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A More Robust Multigrid Algorithm for Diffusion Type Registration Models

Tony Thompson^{*} and Ke Chen^{*†}

Abstract

Registration refers to the useful process of aligning two similar but different intensity image functions 2 in order to either track changes or combine information. Variation 1 models are capable of finding 3 transform maps containing large and non-uniform deformations between such a pair of images. Since 4 finding a transform map is an inverse problem, as with all models suitable regularisation is necessary 5 to overcome the non-uniqueness of the problem. In the case of diff[•] sion type models regularisation 6 terms impose smoothness on the transformation by minimising one gridient of the flow field. The 7 diffusion model also coincides with the basic model for optica. How frameworks of Horn-Schunck 8 (1981, AI). The biggest drawback with variational models is the large computational cost required to 9 solve the highly non-linear system of PDEs; Chumchob-Che. (201', JCAM) developed a non-linear 10 multigrid (NMG) method to address this cost problem. h. vever, a closer look at the analysis of the 11 NMG scheme highlighted omissions which affected the conversion of the NMG scheme. Moreover, 12 the NMG method proposed by Chumchob-Chen did n. • impose any control of non-physical folding 13 which invalidates a map. This paper has proposed several key ideas. First we re-evaluate the analysis 14 of the NMG method to show how the omissions in ¹10₁ ... a noticeable impact on the convergence 15 of the NMG method. In addition, we also provide a way of estimating the convergence rate of a solver 16 on the coarsest grid in order to estimate the **i**, there coarsest has will be required to obtain a 17 solution with appropriate accuracy. Secondly we rop se an extension to the Chumchob-Chen NMG 18 method which controls any folding within a defe mation. Experimental results on the proposed 19 multigrid framework demonstrate improvements in convergence and the accuracy of registrations 20 compared with previous methods. 21

22 Keywords. Variational model, Image re jistrati, n, Fast Multigrid, Mesh folding control

23 1 Introduction

1

Image registration is the process of aligning pairs, or sequences, of similar images. This alignment is 24 achieved by fixing one image called the reference image, and then applying geometric transformations 25 on the remaining images, c .led the template images, such that the template images become similar to 26 the reference image. This a principal is a very powerful tool in many real world applications spanning 27 diverse areas such as computer . naging, weather satellite imaging [19] and especially medical imaging 28 which is of interest to $15 \left[\frac{12}{12}, \frac{12}{12}, \frac{12}{23}, \frac{24}{24} \right]$. However, image registration is also one of the most difficult 29 tasks of image processing "ith .nany challenges to be overcome. Generally image registration models can 30 be classified into tro main categories; parametric and non-parametric models. In parametric models, 31 the transformation is are global and can be described by matching a finite number of features in the 32 images, leading to subscript landmark based registration [31, 33], or the transformations are governed by 33 a small number of parameters such as in the case of affine image registration [3, 15] (with 6 parameters 34 in 2D and 12, aramet rs in 3D). However, the focus of this paper will be on the latter category, namely 35 non-parametric . A.s. 36

Denote respectively a reference and a template image (both given as grey-scale images) $R, T \in \Omega \subset \mathbb{R}^d$. The aim of integer gistration is to transform this T to R such that they become similar to one another,

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or in other words we look to find the transformation $arphi(x):\mathbb{R}^d o\mathbb{R}^d$ such that

$$T \circ \boldsymbol{\varphi}(\boldsymbol{x}) = T(\boldsymbol{\varphi}(\boldsymbol{x})) \approx R(\boldsymbol{x}) \text{ for } \boldsymbol{x} = (x_1, \dots, x_d)^T \in \Omega \subset \mathbb{R}^d.$$
 (1.1)

In variational image registration the transformation $\varphi(x)$ is equivalent to finding the displacement of every pixel x in T to their corresponding pixel in R, and so we can define $\varphi(x)$ by the following

$$\varphi \equiv \varphi(x) = x + u(x) \tag{1.2}$$

where $\boldsymbol{u} \equiv \boldsymbol{u}(\boldsymbol{x}) = (u_1(\boldsymbol{x}), \dots, u_d(\boldsymbol{x}))^T$ denotes the displacement field. Then the problem of determining

 φ is the same as finding u. From this point onward we shall consider only the 2L case, that is d = 2, however all ideas presented in this paper are readily extendible to the 3' λ case. Furthermore we

will also assume that the image domain Ω given by the unit square, that $\Gamma = [0, 1]^2 \subset \mathbb{R}^2$. In order to determine \boldsymbol{u} , the variational minimisation problem will take the following form

$$\min_{\boldsymbol{u}} E(\boldsymbol{u}) = \mathscr{D}(R, T, \boldsymbol{u}) + \alpha \mathscr{R}(\boldsymbol{u})$$
(1.3)

where in the energy functional $\mathscr{D}(R,T,u)$ is a distance measure, $\mathscr{T}(u)$ is the regularisation term and $\alpha \in \mathbb{R}^+$ is a weighting parameter. Note that inclusion of the regularisation term is a necessity as without it the minimisation would be ill-posed in the sense of Hadamard. For the purpose of this paper we shall consider only mono-modal images, that is images taken using the same imaging modality (e.g. CT), this means that image intensities are comparable. In the model on $\mathbb{C}_{2^{-1}}$ case, the typical choice of similarity measure is the sum of squared distances (SSD) measure given by

$$\mathscr{D}(R,T,\boldsymbol{u}) = \frac{1}{2} \int_{\Omega} \left(T(\boldsymbol{x}+\boldsymbol{u}) - R(\boldsymbol{x}) \right)^2 d\Omega .$$
(1.4)

Here SSD is only one of many choices of similar were [34]. Moreover, the choice of regularisation term is less straightforward as there is a large selection to choose from [1,6,17,18,20-22,34-36] and no one is yet the best. In this paper we will only condider one regularisation term, namely the diffusion regulariser and focus on optimal solution. As it numerical implementation, the common approach is to use an optimise-discretise approach, and indeed this is the approach we will adopt throughout this paper.

Solutions of variational models can be omput tionally intensive, but such non-parametric models are 59 worth the effort as they can produce very as a rate results and are able to deal with local deformations 60 effectively; the high computational exprase is due to the need of determining the displacement of every 61 pixel in the image. Multigrid techniques is known fast solvers have been used in previous works [20, 62 21, 25, 27–29, 32, 37, 40 to grea v reduce the computational cost and produce more accurate results, 63 however few of these directly deal with the non-linearity resulting from the similarity measure (1.4). 64 The reason for this is that. "ile multigrid techniques and theories have been established for linear 65 equations for a long time, *achie zing optimal convergence in a non-linear multigrid framework is never* 66 automatic and still poses a g. t challenge. However, the work done by Chumchob-Chen [16] introduced 67 a robust multigrid fram work for liffusion type variational models that treats the non-linearity directly. 68 We propose to improve the sonv rgence problems of the NMG method from [16] through a more in-depth 69 and accurate analysis of ι , r ultigrid framework as well as using an alternate coarsest solver to obtain 70 a more efficient so'ation, 'hus resulting in a better method. Next we address how to overcome mesh 71 folding by incorporating ar additional constraint into the diffusion model presented in [16], this idea can 72 be thought of as $\alpha \sin \beta^{1/2}$ cation of the hyper-elastic model introduced in the work by Burger et al. [11]. 73 The addition c, this c "nstraint imposes that the transformation produced is regular and diffeomorphic i.e. 74 there is no folding. The production of diffeomorphic transformations lead to more physically meaningful 75 results, which is particularly useful in medical imaging. In this paper, we consider one specific (yet widely 76 used) moo ', n... 'y the diffusion model to focus on our main aims: (i) improving the convergence of 77 the NMG me'.od from [16]; (ii) development of a fast NMG method for a refined diffusion model which 78 controls folding. 79

There are, however, many other choices for the regularisation term [1,6,17,18,20–22,34–36], each offering a different model and with their own distinct benefits and drawbacks. In particular, we mention

Total Variation (TV) [20, 21, 35, 36]:
$$\mathscr{R}^{TV}(\boldsymbol{u}) = \sum_{s=1}^{\infty} \int_{\Omega} |\nabla u_s| d\Omega$$
 where $|\cdot|$ denotes the Euclidean

83 norm;

88

Linear Elastic (LE) [1] [6] [22] [34]:
$$\mathscr{R}^{LE}(\boldsymbol{u}) = \int_{\Omega} \frac{\mu}{4} \sum_{s,t=1}^{2} \left(\partial_{x_s} u_t + \partial_{x_t} u_s\right)^2 + \frac{\lambda}{2} \left(\nabla \cdot \boldsymbol{u}\right)^2 d\Omega$$
 where μ, λ

85 are Lamé constants;

Mean Curvature (MC) [17,18]:
$$\mathscr{R}^{MC}(\boldsymbol{u}) = \frac{1}{2} \int_{\Omega} \sum_{s=1}^{2} \nabla \cdot \left(\frac{\nabla u_s}{\sqrt{|\nabla u_s|^2 + \beta}} \right)^2 d\Gamma$$
 where β is some small positive quantity.

87 positive quantity.

While each such models might be solved by a NMG framework, achieving cotine 'officiency would require
further work and development.

The remainder of this paper will be set out as followed. In §2 we will introduce the formulation of the registration model focusing specifically on the diffusion model. Jext in 3 we will discuss the nonlinear multigrid (NMG) framework applied to the diffusion model, long with a detailed analysis to highlight how we can improve the convergence of the Chumchob- hen NMG method. Then in §4 we will formulate our non-folding constraint model, and also present an optimis tion for the implementation of the constraint. §5 will comprise of tests and comparisons with our proposed work, and finally in §6 we

97 will present our conclusions.

³⁸ 2 Review of the registration model and its algorithm of [16]

The model. The diffusion regulariser is a popular croce group variational models [7–10,30], it imposes a simple smoothness constraint upon the displacement field and is given by the following

$$\mathscr{R}^{\mathrm{Diff}}(\boldsymbol{u}) = \frac{1}{2} \int_{\mathcal{M}} \sum_{s=1}^{\infty} \left| \nabla u_s \right|^2 d\Omega .$$
 (2.1)

In fact, the diffusion model is one of the few models that coincides with models from optical flow
frameworks (see [8,9,30] as examples), thich is particularly useful when registering sequences of images.
The diffusion model is given by the folic ting m nimisation problem

$$\min_{\boldsymbol{u}} E^{\text{Diff}}(\boldsymbol{u}) = \mathscr{D}(R, \boldsymbol{\nabla} \boldsymbol{v}) + \mathscr{R}^{\text{Diff}}(\boldsymbol{u}) = \frac{1}{2} \int_{\Omega} (T_{\boldsymbol{u}} - R)^2 + \alpha \sum_{s=1}^2 |\nabla u_s|^2 \, d\Omega$$
(2.2)

where $T_{\boldsymbol{u}} \equiv T(\boldsymbol{x} + \boldsymbol{u})$ and $R \equiv R_{(\boldsymbol{u})}$ The corresponding Euler-Lagrange (EL) equations are derived from the following limits

$$\lim_{\varepsilon_1 \to 0} \frac{E^{\operatorname{Diff}}(u_1 + \varepsilon_1 \phi_1, u_2) - E^{\operatorname{Diff}}(u_1, u_2)}{\varepsilon_1} = 0, \ \lim_{\varepsilon_2 \to 0} \frac{E^{\operatorname{Diff}}(u_1, u_2 + \varepsilon_2 \phi_2) - E^{\operatorname{Diff}}(u_1, u_2)}{\varepsilon_2} = 0 \quad (2.3)$$

which eventually result in the ollowing integrals

$$\int_{\Omega_{\iota}} \phi_m \Big[\delta_{\iota_m} T_{\boldsymbol{u}} \left(T_{\boldsymbol{u}} - R \right) - \alpha \Delta u_m \Big] d\Omega + \alpha \int_{\partial \Omega} \phi_m \left(\nabla u_m \cdot \boldsymbol{n} \right) dS = 0$$
(2.4)

and thus, afte the us, of the fundamental lemma of calculus of variations, yield the EL equations

$$-\alpha\Delta u_m + F_m(\boldsymbol{u}) = 0 \tag{2.5}$$

with Neuman boundary conditions $\nabla u_m \cdot n = 0$ where *n* denotes the outward unit normal and

$$F_m(\boldsymbol{u}) = \partial_{\boldsymbol{u}_m} T_{\boldsymbol{u}} \left(T_{\boldsymbol{u}} - R \right)$$
(2.6)

denote the force terms, for m = 1, 2.

¹¹⁰ 2.1 Optimise-discretise approach for diffusion model

We consider a numerical approximation to the EL equations (2.5) by discretising the image domain Ω into a uniform $n \times n$ mesh with interval width h, using a finite difference (FD) meth d. The size of the mesh is chosen to be equal to the dimension of the image (e.g. 512×512 to coinci. with resolution of given images) and in general need not be square, however in this paper we consider sq. re images as this is common for medical image slices. Using the following central FD approximations

$$(\partial_{u_{1}}T_{\boldsymbol{u}})_{i,j} \approx \frac{1}{2h} \left((T_{\boldsymbol{u}})_{i+1,j} - (T_{\boldsymbol{u}})_{i-1,j} \right), \ (\partial_{u_{2}}T_{\boldsymbol{u}})_{i,j} \approx \frac{1}{2h} \left((T_{\boldsymbol{u}})_{i,j\neg} - (\neg)_{i,j-1} \right)$$
$$(\Delta u_{m})_{i,j} \approx \frac{1}{h^{2}} \left((u_{m})_{i,j-1} + (u_{m})_{i-1,j} - 4(u_{m})_{i,j} + (u_{m})_{i+1,j\neg} (u_{m})_{i,j+1} \right)$$
(2.7)

at a general discrete point (i, j), leads to the following discrete versi ns of the EL equations (2.5)

$$-\alpha \left(\Delta u_m\right)_{i,j} + \left(F_m(\boldsymbol{u})\right)_{i,j} = \boldsymbol{\sigma}$$
(2.8)

117 with

$$(F_m(\boldsymbol{u}))_{i,j} = (\partial_{u_m} T_{\boldsymbol{u}})_{i,j} \left((T_{\gamma})_{i,j} - (\boldsymbol{\iota})_{i,j} \right)$$
(2.9)

118 for m = 1, 2 and $i, j = 2, \ldots, n-1$.

