A parallel goal-oriented adaptive finite element method for 2.5-D electromagnetic modelling

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SUMMARY
We present a parallel goal-oriented adaptive finite element method that can be used to rapidly compute highly accurate solutions for 2.5-D controlled-source electromagnetic (CSEM) and 2-D magnetotelluric (MT) modelling problems. We employ unstructured triangular grids that permit efficient discretization of complex modelling domains such as those containing topography, dipping layers and multiple scale structures. Iterative mesh refinement is guided by a goal-oriented error estimator that considers the relative error in the strike aligned fields and their spatial gradients, resulting in a more efficient mesh refinement than possible with a previous approach based on the absolute errors. Reliable error estimation is accomplished by a dual weighted residual method that is carried out via hierarchical basis computations. Our algorithm is parallelized over frequencies, wavenumbers, transmitters and receivers, where adaptive refinement is performed in parallel on subsets of these parameters. Mesh sharing allows an adapted mesh generated for a particular frequency and wavenumber to be shared with nearby frequencies and wavenumbers, thereby efficiently reducing the parallel load of the adaptive refinement calculations. We demonstrate the performance of our algorithm on a large cluster computer through scaling tests for a complex model that includes strong seafloor topography variations and multiple thin stacked hydrocarbon reservoirs. In tests using up to 800 processors and a realistic suite of CSEM data parameters, our algorithm obtained run-times as short as a few seconds to tens of seconds.

Key words: Numerical solutions; Numerical approximations and analysis; Electromagnetic theory; Magnetotelluric; Marine electromagnetics.

1 INTRODUCTION
Numerical techniques employed to model electromagnetic (EM) geophysical problems commonly utilize a domain discretization based on a structured rectangular grid (e.g. Wannamaker et al. 1987; Mackie et al. 1993; Newman & Alumbaugh 1995). Such grids are appealing since they can be crafted without the need of specialized software and permit a convenient approximation of the spatial derivatives required by the traditional finite difference method. However, EM surveys are often acquired over complex terrains with large topographic or bathymetric variations that can impart a strong influence on the measured data (e.g. Wannamaker et al. 1986; Schwalenberg & Edwards 2004; Li & Constable 2007; Constable et al. 2009; Key & Constable 2011), yet including topography in rectangular modelling grids can be challenging. For example, the fine-scale discretization required to handle topography in the central region of a model leads to thin rows and columns of cells that extend laterally and vertically throughout the entire domain, often resulting in highly elongated cells (e.g. Constable et al. 2009, fig. 9). Such high aspect ratio cells can impart a large condition number to the resulting numerical linear system and often result in poor convergence for iterative Krylov solvers. Similar grid design problems arise for the discretization of known subsurface structures of highly variable shape and those that are inclined with respect to the structured grid.

The finely spaced rows and columns of cells required to represent these features will propagate throughout the entire model domain, resulting in a dense grid that may require more memory than is available and longer than desired run-times. Frequently, one must compromise between fine meshing to accurately represent known structures and coarse meshing to allow the problem to fit into memory or to be solvable in a reasonable time-frame. Certainly future improvements in computational power will allow these problems to be overcome by brute-force fine meshing of the entire model domain; however, for present day computers such profligate meshing schemes can only be implemented for relatively limited model domains.

A well-known solution to these discretization difficulties is to instead use an unstructured grid of triangular elements that can readily conform to highly complex model features. Although unstructured grids have long been used in the applied math and engineering communities, they have only recently been applied to EM geophysical...
problems, as exemplified by finite element codes developed to handle the topographic variations encountered in marine EM surveys (Key & Weiss 2006; Li & Key 2007; Franke et al. 2007; Li & Pek 2008). In part, these recent developments have been enabled by the widespread availability of robust computational geometry tools for creating constrained conforming Delaunay triangulations (e.g. Shewchuk 2002b), thereby lowering a significant software hurdle for the EM geophysicist interested in utilizing unstructured grids. In addition to the ease of handling topography, another key benefit of unstructured grids is the ability to efficiently discretize multiple-scale structures. For example, tiny grid elements required for near-surface features do not propagate into deeper regions when meshed with an unstructured grid. This model representation efficiency results in faster run-times for a given model structure, as well as the ability to handle more complex structures for a given memory capacity.

Another concern for model discretization is the desired solution accuracy. The designer of structured grids must consider how fine the grid spacing should be to ensure a sufficiently accurate numerical computation. Typically, rules-of-thumb for grid density and the user’s prior experience are used to design a static grid. However, such methods often offer little confidence that the grid is indeed accurate for a given set of model and data parameters. A solution to these cumbersome problems is to instead utilize a dynamic approach to grid generation, in which solution accuracy is enforced through the use of automated adaptive refinement methods. Adaptive methods seek to increase the solution accuracy by iteratively refining the grid, where each iteration consists of selecting a subset of elements for refinement based on an estimate of their contribution to the solution error, and then refining the grid by creating new smaller elements in these regions. Such methods offer great potential since they can be used to automatically compute an efficient and accurate model discretization, without requiring user intervention or even expert knowledge about a particular problem.

When the solution behaviour is at least approximately known beforehand, grid refinement can be performed using a local a priori error estimator related to the element shape and size. However, given the complicated nature of electromagnetic coupling between the large conductivity variations encountered in geological structures, a priori methods will only have limited effectiveness for EM geophysical applications. A posteriori methods based on either post-processing of the finite element solution or the computation of an auxiliary solution can overcome this difficulty (e.g. Ainsworth & Oden 2000). For example, gradient recovery methods involve post-processing of the finite element solution to obtain an improved gradient estimate, from which asymptotically exact error estimators can be obtained (e.g. Bank & Xu 2003). Other methods involve examining the residual of the finite element solution (e.g. Ainsworth & Oden 2000). In a relevant example, Franke et al. (2007) applied an error estimator based on the solution residual and the jump in tangential field components across element boundaries to create an adaptive method for magnetotelluric (MT) modelling.

The most commonly used error estimators seek to reduce the global error in the finite element solution. Specifically, the grid is refined anywhere the solution is estimated to be inaccurate. While such global error estimators can be effective for EM geophysics (e.g. Franke et al. 2007), a more efficient alternative approach can be found by noting that geophysical observations are typically made at only a relatively few discrete points, and therefore the solution only needs to be accurate at these locations. Such methods fall under the category of goal-oriented error estimation (cf. Becker & Rannacher 2001; Estep et al. 2005; Ovall 2006), where the goal is to obtain an accurate solution with respect to some functional measure—in our case the measure of interest restricts attention to some subdomain of the model, for example, at the location of EM sensors. Goal-oriented error estimators are typically developed using a sensitivity functional that measures how the error in one portion of the model may corrupt the solution elsewhere, an effect referred to as pollution. The origins of estimating and controlling error in some quantity of interest in the context of finite element computations, as opposed to error in a global norm, can be traced at least back to a series of papers by Babuška & Miller (1984a,b,c). Perhaps the earliest discussion of pollution effects for finite element problems may be found in Nitescu & Schatz (1974), and another very good standard reference for understanding this phenomenon is given in Wahlbin (1991).

We have developed goal-oriented error estimators for 2-D EM geophysics by applying the dual-error weighting approach (DEW, Ovall 2006) to the 2-D MT problem (Key & Weiss 2006) and the 2.5-D controlled-source electromagnetic (CSEM) problem (Li & Key 2007). The DEW approach has proven useful for generating accurate EM responses for simple blocky models but suffers some setbacks as the complexity of the model increases. In particular, the DEW method uses gradient recovery computations that must be performed piecewise over regions of uniform conductivity, and these scale poorly as the complexity of the model increases. Furthermore, the recovered gradients can be highly inaccurate in uniform conductivity regions that are only a single element thick (as is often the case for thin geological strata), and therefore can produce inaccurate error estimates in these regions.

In this work we investigate an alternative goal-oriented error estimator based on the dual weighted residual (DWR) method that was proposed (but not implemented) in Ovall (2006). A fundamental difference between the DWR and DEW methods is that for DWR the errors are determined through the weighting of residuals via hierarchical basis computations, rather than through the gradient recovery operations used by the DEW method. The hierarchical basis approach is advantageous since the underlying physics are applied throughout the entire model domain during the error estimate computations, rather than the piecewise application used for the gradient recovery operations. The dual approach is sourced using a new error functional that is related to the relative error in the strike aligned fields and their spatial gradients, rather than the absolute error functional implemented in previous works (Key & Weiss 2006; Li & Key 2007).

We also consider the practical implementation of our algorithm for modern parallel computers, with the goal of being able to compute accurate 2-D CSEM responses in less than 1 min for complicated models and a full suite of data parameters typical of offshore CSEM surveys. In addition to the straightforward embarrassingly parallel aspects of multifrequency, multitransmitter EM computations, we present a framework for optimized parallel computations by performing adaptive refinement on subsets of the full suite of data parameters. The adaptively refined grids generated for frequency and wavenumber subsets are then shared with nearby frequencies and wavenumbers, thus balancing computational speed with numerical accuracy.

