)\Xi(r)}{r^{4}} \end{pmatrix},$$

$$(2.21)$$

where we have defined  $\Xi(r)$  and  $\mathcal{R}(r)$  as

$$\Xi(r) = \frac{1 - H^2(r)}{\Omega^2(r)}$$
,  $\mathcal{R}(r) = r - R_i + R_i \Omega(r)$ , (2.22)

with  $R_i$  being the interface radius, which we will now briefly discuss.

In some applications, it may be useful to consider a flat slice where most of the dynamics happen, followed by a hyperboloidal slice, which allows us to study the decay of our solution as it approaches future null infinity [7, 8]. That scenario can be achieved using *hyperboloidal layers*, which consist of performing a transition from flat slices to hyperboloidal slices at a chosen interface radius  $R_i$ , making it such that we achieve a hyperboloidal slice after some transition length  $l_t$ . The details of this transition will be further discussed in section 4.2.

#### 2.4 The Wave Equation

The wave equation is a symmetric hyperbolic partial differential equation that describes the propagation of all types of wave-like phenomena. In the study of gravitational waves, this equation is of particular importance due to its structural similarity to the Einstein field equations on the *Generalized Harmonic Gauge (GHG)*. The wave equation, in its simpler form, is given by

$$\Box_a \Psi = S, \tag{2.23}$$

where  $\Box_g$  is the d'Alembert operator,  $\Psi$  is the scalar field, and S is a source term. The Einstein equations in the GHG can be written as

$$\Box_{q}g_{ab} = 2g^{cd}g^{ef}(\partial_{e}g_{ca}\partial_{f}g_{db} - \Gamma_{ace}\Gamma_{bdf}), \tag{2.24}$$

where  $g_{ab}$  is the spacetime metric,  $\Gamma_{abc}$  are the Christoffel symbols, and  $g^{ab}$  are the inverse metric [10]. The similarity between these two equations allows us to study the wave equation as a model for gravitational waves.

It is important to note that when we perform a conformal transformation of the spacetime like the one described in section 2.3, the d'Alembert operator in the new metric  $\tilde{g}_{ab}$  becomes

$$\Box_{\tilde{g}}\tilde{\psi} = \tilde{g}^{ab}\tilde{\nabla}_{a}\tilde{\nabla}_{b}\tilde{\psi} = \Omega^{s-2}g^{ab}\nabla_{a}\nabla_{b}\psi + s\Omega^{s-3}\psi\,g^{ab}\nabla_{a}\nabla_{b}\Omega \tag{2.25}$$

$$+ s(n+s-3)\Omega^{s-4}\psi g^{ab}\nabla_a\Omega\nabla_b\Omega$$
 (2.26)

where s is the conformal weight of the scalar field  $\psi$  (which we can choose freely) and n is the dimension of the spacetime [9]. In this work, we will be working in a four-dimensional spacetime, so n=4. The conformal weight s will be chosen to be s=-1 to simplify the expression of equation (2.26). Thus, we have

$$\tilde{g}^{ab}\tilde{\nabla}_a\tilde{\nabla}_b\tilde{\psi} = \Omega^{-3}g^{ab}\nabla_a\nabla_b\psi - \Omega^{-4}\psi\,g^{ab}\nabla_a\nabla_b\Omega,\tag{2.27}$$

We can now substitute (2.23) into (2.27) and write the resulting equation in terms of the rescaled quantities, obtaining

$$\tilde{g}^{ab}\tilde{\nabla}_{a}\tilde{\nabla}_{b}\tilde{\psi} = \Omega^{-3}S - \Omega^{-1}\tilde{\psi}\bigg(\tilde{g}^{ab}\tilde{\nabla}_{a}\tilde{\nabla}_{b}\Omega - 2\Omega^{-1}\tilde{g}^{ab}\tilde{\nabla}_{a}\Omega\tilde{\nabla}_{b}\Omega\bigg),\tag{2.28}$$

which is the wave equation for the rescaled scalar field  $\tilde{\psi}$  in the compactified spacetime with metric  $\tilde{g}_{ab}$  .