119 2.2 The collective pointwise smoother

The term smoother, which stems from multigric theory is nothing but an iterative solver. In [16] the lexicographic Gauss-Seidel (GS-LEX) method was employed to solve the linear part of the system (2.8) through an inner iteration loop, and a fixed point iteration scheme to solve the non-linear part through an outer iteration loop. In a lexicographical or a ring system, a general discrete point (i, j) as in (2.9) is linked to the global index k = (j-2)(n-1) + (i-1), with n the size of the discrete image dimensions; then for m = 1, 2, we get

$$-\alpha \left(\Lambda u_r \right)_k + \left(F_m(\boldsymbol{u}) \right)_k = 0 \tag{2.10}$$

as illustrated in Figure 1. Now to solve the non-linear part of this system, we employ the following semi-implicit fixed point iteration scheme

$$(\Delta u_m)_k^{(l+1)} + (F_m(\boldsymbol{u}))_k^{(l+1)} = 0$$
(2.11)

128 where

$$(F_{1}(\boldsymbol{u}))_{k}^{(l+1)} = \left(\partial_{u_{1}} r \left(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l)}\right)\right)_{k} \left(\left(T(x_{1} + u_{1}^{(l+1)}, x_{2} + u_{2}^{(l)})\right)_{k} - \left(R(x_{1}, x_{2})\right)_{k}\right)$$
$$(F_{2}(\boldsymbol{u}))_{k}^{(l+1)} = \left(\partial_{u_{2}} T \left(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l)}\right)\right)_{k} \left(\left(T(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l+1)})\right)_{k} - \left(R(x_{1}, x_{2})\right)_{k}\right).$$
(2.12)

The key questing addressed in [16] was how to treat the non-linear terms $(T(x_1 + u_1^{(l+1)}, x_2 + u_2^{(l)}))_k$, ($T(x_1 + u_1^{(l)}, z_2 + u_2^{(l_2^{-1})}))_k$ in a GS-LEX scheme. It proposed to use the first order approximations:

$$\begin{pmatrix} Y_{1}, y_{1}, y_{2} + u_{2}^{(l+1)}, x_{2} + u_{2}^{(l)} \end{pmatrix}_{k} \approx \left(T(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l)}) \right)_{k} \\ + \left((u_{1})_{k}^{(l+1)} - (u_{1})_{k}^{(l)} \right) \left(\partial_{u_{1}} T(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l)}) \right)_{k} \\ \left(T(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l+1)}) \right)_{k} \approx \left(T(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l)}) \right)_{k} \\ + \left((u_{2})_{k}^{(l+1)} - (u_{2})_{k}^{(l)} \right) \left(\partial_{u_{2}} T(x_{1} + u_{1}^{(l)}, x_{2} + u_{2}^{(l)}) \right)_{k}$$

which are substituted back into the discrete force terms (2.10) leading to the following discrete system

$$-\alpha \left(\Delta u_m\right)_k^{(l+1)} + \left(\partial_{u_m} T_{\boldsymbol{u}}\right)_k^{(l)} \left(\left(T_{\boldsymbol{u}}\right)_k^{(l)} + \left(\left(u_m\right)_k^{(l+1)} - \left(u_m\right)_k^{(l)}\right) \left(\partial_{u_m} T_{\boldsymbol{u}}\right)_k^{(l)} - \left(R\right)_k \right) = 0$$
(2.13)

with $(T_{\boldsymbol{u}})_k^{(l)} \equiv (T(\boldsymbol{x} + \boldsymbol{u}^{(l)}))_k$ etc. for m = 1, 2. Using the FD approximations (2.7), we can write (2.13) in the following way

$$-\frac{\alpha}{h^2} \left((u_m)_{k-n}^{(l+1)} + (u_m)_{k-1}^{(l+1)} \right) + \left(\left((\partial_{u_m} T_{\boldsymbol{u}})^2 \right)_k^{(l)} + \frac{4\alpha}{h^2} \right) (u_m)_k^{(l+1)} - \frac{\alpha}{h^2} \left((u_m)_{k+1}^{(l+1)} + (u_m)_{k+n}^{(l+1)} \right) = \left((\partial_{u_m} T_{\boldsymbol{u}})^2 \right)_k^{(l)} (u_m)_k^{(l)} - (\partial_{u_m} T_{\boldsymbol{u}})_l^{(l)} \left((T_{\boldsymbol{u}})_k^{(l)} - (R)_k \right)$$
(2.14)

for m = 1, 2. Then to compute the (l + 1) updates in (2.14), we use a GS-L. X based method.



Figure 1: Illustration of how the domain Ω is discretised by $n \times n$ grid points. The dashed blue line represents the boundary $\partial\Omega$ of the discrete domain, with the boxed points representing the used boundary points, and the black lines show the $(n-2) \times (r-2)$ grid corresponding to the blue interior points. The indexing on the interior points show now the clobal index k is ordered lexicographically.

However, such an iterative methed is not rective as a standalone solver since solving the discrete system

of PDEs (2.10) pixel-wise can .e., ' to a very high computational cost, especially for big images. This

fact is well-known for simpler PDEs such as the Poisson equation (corresponding to $F_m = 0$ and $h \to 0$). One natural way of reducing the cost of calculating the displacement field is a NMG method in which

139 this (slow) iterative method is used as a smoother.

There has already been . lot of work regarding the implementation of NMG methods [21,25,27,28,32] for related models, each h vin, its wn unigrid iterative solver, however most of these works do not address the non-linearity in the sm. 'a ity measure directly, instead linear diagonal terms or augmented systems are used. Chumch o-Che: [16] proposed a robust solver which does directly deal with this non-linearity arising from the S 'D terr, however an inaccurate analysis of the NMG method lead to a less than optimal convergence is nor the NMG method which we will demonstrate in the next section.

146 2.3 The Nix method

There are two neoretical principles driving multigrid methods for linear PDEs. The first is that, although standard iterat. The methods such as the Jacobi and GS methods have poor convergence rates when used independently, they are effective at smoothing out any high frequency error components within a small number of iterations. This property leads to the second key principle of multigrid methods, namely low frequency error components can be well approximated on a coarser grid. Naturally an approximate and accurate solution on a coarser grid can then be interpolated back to the fine grid to approximate the original problem; this two-grid approach is significantly cheaper than working solely on the fine grid.

In fact this strategy allows us to obtain a more accurate approximation efficiently as we can perform a 154 larger number of iterations on the coarser grid in less time when compared with iterating the fine grid 155 alone. This fine-coarse-fine strategy, known as the two-grid V-cycle (see [5] for details), can however be 156 repeated on the coarse grid to interact with even coarser grids until some coarsest grid with few points. 157

While multigrid frameworks are known, and indeed very easy to implement for linea. cases, problems 158 like (2.5) which are highly non-linear prove significantly more difficult to develop a converging NMG 159 method. Now we present the FAS-NMG algorithm of [16] for (2.10) before we hig light the omissions 160 in the analysis which resulted in an overestimated smoothing rate (thus leading to a less optimal NMG 161 method with slower convergence rate), and include our more accurate analysis \neg over \neg one this problem. 162 Here FAS stands for "full approximation scheme" by A. Brandt for solving a _____-line_ " operator equation. 163 First consider a two grid setting where Ω^h denotes a fine grid and Ω^H a coal set grid with $h = \frac{1}{n-1}$, H = 2h. 164 Also denote the system (2.10) by the operator notation on Ω^h

165

$$\mathcal{N}^h(\boldsymbol{u}^h) = \mathcal{G}^h \tag{2.15}$$

with 166

$$\mathcal{N}^{h} = \begin{pmatrix} \left(\mathcal{N}_{1}^{h}\right)_{k} \\ \left(\mathcal{N}_{2}^{h}\right)_{k} \end{pmatrix}, \ \boldsymbol{u}^{h} = \begin{pmatrix} \left(u_{1}^{h}\right)_{k} \\ \left(u_{2}^{h}\right)_{k} \end{pmatrix}, \ \mathcal{G}^{h} \begin{pmatrix} \begin{pmatrix} \prime - h \\ \sqrt{2} \\ \sqrt{2} \end{pmatrix}_{k} \end{pmatrix}.$$
(2.16)

and where $(\mathcal{N}_1^h)_k = (F_1(\boldsymbol{u}^h))_k - \alpha (\Delta^h u_1^h)_k$, $(\mathcal{N}_2^h)_k = (\square(\boldsymbol{u}^h))_k - \alpha (\Delta^h u_2^h)_k$, $(g_1^h)_k = (g_2^h)_k = 0$, $k = 1, 2, \ldots, (n-2)^2$. The main steps of the FAS-NMG at as reliave. 167 168

Smoothing step. Apply the iterative method (2.14) starting from some initial guess. This 169 is the pre-smoothing step required to obtain a smooth app. ximation $\bar{\boldsymbol{u}}^h = (\bar{u}_1^h, \bar{u}_2^h)^T$ which has residual 170 $\boldsymbol{r}^h = \mathcal{G}^h - \mathcal{N}^h(\bar{\boldsymbol{u}}^h).$ 171

To improve this smooth approximation, it remains to ompute the algebraic error (or the residual cor-172 rection) $e^h = (e_1^h, e_2^h)^T = u^h - \bar{u}^h$ which cannot be omputed directly on Ω^h . 173

Restriction. Since only smooth errors can pproximated on a coarser grid, we first solve the 174 FAS coarse grid residual equation 175

$$\mathcal{N}^{H}(\boldsymbol{u}^{H}) \equiv \mathcal{N}^{H^{\prime}-H} + \boldsymbol{e}^{H}) = \boldsymbol{r}^{H} + \mathcal{N}^{H}(\bar{\boldsymbol{u}}^{H}) \equiv \mathcal{G}^{H}$$
(2.17)

where $\bar{\boldsymbol{u}}^{H} = \mathcal{R}_{h}^{H} \bar{\boldsymbol{u}}^{h}$, $\boldsymbol{e}^{H} = \mathcal{R}_{h}^{H} \boldsymbol{e}^{h}$, $\boldsymbol{r}^{H} = \mathcal{R}_{h}^{H} \boldsymbol{r}^{h}$ and \mathcal{R}_{h}^{H} is the restriction operator, which we take to be the full-weighted restriction operator, define \boldsymbol{v} by the following stencil 176 177

$$\mathcal{T}_{h}^{I} = \frac{1}{16} \begin{bmatrix} 1 & 2 & 1 \\ 2 & 4 & 2 \\ 1 & 2 & 1 \end{bmatrix}_{h}^{H}$$
(2.18)

Coarse grid solution. For a t original method (or in a multigrid setting where Ω^H is the coarsest level 178 and computations are inex, 205, ve), the above coarse grid equation must be solved accurately to obtain 179 solutions u^H . Based on his u^{-1} and its initial guess \bar{u}^H , we obtain the residual correction 180

$$\boldsymbol{e}^{H} = \boldsymbol{u}^{H} - \bar{\boldsymbol{u}}^{H}. \tag{2.19}$$

Interpolation. Now we vish to use (2.19) to correct the approximations on the finer grid Ω^h ; we do 181 this by interpolatin, ' the c' rrections using bilinear interpolation. That is we compute 182

$$e^{h} = \mathcal{I}_{H}^{h} e^{H}, \quad \mathcal{I}_{H}^{h} = \frac{1}{4} \begin{bmatrix} 1 & 2 & 1 \\ 2 & 4 & 2 \\ 1 & 2 & 1 \end{bmatrix}_{H}^{h}.$$
 (2.20)

Once the corrections have been interpolated to the next fine grid level, we use them to update the 183 current grid le el approximations via $u^h = \bar{u}^h + e^h$. After the approximations have been corrected, we 184 use a post-smoothing step to remove any interpolation errors. This process of interpolation, correction 185 and smoothing is repeated until the approximations on the original grid level have been corrected and 186 smoothed, thus resulting in our final solutions \boldsymbol{u}^h . 187

- **Remark 2.1.** According to the work done in [26], there are three conditions which need to be satisfied 188
- regarding the orders of the restriction and interpolation methods for a convergent NMG. For an order M 189 PDE, we require 190
- (i) $m_R + m_I \ge M$; (ii) $m_I \ge M$ and $m_R \ge 0$; (iii) $m_R \ge M$ and $m_I \ge 0$ 191

where m_R , m_I denote the high frequency orders of the restriction and interpolation f being respectively. 192

- In our case we have $m_R = 2$, $m_I = 2$, for the full-weighted restriction and bilinear interpol. *ion operators* 193
- respectively, and so all three conditions are satisfied. 194

Below the FAS-NMG algorithm has been summarised 195

 $\overrightarrow{\textbf{Algorithm 1 } \boldsymbol{u}_h^{(k+1)} \leftarrow FASNMG(R^h, T^h, n, h, level, \boldsymbol{u}_h^{(k)}, \mathcal{G}^h, \alpha, \nu_1, \nu_2)}$

 $\frac{\overline{\boldsymbol{u}}_{h}^{(k)}}{\overline{\boldsymbol{u}}_{h}^{(k)}} \underbrace{S \text{ nooth}(R^{h}, T^{h}, \boldsymbol{u}_{h}^{(k)}, \mathcal{G}^{h}, \alpha, \nu_{1})}$ 1: Pre-smoothing step by performing ν_1 steps (relaxation sweeps) 2: Coarse-grid correction

- Compute the residual $\boldsymbol{r}_{h}^{(k)} = \mathcal{G}^{h} \mathcal{N}^{h}(\boldsymbol{u}_{h}^{(k)})$ Restrict residual and smooth approximations $\boldsymbol{r}_{H}^{(k)} = \mathcal{R}_{h}^{H}\boldsymbol{r}_{h}^{(k)}, \, \bar{\boldsymbol{u}}_{H}^{(k)} = \mathcal{R}_{h}^{H}\bar{\boldsymbol{u}}^{(k)}$ Set $level \rightarrow level 1, \, H = 2h, \, nc = \frac{n}{2}$
 - Form RHS of coarse grid PDEs $\mathcal{G}^{H} = r^{H} + \mathcal{N}^{H}(\bar{u}_{H}^{(k)})$
- Solve residual equation on coarse grid to obtain approximatio..... $ilde{u}_{H}^{(k)}$ 3: if level = 1 then
- 3: If level = 1 then Solve to obtain solutions u_H^(k) to high accuracy using a coarsest, rid solver.
 4: else level > 1 Repeat the FAS-NMG procedure recursively to the n xt level i.e. ū_H^(k) ← FASNMG(R^H, T^H, nc, H, level 1, ũ_H^(k), G^H, v₁, v₂)
 5: end if
- 5: end if
 - Compute the correction $oldsymbol{e}_{H}^{(k)}=oldsymbol{u}_{H}^{(k)}-oldsymbol{ar{u}}_{H}^{(k)}$
- Interpolate the correction $e_H = u_H u_H$ Interpolate the correction to next fine grid level $e_h^{(k)} = \tau_T^h e_H^{(k)}$ Update current grid level approximations using converse $\hat{u}_h^{(k)} = \bar{u}_h^{(k)} + e_h^{(k)}$ 6: Post-smoothing step by performing ν_2 steps (relaxation sweeps) $u_h^{(k+1)} \leftarrow Smooth(R^h, T^h, \hat{u}_h^{(k)}, \mathcal{G}^h, \alpha, \nu_2)$

In [16], the coarsest solver that was adopted was a. additive operator splitting (AOS) method. For the diffusion model, it takes the following for $\dots \gamma_m^{(k+1)} = \frac{1}{2} \sum_{s=1}^2 \left[I - 2\tau \alpha L_{x_s} \right]^{-1} \left(u_m^{(k+1)} + \tau g_m - \tau F_m(u) \right)$ where I denotes the identity operator, $\gamma > 0$ t₁ e time-step, g_m the RHS coming from the NMG frame-196 197 198 work, $F_m(\boldsymbol{u})$ the force terms given in (2.2) for i = 1, 2 and $L_{x_s} = \partial_{x_s x_s}$ denote the parts of the discrete 199 Laplace operator in the x_s directions for s = 1, 2 respectively. The above equations are updated along 200 the x_1, x_2 directions separately, this leading to the system 201

$$\begin{cases} \left[I - \gamma \tau \alpha L_{x_1}\right] u_{m,p_1}^{\left(k+\frac{1}{2}\right)} = u_m^{\left(k\right)} + \tau g_m - \tau F_m(\boldsymbol{u}), \\ \left[I - 2\tau \alpha L_{x_2}\right] u_{m,p_2}^{\left(k+\frac{1}{2}\right)} = u_m^{\left(k\right)} + \tau g_m - \tau F_m(\boldsymbol{u}) \end{cases}$$
(2.21)

with the updates $u_m^{(k+1)} = \frac{1}{2} \left(\frac{(k+\frac{1}{2})}{m,p_1} + u_{m,p_2}^{(k+\frac{1}{2})} \right)$ for m = 1, 2. 202

Remark 2.2. In [16], he *i*-elliviticity for the proposed smoother was computed in order to check whether 203 the smoother was suitable for ise in the NMG method. From the resulting calculation, the h-ellipticity 204 was found to have c value of $\frac{16}{16}$, and it was concluded that the smoother was suitable for use in the NMG 205 method. By performing the same calculation for our proposed smoother in §2.2, which is similar to the one used in [16], we also stained a value of $\frac{1}{16}$ and thus reached the same conclusion. 206 207

An improved analysis of the NMG algorithm of [16] 3 208

As mentioned, the above Algorithm 1 as implemented by Chumchob-Chen [16] could still be slow to 209 converge to a solution from new experiments. We found that a major part of this convergence problem 210 was a result of an inaccurate analysis of the smoothing rate, which lead to an overestimation of the rate. 211

- By re-evaluating the analysis of the NMG method, as well as building in some new components, lead to 21 2
- our NMG algorithm with a vastly improved convergence rate. 213

In this section we will outline our more detailed and accurate analysis of the NMG framework. We do
this by analysing two key components of the NMG algorithm (namely the smoothing rate of the smoother
and the coarsest grid solver), which leads to an optimal NMG method.