2 PROBLEM STATEMENT

We begin by presenting the coupled differential equations that govern 2-D electromagnetic geophysics. These well-studied equations (e.g. Stoye & Greenfield 1976; Lee & Morrison 1985; Everett &
However, for low-frequency geophysical applications the imaginary \( \omega \epsilon \)
and the governing equations for the frequency domain electric field \( \mathbf{E} \) and magnetic field \( \mathbf{H} \) are
\[
\nabla \times \mathbf{E} - i \omega \mu \mathbf{H} = \mathbf{M}^f,
\]
\[
\nabla \times \mathbf{H} - \sigma \mathbf{E} = \mathbf{J}^f,
\]
where \( \mu \) is the magnetic permeability and \( \omega \) is the angular frequency. The quantity \( \sigma \) denotes the complex electric conductivity \( \sigma = \sigma - i \epsilon \), where the dielectric permittivity \( \epsilon \) is included for completeness. However, for low-frequency geophysical applications the imaginary term has a much smaller magnitude than the real conductivity \( \sigma \) and therefore is typically neglected (e.g. Ward & Hohmann 1988).

Despite the 2-D conductivity distribution, the sources \( \mathbf{J}^f \) and \( \mathbf{M}^f \) may contain 3-D variations and therefore these equations describe 3-D electromagnetic field variations. They can be transferred into 2-D equations by Fourier transformation with respect to \( x \), written as
\[
\mathbf{F}(k_y, y, z) = \int_{-\infty}^{\infty} \mathbf{F}(x, y, z) e^{-i k_x x} \, dx,
\]
where \( k_x \) is the spatial wavenumber in the strike direction and a hat (\( \hat{\cdot} \)) denotes the quantity in the wavenumber domain \( (k_y, y, z) \).

After Fourier transformation of (1) and (2), the coupled differential equations for the strike parallel components \( \hat{E}_x \) and \( \hat{H}_x \) are found to be
\[
-\nabla \cdot (\sigma \lambda \nabla \hat{E}_x) - \nabla \cdot (i k_y \lambda R \nabla \hat{H}_x) + \sigma \hat{E}_x = f_1,
\]
\[
-\nabla \cdot (i k_y \lambda R \nabla \hat{E}_x) - \nabla \cdot (i \omega \mu \lambda \nabla \hat{H}_x) + i \omega \mu \hat{H}_x = f_2,
\]
where
\[
\lambda^{-1} = k_x^2 - i \omega \mu \sigma = k_x^2 - \omega^2 \mu \epsilon - i \omega \mu \sigma,
\]
\[
f_1 = -\nabla \cdot (\sigma \lambda R \hat{M}_x^f) + \nabla \cdot (i k_y \lambda \hat{J}_y^f) - \hat{J}_y^f,
\]
\[
f_2 = -\nabla \cdot (i \omega \mu \lambda \hat{M}_y^f) + \nabla \cdot (i k_y \lambda \hat{M}_y^f) - \hat{M}_y^f,
\]
\[
R = \begin{pmatrix} 0 & -1 \\ 1 & 0 \end{pmatrix},
\]
\[
\hat{M}_y^f \text{ and } \hat{J}_y^f \text{ are the vector source currents transverse to } \hat{x} [\text{i.e. } \hat{J}_y^f = (J_y^f, \hat{J}_x^f)], \text{ and after the Fourier transformation the } \nabla \text{ operator is only defined in the transverse plane } [i.e. } \nabla = (\frac{\partial}{\partial y}, \frac{\partial}{\partial z})]. \text{ Noting that the rotation matrix } R \text{ rotates vectors by } \pi/2 \text{ counter-clockwise, we can express (4)-(5) in the compact form}
\]
\[
-\nabla \cdot (A \nabla \mathbf{u}) + C \mathbf{u} = \mathbf{f} \text{ in } \Omega \quad \mathbf{u} = 0 \text{ on } \partial \Omega,
\]
where \( A = \lambda \begin{pmatrix} \sigma I & ik_y R \\ ik_y R & i \omega \mu I \end{pmatrix} \) and \( C = \begin{pmatrix} \sigma & 0 \\ 0 & i \omega \mu \end{pmatrix} \),
\[
\nabla \cdot \mathbf{A} \nabla \mathbf{u} = \nabla \cdot \mathbf{w} = (\nabla \cdot \mathbf{w}_1, \nabla \cdot \mathbf{w}_2).
\]

Here and elsewhere, \( \Omega \subset \mathbb{R}^2 \) is our bounded computational domain, and \( I \in \mathbb{R}^{2 \times 2} \) is the identity matrix. We point out that for (4) and (5) to make sense, it is necessary that \( \lambda^{-1} \neq 0 \), as \( \lambda \) appears in both the differential operators and the right-hand sides. We will henceforth assume that \( \lambda^{-1} \neq 0 \), which will always be true for the non-zero frequency and conductivity encountered for EM geophysical applications.

Once \( \hat{E}_x \) and \( \hat{H}_x \) have been obtained via the solution of (10), the transverse field components can be computed using
\[
\hat{E}_y = -i k_y \lambda (\nabla \hat{E}_x - R \hat{M}_x^f) + i \omega \mu \lambda (R \nabla \hat{H}_x + \hat{J}_y^f),
\]
\[
\hat{H}_y = -i k_y \lambda (\nabla \hat{H}_x - R \hat{J}_y^f) + \sigma \lambda (R \nabla \hat{E}_x + \hat{M}_y^f).
\]

2.1 Finite element solution
For the finite element method, the coupled system of partial differential equations (10) is converted to the weak form and the 2-D EM problem is restated as a coupled variational problem: Find \( \mathbf{u} \in \mathcal{H} = [H^1_{0}(\Omega)]^2 \) such that
\[
\int_{\Omega} (A \nabla \mathbf{u}) : \nabla \mathbf{v} + (C \mathbf{u}) \cdot \nabla \mathbf{v} = \int_{\Omega} \mathbf{f} \cdot \nabla \mathbf{v} \text{ for all } \mathbf{v} \in \mathcal{H},
\]
where \( H^1(\Omega) \) denotes the Sobolev space of (complex-valued) functions which are square-integrable and whose first-order weak derivatives are also square-integrable; \( H^1_{0}(\Omega) \subset H^1(\Omega) \) consists of those functions which vanish on the boundary in the sense of trace (cf. Adams & Fournier 2003; Evans 2010). The notation \( \nabla \) and \( \mathbf{v} \) used here and elsewhere denotes the complex conjugate of a scalar or the componentwise complex conjugate of a vector. The space \( \mathcal{H} \) is equipped with the norm \( ||| \cdot ||| \) and semi-norm \( || \cdot ||_1 \) defined by
\[
||| \mathbf{v} ||| = \int_{\Omega} |\nabla \mathbf{v}|^2 + |\mathbf{v}|^2 \, dV, \quad ||\mathbf{v}||_1 = \int_{\Omega} |\nabla \mathbf{v}|^2 \, dV.
\]
Because of a Poincaré–Friedrichs inequality, the semi-norm \( || \cdot ||_1 \) is actually a norm on \( \mathcal{H} \), which is equivalent to the standard norm \( ||| \cdot |||_1 \), that is, there is a constant \( C > 0 \) depending only on the domain \( \Omega \) such that
\[
||| \mathbf{v} ||| \leq C ||\mathbf{v}||_1 \quad \text{for any } \mathbf{v} \in \mathcal{H}.
\]
Setting \( \mathbf{w} = (w_1, w_2) = A \nabla \mathbf{u} \), the double-dot notation should be understood as
\[
(A \nabla \mathbf{u}) : \nabla \mathbf{w} = \nabla \cdot \mathbf{v} = w_1 \cdot \nabla v_1 + w_2 \cdot \nabla v_2.
\]

For our finite element discretization, we consider a family of triangulations \( \mathcal{T}_h \) on \( \Omega \), also called meshes or grids, which is
conforming—the intersection of any two triangles \( \tau, \tau' \in \mathcal{T}_n \) (for any \( n \)) is a common vertex, a common (complete) edge or is empty; and shape-regular—all angles of all triangles in all triangulations are bounded away from 0 and \( \pi \), with bounds independent of \( n \). Such meshes can be highly adapted, and we make no assumptions about relationships between meshes, though one of our aims is that \( \mathcal{T}_n \) will be better suited for approximating \( u \) as \( n \) increases. Given \( \mathcal{T}_n \) we define the corresponding finite element space.

\[ V_n = \{ v \in \mathcal{H} \cap [C(\Omega)]^2 : v \rvert_{\tau} \in [P_1(\tau)]^2 \text{ for each } \tau \in \mathcal{T}_n \}. \]

(18)

Here and elsewhere, \( P_k(S) \) is the space of polynomials of total degree \( k \) or less on \( S \), and \( C(\Omega) \) denotes the set of globally continuous functions on \( \Omega \), meaning \( v \in V_n \) is continuous and both of its components are affine when restricted to a triangle. In other words, \( V_n \) refers to the space of piecewise-linear functions used for the finite element solution. A discrete version of (17) is as follows: Find \( u_n \in V_n \) such that

\[ B(u_n, v) = F(v) \quad \text{for all } v \in V_n. \]

(19)

The solution of (19) is equivalent to the solution of a \( 2N \times 2N \) sparse linear system, where \( N \) is the number of interior vertices in \( \mathcal{T}_n \).

The technical details related to forming the linear systems associated with (19) and (21) (below) are postponed until Appendix A; a discussion of the well posedness of (17), (19) and (21) is given in Appendix B.