To solve the previous equation numerically, it is useful to do a 3+1 decomposition of spacetime and write the system as a set of first-order partial differential equations. To do that, we start by employing the 3+1 split of the metric  $\tilde{g}_{ab}$  described in section 2.2 and define the time reduction variable  $\tilde{\Pi}$  and the spatial reduction variables  $\tilde{\phi}_i$  as

$$\Pi = -\mathcal{L}_n \tilde{\psi} \qquad \qquad \phi_i = \tilde{D}_i \tilde{\psi} , \qquad (2.29)$$

where  $\mathcal{L}_n$  is the Lie derivative along the normal vector to the hypersurface  $n^a$ , and  $\tilde{D}_i$  is the projection of the covariant derivative onto the hypersurface. It is important to note that these definitions introduce new constraints to our system, given by

$$\begin{cases} \mathcal{L}_n \tilde{\psi} + \Pi \stackrel{!}{=} 0 \\ \tilde{D}_i \tilde{\psi} - \phi_i \stackrel{!}{=} 0 \end{cases} , \tag{2.30}$$

where  $\stackrel{!}{=}$  denotes that we may allow for small violations of these constraints due to numerical errors. The second constraint is of particular importance as we will see later in section 4.1. Due to this importance, we will denote it as  $C_i$ .

Using the previous definitions, we can write the wave equation as a set of first-order partial differential equations as

$$\begin{cases} \partial_{t}\tilde{\psi} = -\tilde{\alpha}\tilde{\Pi} + \beta^{i}\tilde{\phi}_{i} \\ \partial_{t}\tilde{\phi}_{i} = -\tilde{\alpha}(\tilde{D}_{i}\tilde{\Pi}) - (\tilde{D}_{i}\tilde{\alpha})\tilde{\Pi} + \beta^{j}\partial_{j}\tilde{\phi}_{i} + \tilde{\phi}_{j}\partial_{i}\beta^{j} \\ \partial_{t}\tilde{\Pi} = -\tilde{\alpha}\tilde{a}^{i}\tilde{\phi}_{i} - \tilde{\alpha}\tilde{\gamma}^{ij}\tilde{D}_{j}\tilde{\phi}_{i} + \tilde{\alpha}\tilde{K}\tilde{\Pi} + \beta^{i}\tilde{D}_{i}\tilde{\Pi} \\ -\tilde{\alpha}\Omega^{-1}\Big(\tilde{g}^{ab}\tilde{\nabla}_{a}\tilde{\nabla}_{b}\Omega - 2\Omega^{-1}\tilde{g}^{ab}\tilde{\nabla}_{a}\Omega\tilde{\nabla}_{b}\Omega\Big)\tilde{\psi} + \tilde{\alpha}\Omega^{-3}S \end{cases}$$

$$(2.31)$$

It is also necessary to calculate the characteristic variables of our system and their respective speeds. To do so, we start by calculating the principal symbol of our system and its eigenvalues, obtaining advection equations for the characteristic variables

$$\begin{cases} \partial_t \tilde{\psi} = 0 \\ \partial_t (\tilde{\Pi} \pm \tilde{\phi}_s) = -(-\beta^s \pm \tilde{\alpha}) \partial_s (\tilde{\Pi} \pm \tilde{\phi}_s) \end{cases} , \tag{2.32}$$

where  $\tilde{\phi}_s = \tilde{\phi}_i s^i$  (it is important to note that this definition does not let  $\tilde{\phi}_i$  and  $s^i$  commute) and similarly for  $\beta^s$ , with  $s^i$  being an arbitrary unit spatial vector. From this, we can identify the characteristic variables as

$$\begin{cases} u_0 = \tilde{\psi} \\ u_{\pm} = \tilde{\Pi} \pm \tilde{\phi}_s \\ (u_2)_i = (\delta_i^j - s_i s^j) \tilde{\phi}_j \end{cases}$$

$$(2.33)$$

with the associated characteristic speeds being

$$\begin{cases} v_0 = 0 \\ v_{\pm} = -\beta^s \pm \tilde{\alpha} \end{cases}$$

$$v_2 = -\beta^s$$
(2.34)