²¹⁷ 3.1 Smoother analysis using Local Fourier Analysis (LFA)

We begin our analysis of the NMG method by showing an improved, and more a curate, LFA of the smoother scheme that was described in [16]. A discrete error (e.g. residual) function on a grid can be written as a sum of two terms:

• high frequency error components (are not visible if the problem is r stri \mathbb{Z}^4 to a coarser grid);

• low frequency error components (that can be accurately represented a coarser grid).

The sole purpose of the smoother, within a MG framework, is to remove any high frequency error components. Local Fourier Analysis (LFA) is used to measure how ether scheme is.

Although LFA was originally designed to analyse discrete linear operator equations, it was extended by 225 A. Brandt (see [38]) to study non-linear operators via a 'freezin, " α localised coefficients. To start we 226 first assume that we are working on an infinite grid, this the. allow us to remove any influence from 227 the boundary conditions. Next we assume that the discrete form α_{3} non-linear operator, with variable 228 coefficients, can be replaced locally by an operator with cons. Int coefficients and extended to the infinite 229 grid. We need to ensure all high frequency error component, are removed prior to restriction to a coarse 230 grid. As a result it is imperative that we know how effective ∞ relaxation scheme is at smoothing out 231 the errors so we can adjust the number of sweeps required for the pre- and post-smoothing steps. Using 232 LFA we obtain a value μ which is defined to be the smoothing factor for a given relaxation scheme. 233

LFA for pointwise smoother from [16]. While the smoother we described in §2.2 is similar to the one used in [16], we found that the smoother and pris in ¹16] contained an omission which lead to a very over-optimistic smoothing rate (practically to a show convergence if using it as a guide). In [16], the discrete system (2.10) was written in the foll show v

$$\mathcal{N}_{+}^{h}\boldsymbol{u}_{new}^{h} + \mathcal{N}_{0}^{h}\boldsymbol{u}_{new}^{h} + \mathcal{N}_{-}^{h}\boldsymbol{u}_{old}^{h} = \mathcal{G}^{h}$$

$$(3.1)$$

where u_{new}^h , u_{old}^h denote the current at d previous approximations of u^h respectively, and

$$\mathcal{N}^{h}_{+} = \begin{pmatrix} -\alpha \mathscr{L}^{h}_{+} & 0\\ 0 & -\epsilon \mathscr{L}^{h}_{+} \end{pmatrix}, \ \mathcal{N}^{h}_{0} = \begin{pmatrix} -\alpha \mathscr{L}^{h}_{0} + \sigma^{h}_{11} & \sigma^{h}_{12} \\ \sigma^{h}_{12} & -\alpha \mathscr{L}^{h}_{0} + \sigma^{h}_{22} \end{pmatrix}$$
$$\mathcal{N}^{h}_{-} = \begin{pmatrix} -\alpha \mathscr{L}^{h}_{-} & 0\\ \cdots & -\alpha \mathscr{L}^{h}_{-} \end{pmatrix}, \ \mathcal{G}^{h} = \begin{pmatrix} g^{h}_{1} - F^{h}_{1} \\ g^{h}_{2} - F^{h}_{2} \end{pmatrix}$$
(3.2)

with $\sigma_{pq}^{h} = \partial_{u_{p}} T_{\boldsymbol{u}}^{h} \partial_{u_{q}} T_{\boldsymbol{u}}^{h}$, g_{m}^{h} '-note the RHS coming from the NMG scheme, F_{m}^{h} are the discrete force terms as given in (2.9) and where \mathscr{L}_{+}^{h} , \mathscr{L}_{0}^{h} , \mathscr{L}_{-}^{h} define the following stencils

$$\mathscr{L}_{-}^{h} = \frac{1}{h^{2}} \begin{pmatrix} 0 & 0 & 0\\ 1 & 0 & 0\\ 0 & 1 & 0 \end{pmatrix}, \ \mathscr{L}_{0}^{h} = \frac{1}{h^{2}} \begin{pmatrix} 0 & 0 & 0\\ 0 & -4 & 0\\ 0 & 0 & 0 \end{pmatrix}, \ \mathscr{L}_{-}^{h} = \frac{1}{h^{2}} \begin{pmatrix} 0 & 1 & 0\\ 0 & 0 & 1\\ 0 & 0 & 0 \end{pmatrix}.$$
(3.3)

for p, q, m = 1, 2. The smoothing rate in [16] was then calculated on a 32×32 grid after a total of 5 241 outer and 5 in versitient tion loops had been performed, thus resulting in an average smoothing rate of 24 2 $\mu_{avg} \approx 0.5$ when using $\alpha = \frac{1}{10}$. However, in the analysis of [16] we notice that the $(u_m)_k^{(l)}$ terms, which 24 3 result from 'ne risation of the SSD term, where not included in the smoothing rate calculation. This 244 omission me. \cdot that the obtained rate of 0.5 was a vast overestimation of the actual smoothing rate, 245 and as a result this lead to an underestimation of the number of pre-smoothing steps required before 246 restriction. This means that when we restrict the problem to a coarser grid, there are still high frequency 247 error components on the fine grid which have not been removed, and so the coarse grid correction that 248 we obtain is much less accurate thus leading to more NMG cycles being required to reach an accurate 249 solution. This omission, as we will now show, has a noticeable effect on the smoothing rate. 250

Revised LFA for pointwise smoother from §2.2. Here we will repeat the analysis of the smoothing rate, with the $(u_m)_k^{(l)}$ terms included, in order to illustrate the impact the addition of these terms have on the smoothing rate. We begin by writing the discrete equations (2.10) in the following form

$$\mathcal{N}^h \boldsymbol{u}^h + \mathcal{M}^h \boldsymbol{u}^h = \mathcal{G}^h \tag{3.4}$$

where \mathcal{G}^h is as in (3.2), and

$$\mathcal{N}^{h} = \begin{pmatrix} -\alpha \Delta^{h} + \sigma_{11}^{h} & 0\\ 0 & -\alpha \Delta^{h} + \sigma_{22}^{h} \end{pmatrix}, \ \mathcal{M}^{h} = \begin{pmatrix} -\sigma_{11}^{h} & 0\\ 0 & -\sigma_{22}^{h} \end{pmatrix}$$
(3.5)

using the following representation of the discrete Laplace operator $\Delta^h \equiv \mathscr{L}_+^i + \mathscr{L}_0^{i-1} \mathscr{L}_-^h$, with $\mathscr{L}_+^h, \mathscr{L}_0^h, \mathscr{L}_-^h$ as defined in (3.3), then we can express (3.4) in the following way

$$\mathcal{N}^{h}_{+}\boldsymbol{u}^{h}_{new} + \mathcal{N}^{h}_{0}\boldsymbol{u}^{h}_{new} + \mathcal{N}^{h}_{-}\boldsymbol{u}^{h}_{old} + \mathcal{M}^{h}\boldsymbol{u}^{h}_{ol} = \mathcal{G}^{h}$$
(3.6)

and subtracting (3.6) from (3.4) yields the local error equation g^{\dagger} in by

$$\left[\mathcal{N}^{h}_{+} + \mathcal{N}^{h}_{0}\right]\boldsymbol{e}^{h}_{new} = -\left[\mathcal{N}^{h}_{-} + \mathcal{M}^{\dagger}_{}\right]_{old}^{\iota}$$
(3.7)

where \mathcal{N}^h_+ , \mathcal{N}^h_0 , \mathcal{N}^h_- are as defined in (3.2) and

$$\boldsymbol{e}_{new}^{h} = \left(e_{1\,new}^{h}, e_{2\,new}^{h}\right)^{T}, \ \boldsymbol{e}_{old}^{h} = \left(e_{1\,old}^{h}, e_{2\,old}^{h}\right)^{T}.$$
(3.8)

Using Fourier components, we can rewrite (3.7) in the tollowing way

$$\left[\hat{\mathcal{N}}^{h}_{+}(\boldsymbol{\theta}) + \hat{\mathcal{N}}^{h}_{0}(\boldsymbol{\theta})\right]\psi^{new}_{\boldsymbol{\theta}}\exp\left(\frac{2i\theta_{1}i\pi}{n} + \frac{2i\theta_{2}j\pi}{n}\right) = -\lfloor\hat{\mathcal{N}}^{h}_{-}(\boldsymbol{\theta}) + \hat{\mathcal{M}}^{h}(\boldsymbol{\theta})\right]\psi^{old}_{\boldsymbol{\theta}}\exp\left(\frac{2i\theta_{1}i\pi}{n} + \frac{2i\theta_{2}j\pi}{n}\right) \quad (3.9)$$

where $i = \sqrt{-1}$, $\theta \in \Theta = [-\pi, \pi)^2$ and ψ_{θ}^* are Fourier coefficients. From here we determine the local smoothing rate μ_{loc} using the following

$$\mu_{\max} = \max_{loc} \mu_{loc}, \qquad \gamma_{\infty} \equiv \mu_{loc}(\boldsymbol{\theta}) = \sup\left\{\rho(\hat{\boldsymbol{S}}^{h}(\boldsymbol{\theta})) \middle| \boldsymbol{\theta} \in \boldsymbol{\Theta}_{high}\right\}$$
(3.10)

where $\Theta_{high} = \Theta \setminus \left[-\frac{\pi}{2}, \frac{\pi}{2}\right)^2$, $\rho(\cdot)$ denotes the spectral radius, and the amplification matrix $\hat{S}^h(\theta)$ is given by

$$\hat{\boldsymbol{S}}^{h}(\boldsymbol{\theta}) = -\left[\hat{\boldsymbol{\mathcal{N}}}_{-}^{h}(\boldsymbol{\theta}) + \hat{\mathcal{N}}_{0}^{h}(\boldsymbol{\theta})\right]^{-1} \left[\hat{\mathcal{N}}_{-}^{h}(\boldsymbol{\theta}) + \hat{\mathcal{M}}^{h}(\boldsymbol{\theta})\right]$$
(3.11)

264 with

$$\hat{\mathcal{N}}_{+}^{h}(\boldsymbol{\theta}) = \begin{pmatrix} -\frac{\alpha}{h^{2}} \left(e^{-i\boldsymbol{\omega}_{+}} + e^{-i\boldsymbol{\omega}_{2}} \right) & 0 \\ 0 & -\frac{\alpha}{h^{2}} \left(e^{-i\boldsymbol{\omega}_{1}} + e^{-i\boldsymbol{\omega}_{2}} \right) \end{pmatrix}, \\ \hat{\mathcal{N}}_{0}^{h}(\boldsymbol{\theta}) = \begin{pmatrix} \frac{4\alpha}{h^{2}} + \sigma_{11}^{h} & 0 \\ 0 & \frac{4\alpha}{h^{2}} + \sigma_{22}^{h} \end{pmatrix}, \\ \hat{\mathcal{N}}_{-}^{h}(\boldsymbol{\theta}) = \begin{pmatrix} -\frac{\alpha}{h^{2}} \left(e^{-i\boldsymbol{\omega}_{+}} + e^{i\boldsymbol{\omega}_{2}} \right) & 0 \\ 0 & -\frac{\alpha}{h^{2}} \left(e^{i\boldsymbol{\omega}_{1}} + e^{i\boldsymbol{\omega}_{2}} \right) \end{pmatrix}, \\ \hat{\mathcal{M}}^{h}(\boldsymbol{\theta}) = \begin{pmatrix} -\sigma_{11}^{h} & 0 \\ 0 & -\sigma_{22}^{h} \end{pmatrix}$$
(3.12)

where $\omega_m = \frac{2\theta_m \pi}{n}$ for m = 1, 2. Implementing the revised local smoothing rate formulae, under the same 265 conditions that were used in [16], we obtained an average and maximum smoothing rate of $\mu_{avg} \approx 0.69854$ 266 and $\mu_{max} \approx (74762)$ espectively. By the smoothing rate of 0.5 in [16] within each outer iteration, 5 267 inner iterations', result in reduction of the error by 0.0313 which appeared satisfactory. However 268 5 inner ite wood reduce only by 0.17 and 0.23 respectively using our new smoothing rates μ_{avg} 269 and μ_{max} . In order to reduce to the level of error claimed in [16], we estimate that we would require up 270 to 12 inner ite, tions. So we see that the original analysis in [16] resulted in the estimated number of 271 pre-smoothing steps being roughly half of the number of steps that would actually be required to reduce 272 the error to quoted level. 273

²⁷⁴ 3.2 Convergence analysis of two coarsest grid solvers by LFA

Next we give a simple solution to the challenging problem of getting the convergence rate of a non-linear
iterative method. Here we remark that this analysis was not performed in [16]. Co sequently, we can
compare methods and guide the number of iterations to be prescribed on the coars and grid. Recall that
the AOS solver (2.21) was used by Chumchob-Chen [16]. Here we shall propose to use a mod point type
solver on the coarsest grid instead.