### 3 Hierarchical Basis Error Estimation

#### 3.1 The global error estimator

We wish to approximate the finite element error \( u - u_n \) as a function in an auxiliary space, \( u - u_n \approx e_n \in W_n \). Defining \( V_n \) to be the set of interior vertices of \( \mathcal{T}_n \), we choose the auxiliary space to be

\[ W_n = \{ v \in \mathcal{H} \cap [C(\Omega)]^2 : v \rvert_{\tau} \in [P_2(\tau)]^2 \text{ for each } \tau \in \mathcal{T}_n \}. \]

(20)

Such spaces are said to consist of quadratic ‘edge-bump’ functions, so-called because the standard basis for this space is naturally associated with the edges in the mesh, and such functions vanish at every vertex. The approximate error function is computed by ‘projecting’ the error onto \( W_n \).

\[ B(e_n, v) = F(v) - B(u_n, v) = B(u - u_n, v) \quad \text{for all } v \in W_n. \]

(21)

We note that the spaces of functions \( X_n = V_n \oplus W_n \), whose functions \( x \) which can be expressed uniquely as \( x = v + w \), where \( v \in V_n \) and \( w \in W_n \), are precisely those which are globally continuous and quadratic on each triangle. The natural bases for the spaces \( V_n \) and \( W_n \) are described in Appendix A. Together they form a so-called ‘hierarchical basis’ for the space \( X_n \)—in contrast to the standard basis for \( X_n \), in which every basis function has degree 2.

The solution of (21) is equivalent to the solution of a \( 2M \times 2M \) linear system, where \( M \) is the number of interior edges in \( \mathcal{T}_n \).

Hierarchical basis error estimation, as outlined in the previous paragraph, is well known, and its properties are fairly well understood, at least in the case of scalar equations (cf. Bank 1996; Ainsworth & Oden 2000; Oval 2007). In the scalar case, a wealth of empirical data strongly suggests, and it can sometimes be proven (cf. Bank 1996) that there are reasonable constants \( c_1, c_2 > 0 \), independent of \( n \), for which \( c_1 \| e_n \| \leq \| u - u_n \| \leq c_2 \| e_n \| \) for \( n \) sufficiently large; also Oval (2007) gives a partial explanation of why one often sees \( \lim_{n \to \infty} \| u_n \| / \| u - u_n \| = 1 \) in practice, with ratios very near 1 even for modest mesh sizes. So we see that the global norm \( \| e_n \| \) can be reliably used to determine if \( \| u - u_n \| \) is below a given tolerance. Local norms on triangles \( \tau \), namely \( \| e_n \|_{1,\tau} \), are used to decide which triangles should be refined to further decrease the error.

#### 3.2 The goal-oriented error estimator

The global error estimator represented by (21) is useful when considering the accuracy of the finite element method over the entire model domain. However, for most practical geophysical applications, the finite element solution only needs to be locally accurate. For example, in EM geophysics an accurate solution is only required at discrete points in the model corresponding to the locations of the electric and magnetic sensors deployed for field surveys. In this case, the goal of obtaining accurate responses at the sensor locations can usually be accomplished with a much sparser mesh than required when a globally accurate solution is desired.

Consider a dual functional \( G(u) \) that is some measure of the solution at the discrete locations of interest in the model domain. For example, \( G(u) \) could be a measure of the absolute or relative size of \( u \) at the EM sensor locations, or could be a quantity derived from \( u \) such as the MT impedance. The purpose of a goal-oriented error estimator is to reduce the size of the functional error \( |G(u - u_n)|\). In other words, the goal is to reduce the error in the quantity of interest \( G(u) \), rather than the global error in \( u \). The dual/adjoint problem associated with the functional \( G \) is as follows: Find \( w \in \mathcal{H} \) such that

\[ B(v, w) = G(v) \quad \text{for all } v \in \mathcal{H}. \]

(22)

We emphasize the reversal of the roles in the arguments between this problem and the primal problem (17), and note that, on the discrete level, the matrices associated with the computation of (approximate) solutions are conjugate-transposes (adjoints) of each other. The corresponding functionals of the primal and dual solutions \( u \) and \( w \) are related by

\[ G(u) = B(u, w) = F(w). \]

(23)

The key relationship for the dual functional error \( G(u - u_n) \) is then

\[ G(u - u_n) = B(u - u_n, w) = F(w) - B(u_n, w). \]

(24)

The middle term shows that the dual solution \( w \) can be thought of as a sensitivity term for the error \( u - u_n \), which serves to weight the error according to its influence on the dual functional \( G(u) \). This idea has been exploited previously in the DEW approach (Oval 2004, 2006) and has proven useful for EM geophysical applications (Key & Weiss 2006; Li & Key 2007; Li & Pek 2008). However, DEW requires a gradient recovery operator to approximate the gradient terms in \( B(u - u_n, w) \), which for EM geophysics applications often suffers from a few shortcomings. To preserve the underlying physical boundary conditions, the gradient recovery operations must be performed piecewise over regions of uniform conductivity, and thus scale poorly as the complexity of the model increases. More importantly, the recovered gradients can be highly inaccurate in uniform conductivity regions that are only a single element thick (as is often the case when meshing thin geological strata), and therefore often produce inaccurate error estimates in these regions. Here
we consider an alternative approach based on the suggestion that the term on the right-hand side of eq. (24) could prove effective for a more robust goal-oriented error estimator (Ovall 2006). The approach described later can be considered an example of the broader class of DWR methods (cf. Becker & Rannacher 2001), in which the (approximate) solution of a related dual/adjoint problem is used to ‘weight’ either the strong or weak form of the residual of the primal/original problem. Those referring to (Ovall 2006) should note that, in that work, we call such methods dual-residual-weighting (DRW) methods—this just reflects a different naming convention.

We begin by solving the discrete dual problem for \( w_n \in V_n \).

\[
B(v, w_n) = G(v) \quad \text{for all } v \in V_n.
\]

(25)

The discrete error in the dual solution, \( w - w_n \approx \delta_n \in W_n \), is then found in a similar manner to the discrete error in the primal problem (eq. 21)

\[
B(v, \delta_n) = G(v) - B(v, w_n) \quad \text{for all } v \in W_n.
\]

(26)

An approximation to the error in the dual functional is then found by inserting \( w \approx w_n + \delta_n \) into (24), giving

\[
G(u - u_n) \approx \mu = F(w_n + \delta_n) - B(u_n, w_n + \delta_n)
\]

(27)

where the final equality in (27) is due to the identity \( F(w_n) = B(u_n, w_n) \). We also define the local element error indicator for triangle \( \tau \)

\[
\mu_\tau = |F(\delta_\tau) - B(u_\tau, \delta_\tau)|.
\]

(28)

A critical part of this method is the choice of an appropriate dual functional \( G(v) \), which in the following context is referred to as the error functional since it is a measure of the error in a specific quantity of interest. Since 2-D EM problems typically require high accuracy in both the strike aligned fields and their spatial gradients, we find the following functional to be effective:

\[
G \equiv G_s(v) = a_0 \int_{\tau_e} \bar{\varepsilon}_e \cdot v \, dV + a_1 \int_{\tau_e} \nabla \bar{\varepsilon}_e : \nabla v \, dV,
\]

(29)

where

\[
a_0 = \frac{1}{\int_{\tau_e} |(u_n + e_\tau)|^2 + c_0^2} \, dV,
\]

(30)

\[
a_1 = \frac{1}{\int_{\tau_e} |(\nabla(u_n + e_\tau)|^2 + c_1^2} \, dV,
\]

(31)

and \( \tau_e \) is the possibly discontinuous subdomain of elements containing the EM receivers. The terms \( a_0 \) and \( a_1 \) serve to weight the integrals so that they approximate the relative errors in \( u \) and \( \nabla u \) rather than their respective absolute errors. This relative weighting is appropriate for EM geophysics applications since the sources of relative error are typically more significant than the absolute errors such as sensor noise-floors. For example, uncertainties in sensor orientations, source–receiver offsets, sensor responses and other data parameters often result in an estimated data relative error of order 1 per cent regardless of the absolute field strength. Furthermore, the fields produced by CSEM transmitters can exhibit several orders of magnitude variation across an array of EM sensors. If our error functional considered only the absolute error, the adaptive refinement method could get bogged down performing excessive refinement in regions with large field amplitudes while neglecting smaller amplitude regions, at least until the absolute errors in the large-amplitude regions have been driven down to similar levels. This could result in catastrophic over-refinement that could possibly consume all available computational memory. The relative error functional given above will be far more parsimonious when guiding adaptive mesh refinement since it will operate somewhat independently of the absolute field amplitude. However, since real EM data at far source–receiver offsets are ultimately dominated by the sensor noise-floor, we do include absolute error cut-off values in our error functional through the constants \( e_0 \) and \( e_1 \), which also ensure the denominators do not equal zero. For example, \( e_0 \) could be set to the sensor noise-floor so that the dual functional will only be large when the error is significantly above the noise-floor. Such a cut-off would result in a more sparsely refined mesh than if, for example, \( e_0 \) was set to a value a few orders of magnitude smaller.

In summary, the DWR error-estimator aims to find a suitable approximation of the error functional \( G(u - u_n) \) as follows. The primal solution \( u_n \) and its estimated global error \( e_n \) are computed using (19) and (21). These quantities are then inserted into the error functional in (29). Eqs (25) and (26) are then solved for the dual solution \( w_n \) and its estimated error \( \delta_n \). Finally, the local error indicator for each triangular element is found with (28).