In order to keep constraint violations from growing exponentially, it may be useful to add multiples of the constraint  $C_i$  to the evolution equations of our system since doing so does not alter the physical behavior of the system, but it improves numerical stability. Doing so, we obtain the final form of our

system as

$$\begin{cases} \partial_{t}\tilde{\psi} = -\tilde{\alpha}\tilde{\Pi} + \beta^{i}\tilde{\phi}_{i} + (1+\gamma_{1})\beta^{i}(\tilde{D}_{i}\tilde{\psi} - \tilde{\phi}_{i}) \\ \partial_{t}\tilde{\phi}_{i} = -\tilde{\alpha}(\tilde{D}_{i}\tilde{\Pi}) - (\tilde{D}_{i}\tilde{\alpha})\tilde{\Pi} + \beta^{j}\partial_{j}\tilde{\phi}_{i} + \tilde{\phi}_{j}\partial_{i}\beta^{j} + \gamma_{2}\tilde{\alpha}(\tilde{D}_{i}\tilde{\psi} - \tilde{\phi}_{i}) \\ \partial_{t}\tilde{\Pi} = -\tilde{\alpha}\tilde{a}^{i}\tilde{\phi}_{i} - \tilde{\alpha}\tilde{\gamma}^{ij}\tilde{D}_{j}\tilde{\phi}_{i} + \tilde{\alpha}\tilde{K}\tilde{\Pi} + \beta^{i}\tilde{D}_{i}\tilde{\Pi} \\ -\tilde{\alpha}\Omega^{-1}\Big(\tilde{g}^{ab}\tilde{\nabla}_{a}\tilde{\nabla}_{b}\Omega - 2\Omega^{-1}\tilde{g}^{ab}\tilde{\nabla}_{a}\Omega\tilde{\nabla}_{b}\Omega\Big)\tilde{\psi} + \tilde{\alpha}\Omega^{-3}S + \gamma_{3}\beta^{i}(\tilde{D}_{i}\tilde{\psi} - \tilde{\phi}_{i}) \end{cases}$$

$$(2.35)$$

where  $\gamma_1$ ,  $\gamma_2$ , and  $\gamma_3$  are free parameters which quantify the constraint violation we allow in our system. This addition will also change the characteristic variables and their respective speeds. Choosing  $\gamma_3=\gamma_1\gamma_2$  for convenience, we obtain the characteristic variables as

$$\begin{cases} u_0 = \tilde{\psi} \\ u_{\pm} = \tilde{\Pi} \pm \tilde{\phi}_s - \gamma_2 \tilde{\psi} \\ (u_2)_i = (\delta_i^j - s_i s^j) \tilde{\phi}_j \end{cases}$$

$$(2.36)$$

with the respective characteristic speeds being

$$\begin{cases} v_0 = -(1+\gamma_1)\beta^s \\ v_{\pm} = -\beta^s \pm \tilde{\alpha} \\ v_2 = -\beta^s \end{cases}$$
 (2.37)

#### 2.5 The Cubic Wave Equation

# **Chapter 3**

# **Numerical Setup**

In this chapter, we will describe the numerical setup of the bamps code. We will start by discussing the grid setup in section 3.1, followed by the numerical method used to solve the partial differential equation systems in section 3.2 and a brief description of how bamps handles patch boundaries in section 3.3. Finally, we will discuss the Adaptive Mesh Refinement (AMR) strategy employed by bamps in section 3.4.