Our coarsest grid solver. From §2.2 we have the following lexicographically viered discrete system
 of linear equations

$$-\frac{\alpha}{H^2} \left((u_m)_{k-n}^{(l+1)} + (u_m)_{k-1}^{(l+1)} \right) + \left(\left((\partial_{u_m} T_{\boldsymbol{u}})^2 \right)_k^{(l)} + \frac{4\alpha}{H^2} \right) (u_m)_k^{(l+1)} - \frac{\alpha}{H^2} \left((u_m)_{k+1}^{(l+1)} + (u_m)_{k+n}^{(l+1)} \right) = \left((\partial_{u_m} T_{\boldsymbol{u}})^2 \right)_k^{(l)} (u_m)_k^{(l)} - (\partial_{u_m} T_{\boldsymbol{u}})_k^{(l)} \left((T_{\boldsymbol{u}})_k^{(l)} - (R)_k \right)$$
(3.13)

for m = 1, 2. In matrix notation, we can express these equations a. matrix equations $A_m u_m = f_m$, where $u_m, f_m \in \mathbb{R}^{(n-2)^2 \times 1}$ are column vectors and $A_m \in \mathbb{R}^{(\gamma-1)^2 \times (n-2)^2}$ are the block tridiagonal system matrices with the following structure

$$\boldsymbol{A}_{m} = \begin{pmatrix} A_{m_{2}} & I_{1} & & \\ I_{1} & \ddots & \ddots & \\ & \ddots & \ddots & I_{1} \\ & & & I_{1} & A_{m_{n-1}} \end{pmatrix}, \boldsymbol{u}_{m} = \begin{pmatrix} (u_{m})_{k_{2}(2)} \\ \vdots \\ (u_{m})_{k_{i}(z)} \\ \vdots \\ (u_{m})_{k_{n-2}(z-2)} \end{pmatrix}, \boldsymbol{f}_{m} = \begin{pmatrix} (f_{m})_{k_{2}(2)} \\ \vdots \\ (f_{m})_{k_{i}(j)} \\ \vdots \\ (f_{m})_{k_{n-2}(n-2)} \end{pmatrix}$$
(3.14)

where A_{m_i} , $I_1 \in \mathbb{R}^{(n-2) \times (n-2)}$ are matrices with stru +v.e

with $(a_m)_{k_i(j)} = \left(\left(\partial_{u_m} T_u \right)^2 \right)_{k_i(j)} + \frac{4\omega}{H^2}$ and where $k_i(j) = (j-2)(n-1) + (i-1)$ denotes a general lexicographically ordered discrete point (i, j), as shown in Figure 1. Also

$$(f_m)_{k_i(j)} = \left(\left(\partial_{u_m} \mathcal{D}_{\boldsymbol{u}} \right)^2 \right)_{k_i(j)} (u_m)_{k_i(j)} - \left(\partial_{u_m} T_{\boldsymbol{u}} \right)_{k_i(j)} \left(\left(T_{\boldsymbol{u}} \right)_{k_i(j)} - \left(R \right)_{k_i(j)} \right)$$
(3.16)

for m = 1, 2 and i, j = 2, ..., n - 1. Then our proposed algorithm is as shown in Algorithm 2

In order to demonstrate the improvement in convergence rate of our proposed coarsest grid solver over 289 the AOS scheme used in [1,1] we first need a way to measure the convergence rate. To do this we 290 shall employ LFA to est mate the convergence rates of both of our proposed solver and the AOS solver. 291 The purpose is to dis rim late these two estimations. Unfortunately due to the non-linearity of the 292 problem we are unable to the in a sharp measure of the convergence rate, and so using LFA to obtain 293 an approximation i, the bast option. It should be remarked that LFA used for this convergence analysis 294 is only viable on a coarse rid (e.g. 8×8 mesh) as the rate is not sharp especially on a fine grid (e.g. 295 $128 \times 128 \text{ mesh}$ 296

Analysis of the proposed coarsest grid solver. To estimate the convergence rate \mathcal{P} of a given 297 solver, we foll r a si nilar method to that in the smoother analysis shown in §3.1. That is we must 298 evaluate t¹ complimation matrix $\hat{S}^{H}(\theta)$ at every discrete interior point (i, j) for $i, j = 2, \ldots, n-1$ and 299 where n der ste stress the size of the image dimensions. However, where we restricted θ to only consider the 300 high frequency range Θ_{high} in the smoother analysis, now we consider θ over the entire Fourier domain 301 Θ . Since our proposed direct solver is based upon the pointwise smoother shown in §2.2, the derivation 302 of the amplification matrix $\hat{S}^{H}(\theta)$ is very similar to that shown in §3.1. Then, the convergence rate for 303 our proposed direct solver can be estimated locally by the following 304

$$\mathcal{P}_{D \max} = \max_{\text{loc}} \mathcal{P}_{D \log}, \qquad \mathcal{P}_{D \log} \equiv \mathcal{P}_{D \log}(\boldsymbol{\theta}) = \sup \left\{ \rho(\hat{\boldsymbol{S}}^{H}(\boldsymbol{\theta})) \middle| \boldsymbol{\theta} \in \boldsymbol{\Theta} \right\}$$
(3.17)

where $\Theta \in [-\pi,\pi)^2$, $\rho(\cdot)$ denotes the spectral radius and $\hat{S}^H(\theta)$ is the amplification matrix as given by

$$\hat{\boldsymbol{S}}^{H}(\boldsymbol{\theta}) = - \big[\hat{\mathcal{N}}_{+}^{H}(\boldsymbol{\theta}) + \hat{\mathcal{N}}_{0}^{H}(\boldsymbol{\theta}) + \hat{\mathcal{N}}_{-}^{H}(\boldsymbol{\theta})\big]^{-1}\hat{\mathcal{M}}^{H}(\boldsymbol{\theta})$$

with $\hat{\mathcal{N}}^H_+(\boldsymbol{\theta}), \hat{\mathcal{N}}^H_0(\boldsymbol{\theta}), \hat{\mathcal{N}}^H_-(\boldsymbol{\theta}), \hat{\mathcal{M}}^H(\boldsymbol{\theta})$ as in (3.12) and H = 2h.

Algorithm 2 $\boldsymbol{u}_{H}^{(l+1)} \leftarrow DirectSolve(R^{H}, T^{H}, \boldsymbol{u}_{H}^{(k)}, \mathcal{G}^{H}, \alpha, IMAX, Tol)$

1: Initialise $\boldsymbol{u}_{H}^{(l)} = \boldsymbol{u}_{H}^{(k)}$ Construct discrete Laplacian parts of sparse matrices \boldsymbol{A}_{m} 2: for $l = 1, \cdots, IMAX$ do Deform template image using $\boldsymbol{u}_{H}^{(l)} \to T_{\boldsymbol{u}}^{H}$ Compute FD approximations for derivatives of $T_{\boldsymbol{u}}^{H} \to \partial_{u_{1}}T_{\boldsymbol{u}}^{H}, \partial_{u_{2}}T_{\boldsymbol{u}}^{H}$ Compute RHS f_{m} (matrices) and then convert to column vectors \boldsymbol{f}_{m} Add remaining diagonal parts to \boldsymbol{A}_{m} Compute $\boldsymbol{u}_{mH}^{(l+1)} \to \boldsymbol{u}_{mH}^{(l+1)} = \boldsymbol{A}_{m}^{-1}\boldsymbol{f}_{m}$ Reshape $\boldsymbol{u}_{mH}^{(l+1)}$ to matrices $\boldsymbol{u}_{mH}^{(l+1)}$ 3: if $\|\boldsymbol{u}_{1H}^{(l+1)} - \boldsymbol{u}_{1H}^{(l)}\|_{2}^{2} < Tol$ and $\|\boldsymbol{u}_{2H}^{(l+1)} - \boldsymbol{u}_{2H}^{(l)}\|_{2}^{2} < Tol$ then Exit for loop 4: end if 5: end for

Analysis of the block formulation of our proposed parsest grid solver. Previously in order to estimate the convergence rate for the pointwise case we we will have a single equation of the form shown in (3.9) for each discrete interior point from which we would determine the amplification matrix, now however we construct the amplification matrix from a single system of equations with the following structure

$$B\Psi_{\theta}^{n, \circ} = C\Sigma_{\theta}^{\circ old}$$
(3.18)

where $B, C \in \mathbb{R}^{2(n-2)^2 \times 2(n-2)^2}$ and $\Psi_{\theta}^* \in \mathbb{R}^{2(n-2) \times 1}$ are block matrices and block column vectors respectively with structure

$$\boldsymbol{B} = \begin{pmatrix} \boldsymbol{B}_1 & \boldsymbol{\mathcal{J}} \\ \boldsymbol{D} & \boldsymbol{\mathcal{P}}_2 \end{pmatrix}, \, \boldsymbol{C} = \begin{pmatrix} \boldsymbol{C}_1 & \boldsymbol{D} \\ \boldsymbol{D} & \boldsymbol{C}_2 \end{pmatrix}, \, \boldsymbol{\Psi}_{\boldsymbol{\theta}}^* = \begin{pmatrix} \boldsymbol{\psi}_{\boldsymbol{\theta}}^* \\ \boldsymbol{\psi}_{\boldsymbol{\theta}}^* \end{pmatrix}$$
(3.19)

with $B_m, C_m, D \in \mathbb{R}^{(n-2)^2 \times (n-2)^2}$ and $\psi_{\theta}^* \in \mathbb{R}^{(n-2)^2 \times 1}$ given by

$$\boldsymbol{B}_{m} = \begin{pmatrix} B_{m_{2}} & J_{1} & & \\ J_{2} & \ddots & \ddots & \\ & \ddots & \ddots & J_{-} \\ & & J_{2} & B_{\gamma_{n-1}} \end{pmatrix}, \boldsymbol{C} = \begin{pmatrix} C_{m_{2}} & & \\ & \ddots & \\ & & C_{m_{n-1}} \end{pmatrix}, \boldsymbol{D} = \begin{pmatrix} D_{2} & & \\ & \ddots & \\ & & D_{n-1} \end{pmatrix}, \boldsymbol{\psi}_{\boldsymbol{\theta}}^{*} = \begin{pmatrix} (\psi_{\boldsymbol{\theta}}^{*})_{1} \\ \vdots \\ (\psi_{\boldsymbol{\theta}}^{*})_{k} \\ \vdots \\ (\psi_{\boldsymbol{\theta}}^{*})_{(n-2)^{2}} \end{pmatrix}$$
(3.20)

and where $B_{m_j}, C_{m_j}, I_{j}, I_n \in \mathbb{R}^{(n-2) \times (n-2)}$ are given by

$$B_{m_{j}} = \begin{pmatrix} (b_{m})_{k_{2}(j)} & -\frac{c}{H^{2}} i^{i\omega_{j}} \\ -\frac{\alpha}{H^{2}} e^{-\omega_{1}} & \ddots \\ & & & \\ &$$

with $(b_m)_{k_i(j)} = \left(\left(\partial_{u_m} T_{\boldsymbol{u}} \right)^2 \right)_{k_i(j)} + \frac{4\alpha}{H^2}, (c_m)_{k_i(j)} = \left(\left(\partial_{u_m} T_{\boldsymbol{u}} \right)^2 \right)_{k_i(j)}, (d)_{k_i(j)} = \left(\partial_{u_1} T_{\boldsymbol{u}} \right)_{k_i(j)} (\partial_{u_2} T_{\boldsymbol{u}})_{k_i(j)},$ $\omega_m = \frac{2\theta_m \pi}{n} \text{ and } k_i(j) = (j-2)(n-1) + (i-1) \text{ for } m = 1, 2 \text{ and } i, j = 2, \dots, n-1.$ Then the convergence rate \mathcal{P}_B for the block formulation of our direct solver is estimated from the following

$$\mathcal{P}_{B} \equiv \mathcal{P}_{B}(\boldsymbol{\theta}) = \sup\left\{\rho\left(\hat{\boldsymbol{S}}^{H}(\boldsymbol{\theta})\right) \middle| \boldsymbol{\theta} \in \boldsymbol{\Theta}\right\}$$
(3.22)

with amplification matrix $\hat{S}^{H}(\theta) = B^{-1}C$. On this coarsest grid, n is small so estimating \mathcal{P}_{B} is feasible.

Convergence analysis for AOS solver. We again remark that an analysis to estimate the convergence of the coarsest solver in [16] was not performed. From [16], the AOS scheme for the diffusion model is shown in (2.21) for m = 1, 2. We use a similar method to the one shown in §3.1 to derive the amplification matrix for the AOS method. However, since the AOS scheme solves along the $z_1 \, \ldots \, d \, x_2$ directions separately, we will obtain two convergence rates $\mathcal{P}_{A_1}, \mathcal{P}_{A_2}$ for these directions respectively. We start by expressing the discrete versions of (2.21) by the following system

$$\mathcal{N}_m^H \boldsymbol{u}_{p_m}^H + \mathcal{M}_m^H \boldsymbol{u}_{p_m}^H = \mathcal{G}_m^H \tag{3.23}$$