4 ADAPTIVE REFINEMENT METHOD FOR NUMERICAL ACCURACY

Our adaptive refinement procedure starts with a coarse Delaunay triangulation of the model domain. We apply the goal-oriented error estimation method as described to produce the local error indicator \( \mu_\tau \) for each element. We then identify the fraction \( \alpha \) of elements with the largest errors, marking them for refinement. After a new refined mesh has been created, and eqs (19), (21), (25) and (26) have been solved on this mesh, \( \mu_\tau \) is computed and again the \( \alpha \) elements with largest error estimates are marked for refinement. This process is repeated until the functional error is less than some user-specified tolerance; for example, less than 1 per cent relative error is typically desired. A downside to this approach is that computing \( \mu_\tau \) for each iteration requires the relatively expensive solutions of the finite element systems in eqs (19), (21), (25) and (26). A shortcut around this inefficiency utilizes the special structure of the error estimator, which allows it to be forward projected onto the refined mesh, thereby yielding a reasonable approximation of \( \mu_\tau \) on the refined grid without requiring a new solution of the finite element systems (e.g. Bank & Xu 2003). In practice, we use a low-order quadrature integration to compute \( \mu_\tau \) on the refined elements. This method can be re-applied for a few refinement iterations before the approximation breaks down and a new computation of the error estimator is required. We find that five iterations of refining the 6 per cent worst elements by approximately subdividing them in half works well, with the mesh vertex count doubling in size, after which the finite element systems for the error estimator need to be solved for the new mesh.

5 PARALLEL IMPLEMENTATION DETAILS

We have implemented this goal-oriented adaptive refinement methodology into a parallel Fortran code named MARE2DEM (Modeling with Adaptively Refined Elements for 2-D EM) that currently supports both MT and CSEM problems. Parallel constructs were implemented using the Message Passing Interface standard. For CSEM modelling, zero Dirichlet conditions are applied to the model boundaries and the dipole sources are approximated using point delta functions, rather than the discrete pseudo-delta-function approach used in fixed-grid modelling approaches (e.g. Mitsuhata.
Dipole sources are obtained by the vector superposition of the fields computed at discrete $k_x$ values. The fields from arbitrarily oriented dipole sources are obtained by the vector superposition of the fields from a source oriented in the $\hat{x}$ direction and another in the $(\hat{y}, \hat{z})$ plane, therefore requiring at most one additional source computation. We use 401 point digital filters designed in a manner similar to Guptaarma & Singh (1997). For the MT method, the plane-wave source field is implemented through 1-D boundary conditions applied to the left- and right-hand sides of the model domain with a cosine taper of these solutions applied along the top and bottom boundaries (Wannamaker et al. 1987).

5.1 Mesh generation and refinement

The input conductivity model consists of segment bound polygonal regions with an assigned constant conductivity for each region. The model outer boundaries are placed at least several skin depths away from the region of interest so that anomalous fields generated in the centre of the model are sufficiently attenuated by the model boundaries, in accordance with the imposed Dirichlet conditions. Mesh generation and subsequent refinements are accomplished by direct calls from Fortran to routines in the C-code Triangle, a program for robust and rapid generation of constrained, conforming Delaunay triangulations (Shewchuk 1996). Triangle uses a minimum interior angle constraint to generate high-quality triangulations that are suitable for numerical accuracy and stability considerations (e.g. Shewchuk 2002a). This angle constraint presents the practical issue that the input model structure should not have narrow ‘slivers’, where two segments intersect with an angle smaller than the constraint. A typical minimum angle constraint in the range of 20°–30° serves to balance mesh quality with flexibility in the input model structure. Mesh refinement is performed in Triangle through the use of element area constraints. In MARE2DEM, elements that have been marked for refinement are given area constraints that are half their current area.

The forward projection of the error estimate from a coarse mesh onto a refined mesh requires knowledge of which parent element in the coarse mesh contains the centroid of each child element in the refined grid. Since refinement in Triangle results in a sequence of meshes containing a hierarchy of vertices rather than a hierarchy of triangles, identifying the parent element is more costly than for algorithms that refine simply through triangle bisection. Because the error needs to be projected onto all elements of the refined grid, computing brute-force in-triangle point location queries for each element in the refined grid would scale very poorly as the mesh size increases. Instead, MARE2DEM performs efficient point location queries by using a variant of the jump-and-walk method that uses a 2-D tree structure (e.g. Kennel 2004) for rapid location of the nearest vertex (the jump) before traversing adjacent triangles to find which one contains the query point (the walk), resulting in a very favourable expected time dependence of $\Theta(n^{0.056})$ (Devroye et al. 2004).

5.2 Efficient solution of the linear systems

The four linear systems required for the error estimator may seem like an expensive burden that limits its practicality. However, the sparse matrices for these systems are identical between (19) and (25), and between (21) and (26); therefore, they can be solved efficiently using the LU decomposition (e.g. Golub & Van Loan 1996).

$$Ax = LUx = b.$$  \hspace{1cm}  (32)

where $L$ and $U$ are lower and upper triangular matrices. Once $L$ and $U$ have been computed, the solution for any right-hand side vector $b$ is found using forward and backward substitutions for the triangular systems

$$y = L^{-1}b, \quad x = U^{-1}y.$$  \hspace{1cm}  (33)

MARE2DEM solves for a single right-hand side vector $b$ of source vectors can be added to the linear system with only a small increase in the computational time. Since the system matrices for the primal and dual problems are identical, MARE2DEM stores the LU factors computed for the primal problem and then reuses them for the dual problem (both for the linear and bump system matrices). The effective cost of the goal-oriented error estimator is therefore the additional time required to factorize the bump system matrix and the time for the triangular solutions of the three additional systems, totaling to roughly a factor of two increase over the time required for the primal problem alone. Additionally, as we show in the Appendix, the coupling terms between $\tilde{E}_i$ and $H_i$ integrate to zero for the hierarchical basis system; hence MARE2DEM solves two smaller $M \times M$ systems rather than a single $2M \times 2M$ system when computing the hierarchical basis.

![Figure 1. Scaling of the SuperLU solver for a typical adaptive refinement problem as a function of the number of mesh vertices. Triangles and diamonds show the time to compute the LU factorization for the sparse linear systems associated with the linear basis of the space $V_0$ and the quadratic bump basis of the space $W_0$. Dots show the time to compute the triangular solves for a single right-hand side vector. Computations were performed serially on a MacPro desktop computer containing two 2.93 GHz Quad-core Intel Xeon processors.](image)
residuals. Since \( k_z = 0 \) for the MT method, all the linear systems are decoupled and the primal and dual problems are solved rapidly as two uncoupled \( N \times N \) systems rather than the \( 2N \times 2N \) coupled system required for the CSEM method.

Another efficiency of the LU decomposition is found for CSEM modelling. Multiple transmitter sources \( \mathbf{J}_t \) and \( \mathbf{M}_t \) in (4) and (5) result in multiple right-hand side vectors in the linear systems that can be solved rapidly using the LU method described. For mesh \( i \) with \( N_i \) vertices and \( n_i \) transmitters on the right-hand sides of the linear systems, the time to solve the finite element systems and compute the error estimator is approximately

\[
t_i = a N_i^0 + 2c N_i^d + 4n_i e N_i^f,
\]

where the first term on the right-hand side represents LU factorization time for the \( 2N_i \times 2N_i \) sparse matrix used for both the primal and dual problems in (19) and (25), the second term is the LU factorization time for the two decoupled bump systems in (21) and (26), and the third term is the triangular solution time for the two linear and two bump systems for \( n_i \) sources. Table 1 lists values for the scaling coefficients obtained by least squares fits to the performance data shown in Fig. 1.

Fig. 2 shows \( t_i \) as a function of the number of sources for meshes with 1000, 10 000 and 100 000 vertices. We can see that about 10–20 sources can be put on the right-hand side before the run-time doubles, and that further additional sources produce a run-time that is directly proportional to the number of sources. This suggests that for problems with multiple transmitters, some efficiency can be gained lumping sources together into groups of about 10 for the adaptive refinement computations, rather than modelling each transmitter as an independent modelling task. A trade-off in this approach is that the resulting mesh will require more vertices (and hence memory and run-time) to more accurately capture the electromagnetic fields produced by each transmitter. However, for the CSEM problems we have examined this is typically far offset by the time-efficiency gained from the multiple right-hand side approach.

5.3 Parallelization

Many aspects of the 2.5-D EM problem are embarrassingly parallel and readily allow for large savings in the computational run-time. Eqs (4) and (5) show that independent computations are required for each wavenumber \( k_x \) and frequency \( \omega \), as well as for each source in \( \mathbf{J}_t \) and \( \mathbf{M}_t \). A typical 2-D CSEM data set contains around 100 transmitter locations and 10 transmission frequencies, while about 30 wavenumbers are required for an accurate 2.5-D computation. Thus, the fastest run-time would be obtained by running each of the 30 000 independent tasks on a cluster of the same size. Such massive clusters exist but represent the pinnacle of current technology and are therefore of limited availability; more typical clusters consist of a few hundred processors, while desktop machines routinely ship with eight or more processing cores. Thus, for ordinary CSEM modelling problems there will be significantly more parallel modelling tasks than available processors, although the number of tasks can be reduced by an order of magnitude through the transmitter grouping method mentioned in the previous section.