#### 3.1 Grid Setup

The bamps code uses a multi-patch grid setup to cover the computational domain. Even though bamps is capable of supporting both a cubed-ball grid and a cubed-sphere grid, in this work we will only use the cubed-ball grid. The grid is composed of several patches which are shaped like deformed cubes that fit together to cover the domain. Each patch is mapped by two different coordinate systems: the local patch coordinates  $(\bar{x}, \bar{y}, \bar{z})$  and the global Cartesian coordinates (x, y, z).

The cubed-ball grid consists of a central cube surrounded by six deformed cube patches that form a spherical shell around the central cube forming a transition shell, which is itself surrounded by six more deformed cube patches that form the outer spherical shell. In the central cube, the local patch coordinates and the global Cartesian coordinates are identical, i.e.,  $(\bar{x}, \bar{y}, \bar{z}) = (x, y, z)$ . However, in the other patches, a coordinate transformation must be applied to map the local patch coordinates to the global Cartesian coordinates. This transformation is performed in two steps: first, the local coordinates are transformed to temporary global coordinates  $(\tilde{x}, \tilde{y}, \tilde{z})$ , which are then transformed to the final global Cartesian coordinates (x, y, z) by performing a rotation of the patch to their correct placement on the sphere. The transformation from local patch coordinates to temporary global coordinates is given by

$$x_t = \frac{\bar{x}}{\bar{s}}, \qquad y_t = \frac{\bar{x}}{\bar{s}}\bar{y}, \qquad z_t = \frac{\bar{x}}{\bar{s}}\bar{z},$$
 (3.1)

where, for the outer patches,

$$\bar{s} = \sqrt{1 + \bar{y}^2 + \bar{z}^2} \tag{3.2}$$

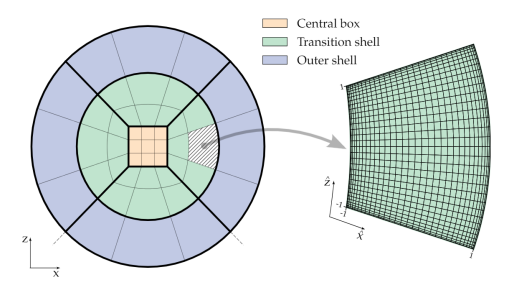


Figure 3.1: Two-dimensional sketch of the bamps cubed-ball grid layout. The grid is composed of several patches shaped like deformed cubes. These patches can be further divided into subpatches. Each subpatch is covered by Gauss-Lobatto grids. [11]

and for the inner patches,

$$\bar{s}(\lambda) = \sqrt{\frac{1+2\lambda}{1+\lambda(\bar{y}^2+\bar{z}^2)}}, \qquad \lambda = \frac{\bar{x}^2-\bar{x}_0^2}{\bar{x}_1^2-\bar{x}_0^2},$$
 (3.3)

where  $\bar{x}_0$  and  $\bar{x}_1$  are the boundaries of the patch in the local coordinates [11].

Each patch can be further divided into several subpatches, which are smaller cubes that allow for a more refined grid in specific regions of the domain. When dividing a patch into subpatches, the division is done such that subgrids of two neighboring patches match, and that neighboring patches and subpatches have the same grid-point positions on their respective boundaries. In figure 3.1, we show an example of a cubed-ball grid divided into subpatches.

#### 3.2 Numerical Method

bamps uses the method of lines to solve the system of partial differential equations, meaning that it treats time as a continuous variable and space as discrete. In order to integrate fields forward in time, a 4th order Runge-Kutta scheme is used. The spatial discretization is performed using Gauss-Lobatto collocation points, which is done by mapping the local patch coordinates  $(\bar{x}, \bar{y}, \bar{z})$  to the reference cube coordinates  $(\tilde{x}, \tilde{y}, \tilde{z}) \in [-1, 1]^3$  and then using the Gauss-Lobatto points in each direction. The Gauss-Lobatto points in the  $\tilde{x}$  direction are given by

$$\tilde{x}_{\alpha} = -\cos\left(\frac{\pi\alpha}{N_x - 1}\right) , \qquad (3.4)$$

with  $\alpha=0,1,\ldots,N_x-1$  and  $N_x$  being the number of grid points in the  $\tilde{x}$  direction. The collocation points in the other two directions are defined similarly. It is important to note that the number of angular grid

points must be the same in all patches to ensure that the grid points on the boundaries of neighboring patches match [11]. A grid generated using Gauss-Lobatto points can be seen in figure 3.1.