326 with

$$\mathcal{N}_{m}^{H} = \begin{pmatrix} 1 - 2\tau\alpha\partial_{x_{m}x_{m}}^{H} & 0\\ 0 & 1 - 2\tau\alpha\partial_{x_{m}x_{m}}^{H} \end{pmatrix}, \ \mathcal{M}_{m}^{H} = \begin{pmatrix} -1 & 0\\ 0 & -1 \end{pmatrix}, \ \mathcal{G}_{m}^{H} = \begin{pmatrix} \tau g_{1}^{H} - \tau F_{1}^{H}(\boldsymbol{u})\\ \tau g_{2}^{T} - \tau F_{2}^{H}(\boldsymbol{u}) \end{pmatrix}$$
(3.24)

where g_m^H are the discrete RHS coming from the NMG method and $\Gamma^H(\boldsymbol{u})$ is the discrete force terms given in (2.9). The x_1, x_2 directions of the discrete Laplace optimizer can be represented by $\partial_{x_m x_m}^H = \mathcal{L}_{m+}^H + \mathcal{L}_{m0}^H + \mathcal{L}_{m-}^H$, where $\mathcal{L}_{m+}^H, \mathcal{L}_{m0}^H, \mathcal{L}_{m-}^H$ define the folloting step cills

$$\mathcal{L}_{1+}^{H} = \frac{1}{H^{2}} \begin{pmatrix} 0 & 0 & 0 \\ 1 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}, \ \mathcal{L}_{10}^{H} = \frac{1}{H^{2}} \begin{pmatrix} 0 & 0 & 0 \\ 0 & -2 & 0 \\ 0 & 0 & 0_{f} \end{pmatrix}, \ \mathcal{L}_{-}^{H} = \frac{1}{H^{2}} \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & 1 \\ 0 & 0 & 0 \end{pmatrix}$$
$$\mathcal{L}_{2+}^{H} = \frac{1}{H^{2}} \begin{pmatrix} 0 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 1 & 0 \end{pmatrix}, \ \mathcal{L}_{20}^{H} = \frac{1}{H^{2}} \begin{pmatrix} 0 & 0 & 0 \\ 0 & -2 & 0 \\ 0 & 0 & 0 \end{pmatrix}, \ \mathcal{L}_{2-}^{H} = \frac{1}{H^{2}} \begin{pmatrix} 0 & 1 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{pmatrix}$$
(3.25)

then we can write (3.23) in the following way

$$\mathcal{N}_{m+}^{H}\boldsymbol{u}_{p_{m}\,new}^{H} + \mathcal{N}_{m0}^{H}\boldsymbol{u}_{p_{m}\,ne.}^{H} + \mathcal{N}_{m-}^{H}\boldsymbol{u}_{p_{m}\,old}^{H} + \mathcal{M}_{m}^{H}\boldsymbol{u}_{p_{m}\,old}^{H} = \mathcal{G}_{m}^{H}$$
(3.26)

where $u_{p_m new}^H$, $u_{p_m old}^H$ denote the current and previous approximations of $u_{p_m}^H$ in the x_m directions respectively, and

$$\mathcal{N}_{m+}^{H} = \begin{pmatrix} -2\tau\alpha\mathscr{L}_{m+}^{H} & 0\\ 0 & -2\tau\alpha\mathscr{L}_{n+}^{H} \end{pmatrix}, \ \mathcal{N}_{m0}^{H} = \begin{pmatrix} 1 - 2\tau\alpha\mathscr{L}_{m0}^{H} & 0\\ 0 & 1 - 2\tau\alpha\mathscr{L}_{m0}^{H} \end{pmatrix}$$
$$\mathcal{N}_{m-}^{H} = \begin{pmatrix} -2\tau\alpha\mathscr{L}_{n-} & 0\\ 0 & -2\tau\alpha\mathscr{L}_{m-}^{H} \end{pmatrix}, \ \mathcal{M}_{m}^{H} = \begin{pmatrix} -1 & 0\\ 0 & -1 \end{pmatrix}$$
(3.27)

for m = 1, 2. Using a sir ilar process to that shown in §3.1, for computing the smoothing rate, we stimate the convergence rate rom the following

$$\mathcal{P}_{\max} = \max_{\text{loc}} \mathcal{P}_{A \, loc}, \qquad \mathcal{P}_{A \, loc} = \frac{1}{2} \left(\mathcal{P}_{A_1 \, loc} + \mathcal{P}_{A_2 \, loc} \right), \\ \mathcal{P}_{A_m \, loc} \equiv \mathcal{P}_{A_m \, loc}(\boldsymbol{\theta}) = \sup \left\{ \rho \left(\hat{\boldsymbol{S}}_m^H(\boldsymbol{\theta}) \right) \big| \boldsymbol{\theta} \in \boldsymbol{\Theta} \right\}$$
(3.28)

where $\rho(\cdot)$ aga: α enotes the spectral radius, and $\hat{S}_m^h(\theta)$ denote the amplification matrices given by

$$\hat{\boldsymbol{S}}_{m}^{H}(\boldsymbol{\theta}) = -\left[\hat{\mathcal{N}}_{m+}^{H}(\boldsymbol{\theta}) + \hat{\mathcal{N}}_{m0}^{H}(\boldsymbol{\theta})\right]^{-1} \left[\hat{\mathcal{N}}_{m-}^{H}(\boldsymbol{\theta}) + \hat{\mathcal{M}}_{m}^{H}(\boldsymbol{\theta})\right]$$
(3.29)

336 and where

$$\mathcal{N}_{m+}^{H}(\boldsymbol{\theta}) = \begin{pmatrix} -\frac{2\tau\alpha}{H^{2}}e^{-i\omega_{m}} & 0\\ 0 & -\frac{2\tau\alpha}{H^{2}}e^{-i\omega_{m}} \end{pmatrix}, \\ \hat{\mathcal{N}}_{m0}^{H}(\boldsymbol{\theta}) = \begin{pmatrix} 1 + \frac{4\tau\alpha}{H^{2}} & 0\\ 0 & 1 + \frac{4\tau\alpha}{H^{2}} \end{pmatrix}$$
$$\hat{\mathcal{N}}_{m-}^{H}(\boldsymbol{\theta}) = \begin{pmatrix} -\frac{2\tau\alpha}{H^{2}}e^{i\omega_{m}} & 0\\ 0 & -\frac{2\tau\alpha}{H^{2}}e^{i\omega_{m}} \end{pmatrix}, \\ \hat{\mathcal{M}}_{m}^{H}(\boldsymbol{\theta}) = \begin{pmatrix} -1 & 0\\ 0 & -1 \end{pmatrix}$$
(3.30)

337 Comparison of convergence rates for two coarsest grid solvers. Once we have an estimate of

the convergence rate \mathcal{P} , we can compute the number of iterations l required to reach a desired tolerance 10^{-k} using the following

$$l = -\frac{k\ln(10)}{\ln(\mathcal{P})} \tag{3.31}$$

			AOS Solver	Direct	Solver (Pointwise)	Direc	To' er (Block)
Grid Size	α	\mathcal{P}_A	Tol $10^{-1}/10^{-2}/10^{-3}$	\mathcal{P}_D	Tol $10^{-1}/10^{-2}/10^{-3}$	\mathcal{P}_B 1	$10^{-1}/10^{-2}/10^{-3}$
	$\frac{1}{10}$	0.99915	2709/5417/8124	0.40511	3/6/8	0.14573	2/3/4
4×4	$\frac{1}{20}$	0.99957	5355/10708/16062	0.51635	4/7/11	0.2 IJU	2/4/6
	$\frac{1}{30}$	0.99971	7940/15879/23817	0.61297	5/10/15	5084 0	3/5/7
	$\frac{1}{10}$	0.99937	3655/7309/10962	0.82924	13/25/37	0. 41	3/6/8
8×8	$\frac{1}{20}$	0.99968	7195/14390/21584	0.90661	24/47/71	0.6300.	5/10/15
	$\frac{1}{30}$	0.99979	10965/21928/32892	0.93578	35/70/105	0.76812	9/18/27
	$\frac{1}{10}$	0.99947	4344/8688/13031	0.97391	88/175/262	0.9965	632/1262/1894
16×16	$\frac{1}{20}$	0.99973	8528/17055/25582	0.98679	174/647/520	1.0000	-
	$\frac{1}{30}$	0.99982	12792/25583/38374	0.99116	$260/519/77^{\circ}$	J0	-

Table 1: Comparison 2 of convergence rates (averaged over 5 FA. $\$ MG ycles) for the Chumchob-Chen AOS solver and our direct solver. For each solver the convergence rate and number of iterations required to reach tolerances of 10^{-1} , 10^{-2} , 10^{-3} are shown for multiple α values on various coarsest grid sizes for the lung CT example (Example 2 in Figure 3).

Crid Size			AOS Solver	Direct	So. r (Pointwise)	Dire	ect Solver (Block)
GIIG Size		\mathcal{P}_A	Tol $10^{-1}/10^{-2}/10^{-3}$	\mathcal{P}_D	Tol 10 $^{1}/10^{-2}/10^{-3}$	\mathcal{P}_B	Tol $10^{-1}/10^{-2}/10^{-3}$
	$\frac{1}{10}$	0.99915	2708/5416/8123	0.6547	6/11/17	0.32791	3/5/7
4×4	$\frac{1}{20}$	0.99957	5355/10708/16061	0.79307	10/20/30	0.51094	4/7/11
	$\frac{1}{30}$	0.99971	7940/15879/23817	0.85177	15/29/44	0.62553	5/10/15
	$\frac{1}{10}$	0.99937	3655/7309/10962	0.9. 10	39/77/115	0.70146	7/13/20
8×8	$\frac{1}{20}$	0.99968	7195/14390/21584	0.969.5	74/148/222	0.88868	20/40/59
	$\frac{1}{30}$	0.99979	-10965/21928/32892	0.07894	109/217/325	0.97361	87/173/259
	$\frac{1}{10}$	0.99947	4344/8688/13031	₹ 18925	214/427/640	1.00000	=
16×16	$\frac{1}{20}$	0.99973	8528/17055/25582	0.994、	428/856/1283	1.00000	-
	$\frac{1}{30}$	0.99982	12792/25583/38374	0.99643	644/1288/1932	1.00000	-

Table 2: Comparison 1 of convergence ates (av raged over 5 FAS-NMG cycles) for the Chumchob-Chen AOS solver and our direct solver. For each only r the convergence rates and number of iterations required to reach tolerances of 10^{-1} , 10^{-2} , $1'^{-3}$ re shown for multiple α values on various coarsest grid sizes for the hand example (Example 3 in Figure 3)

From Tables 1 and 2 we see that our Crect solver converges much faster than the Chumchob-Chen AOS solver on several different coarset grid sizes for both Hand and Lung CT examples (Examples 1 and 2 in Figure 3) respectively, especial on the 4×4 and 8×8 grids; this improvement has a significant impact on the number of iterations required to reach a desired tolerance, which in turn will have a noticeable effect on the number of FAS NMG cycles needed to obtain a good registration result as well as the time taken. As is also clear if y i both tables, the rates are too high and both solvers are not effective on the less coarse 16×16 cml, posibly due to limitation of the analysis; we would conclude that the coarsest grid is kept as 8×3 .

Hence the improved IMC method, to be denoted by unconstrained INMG, is taken as Algorithm 1 equipped with the consect grid solver by Algorithm 2 and the predicted number of smoothing steps of $\nu_1, \nu_2 \ge 8$ sinc $\mu_{\text{max}}^8 = 0.74762^8 < 0.1$ is believed to be small enough.

351 4 Non-Olding constraint model

We now present another model to deliver diffeomorphic transforms. Folding in the transformation is a problem which can occur in image registration, unless it is specifically controlled. In real applications the presence of folding would suggest an inaccurate registration result as such transformations are nonphysical. In this section we will first introduce our proposed improved diffusion model, which removes any folding that may occur in the transformation φ , as well as including a NMG scheme (Algorithm 1). Then we will extend this model to increase robust with respect to the weighting parameter α .

358 4.1 Improved diffusion model formulation and optimise-discretise approach

In the work by Burger et al. [11], it was explained that the sign of the determine \dots det $\nabla \varphi$ can indicate the presence of any folding in the transformation $\varphi = x + u$, or more specifically the sign of

$$\det \nabla \varphi = (1 + u_{1_{x_1}}) (1 + u_{2_{x_2}}) - u_{1_{x_2}} u_{2_{x_1}}.$$
(4.1)

If det $\nabla \varphi \leq 0$ then this indicates that folding in the transformation is pre-ent while if det $\nabla \varphi > 0$ then no folding occurs in the transformation. In [11] this information was used to τ d an additional term into the diffusion energy functional (2.2) which penalises this determinant in order to produce diffeomorphic image registrations, thus resulting in the following 2D hyper-elastic energy functional

$$E^{\text{Hyper}}(\boldsymbol{u}) = \frac{1}{2} \int_{\Omega} \left(T_{\boldsymbol{u}} - R \right)^2 + \alpha \sum_{s=1}^2 \left| \nabla u_s \right|^2 + \beta \left(\frac{\left(\det \nabla \cdot - 1 \right)^2}{\operatorname{us}^4 \nabla \varphi} \right)^2 d\Omega, \tag{4.2}$$

where $\alpha \in \mathbb{R}^+$, $0 \le \beta \in \mathbb{R}$ are weighting parameters. Although α may be possible to develop an effective smoother for solving (4.2), which has a strong non-linearity. in this paper however we instead propose an extension to the diffusion model (2.2) as a simplification of α , by r-elastic model (4.2) to control any folding. We propose to introduce a constraint into the diffusion model which ensures a positive value of the determinant (4.1). In other words, we aim to solve the falling minimisation problem

$$\min_{\boldsymbol{u}} E^{\text{Diff}}(\boldsymbol{u}), \quad c \neq \det \forall \boldsymbol{\varphi} > 0$$
(4.3)

or equivalently, using an optimise-discretise approach, ve look to solve the following EL equations

$$-\alpha \Delta u_m + F_m(\boldsymbol{u}) = \boldsymbol{\gamma} \quad \text{i. } \det \nabla \boldsymbol{\varphi} > 0 \tag{4.4}$$

with Neumann boundary conditions $\nabla u_m \cdot \boldsymbol{n} = 1$ and where $F_m(\boldsymbol{u})$ are as in (2.6) for m = 1, 2.

4.2 Estimating the determinant using finite elements

In order for us to be able to impose the constraint in (4.4), we must first obtain an approximation of the determinant at every discrete interprovide of Ω^h , that is we need to compute

$$\boldsymbol{Q} \equiv (Q_{ij}) = (\det {}^{,7} \boldsymbol{\varphi})_{i,j} = (1 + (u_{1_{x_1}})_{i,j}) (1 + (u_{2_{x_2}})_{i,j}) - (u_{1_{x_2}})_{i,j} (u_{2_{x_1}})_{i,j}$$
(4.5)

where $Q \in \mathbb{R}^{(n-2)\times(n-2)}$ is the matrix consisting of determinant values at the discrete interior points 375 (i,j) for $i, j = 2, \ldots, n - \ldots$ I) compute the entry (Q_{ij}) , we need to determine the discrete partial 376 derivatives $(u_{m_{x_1}})_{i,j}$, $(u_{m_{x_2}})_{i,j}$ for m = 1, 2. We do this by splitting our discrete domain Ω^h into a mesh 377 of finite elements consisting of pix ewise linear triangular basis functions as shown in Figure 2(a). In fact 378 for each interior point i, j we need to compute the determinant in each of the four triangles T_1, \ldots, T_4 379 as shown in Figure 2(b). Doi g this gives us a clearer picture of the local geometry surrounding the 380 (i, j) point, thus a' owing us to better detect any mesh folding of the transformation. Once we have 381 determinant value for each of the triangles, we assign the smallest value to be our (Q_{ij}) entry, this 382 in essence considers be orst possible case for each (i, j) allowing us to better detect and correct all 383 potential foldi .g in the transformation. Now for linear triangular basis functions, we can approximate 384 385 $u_m(\boldsymbol{x})$ by the following linear functions

$$L_m(\mathbf{x}) = a_{u_m} + b_{u_m} x_1 + c_{u_m} x_2 \tag{4.6}$$

where a_{u_m} , b_{ι} , $c_{u_m} \in \mathbb{R}$ are coefficients to be determined for m = 1, 2. From (4.6) we see that the partial derivatives $u_{m_{x_1}}$, $u_{m_{x_2}}$ are given by the coefficients b_{u_m} , c_{u_m} respectively. Then looking at the first triangle T_1 , at a general discrete interior point (i, j), we have the following system

$$\underline{\text{Triangle } T_1}: \begin{pmatrix} 1 & x_i & y_j \\ 1 & x_{i+1} & y_j \\ 1 & x_i & y_{j+1} \end{pmatrix} \begin{pmatrix} a_{1\,u_1} \\ b_{1\,u_1} \\ c_{1\,u_1} \end{pmatrix} = \begin{pmatrix} (u_1)_{i,j} \\ (u_1)_{i+1,j} \\ (u_1)_{i,j+1} \end{pmatrix}, \begin{pmatrix} 1 & x_i & y_j \\ 1 & x_{i+1} & y_j \\ 1 & x_i & y_{j+1} \end{pmatrix} \begin{pmatrix} a_{1\,u_2} \\ b_{1\,u_2} \\ c_{1\,u_2} \end{pmatrix} = \begin{pmatrix} (u_2)_{i,j} \\ (u_2)_{i+1,j} \\ (u_2)_{i,j+1} \end{pmatrix};$$

389



Figure 2: Finite element splitting of the discrete domain Ω^h using linear triangle basis functions

we obtain similar systems for each of the remaining triangles T_2 , T_3 and T_4 . Then, to compute the coefficients $a_{l u_m}$, $b_{l u_m}$, $c_{l u_m}$, we solve

$$\boldsymbol{s}_l = \boldsymbol{A}_l^{-1} \boldsymbol{v}_{1l}, \quad \boldsymbol{t}_l = \boldsymbol{A}_l^{-1} \boldsymbol{v}_{2l} \tag{4.7}$$

where $s_l = (a_{l\,u_1}, b_{l\,u_1}, c_{l\,u_1})^T$, $t_l = (a_{l\,u_2}, b_{l\,u_2}, c_{l\,u_2})^T$ are the column vectors of coefficients for $(u_1)_{i,j}$, $(u_2)_{i,j}$ respectively, A_l^{-1} are the inverses of the matrices corresponding to the edges of the triangles T_l and $v_{m\,l} = (u_{m\,1}, u_{m\,2}, u_{m\,3})^T$ are the values of u_m as each vertex of the triangles T_l for $l = 1, \ldots, 4, m = 1, 2$. Then, once all elements of Q have been computed, we take the minimum value of the matrix Q to be used to see if the constraint has been satismed. This method can be summarised by Algorithm 3. Once we have a value for Q_{min} , we use Algorithm 4 to impose the constraint and determine whether we accept the updated transformation or not.