To demonstrate how MARE2DEM might scale on a large parallel system, we can create a model for the expected run-time, which is dominated by the solution of the linear systems. Assuming that the refinement iterations double the vertex count at each iteration (i.e. \( N_i = 2^{n_i-1} N_1 \)), the total time to perform \( n \) refinement iterations is

\[
T_n = \sum_{i=1}^{n} t_i = a N_1^0 \left( \frac{2^n - 1}{2^1 - 1} \right) + 2c N_1^d \left( \frac{2^n - 1}{2^1 - 1} \right) + 4n e N_1^f \left( \frac{2^n - 1}{2^1 - 1} \right).
\]

If there are \( n_s \) sources used for LU factorization speedup, the total number of independent tasks is then \( n_{\text{tasks}} = n_n / n_s \). If we assume that each task requires roughly the same adaptive refinement time and \( n_{\text{proc}} \) is the number of parallel processors available, the expected total run-time required to complete all tasks at a given wavenumber and frequency is then

\[
T_{\text{total}} = \frac{n_{\text{tasks}}}{n_{\text{proc}}} T_n.
\]

As an example of the implications of this equation, Fig. 3 shows \( T_{\text{total}} \) as a function of the number of processors used, where the initial mesh contained \( N_1 = 5000 \) vertices and \( n = 4 \) and \( n_s = 1024 \). Although the fastest run-time is achieved by using a single processor for each task (i.e. \( n_s = 1 \)), a reasonable run-time can be achieved with a factor of 16 fewer processors by grouping the sources together into sets of \( n_s = 16 \). On a single processor, grouping all sources together (\( n_s = 1024 \)), while not very efficient, is still over 10 times faster than solving for each source separately (\( n_s = 1 \)).

For the final component of the parallel implementation, we introduce the concepts of mesh sharing and receiver grouping. Traditional methods for EM modelling utilize a single fixed mesh that is designed to be suitable for the entire range of parameters (i.e. all transmitters, frequencies, receivers and wavenumbers), and therefore such a mesh is likely to be overmeshed for at least some of the data parameters, leading to an inefficiency in the run-time. Conversely, the adaptive finite element technique allows MARE2DEM...
to generate a unique mesh that is optimized for a given set of parameters, resulting in a very efficient use of mesh vertices. Since the EM solutions typically vary smoothly as a function of frequency and wavenumber, a mesh refined for a specific pair of these values is usually accurate for nearby values, as shown previously by examples in Key & Weiss (2006) and Li & Key (2007). Therefore, another efficiency is to share an adaptively refined mesh with neighbouring frequencies and wavenumbers so that the refinement iterations only need to be computed once per mesh sharing group. Fig. 4 provides a visual interpretation of the mesh sharing concept, illustrating how various data parameters can be grouped together while the adaptive refinement iterations are performed only for the centre parameter value. Since the refinement iterations require time $T_n$ and the computations for the nearby frequency–wavenumber pairs require time $t_n$, mesh sharing in the frequency–wavenumber space typical results in about 2–4 times faster run-times than for a full refinement sequence of every frequency–wavenumber pair.

Finally, the receivers can also be decomposed into small groups so that a given adaptive refinement task considers only a few receivers and therefore will run faster since a few receivers can be adaptively meshed with a more limited amount of mesh refinement than when all receivers are considered. We note that there can sometimes be a negative trade-off in performance for receiver grouping since each receiver group increases the number of parallel tasks to be performed, while the change in run-time for having fewer receivers in a given group may only be marginal. The best increase in performance will be found for very simple starting models with low mesh vertex counts to begin with, and always when the number of available processors is equal to or greater than the number of parallel tasks.

### 6 Validation

Here we present examples that demonstrate the reliability and effectiveness of MARE2DEM for CSEM and MT modelling. The first example considers marine CSEM responses for the canonical 1-D reservoir model, consisting of a 100-m-thick, 100-ohm-m reservoir buried 1 km below the seafloor in 1 ohm-m sediments. The usefulness of the canonical 1-D model resides in its well-studied solutions and semi-analytic solutions are available (Weaver et al. 1985; Weaver 1986). The effectiveness of MARE2DEM for CSEM and MT modelling. The first example consists of a 0.25 Hz horizontal electric dipole transmitter located 50 m above the seafloor while receivers are positioned along the seafloor every 500 m from 0 to 15 km range. Computations were performed on a MacPro Desktop containing two 2.93 GHz quad-core Intel CPUs, for a full capability of eight parallel processors. The MARE2DEM computations used 30 wavenumbers spaced logarithmically from $10^{-6}$ to $10^{-3}$ m$^{-1}$, with six continuous subsets of five wavenumbers used for mesh sharing. The mesh sharing subsets reached the 1 per cent requested error tolerance within five–six adaptive refinement iterations. The starting mesh contained 4584 vertices, while the final meshes contained around 18,000–19,000 vertices, depending on the specific wavenumber.

The accuracy of the MARE2DEM solutions is measured by comparisons with the semi-analytical 1-D responses from the open-source code Dipole1D (Key 2009). Fig. 5 shows that all field components for inline receivers are below the requested 1 per cent error tolerance, demonstrating the validity of the MARE2DEM solutions for this model and data parameters. Fig. 6 extends this test to illustrate the code’s effectiveness for a broad range of requested error tolerances. The true errors for all field components are slightly below the requested error tolerances, indicating that the goal-oriented error estimator is reliable for achieving a given tolerance. Furthermore, the increase in run-time that accompanies the significant reduction in the solution error is very modest—less than a factor of 10 despite the 1000-fold decrease in error. This demonstrates the efficiency in mesh refinement possible with goal-oriented error estimation.

Fig. 7 shows the accuracy of MARE2DEM for an example land MT problem consisting of the segmented slab model, for which semi-analytic solutions are available (Weaver et al. 1985; Weaver et al. 1986). The segmented slab is composed of a 100-km-thick
slab that is divided into a 5-km-wide, 1 ohm-m region in the middle, and is bound on the left- and right-hand side by 10 and 50 ohm-m regions. The analytic solution has a perfect conductor underneath the slab, which we approximate in the finite element model by setting the basal resistivity to $10^{-8}$ ohm-m. MT responses at 100 s period were computed for 197 sites atop the slab at lateral positions between $-100$ and 100 km. For the MARE2DEM computations, the receivers were grouped into subsets of 20 receivers for each adaptive mesh refinement computation. On the MacPro computer described earlier, the entire process of adaptive refinement computations for all groups took about 1 s. Fig. 7 shows that the relative error in strike aligned electric field ($E_x$) is mostly below the 1 per cent requested tolerance, although many of the receivers over the conductive middle segment have errors of up to 1.4 per cent. Fig. 8 shows the coarse starting mesh consisting of 139 vertices and two examples of adaptively refined meshes for two different receiver groups that required around 3400 and 2600 vertices to meet the requested 1 per cent error tolerance. The adaptively refined meshes are substantially denser around the particular receiver groups. They also show significant refinement in regions distal to the receivers, indicating where the mesh required refinement to mitigate pollution effects.

The fundamental datum of a MT survey is the impedance tensor, which for 2-D conductivity structures can be decomposed into the TE and TM mode impedances

$$Z_{TE} = \frac{E_x}{H_y}, \quad Z_{TM} = \frac{E_y}{H_x}. \quad (37)$$

Figure 5. Validation of the magnetic and electric field responses computed by MARE2DEM for the canonical 1-D reservoir model. Electric and magnetic field responses are shown in the top panel and the bottom panel shows their relative errors compared to solutions obtained with the Dipole1D modelling code.

Figure 6. Effectiveness of the goal-oriented error estimator. The top plot shows the true error of the MARE2DEM solutions as a function of the requested error tolerance. The bottom plot shows the accompanying runtime required for the full adaptive refinement computations.

Figure 7. TE electric field response at 100 s period for the segmented slab model (top panel) and the relative error (bottom panel). Vertical lines delineate groups of receivers used for separate adaptive refinement computations.
The previous simple examples provided a validation of MARE2DEM and illustrated the reliability of the goal-oriented error estimator. Here we demonstrate how MARE2DEM performs for a significantly more complicated model and a much broader suite of data parameters. We designed a synthetic model that is representative of typical marine CSEM surveys for hydrocarbon exploration on the continental shelves (Fig. 10). The complex model contains many features that cannot be discretized efficiently on a structured rectangular grid: realistic seafloor topography, tilted geological strata and closely spaced thin resistive layers simulating a stacked reservoir sequence. The three dipping reservoir layers are only 10 m thick and have a resistivity of 20 ohm-m. All of these features are readily accommodated by the unstructured triangular grid using just under 7000 mesh vertices. The stratigraphic intervals might be known \textit{a priori} from seismic exploration and exploratory wells. The purpose of conducting a CSEM survey would be to test whether a seismically imaged geological trap structure is filled with resistive hydrocarbons or conductive sea water (e.g. Eidesmo et al. 2002), or to estimate the saturation of a reservoir as a function of time during production (e.g. Orange et al. 2009). The entire model domain is 200 km wide and 200 km high, vertically including 100 km of air
We modelled the CSEM responses at 12 frequencies spanning inline electric dipole transmitters deep-towed 50 m above the domain horizontally. The unstructured grid used as the starting mesh (bottom panel). Inverted stacked reservoir layers (middle panel, note the vertical exaggeration) and intervals. The three panels show the model (top panel), a close-up of the slope, tilted regional strata and closely spaced thin and dipping reservoir layers. The requested error tolerance was set to 1 per cent. Computations were performed on the Triton Compute Cluster (TCC) maintained by the San Diego Supercomputer Center at the University of California, San Diego. The TCC system is composed of 256 nodes containing two quad-core Intel Nehalem 2.4 GHz CPUs, representing a parallel capacity of 2048 processors.