On each of the collocation points, all the evolution fields u are expanded in each dimension in terms of Chebyshev polynomials  $T_n(\tilde{x})$  as

$$u_{\alpha\beta\delta} = u(\tilde{x}_{\alpha}, \tilde{y}_{\beta}, \tilde{z}_{\delta}) = \sum_{l=0}^{N_z - 1} c_n^x(\tilde{y}_{\beta}, \tilde{z}_{\delta}) T_n(\tilde{x}_{\alpha}) , \qquad (3.5)$$

and similarly for the  $\tilde{y}$  and  $\tilde{z}$  directions. This expansion allows for a spectral representation of the fields, which is advantageous since it allows for the analytical differentiation of our expansion through the analytical derivatives of the Chebyshev polynomials. Additionally, since for many applications the contribution of higher order polynomials decays exponentially, the numerical error in spectral methods usually decays exponentially with the resolution [6]. bamps uses the pseudospectral approach, meaning that instead of storing the coefficients  $c_n^x(\tilde{y}_\beta,\tilde{z}_\delta)$ , it stores the values of the fields at the collocation points. The spatial derivatives of the fields are computed using matrix multiplication. For example, the derivative in the  $\tilde{x}$  direction is computed as

$$(\partial_{\tilde{x}}u)_{\alpha\beta\delta} = \sum_{n=0}^{N_x-1} D_{\alpha n} u_{n\beta\delta} , \qquad (3.6)$$

where  $D_{\alpha n}$  is the Gauss-Lobatto differentiation matrix

$$D_{\alpha\beta} = \begin{cases} -\frac{2(N_x - 1)^2 + 1}{6} & \alpha = \beta = 0\\ \frac{q_{\alpha}}{q_{\beta}} \frac{(-1)^{\alpha + \beta}}{\tilde{x}_{\alpha} - \tilde{x}_{\beta}} & \alpha \neq \beta\\ \frac{-\tilde{x}_{\beta}}{2(1 - \tilde{x}_{\beta}^2)} & \alpha = \beta = 1, \dots, N_x - 2 \end{cases}, \tag{3.7}$$

$$\frac{2(N_x - 1)^2 + 1}{6} & \alpha = \beta = N_x - 1$$

where  $q_{\alpha}=2$  on the boundaries and  $q_{\alpha}=1$  otherwise [6, 11].

In order to ensure numerical stability, bamps uses a filtering method to remove high-frequency modes. The filtering is applied after each full time step and consists of multiplying the fields with a filter matrix in each direction

$$(\mathcal{F}u)_{\alpha\beta\delta} = \sum_{n=0}^{N_x - 1} \mathcal{F}_{n\alpha} u_{n\beta\delta} , \qquad (3.8)$$

where the filter matrix  $\mathcal{F}_{n\alpha}$  is given by

$$\mathcal{F}_{\alpha\beta} = \sum_{n} S_{\alpha n} e^{-36(n/(N_x - 1))^{64}} A_{n\beta} , \qquad (3.9)$$

where  $S_{\alpha\beta}$  is the Chebyshev synthesis matrix and  $A_{n\beta}$  is Chebyshev analysis matrix [6, 11].

The bamps code also supports both 1 dimensional and 2 dimensional reductions, called the cartoon method. TIt is also able to handle mirror symmetries along the x, y and z directions [8].

#### 3.3 Patching Boundary Conditions

Since bamps separates our domain into different patches, we need to impose boundary conditions between adjacent patches so that they can communicate data to each other. An effective way to impose these boundary conditions is to add appropriate penalty terms to the equations on these boundaries. This method, which we call *Penalty Method*, imposes the needed boundary conditions while keeping the evolution equations at the boundary almost unchanged [11, 12].