In practice, Algorithm 3 can be cor .put .tionally expensive on larger grid sizes owing to the fact that we must solve eight inverse problems at erry \dot{c} .screte interior point in the discrete domain Ω^h , consequently this has a severe impact on the CPU time of the NMG scheme for our constrained model. In Appendix A we demonstrate how Algorithr . 3 c. γ be optimised to significantly decrease CPU cost for each iteration of the determinant computation. The method outlined in Algorithm 8 is how we actually compute the determinant in practice, ar 1 the results shown in §5.2 are also obtained using this algorithm.

4.5 4.3 Numerical solution and NMG algorithm for a constrained diffusion model

Based on our NMC framework unconstrained INMG, we will solve our constrained diffusion model by NMG. Adding a constraint, the same pointwise smoother as the one shown in §2.2 and the same coarsest grid solver c_{thc} one described in §3.2 are used. Then our proposed NMG algorithm is shown in Algorithm ℓ , which we denote constrained INMG.

410 4.4 Ar dentive α constrained diffusion model

While our contrained INMG does ensure that the deformations obtained are non-folding, in cases where folding is severe the deformation field u can be penalised so heavily that the deformed template image T_u may have moved very little when compared with the original template image T. To overcome this problem we propose an extension to our constrained INMG model, whereby we re-initialise the NMG method using a larger value of α if the constraint has not been satisfied within a small number of iterations. To construct this adaptive α scheme, we modify the determinant check shown in Algorithm

4 as seen in Algorithm 5. From Algorithm 5 we see that if we reach the iteration limit LMAX, we 417 exit out of the FAS-NMG algorithm and this is when we re-initialise the NMG with a larger weighting 418 parameter α . This process can be summarised by Algorithm 7, and where the algorithm AdaptFASNMG 419 is the same as Algorithm 6 except now Algorithm 5 is used to check the constraint in tead of Algorithm 420 4. Another advantage of the adaptive α scheme shown in Algorithm 7 is its robuliness to the choice 421 of parameter α . Even if the initial α is set too small such that severe folding would n "mally occur, because we keep re-initialising the problem with new values of α , we automatically in d a pseudo-optimal 423 α value where folding is avoided. This will be shown in the next section. Using the pointwise smoother 424 from §2.2, and the coarsest grid solver from §3.2 along with Algorithm 7, then we denote our adaptive 425 α model by **adaptive INMG**. 426

Algorithm 3 $Q_{min} \leftarrow ComputeQ(u^h, n, h)$

1: for i = 2, ..., n - 1 do for j = 2, ..., n - 1 do 2: 3: for l = 1, ..., 4 do Compute the vectors s_l , t_l using (4.7) Compute determinant for triangle $T_l \to \tilde{Q}_l = (1 + b_{l u_1})(1 + c_{l u_2}) \cdot \dots \cdot u_1 b_{l u_2}$ 4:end for Assign minimum \tilde{Q} to be entry $(Q_{ij}) \to (Q_{ij}) = \min \{\tilde{Q}_1, \ldots, Q_4\}$ 5:end for 6: end for Take minimum entry in Q to be minimum determinant v. " $\to Q_{min} = \min \{Q\}$

Algorithm 4 $u_h^{(k+1)} \leftarrow ConstrainU(u_h^{(k)}, h, \lambda, LMAX)$

1: for $l = 1, \cdots, LMAX$ do

2: Compute minimum value of determinant Q_m usin, Algorithm 3

 $\begin{array}{l} \text{if } Q_{min} > 0 \ and \ l \leq LMAX \ \text{then} \\ \text{Accept update } \boldsymbol{u}_h^{(k+1)} = \boldsymbol{u}_h^{(k)} \end{array}$ 3:

else if $Q_{min} \leq 0$ and l < LMAX then Reject update and set $\boldsymbol{u}_h^{(k)} = \lambda \boldsymbol{u}_h^{(k)}, \lambda \in (0, 1)$ else if $Q_{min} \leq 0$ and l = LMAX then 4:

5:

- $\operatorname{Error} \to \operatorname{Constraint}$ failed
- 6: end if

```
7: end for
```

Algorithm 5 $\left[u_{h}^{(k+1)}, c, done \right] \leftarrow AdaptiveU(u_{h}^{(k)}, h, \lambda, LMAX)$

1: Save current 'good' approximation $\rightarrow \hat{u}_h^{(k)} = u_h^{(k)}, c = 0$

2: for $l = 1, \dots, LMAX$ do

Compute minimum v ine i determinant Q_{min} using Algorithm 3 3:

4:

if $Q_{min} > 0$ and $l \leq LM$. 'X then Accept update $\boldsymbol{v}_{h}^{(c+1)} = \boldsymbol{u}_{h}^{(c)}, \, \hat{\boldsymbol{u}}_{h}^{(k)} = \boldsymbol{u}_{h}^{(k)}, \, c = c+1, \, done_alpha = 1, \, break$ else if $Q_{min} \leq 0$ and $l \leq MAX$ then Reject update and so $\boldsymbol{v}_{h}^{(k)} = \lambda \boldsymbol{u}_{h}^{(k)}, \, \lambda \in (0, 1), \, c = c+1$ else if $Q_{min} \geq 0$ and l = LMAX then 5:

- 6:
- Reset to 'good' appr ximation $\rightarrow c = LMAX$, $\boldsymbol{u}_{h}^{(k+1)} = \hat{\boldsymbol{u}}_{h}^{(k)}$, done_alpha = 0 end if 7:
- 8: end for

Algorithm 6 $u_h^{(k+1)} \leftarrow ConstFASNMG(\mathbb{R}^h, \mathbb{T}^h, n, h, level, u_h^{(k)}, \mathcal{G}^h, \alpha, \nu_1, \nu_2)$ $\bar{\boldsymbol{u}}_{h}^{(k)} \leftarrow Smooth(R^{h}, T^{h}, \boldsymbol{u}_{h}^{(k)}, \mathcal{G}^{h}, \alpha, \nu_{1})$ 1: Pre-smoothing step by performing ν_1 steps (relaxation sweeps) 2: Coarse-grid correction Compute the residual $\boldsymbol{r}_{h}^{(k)} = \mathcal{G}^{h} - \mathcal{N}^{h}(\boldsymbol{u}_{h}^{(k)})$ Restrict residual and smooth approximations $\boldsymbol{r}_{H}^{(k)} = \mathcal{R}_{h}^{H} \boldsymbol{r}_{h}^{(k)}, \ \bar{\boldsymbol{u}}_{H}^{(k)} = \mathcal{R}_{h}^{H} \bar{\boldsymbol{u}}_{h}^{(k)}$ Set $level \rightarrow level - 1$, H = 2h, $nc = \frac{n}{2}$ Form RHS of coarse grid PDEs $\mathcal{G}^{H} = \boldsymbol{r}^{H} + \mathcal{N}^{H}(\bar{\boldsymbol{u}}_{H}^{(k)})$ Solve residual equation on coarse grid to obtain approximations $ilde{m{u}}_H^{(k)}$ 3: if level = 1 then Solve to obtain high accuracy solutions $oldsymbol{u}_{H}^{(k)}$ using a coarsest grid solver Use Algorithm 4 to determine whether update is accepted 4: 5: else level > 1 Repeat the FAS-NMG-CONST procedure recursively to t' \circ ne t level i.e. $\bar{\boldsymbol{u}}_{H}^{(k)} \leftarrow ConstFASNMG(R^{H}, T^{H}, nc, H, level - 1, \tilde{\boldsymbol{u}}_{H}^{(k)}, \mathcal{G}^{H}, \alpha, \nu_{1}, \nu_{2})$ 6: end if Compute the correction $\boldsymbol{e}_{H}^{(k)} = \boldsymbol{u}_{H}^{(k)} - ar{\boldsymbol{u}}_{H}^{(k)}$ Interpolate the correction to next fine grid level $e_h^{(k)} = \mathcal{I}_H^h e_{H_{-}}^{(k)}$ Update current grid level approximations using correction $\hat{u}_{l}^{(\prime)} = \bar{v}^{(k)} + e_{h}^{(k)}$ 7: Post-smoothing step by performing ν_2 steps (relaxation sweeps) $\nu_{a}^{(r+1)} - Smooth(R^h, T^h, \hat{u}_h^{(k)}, \mathcal{G}^h, \alpha, \nu_2)$ Computes $u_h^{(k+1)}$ by performing ν_2 relaxation sweeps of sn. other 8: Use Algorithm 4 to determine whether update is accepted if on fines grid level Ω^h

$\overline{\text{Algorithm 7 } \boldsymbol{u}_h^{(k+1)} \leftarrow Adaptive\alpha\left(R^h, T^h, n, h, \boldsymbol{u}_h^{(k)}, \cdots, \boldsymbol{u}_{max}^{(k)}\right)}$

1: Set done NMG = 0, done alpha = 02: while $done_NMG \neq 1$ do if $i^{\alpha} = i^{\alpha}_{max}$ then 3: LMAX = 100end if 4: while $done_NMG \neq 1$ do 5:Set previous 'good' approximation $\rightarrow \boldsymbol{u}_h^{(\kappa)} = \hat{\boldsymbol{u}}_h^{(\kappa)}$ 6: $\text{Perform FAS-NMG} \rightarrow \left[\boldsymbol{u}_{h}^{(k+1)}, c \right] \leftarrow \text{Adapt FASNMG} \left(R^{h}, T^{h}, n, j, level, \hat{\boldsymbol{u}}_{h}^{(k)}, \mathcal{G}^{h}, \alpha, \nu_{1}, \nu_{2} \right)$ 7: if $c \leq LMAX$ and done_alpha $\neq 1$ th. **n** 8: break 9: end if if NMG convergence criteri sat² fied then 10: $done \ NMG = 1$ end if 11:12:end while if $c \leq LMAX$ and done $\mathfrak{slp}_{l} \neq 1$ then 13: Set $\alpha = 2\alpha$, $i^{\alpha} = i^{\alpha} - 1$, $\boldsymbol{u}_{h}^{(k)} = \hat{\boldsymbol{u}}_{h}^{(k)}$ 14:end if 15:16: end while

Experimenta. v.sults $\mathbf{5}$ 427

Here we will presen and compare the results of four models 428

- M1 he NNG method CCNMG from [16] i.e. Algorithm 1; 429
- M2 t. imr.oved NMG method unconstrained INMG of $\S3.2$; 430
- $-v_{\rm c}$ MG method **constrained INMG** of §4.3 i.e. Algorithm 6; • M3 431
- M4 t. e NMG method adaptive INMG of §4.4 i.e. Algorithm 7. 432

Firstly we will demonstrate how our more accurate analysis of the smoothing rate, along with our 433 new coarsest grid solver, impact the number of NMG cycles required for the method to converge when 434 compared with M1. In addition we will also show how this improved convergence of our NMG method 435 M2 results in a significant decrease in CPU time, as well as an improvement in the accuracy of the 436 registration, when compared with M1. 437

438 Secondly, we will show how our method M3 overcomes the issue of transformation folding while still main-

taining good accuracy and CPU times compared with our unconstrained model M2 and the ChumchobChen model M1.

Thirdly we will show how our method **M4** not only overcomes the problem of mesh feiding while keeping a good level of accuracy and CPU times, but also how it can maintain these good trans. "ms while being robust to parameter choice when compared with the other models.

To gain a quantitative measure of the accuracy of the NMG methods, we se S ructural Similarity (SSIM) [39] as well as the relative error given by $\operatorname{Err} = \frac{\|T_u - R\|_2^2}{\|R\|_2^2}$. Moreover, in or ler to highlight the convergence problem of the **M1**, and for fairness, we will consider a method to nove converged only if any of the following stopping criteria has been satisfied:

• The average relative residual of the EL equations reaches a tolerance $\epsilon = 10^{-2}$

- The maximum relative residual of the EL equations reaches a plerance of $\varepsilon_2 = 10^{-2}$
- The number of NMG cycles reaches the maximum number of ε_3 25

451 We shall take 3 pairs of test images (shown in Fig.3) to experiment and compare registrations:

- **Example 1** a pair of CT images from Fig.3(a, d),
- **Example 2** a pair of CT images from Fig. 3(b, e),
- **Example 3** a second pair of Hand images from $F^* 3(c, f)$.

Moreover, in Tables 5-6 we indicate whether a test has bee. 'successful' (results highlighted in green) or whether it has 'failed' (results highlighted in red). We say that ε est has 'failed' if the maximum number of NMG cycles ε_3 has been reached, or if there is folding in the result (i.e. $Q_{min} < 0$). Additionally bold

 $_{\tt 458}$ $\,$ values indicate the results which give the best SSIM and relative error values for each test.

459 5.1 Comparative results of models $\sqrt{1}$ and M2

Here we will demonstrate the improvement on the non M2 over M1. As mentioned in §3, our improvement is to overcome the convergence problem that the present in the former method.