We performed a series of tests to examine the scaling of the MARE2DEM run-time as a function of the number of parallel processors. If each unique transmitter, receiver, wavenumber and frequency combination were considered separately for adaptive mesh refinement, the total number of tasks would be just under one million. Although this would yield the fastest run-time on a parallel system of the same size, this is far in excess of the number of available processors on the TCC system. Instead we use mesh sharing and parameter grouping to reduce the number of tasks significantly. We grouped all receivers together and then applied mesh sharing to groups of three consecutive frequencies and five consecutive wavenumbers. We then varied the number of transmitters grouped together into sets of 1, 11 and 51 transmitters, as shown in Table 2. We also ran a test of only a single frequency (0.3 Hz) to study how fast MARE2DEM performs for a reduced task. Depending on the number of frequencies under consideration and the number of transmitters grouped together, the number of total tasks varies from 30 (for one frequency and all 51 transmitters grouped together) to 18360 (for 12 frequencies and each transmitter considered separately). Due to mesh sharing, the number of refinement tasks is much lower, spanning from six to 1224. We ran MARE2DEM for these parametrizations using from one to 800 parallel processors.

Fig. 11 shows the corresponding wall clock run-times for the scaling tests. All runs exhibit a linear decrease in run-time that is proportional to the number of processors, up until the number of processors exceeds the number of refinement tasks, after which the run-time flattens out regardless of the number of processors used. The linear decrease in run-time is due to the embarrassingly parallel nature of the problem, where the low communications overhead allows for highly efficient parallel scaling. The flattening in run-time with additional processors occurs because the refinement tasks and mesh sharing tasks have been completed and no further speed-ups are possible with additional processors. For example, consider run (b), which required 120 refinement tasks and an additional 1680 computations when the final grids are shared with nearby parameters. The scaling tests show a linear decrease in run-time up until 120 processors were used, after which the increase in the number of processors does not further reduce the run-time, despite the additional 1680 mesh sharing tasks. This is simply explained by the variability of run-time required for each refinement task, which depends on the particular data parameters of the task. When a processor running a refinement task finishes, it then becomes available to compute any mesh sharing tasks. For run (b), when the last (120th) adaptive refinement task completes, all other mesh sharing tasks have already been handled by the other nodes and the only remaining tasks are to compute its own mesh sharing computations. Therefore, no further speed improvements are possible from additional processors.

The scaling tests point out a few different ways that MARE2DEM can be used to model large 2-D CSEM problems in as short a time as possible. For very large clusters containing hundreds of processors or more, the fastest run-times can be obtained by using small groups of transmitters, producing run-times of under a minute for typical CSEM survey parameters when using a few hundred nodes. Conversely, on smaller multicore desktop computers that a typical working group might maintain (of order 10 processors), larger transmitter groups can be used to get run-times as short as a few minutes. In our tests here, the fastest run-time of only

<table>
<thead>
<tr>
<th>Run</th>
<th># Frequencies</th>
<th># Tx grouped</th>
<th># Total tasks</th>
<th># Refinement tasks</th>
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<tr>
<td>a</td>
<td>12</td>
<td>1</td>
<td>18360</td>
<td>1224</td>
</tr>
<tr>
<td>b</td>
<td>12</td>
<td>11</td>
<td>1800</td>
<td>120</td>
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<td>c</td>
<td>12</td>
<td>51</td>
<td>360</td>
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<td>1530</td>
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<td>1</td>
<td>51</td>
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23 s was obtained using 800 processors when only one frequency was modelled and each transmitter was grouped separately. We can predict that on a larger parallel system of at least 9600 processors, the full problem containing all 12 frequencies could be computed in as short as 23 s.

Fig. 12 shows an example of the error estimator $\mu_{\tau}$ computed for the starting mesh (6974 vertices) and the final refined grid (26 065 vertices) computed for a transmitter at 300 km position, $k_x = 0.000016$ m$^{-1}$ and $f = 0.3$ Hz. The bottom panel shows the mesh refinement for the final refined grid. Inverted white triangles show the location of the seafloor EM receivers. Only the central region of the modelling domain is shown.

Table 3 provides a representative example of the adaptive refinement performance when the starting mesh is refined for a single transmitter at 0.3 Hz and wavenumber 0.001 m$^{-1}$. Five iterations of grid refinement were required to reach the target 1 per cent error estimate, requiring a total run-time of 9.2 s. The adaptive refinement iterations dominated about 80 per cent of the run-time, whereas the final grid required an additional 2.0 s, or roughly 20 per cent additional time. When the final grid is shared with nearby frequencies and wavenumbers, only the final run-time of 2.0 s is required for these computations, which illustrates how mesh sharing can lead to significantly faster run-times. The table also documents the effectiveness of the error estimator, as noted by the significant decrease in $\max\{\mu_{\tau}\}$ with each iteration.

Fig. 13 shows an example of the inline electric field responses computed at 0.3 Hz. The responses for all 51 receivers and transmitters are shown for computations that included the three resistive
stacked reservoir layers and for computations where the reservoir layers were removed. The responses show a significant amount of structure, particularly at amplitudes around $10^{-15}$ to $10^{-13}$ V m$^{-2}$, where a large change in slope occurs with the arrival of the air-wave. This typically occurs at large source–receiver offsets, where the responses become dominated by energy that has travelled along the air–sea interface (e.g. Constable & Weiss 2006).

Due to the strong attenuation of the fields with distance from the transmitter, the differences between the responses with and without the reservoir are difficult to identify when viewed on an absolute scale. Fig. 14 shows the relative differences between these responses in a midpoint-offset section, as shown for four of the 12 frequencies computed. These sections are generated by plotting the relative difference between the reservoir and no-reservoir responses.

Figure 13. Horizontal electric field amplitude responses for the complex model with a 0.3 Hz inline electric field transmitter. Solid lines show the responses for the complex model in Fig. 10 and dashed lines show the responses for the same model after removal of the three stacked reservoir layers.

Figure 14. Electric field anomaly midpoint-offset sections for the complex model shown for four frequencies. The relative difference between the complex model response and the response without the reservoir layers is plotted horizontally at the source–receiver midpoint and vertically at the source–receiver offset. Since the responses for the sources and receivers are nearly reciprocal, only positive offsets are shown. Black contour lines show the absolute magnitude of the responses in units of log$_{10}$ V m$^{-2}$. The shaded area for the 4.7 Hz solution covers the region where the solution is below the computational noise floor.
horizontally along the transmitter–receiver midpoint, and vertically at the absolute transmitter–receiver offset. Since longer offsets have a broader and deeper sensitivity, these plots are sometimes referred to as anomaly pseudo-sections. We can now see that the peak signal from the reservoir occurs at transmitter–receiver offsets of about 4–8 km and is largely confined to source–receiver midpoints located over the reservoir. Although the largest anomaly is observed at 1.1 Hz, most of the signal at this frequency is below the typical marine CSEM system noise-floor of about $10^{-15}$ V A$^{-1}$ m$^{-2}$. This is due to the rapid inductive attenuation of high-frequency data with source–receiver offset. Conversely, the lower frequency of 0.3 Hz experiences less attenuation yet still contains a measurable 20–30 per cent anomaly, suggesting that this frequency would be optimal for imaging the reservoir.

8 CONCLUSIONS

The parallel adaptive finite element algorithm presented here can be used for rapid and accurate simulations of 2-D EM problems. The unstructured triangular grid readily incorporates complex structures with an efficient use of mesh vertices and without the discretization difficulties and cumbersome approximations experienced with rectangular grids. Our use of an automated adaptive refinement method based on a goal-oriented error estimator allows the MARE2DEM program to generate reliable EM responses, even for finite element novices who may have little or no experience with mesh design. The LU matrix factorization results in rapid solutions of the additional systems associated with the error estimator, and also allows for efficient computations when multiple transmitters are modelled. We showed that the parallelization over subsets of transmitters, receivers, wavenumbers and frequencies scales very efficiently for clusters of up to at least 1000 processors. Mesh sharing for nearby frequencies and wavenumbers can be used to reduce the parallel load and offers a method for fast solutions when only a smaller number of parallel processors are available.

Our final example demonstrated how MARE2DEM can automatically compute EM responses for a realistic offshore model with realistic data parameters in run-times of under 1 min when run on a large cluster. Further speed improvements may be possible by incorporating a parallel algorithm for the LU factorizations, although the performance increase from this would be modest compared to the high efficiency of the independent parallel EM computations described here. However, if a significantly large parallel system is available, such that all embarrassingly parallel aspects of the problem are exploited and additional processors are still available, then parallel matrix factorizations may be a worthwhile consideration.