It consists of adding to the equation of motion of the incoming characteristic variable a term of the form  $(u_-^{BC}-u_-)$ , where  $u_-$  is the incoming characteristic variable. Thus, if the boundary condition  $u_-^{BC}=u_-$  is satisfied, the penalty terms vanish. We can then determine the appropriate penalty terms to add to the equation when written in their fundamental form by rewriting our evolution equations on the boundaries as functions of the characteristic variables, incorporating the penalty terms, and then transforming our system back [11, 12]. That way, the evolution equations (2.31) on the boundary will get modified to

$$\begin{cases}
(\partial_{t}\tilde{\psi})^{BC} = \partial_{t}\tilde{\psi} \\
(\partial_{t}\tilde{\Pi})^{BC} = \partial_{t}\tilde{\Pi} + \frac{1}{2}p \left(v_{+}(u_{+}^{BC} - u_{+}) + v_{-}(u_{-}^{BC} - u_{-})\right) \\
(\partial_{t}\tilde{\phi}_{i})^{BC} = \partial_{t}\tilde{\phi}_{i} + \frac{1}{2}p n_{i} \left(v_{+}(u_{+}^{BC} - u_{+}) - v_{-}(u_{-}^{BC} - u_{-})\right) + p v_{2} \left((u_{2}^{BC})_{i} - (u_{2})_{i}\right)
\end{cases} ,$$
(3.10)

where p is the penalty coefficient, which can determined from a semi-discrete energy analysis, and  $n^i$  is the unit outgoing normal vector to the boundary. Even though in equation (3.10) the terms regarding every characteristic variable are included, it is important to note that only incoming characteristic variables are taken into account in this method, thus if any of the characteristic speeds are greater than 0, the penalty term associated will be null.

If we allow for constraint violations in our system (as we did in equation (2.35)), the characteristic variables and their respective speeds will change, and we will need to modify the penalty terms accordingly, obtaining

$$\begin{cases} (\partial_{t}\tilde{\psi})^{BC} = \partial_{t}\tilde{\psi} - p \, v_{0}(u_{0}^{BC} - u_{0}) \\ (\partial_{t}\tilde{\Pi})^{BC} = \partial_{t}\tilde{\Pi} + \frac{1}{2}p \, \left(v_{+}(u_{+}^{BC} - u_{+}) + v_{-}(u_{-}^{BC} - u_{-})\right) - 2 \, \gamma_{2} \, p \, v_{0}(u_{0}^{BC} - u_{0}) \\ (\partial_{t}\tilde{\phi}_{i})^{BC} = \partial_{t}\tilde{\phi}_{i} + \frac{1}{2}p \, n_{i} \, \left(v_{+}(u_{+}^{BC} - u_{+}) - v_{-}(u_{-}^{BC} - u_{-})\right) + p \, v_{2} \, \left((u_{2}^{BC})_{i} - (u_{2})_{i}\right) \end{cases}$$
(3.11)

#### 3.4 Adaptive Mesh Refinement

bamps has the capability of generating new grids during the evolution in order to supply additional resolution where needed to adequately represent the solution, and consolidate these fine grids back into fewer coarser ones once the additional resolution is no longer required (h-refinement). It is also able to adjust the resolution in any individual grid (p-refinement). This process is called Adaptive Mesh Refinement (AMR).

The AMR procedure in bamps consists in:

1. Evaluate the h-refinement indicator on each subpatch;

- 2. Generate a set of initial h-refinement flags;
- 3. Modify the h-refinement flags to satisfy the constraints for a 'legal' bamps grid;
- 4. Apply the h-refinement operations;
- 5. Perform load balancing to consolidate grids that are marked to be coarsened together;
- 6. Apply h-coarsening operations;
- 7. Evaluate the p-refinement indicator function on each grid;
- 8. Apply p-refinement and p-coarsening operations;
- 9. Apply final load balancing.