Test on Example 1. From Figures 8 and 2 we see that our M2 produces visually similar deformed 462 template images $T_{\boldsymbol{u}}$ and final error images $|T_{\boldsymbol{u}} - R|$ when compared with those obtained from M1. The 463 first two columns of Table 5 show several \cdot st results of varying resolutions and parameters α . There, 464 abbreviations 'SSIM', 'Err', 'NMG', 'C^TU' represent the final structural similarity, final relative error, 465 number of multigrid cycles perforn. ⁴ and CPU time respectively. When we look at the table we see 466 that our M2 requires consistent'y fewer `MG cycles to produce these accurate results. In fact, the M1 467 method almost always fails to co. erge within the allowed number ε_3 of NMG cycles to the required 468 tolerances. This confirms our statements earlier on the convergence problem of M1. Moreover, this also 469 leads to a drastic improvem int i CPU time, especially in the 512^2 and 1024^2 cases where the **M1** model 470 requires a much larger number of NMG cycles. 471

Test on Example 2. Alt' ough visual differences between the models are small in Figures 6 and 7, in Table 4, we see that \mathbf{M}' , is ' etter than $\mathbf{M1}$ (in all indicators: SSIM/Err/NMG cycles/CPU) for the first α value, but for the c b r two cases of α both models failed to give diffeomorphic maps due to det $\nabla \varphi < 0$.

Test on Example For the second lung CT example visual differences between the models are small in Figures 4 at a 5. We can see that, from Table 3, M2 is successful for all cases of α but M1 failed in several cases. On convergence alone, M1 is not as fast as M2 because it takes many NMG cycles.

We remark that, in the M1 method tested above, we have used the original CCNMG AOS solver on 479 the coarses graduate the (new) updated smoothing rates to predict the number of smoothing steps 480 required on the grids; that is to say, the NMG cycles displayed are better than the original work. To 481 illustrate the in portance of our re-analysis in LFA, we will give a brief comparison using the old and 482 new smoothing rates for a specific test. Considering **Example 1** from Figure 8 of size 128^2 with $\alpha = \frac{1}{10}$, 483 we obtained SSIM/Err(%)/NMG/CPU(s) values of 0.774/1.48/21/1.169 using the M1 method with 484 smoother steps based upon the rate $\mu = 0.5$. However if we perform the same test with smoother steps 485 based upon our re-calculated rate $\mu = 0.74762$, we obtain values of 0.775/1.46/10/0.959. Clearly there 486

is a vast improvement (reduction) in the number of NMG cycles required with small improvements in
the other three values and the overall improvement of M2 over M1 is also due to the new coarsest grid
solver.

⁴⁹⁰ 5.2 Comparative results of models M2 and M3

In §4 we introduced our constrained version M3 in order to prevent any folding from occurring in the transformation. This was achieved by ensuring det $\nabla \varphi > 0$ for every discrete interproduct or point in Ω^h . Here we will present results comparing M2 and M3 to show how this constraint dochindeed prevent folding while still maintaining good accuracy and CPU time using the same three manples from §5.1. The abbreviation Q_{min} represents the minimum determinant value det $\nabla \varphi$. There small 'Err' means a small fitting error while $Q_{min} > 0$ implies a correct registration transformation.

Test on Example 1. From columns 2 and 3 of Table 5 we see that our N. 2 always produces positive Q_{min} values; as a result we obtain the exact same results with our M. method with very small increases in CPU times owing to the constraint checking. This also transitiones to regures 8 and 9 where we see that all images look very similar visually.

Test on Example 2. From Table 4 we see that M3 has over $met^{1/2}$ mesh folding problems of M2 by positive Q_{min} values in all cases. In achieving this convergent non-folding result, the number of NMG cycles taken by M3 is more than M2. Although the CPU of pes in these cases also increase noticeably, we do however still see a reduction and consistency in the number of NMG cycles when compared with the M1 method. The CPU time increase could be reduced by a more computationally efficient implementation of our smoother code to penalise the transformation only in regions where folding is present.

Test on Example 3. Here we see the exact same protern as in Example 1 since our M3 produces positive determinant values in all cases and iden. The ults to M2 with small increases in CPU times as shown in Table 3, with improvements in all cate on roover the M1 method especially in convergence and CPU times.

512 5.3 Comparative results of rous 1s M3 and M4

Additionally in §4 we introduce and (xten. on .o our M3 model to be robust to parameter choice while maintaining a non-folding transformatio. Here we will consider a case where severe folding would occur and our M3 model, while producing a non-folding deformation, performs poorly in terms of registration accuracy whereas our M4 mode' also a, its folding while producing good registration accuracy.

From Table 6 we see that alt_lough ve obtain very good accuracy from our M2 model, we also have 517 severe folding in the transform tions in all tests as indicated by the negative Q_{min} values. Looking at 518 the results for our M3 model we see that the folding problem has been overcome and all Q_{min} values are 519 now positive, however we also ee that we have lost the accuracy of the result with regard to error when 520 compared with the M2 res lts, especially on the 127^2 and 256^2 images. Our M4 model on the other 521 hand no only produce no -fol ing results like with our M3 model, but also maintains a similar level 522 of accuracy when compared \mathbf{v} th the results from our M2 model. In addition we also see that our M4 523 model achieves this with only a slight increase in CPU time when compared with those from the M2 524 model, and is over wice as fast as our M3 model. From Figures 10 and 11 we see that visually there is 525 a noticeable difference, 'ween the deformed template from our M3 model compared with those from 526 our M2 and Γ_{14} models, especially in the error images. 527

528 5.4 Te. v. o. MG efficiency and parameter robustness

NMG efficiency. In this work, we are concerned with transforms' quality and fast solution by a NMG. For the latter, we expect the optimal efficiency of $O(N \log N)$ complexity in achieving a fixed accuracy (with $N = n^2$ for $n \times n$ images). Let t_n denote the CPU times required by registering two $n \times n$ images. Then for an optimal NMG, we expect the CPU increase to be of ratio $t_n/t_{n/2} = Cn^2 \log n^2/(C(n/2)^2 \log(n/2)^2) = 4 + 4 \log 4/\log(n/2)^2 \approx 4.5$. In Table 7, we show test results of all four NMG methods for varying resolutions, where in M1 we use the original analysis of [16]

to set the number of smoothing steps. Clearly M2, M3 and M4 exhibit nearly optimal complexity but M1 shows irregular patterns, which justify our re-analysis for Algorithm 1.

Finally to give an indication of the convergence history of M1 and M2, we plot in Fig.12 the residuals for more NMG cycles. Evidently M2 has faster convergence plot than M1.

Parameter robustness. In the diffusion model, the weighting parameter α indicates n, \forall strongly we wish to enforce smoothness on the deformation from the regularisation term. Specifically, a larger value 540 of α will impose a strong penalisation on non-smooth deformations leading to no folding, however this 541 also leads to a less accurate registration with regards to error. On the other ha.¹ a smaller value of 542 α will lead to a more accurate registration in terms of error, but will also in ease the likelihood of 543 folding occurring. Moreover, selecting a 'good' value for α can be very time onsuming as in general a 544 pre-multigrid routine is usually required to find this 'best' α (for exam 'e t' e cooling process in [16]), 545 which can noticeably increase the computational work and CPU time. For this reason, having a model 546 which is robust to the choice of weighting parameter is very useful as 'ne nee⁴ for finding the 'best' value for α is less important. Here we will compare how the value of α inpacts the relative error (denoted 548 'Err') and minimum determinant value (denoted ' Q_{min} ') for models 1. '2 and M4. From Figure 13(a) 549 we see that as α gets smaller the error also decreases, however 'ooki' out Figure 13(b) we see that the 550 value of Q_{min} is also decreasing to a point where it is always gative as highlighted by the dotted 551 line. This suggests that our model M2 has a limit where it n. intains physically accurate non-folding 552 deformations, and once past this point folding always occurs. Look ng at Figure 14(a) we see that our 553 M4 model follows a similar pattern with regard to a decrealing er or as α decreases like with our M2 model, however from Figure 14(b) we see that our M4 mod ¹ always maintains the physical integrity of 555 the deformation with $Q_{min} > 0$ for all tested values of α From his we can conclude that our adaptive α 556 model M4 is very robust to the initial value of α , even to small values, while maintaining a consistently 557 good registration accuracy in terms of error. 558



Figure 3: Three Pairs of Test Images.

559 6 Conclusions

In this paper w have first presented an improved NMG method, with regard to convergence and accuracy, over that proposed by Chumchob-Chen through a more detailed and accurate analysis of the multigrid method, as well as a different coarsest grid solver. Secondly we proposed an extension to our NMG method with the aim of producing non-folding transformations, which was achieved by imposing an additional constraint into our improved NMG method. Next we extended our **constrained INMG** to be more robust to parameter choice while keeping non-folding deformations and good registration accuracy. We then used three examples to demonstrate the improvement in accuracy and NMG cycles required for convergence over the Chumchob-Chen NMG, as well as how our constrained INMG and adaptive INMG overcame folding by ensuring det $\nabla \varphi > 0$.

Image Cine m2		M1	M2	M3
Image Size n	α	$SSIM/Err$ (%)/NMG/CPU (s)/ Q_{min}	SSIM/Err (%)/NMG/CPU (s)/ Q_{min}	SSIM/Err \wedge)/1, "G/CPU (s)/ Q_{min}
128 ²		0.930/0.54/2/0.391/0.797	0.943/0.41 /1/0.333/0.819	0.943/0.41 /1/0. 39/0.819
256^{2}	1	0.943/0.45/5/1.512/0.715	0.951/0.42 /2/1.927/0.803	0.^c JI / 42 /2/2.051/0.803
512^{2}	5	0.959/0.44/13/22.387/0.854	0.964/0.43/2/9.426/0.801	6 964// 43 /2/9.721/0.801
1024^{2}		0.972/0.44/25/196.585/0.872	0.975/0.43 /3/66.178/0.822	u. 75 / .43 /3/69.500/0.822
128 ²		0.931/0.52/1/0.316/0.612	0.945/0.39 /1/0.425/0.694	0.94 5, 39 /1/0.437/0.694
256^{2}	1	0.945/0.43/25/6.887/0.464	0.953/0.40/1/1.090/0.660	υ. 53/0.4 J/1/1.164/0.660
512^{2}	10	0.961/0.43/10/17.204/0.734	0.965/0.41 /1/5.057/0.668	0.96 , 41 /1/5.250/0.668
1024^{2}		0.974/0.43/23/180.785/0.745	0.976/0.42 /1/22.972/0.685	*~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
128^{2}		0.937/0.45/25/1.919/0.619	0.947/0.38/3/0.976/0.559	0.94, /0.38/3/1.010/0.559
256^{2}	1	0.948/0.40/25/6.820/0.230	0.954/0.39/1/1.080/0.511	0.954/0.39/1/1.146/0.511
512^{2}	15	0.962/0.41/12/20.657/0.631	0.966/0.40/1/4.886/0.526	966/0.40 /1/5.150/0.526
1024^{2}		0.975/0.41/18/141.395/0.644	0.977/0.40 /1/24.642/0.55	0.977/0.40 /1/25.546/0.554

Table 3: Example 2 – Registration comparison of 3 methods on multiple image sizes for different α values, with an initial relative error of 0.60% and initial SSIM values of 0.933, 0.942, 0.957, 0.972 for the 128^2 , 256^2 , 512^2 , 1024^2 images respectively.

, , ,		0 1		
Imaga Siza n2	0	M1	M2	M3
mage size n		$SSIM/Err$ (%)/NMG/CPU (s)/ Q_{min}	$SSIM/Err (\%)/NMG/CP (^)/Q_{min}$	$SSIM/Err (\%)/NMG/CPU (s)/Q_{min}$
128 ²		0.750/1.28/4/0.530/0.642	0.764/1.17/2/0.586/0.60	0.764/1.17/2/0.603/0.664
256^{2}	1	0.752/1.35/11/3.102/0.640	0.786/1.14/o, 926/0.6	0.786/1.14 /3/3.015/0.645
512^{2}	5	0.806/1.32/25/42.794/0.618	0.832/1.1 /4/18.5 10.83	0.832/1.18/4/19.188/0.683
1024^{2}		0.860/1.34/25/199.920/0.640	0.883/1.20/4/ 853/0.701	0.883/1.20/4/94.397/0.701
128^{2}		0.766/1.04/3/0.456/0.406	0.783/0.05/0/0.070	0.783/0.95 /2/0.715/0.070
256^{2}	1	0.768/1.11/7/2.038/0.344	0.803/0. ¹ /3/2.879/ − 0.028	0.800/0.95/6/6.251/0.027
512^{2}	10	0.819/1.07/20/34.047/0.280	0.847/0.95 / '14.244/0.091	0.847/0.95 /3/14.784/0.091
1024^{2}		0.873/1.06/25/195.431/0.271	0. 06 /4/68.196/0.145	0.893/0.96 /4/71.186/0.145
128^{2}		0.774/0.89/3/0.488/0.080	0.79 $\sqrt{0.8}$ $\sqrt{3}/\sqrt{920} - 0.687$	0.757/1.18/8/3.424/0.015
256^{2}	1	0.802/0.77/6/1.786/-0.165	0.811 $6/2/1.952/-0.862$	0.772/1.23/5/8.047/0.024
512^{2}	15	0.826/0.91/15/25.598/ - 0.122	54/0.1/3/13.750/-0.680	0.827/1.18/6/40.789/0.012
1024^2		0.880/0.89/25/195.370/-0.156	0. `95 , ` 80 /3/69.076/ − 0.584	0.881/1.16/6/182.460/0.011

Table 4: Example 2 – Registration comparis γ or C methods on multiple image sizes for different α values, with an initial relative error of 1.99% and minimal SSIM values of 0.667, 0.704, 0.769, 0.838 for the 128², 256², 512², 1024² images respective? Clearly although **M2** does converge quickly, the **M3** offers both speed and correct transforms.

Imaga Siza n ²		M1	M2	M3
Image Size n		$SSIM/Err (\%)/NMG/CPU (s)/Q_{min}$	$SIM/Err (\%)/NMG/CPU (s)/Q_{min}$	$SSIM/Err$ (%)/NMG/CPU (s)/ Q_{min}
128 ²		$0.742/2.42/16/1.4\epsilon$ /0.6 ϵ	0.717/3.30/2/0.633/0.554	0.717/3.30/2/0.644/0.554
256^{2}	1	0.743/2.61 /25/7. 7/0. 1	0.725/3.24/2/1.959/0.517	0.725/3.24/2/2.093/0.517
512^{2}	5	$0.748/3.68/25/4^{\circ}542/717$	0.750/3.24 /2/9.397/0.498	0.750/3.24 /2/9.691/0.498
1024^2		0.747/6.85/25/1 $5.731/0.6$.	0.784/3.24 /2/45.445/0.486	0.784/3.24 /2/47.728/0.486
128 ²		0.775/1.46 /1/ 059/0.600	0.758/1.89/3/0.868/0.420	0.758/1.89/3/0.892/0.420
256^{2}	1	0.776/1.46, 25/6.78., 10 639	0.760/1.87/2/1.984/0.376	0.760/1.87/2/2.118/0.376
512^{2}	10	0.778/2.02 /05/42.149/0.o02	0.778/1.86/2/9.350/0.348	0.778/1.86/2/9.706/0.348
1024^2		0.780/3.6'/25/) $5.403/0.532$	0.807/1.87/2/45.620/0.332	0.807/1.87/2/48.026/0.332
128 ²		0.790 .13 /8 0.814/0.563	0.783/1.33/3/0.891/0.324	0.783/1.33/3/0.922/0.324
256^{2}	1	0.791/1.1 2/5.992/0.561	0.781/1.31/3/2.907/0.266	0.781/1.31/3/3.086/0.266
512^2	15	0.78 / 1.40 / 25 / 225 / 0.539	0.794/1.31/3/13.786/0.246	0.794/1.31/3/14.526/0.246
1024^{2}		$0.7 \ \mathfrak{z}/2.3 \ /25/194.026/0.390$	0.819 / 1.31 /3/66.949/0.235	0.819 / 1.31 /3/69.405/0.235

Table 5: Example $3 - \text{Re}_{c}$'s' ation comparison of 3 methods on multiple image sizes for different α values, with an initial relative error of 13.25% and initial SSIM values of 0.551, 0.587, 0.639, 0.693 for the 128^2 , 256^2 , 512^2 , 1024^2 images respectively.