Although the fast performance of MARE2DEM will be beneficial for the geophysicist interested in forward modelling highly detailed 2-D problems, we anticipate that this algorithm will prove more useful as a kernel routine for non-linear inversions. Gradient-based inversions for 2-D EM have been around for some time (e.g. Unsworth & Oldenburg 1995), yet Bayesian methods have long been considered impractical since they can require tens of thousands of forward computations. The run-times of our complex test model show that Bayesian methods for realistic 2-D EM problems are now tractable on large clusters. In addition to inversion performance improvements from the parallel implementation, the automated adaptive meshing may lead to stability improvements, where an accurate mesh is automatically generated as model structure evolves throughout the inversion process, thereby insuring numerical accuracy is not compromised. An initial test of using MARE2DEM for a stochastic sharp boundary inversion for 2-D MT problems is given in Chen et al. (2010). Finally, the adaptive methods we proposed here could be extended to finite element methods developed for 3-D electromagnetic problems (e.g. Monk 2003).

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APPENDIX A: FINITE ELEMENT DETAILS

This brief appendix is intended as an aid for those interested in the finite element details required for the implementation and testing of our goal oriented adaptive method. Because the solution of the discrete dual/adjoint problems (25) and (26) involves the adjoints of the matrices needed for the solution of the discrete primal problems (19) and (21), we only describe those for the former.
Let $T$ be a given triangulation, having vertices $V$ and edges $E$. For any vertex $z \in V$ we define a function $l_z$, which is linear on every triangle, and which has value 1 at $z$ and value 0 at every other vertex—in brief, $l_z(z') = \delta_{zz'}$ for all $z, z' \in V$. From this relation, it is clear that $l_z$ is identically 0 on any triangle which does not have $z$ as a vertex. If $\tau$ is a triangle which has $z$ as a vertex, and we number the vertices of $\tau$ counter-clockwise starting at $z$, then the formula for $l_z$ on $\tau$ is given in eqs (A3)-(A7)—the link is that $l_z = l_1$ on $\tau$. Because of the key relation $l_z(z') = \delta_{zz'}$, any continuous, piecewise-linear scalar function $v$ on $\Omega$ can be described uniquely in terms of the basis $\{l_z\}$ by $v = \sum_{z \in V} v_z l_z$, where $v_z = v(z)$. More specifically, the finite element solution $u = (u_{a,1}, u_{a,2})$ of (19) can be expressed in the form

$$u_{a,1} = \sum_{z \in V_i} u_{a,1} l_z, \quad u_{a,2} = \sum_{z \in V_i} u_{a,2} l_z,$$

where $V_i$ are the ‘interior’ vertices in the triangulation—our function vanishes on the boundary. After choosing a global ordering of the interior vertices, the finite element solution is found by computing its coefficients $\{u_{a,1} : z \in V_i\}$ and $\{u_{a,2} : z \in V_i\}$ via solving the $(2 \times 2)$ block linear system

$$
\begin{pmatrix}
A_{11} & A_{12} \\
A_{21} & A_{22}
\end{pmatrix}
\begin{pmatrix}
u_1 \\
u_2
\end{pmatrix}
=
\begin{pmatrix}
f_1 \\
f_2
\end{pmatrix},
$$

where $\{\nu_1, \nu_2\}$ is the vector of coefficients for $u_{a,1}$, $\{\nu_1, \nu_2\}$ is the vector of coefficients for $u_{a,2}$, and

(i) the entries of $A_{11}$ are of the form: $\int_{\Omega} \sigma(\lambda) \nabla l_1 \cdot \nabla l_1 + l_1 l_2) \, dV$,

(ii) the entries of $A_{22}$ are of the form: $\int_{\Omega} \sigma(\lambda) \nabla l_2 \cdot \nabla l_2 + l_1 l_2) \, dV$,

(iii) the entries of $A_{12} = A_{21}$ are of the form: $\int_{\Omega} l_1 l_2 \cdot \nabla l_1 \cdot \nabla l_2 \, dV$,

(iv) the entries of $f_1$ are of the form: $\int_{\Omega} f l_1 \cdot dV$ for $k = 1, 2$.

If $N$ is the number of interior vertices, we see that this system is $2N \times 2N$. The linear system A2 can be made complex symmetric simply by multiplying the bottom row by a negative constant.

Similarly, for any edge $e \in E$, let $z \in V$ and $z' \in V$ be its vertices. We define $q_e = 4l_1 l_2$ for all $e \in E$ and $q_e$ is only non-zero on one or (generally) two triangles having $e$ as an edge. Again, if we pick one of these triangles and number its edges and vertices counter-clockwise as before, we have a formula for $q_e$ (A8) on that triangle. Our chosen basis for $W_e$ is $\{q_e : e \in E_i\}$, where $E_i$ is the set of interior edges. The computation of $e_n$ in terms of this basis involves the solution of a $2M \times 2M$ system analogous to (A2), where $M$ is the number of interior edges. As we see in (A15), for this system the analogues of $A_{11}$ and $A_{22}$ are identically 0—that is, the system naturally decouples. Though we do not explicitly give the right-hand side for this system, it is easily derived from (21), and is constructed by computing integrals on triangles of the forms (A16)–(A18). Below we outline the rudimentary aspects of how the integrations are carried out for each element.

For a triangular element $\tau$ with vertices $i, j, k$ and edges $e_i, e_j, e_k$ opposite those vertices (Fig. A1), the linear basis function $l_i$ associated with vertex $i$ varies from value 1 at vertex $i$ to zero at the other two vertices according to

$$l_i = \frac{1}{2|\tau|}(a_i + b_i y + c_i z),$$

where the area of the triangle $|\tau|$ is defined by

$$2|\tau| = \begin{vmatrix} 1 & 1 & 1 \\ y_i & y_j & y_k \\ z_j & z_j & z_k \end{vmatrix} = (a_i + b_i y_i + c_i z_i),$$

and

$$a_i = y_j z_k - y_k z_j, \quad (A5)$$

$$b_i = z_j - z_k, \quad (A6)$$

$$c_i = y_k - y_j, \quad (A7)$$

with cyclic permutation of $i, j, k$. The quadratic bump basis function associated with edge $i$ is

$$q_i = 4l_1 l_2,$$

again with cyclic permutation of $i, j, k$. The bump function $q_i$ is zero at all three vertices and has a quadratic shape along edge $e_i$.

The details of forming the linear systems follow the standard approaches for the finite element method (e.g. Zhu & Cangellaris 2006), so here we only provide a few details that are relevant to our implementation. Using the well-known formula for polynomial integration on triangles (e.g. Eisenberg & Malvern 1973)

$$\int_{\tau} l_i l_j d\tau = \frac{2|\tau|}{(p + q + r + 2)!} l_i l_j,$$

we find the following formulae for the integrals required by the finite element equations. For the linear basis functions $l_i$, these are

$$\int_{\tau} \nabla l_i \cdot \nabla l_j = \frac{1}{4|\tau|} (b_i b_j + c_i c_j),$$

$$\int_{\tau} l_i l_j = \begin{vmatrix} 1 \frac{|e_j|}{|\tau|} \\ |e_j| \frac{|e_i|}{|\tau|} \end{vmatrix}, \quad \begin{array}{c} \quad i = j \\ \quad i \neq j \end{array},$$

$$\int_{\tau} (R \nabla l_i) \cdot \nabla l_j = \frac{1}{4|\tau|} (b_i c_j - b_j c_i).$$

Here, we recall the rotation matrix $R$ described in Section 2. For the quadratic basis functions $q_i$, we have

$$\int_{\tau} \nabla q_i \cdot \nabla q_j = \begin{vmatrix} 1 \frac{|e_j|}{|\tau|} (b_i b_j + c_i c_j) \quad i = j \\ 1 \frac{|e_j|}{|\tau|} (b_i b_j + c_i c_j) \quad i \neq j \end{vmatrix}.$$
The existence of a solution for the infinite dimensional problem such that

\[ \int q_i q_j = \begin{cases} \frac{4}{\pi |r|}, & i = j \\ \frac{4}{\pi^2 |r|}, & i \neq j \end{cases}, \]  

\[ \int (R \nabla q_i) \cdot \nabla q_j = 0. \]  

This last integral is particularly useful since it eliminates the cross coupling between \( E_\alpha \) and \( H_\alpha \) in the linear systems using the quadratic bump basis, and hence each field can be solved separately using \( M \times M \) sparse symmetric linear systems, rather than the larger \( 2M \times 2M \) systems required by the coupled problem. The right-hand side of (21), (26) and (29) require the integral formulae

\[ \int \nabla l_i \cdot \nabla q_j = -\frac{1}{3 |r|} (b_i b_j + c_i c_j). \]  

\[ \int l_i q_j = \begin{cases} \frac{4}{\pi^2 r}, & i \neq j \\ \frac{4}{\pi r}, & i = j \end{cases}, \]  

\[ \int (R \nabla l_i) \cdot \nabla q_j = \frac{1}{3 |r|} (b_i c_j - c_i b_j). \]  

Of course, \((R \nabla q_i) \cdot \nabla l_i = - (R \nabla l_i) \cdot \nabla q_j\), so this last identity can be used with linear and quadratic functions reversed by changing sign.