Several indicators can be used to determine where refinement is needed. In this work, we will use the *truncation error indicator*, which estimates the truncation error of our spectral expansion. This indicator is calculated by performing the spectral decomposition of the evolution fields as described in section 3.2 and then constructing the sequence

$$\tilde{c}_i = \sqrt{\frac{1}{N} \sum_{k=1}^{N} c_i^k}, \qquad i = 1, 2, \dots, \tilde{n}$$
 (3.12)

where  $\tilde{n}$  is the highest mode which is not affected by the filter described in section 3.2, and k enumerates all N lines of grid points. Then, an exponential is fitted to the sequence  $\tilde{c}_i$  obtaining the indicator function

$$\epsilon = 10^{a\tilde{n}+b} \,, \tag{3.13}$$

where a and b are the slope and the offset obtained from the least squares fit on the logarithm of the sequence  $\tilde{c}_i$  [13].

It is important to note that the flags generated in the AMR procedure are then modified to ensure that the end state satisfies the constraints for a 'legal' bamps grid, as previously mentioned. This modification can can be visualized in figure 3.2. In a 'legal' bamps grid, each subpatch is not allowed to be more than one refinement level apart from any of its neighbors, and the grid structure must satisfy a 1:2 condition, meaning that if a subpatch is refined, all its neighbors must either be at the same refinement level or one level coarser. This is done to ensure that the grid structure remains consistent and that the numerical methods used in bamps can operate correctly across patch boundaries [13].

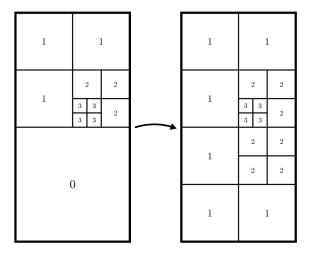


Figure 3.2: Example of a grid that does not satisfy the constraints for a 'legal' bamps grid (left) and the same grid after the modification of the h-refinement flags to satisfy these constraints (right). In a 'legal' bamps grid, each subpatch is not allowed to be more than one refinement level apart from any of its neighbors, and the grid structure must satisfy a 1:2 condition, meaning that if a subpatch is refined, all its neighbors must either be at the same refinement level or one level coarser. This is done to ensure that the grid structure remains consistent and that the numerical methods used in bamps can operate correctly across patch boundaries [13].

## **Chapter 4**

### Results

Insert your chapter material here.

#### 4.1 Wave Equation in Hyperboloidal Slices

The first step towards our goal is to implement the wave equation from equation (2.31) in hyperboloidal slices. We will construct our hyperboloidal slices using

$$H(R) = \frac{2R^2 + s^2 - \sqrt{4R^2s^2 + s^4}}{2\sqrt{R^2 + 1}}, \qquad \qquad \Omega(r) = 1 - \frac{r^2}{s^2}, \tag{4.1}$$

as heigh and compress functions respectively, where s is a free parameter that controls where null infinity is located. The details of this construction were discussed in section 2.3. This particular choice of heigh and compress functions is of particular interest since it leads to a constant outgoing characteristic speed, which has the advantage of keeping the shape of the waves as they propagate [7, 8]. This, however comes with the disadvantage of having an incoming characteristic speed that goes to zero as we approach null infinity. This is an acceptable trade-off since we are not expecting incoming waves in our setup. In figure  $\ref{eq:constraint}$  we can see how the incoming and outgoing light speeds vary along our computational domain.

#### INSERT FIGURE WITH SPEEDS HERE

Giving the initial conditions

$$sjdjf$$
ç $lsdg$  (4.2)

and setting s=30, we obtain the evolution shown in figures ?? and ??, which satisfies the

- 4.2 Wave Equation in Hyperboloidal Layers
- 4.3 Cubic Wave Equation

# **Chapter 5**

# **Conclusions**

Insert your chapter material here.

#### 5.1 Achievements

The major achievements of the present work.

#### 5.2 Future Work

A few ideas for future work.

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