Image Cine m2		M2	M3	M4
Image Size n	1	$SSIM / Err (\%) / NMG / CPU (s) / Q_{min}$	$SSIM/Err$ (%)/NMG/CPU (s)/ Q_{min}	$SSIM/Err$ (%)/NMG/CPU (s)/ Q_{min}
128 ²		0. $12/0.95/2/0.686/ - 3.078$	0.630/7.56/6/2.676/0.032	0.758/1.91/3/0.711/0.554
256^{2}		0 $16/0.74/2/2.458/-0.463$	0.630/9.59/3/5.076/0.060	0.815/0.82/2/2.178/0.168
512^{2}	40	J.824/0.82/2/9.729/-0.132	0.805/1.10/4/27.558/0.025	0.824/0.74 /2/10.318/0.351
1024^2		0.832/0.78/2/45.762/-0.163	0.812/1.64/4/121.546/0.086	0.842/0.73/2/58.604/0.358

Table 6: Exal ble 3 - Registration comparison of 3 methods on multiple image sizes for a 'bad' choice of α , with an initial relative error of 13.25% and initial SSIM values of 0.551, 0.587, 0.639, 0.693 for the 128^2 , 256^2 , 512^2 , 1024^2 images respectively.



(a) **M1** $T_u \alpha = \frac{1}{10} Err = 0.43\%$

(b) **M2** $T_{\boldsymbol{u}}$ $\alpha = \frac{1}{10}$ Err = 0.41%



Figure 4: Example 1 – Registration of 3(a) R and 3(d) T of size 512 > 512 or 5 methods. Image (a) shows the deformed template image T_u obtained using the **M1**, while image (b) shows the deformed template image T_u for our **M2** and image (c) shows the deformed template image T_u for our **M3** for the parameter value $\alpha = \frac{1}{10}$.



Figure 5: Example 1 – Difference images corresponding to registrations of Fig.4. Image (a) shows the initial error between T and R, while images $(\underline{f}, \underline{f}, \underline{f})$ show the final errors between T_u and R for **M1**, our **M2** and our **M3** respectively.



Figure 6: Example 2 – Reg. ^{*r} tion of 3(b) R and 3(e) T of size 512×512 by 3 methods. Image (a) shows the deformed temp ate in. ge T_u obtained using the **M1**, while image (b) shows the deformed template image T_u for our 1 **12** and mage (c) shows the deformed template image T_u for our constrained NMG for the parameter value $a = \frac{1}{10}$.



Figure 7: Example 2 – Difference images corresponding to registrations of Fig. 6. Image (a) shows the initial error between T and R, while images (b), (c), (d) show the final ϵ rors between T_u and R for the **M1**, our **M2** and our **M3** respectively.



Figure 8: Example 3 – Registration of $3(c) R \approx 3(f) T$ of size 512×512 by 3 methods. Image (a) shows the deformed template image T_u obtained using the **M1**, while image (b) shows the deformed template image T_u for our **M2** and image (c) using the **M1**, while image T_u for our **M3** for the parameter value $\alpha = \frac{1}{10}$.



(a) |T - R|

(b) M1 $|T_{\boldsymbol{u}}-R|~\alpha=\frac{1}{10}$

(c) M2 $|T_{\boldsymbol{u}}-R|\;\alpha=\frac{1}{10}$

(d) **M3** $|T_u - R| \alpha = \frac{1}{10}$

Figure 9: Example 3 - \mathcal{D} ; erer ce images corresponding to registrations of Fig.8. Image (a) shows the initial error between \mathcal{T} and \mathcal{P} while images (b), (c), (d) show the final errors between T_u and R for the **M1**, our **M2** and ur **M5** respectively.



(a) **M2** $T_{\boldsymbol{u}} \alpha = \frac{1}{40} Err = 0.82\%$

(b) **M3** $T_{\boldsymbol{u}} \alpha = \frac{1}{40} Err = 1.10\%$

 $(- M4 T_u - \frac{1}{40} Err = 0.74\%$

Figure 10: Example 3 – Registration of 3(c) R and 3(f) T of size $512 \div 512$ by 5 methods. Image (a) shows the deformed template image T_u obtained using the **M2**, while image (b) shows the deformed template image T_u for our **M3** and image (c) shows the deformed template image T_u for our **M4** for the 'bad' parameter value $\alpha = \frac{1}{40}$.



Figure 11: Example 3 – Difference images corresponding to registrations of Fig.10. Image (a) shows the initial error between T and R, while images (a, b, d) show the final errors between T_u and R for our **M2**, **M3** and **M4** respectively.



Figure 12: Co. parison of the number of NMG cycles required for the maximum relative residual to reach a tolerance of 10^{-10} between our **M2** method and the **M1** method





(a) Plot of relative error vs parameter α of model ${\bf M2}$ for Example 1

(b) Plot f mir.... 'n determinant value vs parameter α of model \mathbf{M}_{\star} '. Exa, ple 1





(a) Plot of relative error vs parameter α $\$ f model M4 for Example 1

(b) Plot of minimum determinant value vs parameter α of model **M4** for Example 1

Figure 14: Test of 1. ustness of model M4 to the choice of parameter α (50 values).

Imaga Siza m2			M	1	M2	2	M3	1	M4	L D
Image Size n-	nage E. Aple o		CPU(s)	Ratio	CPU(s)	Ratio	CPU(s)	Ratio	CPU(s)	Ratio
128^2			0.316	-	0.425	-	0.437	-	0.452	-
256^{2}			6.887	21.794	1.090	2.565	1.164	2.666	1.304	2.885
512^{2}	le 1 (CT)	$\frac{1}{10}$	17.204	2.498	5.057	4.639	5.250	4.510	6.202	4.756
102		10	180.785	10.508	22.972	4.543	24.182	4.606	29.072	4.688
12 2			0.456	-	0.636	-	0.715	-	0.831	-
25t `			2.038	4.469	2.879	4.527	6.251	8.743	3.874	4.662
512^{2}	لاً ي Axample 2 (CT)	$\frac{1}{10}$	34.047	16.706	14.244	4.948	14.784	2.365	18.768	4.845
	[10	195.431	5.740	68.196	4.788	71.186	4.815	87.203	4.646
128^2			0.959	-	0.868	-	0.892	-	0.845	-
2.1			6.787	7.077	1.984	2.286	2.118	2.374	2.582	3.059
512	Example 3 (Hand)	$\frac{1}{10}$	42.149	6.210	9.350	4.713	9.706	4.089	12.340	4.779
1024^{2}		10	195.403	4.636	45.620	4.879	48.026	4.948	58.466	4.738

Table 7: Test on optimal complexity in CPU time ratio for 4 NMG methods. The optimal ratio is 4 for an O(N) method (with $N = n^2$). Clearly the newer NMGs are better.

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655 A Opth aised version of Algorithm 3

In our constrained NMG, we check to see whether the constraint in (4.4) has been satisfied after the final post-smoothing step and solver step. While checking the constraint after the coarsest solver step is inexpensive computationally owing to the very small grid size, this is not the case when checking after the post-smoothing step. For each interior point Algorithm 3 needs to solve eight inverse problems which, even though we are only using 3×3 matrices, become very expensive on larger grids thus leading to a significant increase in CPU time. We will now look to exploit the structure and commonality between different interior points, of the matrices A_l , to create an optimised version of Algorithm 3. First we will look at the relation of the matrices A_l at the first interior point (2, 2) and a general ; iterior point (i, j). Looking at the matrix A_1 , we see that

$$\underline{\operatorname{At}\ (2,2)}:\ A_1 = \begin{pmatrix} 1 & h & h \\ 1 & 2h & h \\ 1 & h & 2h \end{pmatrix},\ \underline{\operatorname{At}\ (i,j)}:\ \tilde{A}_1 = \begin{pmatrix} 1 & (i-1)h & (\ \ 1)h \\ 1 & ih & (j-1)h \\ 1 & (i-1)h & h \end{pmatrix}$$

since $((x_1)_2, (x_2)_2) = (h, h)$ and $((x_1)_i, (x_2)_j) = ((i-1)h, (j-1)h)$, then \tilde{A}_1 can be written in the following way

$$\tilde{A}_{1} = \begin{pmatrix} 1 & (x_{1})_{2} + (i-1)h & (x_{2})_{2} + (j-1)h \\ 1 & (x_{1})_{3} + (i-1)h & (x_{2})_{2} + (j-1)h \\ 1 & (x_{1})_{2} + (i-1)h & (x_{2})_{3} + (j-1)h \end{pmatrix} = A_{1} + \begin{pmatrix} 1 \\ 1 \end{pmatrix} \begin{pmatrix} 0, (i-1)h, (j-1)h \end{pmatrix} = A_{1} + \begin{pmatrix} 1 \\ 1 \end{pmatrix} \begin{pmatrix} 0, (i-1)h, (j-1)h \end{pmatrix} = A_{1} + \begin{pmatrix} 1 \\ 1 \end{pmatrix} \begin{pmatrix} 0, (i-1)h, (j-1)h \end{pmatrix}$$

with $\boldsymbol{p} = (1,1,1)^T$, $\boldsymbol{q} = (0,(i-1)h,(j-1)h)^T$. The metrices \hat{A}_l for the remaining triangles can be written in similar ways to (A.1), then we have

$$\tilde{A}_l = A_l + \frac{T}{4} \tag{A.2}$$

with p, q as before, and so the inverse $\tilde{A}_l^{-1} = (A_l + z^T)^{-1}$, at a general discrete interior point, can be computed using the Sherman-Morrison formula [2] given by the following theorem

Theorem A.1. (Sherman-Morrison) Suppose $A \in \mathbb{T}^{n \times 1}$ is an invertible matrix, and $p, q \in \mathbb{R}^{n \times 1}$ are column vectors. Then $(A + pq^T)$ is invertible $\iff 1 + q^T A^{-1} p \neq 0$. If $(A + pq^T)$ is invertible, then its inverse is given by

$$\left(A + \gamma \, \boldsymbol{q}^{*}\right)^{-1} = A^{-1} - \frac{A^{-1} \boldsymbol{p} \boldsymbol{q}^{T} A^{-1}}{1 + \boldsymbol{q}^{T} A^{-1} \boldsymbol{p}} \tag{A.3}$$

where pq^T denotes the outer product of the stors p, q.

It can be shown that the multiplication $q^T \mathbf{i}_l^{-1} \mathbf{p} = 0 \forall l = 1, ..., 4$, therefore the invertibility condition from Theorem A.1 holds for ever interval, j for i, j = 2, ..., n-1 and thus the matrices $(A_l + pq^T)^{-1}$ are invertible for each $l = 1, ..., Then we can use Theorem A.1 to rewrite the inverses <math>(A_l + pq^T)^{-1}$ as

$$(A_l + pq^T)^{-1} = A_l^{-1} - \frac{A_l^{-1} pq^T A_l^{-1}}{1 + q^T A_l^{-1} p} .$$
(A.4)

Next we use the fact that $r \in n \in d$ only determine the $b_{l u_m}$, $c_{l u_m}$ coefficients where m = 1, 2, and so our original inverse problem (4.7) reduces to the following scalar equations

$$b_{l u_{1}} = \omega_{u_{1} l}(2) - \mu_{l} \omega_{u_{1} l}(2), \qquad c_{l u_{1}} = \omega_{u_{1} l}(3) - \mu_{l} \omega_{u_{1} l}(3),$$

$$v_{l u_{2}} = \omega_{u_{2} l}(2) - \mu_{l} \omega_{u_{2} l}(2), \qquad c_{l u_{2}} = \omega_{u_{2} l}(3) - \mu_{l} \omega_{u_{2} l}(3), \qquad (A.5)$$

where $\mu_l = \frac{(c_{jl}(2)q_2 + \frac{1}{p_l}(3)q_3)}{1 + (\omega_l^{-1}(2)\sigma_l + \omega_{pl}(3)q_3)}$ and $\omega_{pl}(2)$, $\omega_{pl}(3)$, q_2 , q_3 , $\omega_{u_ml}(2)$, $\omega_{u_ml}(3)$ denote the second and third component $c_{ll} = A_l^{-1} p$, q^T and $\omega_{u_ml} = A_l^{-1} v_{ml}$ respectively.

Therefore the key message is that per checking step across the entire grid only simple matrix-vector products are needed, if we invert matrices A_l^{-1} at the first pixel and then re-use them. Hence our optimised version of Algorithm 3 can be expressed by the following Algorithm 8 $Q_{min} = FEMOpt(u^h, n, h)$ 1: for l = 1, ..., 4 do Compute matrices A_l corresponding to first interior point (2,2)Compute inverse matrices A_l^{-1} Compute second and third components of $A_l^{-1} \mathbf{p} \to \omega_{pl}(2), \, \omega_{pl}(3)$ 2: **end for** 3: for i = 2, ..., n - 1 do for j = 2, ..., n - 1 do 4:Compute second and third components of $q^T \rightarrow q_2 = (i-1)h, q_3 = (j-1)h$ for l = 1, ..., 4 do 5:Compute μ_l Compute second and third components of $\omega_{u_1 l}, \omega_{u_2 l} \rightarrow \omega_{u_1 l}(2), \omega_{u_1 l}(2), \omega_{u_2 l}(3)$ Determine coefficients $b_{l u_1}, c_{l u_1}, b_{l u_2}, c_{l u_2}$ using (A.5) Compute determinant for triangle $T_l \rightarrow \tilde{Q}_l = (1 + b_{l u_1})(1 + c_{l u_2}) - c_{l u_1}$ 6: end for Assign minimum \tilde{Q} to be entry $(Q_{ij}) \to (Q_{ij}) = \min \{\tilde{Q}_1, \dots, \tilde{Q}_4\}$ 7:end for 8: end for Take minimum entry in Q to be minimum determinant value $\rightarrow O$ = min $\{Q\}$

Finally we show in Table 8 how much speed up can be achieved for . simple example. Clearly Algorithm
8 uses up to 30 times less CPU than Algorithm 3.

Image Size n	Unoptimised Time (s)	• timised Time (s)
256^{2}	4.46	0.17
512^{2}	17.87	0.61
1024^{2}	71.53	2.40
2048^{2}	306 23	9.90

Table 8: Table showing the comparison of CPU tim's per iteration between old unoptimised FEM code and new optimised FEM code.