**APPENDIX B: WELL POSEDNESS OF THE VARIATIONAL PROBLEMS**

In low-frequency EM geophysics applications it is legitimate to take \( \epsilon = 0 \), and we have done so in our numerical experiments. For the analysis below, we make this assumption as well. In this case, \( \lambda^{-1} = k_0^2 - \omega \mu \sigma \), and the ‘diffusion matrix’ \( A \) in our bilinear form \( B \) is given by \( A = A_1 + i A_2 \), where

\[ A_1 = \frac{1}{k_0^2 + (\omega \mu \sigma)^2} \begin{pmatrix} k_0^2 \sigma I & ik_0^2 R \\ ik_0^2 R & -(\omega \mu \sigma)^2 I \end{pmatrix} \]  

and

\[ A_2 = \frac{1}{k_0^2 + (\omega \mu \sigma)^2} \begin{pmatrix} \omega \mu \sigma^2 I & ik_0 \omega \mu \sigma R \\ ik_0 \omega \mu \sigma R & k_0^2 \omega \mu \sigma I \end{pmatrix}. \]  

We will first argue that the problems (17), (19) and (21) can have at most one solution, by showing that the only solution of ‘Find \( u \in \mathcal{H} \) such that \( B(u, v) = 0 \) for all \( v \in \mathcal{H} \) is \( u = 0 \). In particular, we will show that \( B(u, u) = 0 \) only if \( u = 0 \). For the finite dimensional problems (19) and (21), the fact that at most one solution can exist automatically implies that precisely one solution exists. To prove the existence of a solution for the infinite dimensional problem (17), we must also establish a Gårding inequality (cf. Ihlenburg 1998; Hackbusch 2010), which we will describe after proving the uniqueness of solutions.

For \( a, b, c \in \mathbb{R} \), the Hermitian matrix

\[ M = \begin{pmatrix} aI & ibR \\ ibR & cI \end{pmatrix} \]  

has eigenvalues

\[ \lambda_1 = \lambda_2 = \frac{1}{2} \left( a + c - \sqrt{(a-c)^2 + 4b^2} \right), \]  

\[ \lambda_3 = \lambda_4 = \frac{1}{2} \left( a + c + \sqrt{(a-c)^2 + 4b^2} \right). \]  

and corresponding eigenvectors

\[ \psi_1 = \{ (2ib, 0, 0, a - c + \sqrt{(a-c)^2 + 4b^2}) \}. \]  

\[ \psi_2 = \{ (0, -2ib, a - c + \sqrt{(a-c)^2 + 4b^2}, 0) \}. \]  

\[ \psi_3 = \{ (a - c + \sqrt{(a-c)^2 + 4b^2}, 0, 0, 2ib) \}. \]  

\[ \psi_4 = \{ (a - c + \sqrt{(a-c)^2 + 4b^2}, 0, 0, 2ib) \}. \]  

In the case of \( A_2 \), \( \lambda_1 = \lambda_2 = 0 \), \( \lambda_3 = \lambda_4 = \omega \mu (k_0^2 + \sigma^2)/k_0^4 > 0 \), and its eigenvectors (after rescaling) are

\[ \psi_1 = \{ (i k_x, 0, 0, \sigma) \}, \quad \psi_2 = \{ (0, -i k_x, \sigma, 0) \}. \]  

\[ \psi_3 = \{ (\sigma, 0, 0, i k_x) \}, \quad \psi_4 = \{ (0, \sigma, -i k_x, 0) \}. \]  

For \( B(u, u) = 0 \) to hold, it must be the case that

\[ \Re[B(u, u)] = \int_{\Omega} A_1 \nabla u \cdot \nabla u + \omega \mu |u|^2 dV = 0. \]  

Since \( A_2 \nabla u \cdot \nabla u \geq 0 \), this implies that \( u_2 = 0 \). So \( \nabla u = (\partial_x u_1, \partial_y u_1, 0, 0) \). The only way that \( A_2 \nabla u \cdot \nabla u = 0 \) is if \( \nabla u \in \text{span} (\psi_1, \psi_2) \), but \( \sigma > 0 \) (throughout the domain), so this implies that \( \nabla u = 0 \). Finally, the Dirichlet boundary conditions force \( u = 0 \). Therefore, our variational problems can have at most one solution.

To see that they have precisely one solution, we will make an argument based on a Gårding inequality. It is clear that multiplying the eqs (17), (19) and (21) by a non-zero constant \( \alpha - i \beta \) has no effect on whether or not these problems are well posed, and doing so will make the analysis cleaner. We define \( \tilde{B}(v, w) = (\alpha - i \beta) B(v, w) \) and \( \mathcal{F}(w) = (\alpha - i \beta) F(w) \). Recalling our (physically realistic) assumption that \( \lambda^{-1} \neq 0 \), it is clear that both the sesquilinear form \( \tilde{B} \) and the functional \( \mathcal{F} \) are ‘bounded’: there are constants \( L, M > 0 \) such that \( |\mathcal{F}(v)| \leq L \|v\|_1 \) and \( |\tilde{B}(v, w)| \leq M \|v\|_1 \|w\|_1 \) for all \( v, w \in \mathcal{H} \). We will argue that \( \tilde{B} \) satisfies a Gårding inequality as well: there are constants \( m_0, m_1 > 0 \) for which

\[ \Re[B(v, v)] + m_0 \|v\|_0^2 \geq m_1 \|v\|_1^2 \]  

for all \( v \in \mathcal{H} \).  

Here, and below, \( \|v\|_0^2 = \int_\Omega |v|^2 dV \) is the \( L^2 \)-norm (squared). When \( \tilde{B} \) satisfies a Gårding inequality, existence of a solution is equivalent to uniqueness of a solution—and we have already proven the latter of these. Therefore, establishing (B8) will prove that our variational problems are well posed.

For \( \alpha, \beta \in \mathbb{R} \), we have \( \tilde{A} = (\alpha - i \beta) A = \tilde{A}_1 + i \tilde{A}_2 \), where

\[ \tilde{\lambda}_1 = \left( \frac{1}{k_0^4 + (\omega \mu \sigma)^2} \right) \begin{pmatrix} (ak_0^2 \sigma + \beta \omega \mu \sigma^2) I & i (ak_0^2 + \beta k_0 \omega \mu \sigma) R \\ i (ak_0^2 + \beta k_0 \omega \mu \sigma) R & (\beta k_0^2 \omega \mu + \alpha (\omega \mu \sigma^2) \sigma I) \end{pmatrix}; \]  

\[ \tilde{\lambda}_2 = \left( \frac{1}{k_0^4 + (\omega \mu \sigma)^2} \right) \begin{pmatrix} (\omega \mu \sigma^2 - \beta k_0^2 \sigma) I & i (ak_0 \omega \mu \sigma - \beta k_0^2) R \\ i (ak_0 \omega \mu \sigma - \beta k_0^2) R & (ak_0^2 \omega \mu \sigma + \beta (\omega \mu \sigma^2) I) \end{pmatrix}. \]  

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It holds that
\[ \Re[\tilde{B}(v, v)] = \int_{\Omega} \tilde{A}_1 \nabla v : \nabla v + a \hat{\sigma} |v_1|^2 + \beta \omega |v_2|^2 \, dV, \]
\[ \Im[\tilde{B}(v, v)] = \int_{\Omega} \tilde{A}_2 \nabla v : \nabla v + a \omega |v_2|^2 - \beta \hat{\sigma} |v_1|^2 \, dV. \]

We now argue that \( \alpha \) and \( \beta \) may be chosen so that \( \tilde{A}_2 \) is uniformly positive definite on \( \Omega \). Referring back to (B2)–(B3), it holds that
\[ \lambda_1 = \lambda_2 = \frac{X - \sqrt{X^2 - 4Y}}{2}. \]
where \( X = a + c \) and \( Y = ac - b^2 \); it is apparent that \( X^2 \geq 4Y \). In any (bounded) rectangle \( Q = [X_1, X_2] \times [Y_1, Y_2] \) in the domain of \( f(X, Y) = (X - \sqrt{X^2 - 4Y})/2 \), with \( X_1, Y_1 > 0 \), the minimum value of \( f \) on \( Q \) is \( f(X_1, Y_2) > 0 \). For \( \tilde{A}_2 \) we have
\[ X = \frac{(a \omega \mu (k_x^2 + \hat{\sigma}^2) - \beta (k_x^2 - (\omega \mu)^2) \hat{\sigma})}{k_x^4 + (\omega \mu \hat{\sigma})^2}, \]
\[ Y = \frac{\beta (a \omega \mu \hat{\sigma} - \beta k_x^2)}{k_x^4 + (\omega \mu \hat{\sigma})^2}, \]
and it is clear that we can fix \( \alpha, \beta > 0 \) such that \( X \) and \( Y \) are contained in such a rectangle \( Q \) for all points in \( \Omega \). If we set \( \tilde{m}_1 = f(X_1, Y_2) \), it holds that \( \lambda_1 = \lambda_2 \geq \tilde{m}_1 > 0 \), so \( \tilde{A}_2 \) is uniformly positive definite on \( \Omega \). Furthermore,
\[ \Im[\tilde{B}(v, v)] + \beta \hat{\sigma}_{\text{max}} \|v\|_0^2 \geq \tilde{m}_1 \|v\|_1^2 \geq \frac{\tilde{m}_1}{C} \|v\|_1^2, \]
where \( \hat{\sigma}_{\text{max}} \) is the largest value that \( \hat{\sigma} \) attains on \( \Omega \) and \( C \) is the optimal constant in the norm equivalence \( |v| \leq \|v\|_1 \leq C |v|_1 \). Taking \( m_0 = \beta \hat{\sigma}_{\text{max}} \) and \( m_1 = \tilde{m}_1/C \) establishes the Gårding inequality (B8), which completes the proof.