Characterization of an Adaptive Refinement Algorithm for a Meshless Eigenvalue Solver Based on Radial Basis Functions

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Abstract—A meshless method based on a radial basis collocation approach is presented to calculate eigenvalues for the second-order wave equation. Instead of an explicit mesh topology only a node distribution is required to calculate electric fields, thus facilitating dynamic alteration of the discretization of an electromagnetic problem. An algorithm is presented that automatically adapts an initially very coarse discretization by adding points where higher accuracy is required by the physics of the problem. The algorithm is applied to a cylindrical cavity resonator and the rate of convergence is compared to uniform refinements with the radial basis method and to a regular grid-based finite-difference approach.

I. INTRODUCTION

In computational electromagnetics, commonly used domain-discretization solvers are based on finite-differences in time-domain (FDTD), or the finite elements method (FEM) in frequency-domain. FDTD is a second-order algorithm, i.e. the errors decrease at most with the the square of the grid spacing. FEM is usually used with low-order basis functions of polynomial degree 1 or 2. The simulation error for a structured mesh is then comparable to FDTD for similar discretizations. Meshless methods based on a radial basis collocation approach are an emerging class of numerical schemes with advantages in accuracy over low order methods such as FEM/FDTD, but sharing the ease of implementation and flexibility of these methods. No explicit mesh configuration is required to solve physical problems with a meshless method. Instead, a set of nodes is used in a collocation approach, thus avoiding the computational overhead of dealing with mesh elements. This advantage is especially attractive for adaptive node refinement. Due to the unstructured nature of the interpolation algorithm, nodes can be placed in an arbitrary manner, i.e. complex and multi-scale geometries can be modeled accurately by an appropriate node distribution.

The approach with radial basis functions (RBFs) is sometimes referred to as the Radial Point Interpolation Method (RPIM) [1] or the Kansa method [2]. The collocation approach leads to a true meshless method where no volume integration is required, in contrast to a Galerkin approach [3]. Here, only the position of the collocation nodes is required. It has been estimated [4], that this type of methods can offer convergence at much higher rates than FDTD or low-order finite elements (FEM). This means that, in order to solve problems in electromagnetics, much coarser discretizations can be used to achieve the required accuracy, which results either in a shorter computation time or in an increase of the accuracy of a simulation at the same cost.

The goal of our research is to develop a versatile full-wave solver for 3D problems. Steps in that direction were the presentation of a RBF collocation time-domain method in two dimensions [5], as well as the development of an alternate direction implicit (ADI) approach for overcoming the CFL limit [6] and an implementation in three dimensions [7] by Yu and Chen.

This publication is structured as follows. Sec. II summarizes the interpolation method with radial basis functions. The eigenvalue solver for the Helmholtz equation is presented in Sec. III. In the subsequent Sec. IV, the adaptation algorithm is introduced. A numerical experiment illustrates the abilities of the algorithm in Sec. V and the paper is concluded in Sec. VI.

II. RADIAL BASIS FUNCTIONS

RBFs are an elaborate type of basis functions for highly accurate interpolation of scattered data. This kind of basis function is becoming more popular in many fields of science such as statistics and neural networks. They are also an interesting choice of basis functions to solve partial differential equations. The interpolation scheme is explained in detail in [1]. Hence, only a short summary is given here. A field component \( u(x) \) at position \( x \) in a computational domain containing \( N \) nodes is interpolated as

\[
\langle u(x) \rangle = \sum_{n=1}^{N} a_n r_n(x) = r(x)^T a,
\]

with the radial basis functions

\[
r_n(x) = \exp\left(-\alpha^2 \frac{|x_n - x|^2}{d_c^2}\right)
\]

of Gaussian type. The shape parameter \( \alpha \) is used to control the flatness of the basis function (Fig. 1). The normalization
factor \( d_n \) denotes the average node distance inside the computational domain. Here, we interpolate values in 2D with \( x = (x, y) \).

The interpolation parameter vector \( \mathbf{a} \) is calculated in a preprocessing step as follows. A matrix is set up to interpolate the field values in the nodes \( \mathbf{U}_0 = [u(x_1), u(x_2), \ldots, u(x_N)]^T \).

This system can be written in the form

\[
\begin{pmatrix}
   u(x_1) \\
   \vdots \\
   u(x_N)
\end{pmatrix} =
\begin{pmatrix}
   r_1(x_1) & \cdots & r_1(x_N) \\
   \vdots & \ddots & \vdots \\
   r_N(x_1) & \cdots & r_N(x_N)
\end{pmatrix}
\begin{pmatrix}
   a_1 \\
   \vdots \\
   a_N
\end{pmatrix},
\]

which we denote in compact notation as

\[
\mathbf{U}_s = \mathbf{R}_0 \mathbf{a}.
\]

When the interpolation matrix \( \mathbf{R}_0 \) is non-singular, the interpolation parameter \( \mathbf{a} \) can be calculated as

\[
\mathbf{a} = \mathbf{R}_0^{-1} \mathbf{U}_s.
\]

\( \mathbf{R}_0 \) can become singular when two nodes coincide, or when the basis functions are too flat, leading to a unitary matrix. The shape functions \( \Psi(x) \) and their derivatives are subsequently obtained as

\[
\begin{align}
\langle u(x) \rangle &= r^T(x) \mathbf{a} = r^T(x) \mathbf{R}_0^{-1} \mathbf{U}_s \\
\Psi(x) &= \sum_{j=1}^{N} \Psi_j(x) u(x_j), \\
\langle \partial_\kappa u(x) \rangle &= \partial_\kappa r^T(x) \mathbf{a} = \partial_\kappa r^T(x) \mathbf{R}_0^{-1} \mathbf{U}_s \\
\partial_\kappa \Psi(x) &= \sum_{j=1}^{N} \partial_\kappa \Psi_j(x) u(x_j), \\
\langle \partial_\kappa^2 u(x) \rangle &= \partial_\kappa^2 r^T(x) \mathbf{a} = \partial_\kappa^2 r^T(x) \mathbf{R}_0^{-1} \mathbf{U}_s \\
\partial_\kappa^2 \Psi(x) &= \sum_{j=1}^{N} \partial_\kappa^2 \Psi_j(x) u(x_j)
\end{align}
\]

with \( \partial_\kappa \) the spatial derivative in \( \kappa = x, y \) direction. The shape function \( \Psi(x) = [\Psi_1(x), \ldots, \Psi_N(x)] \) fulfills the delta property

\[
\Psi_j(x_i) = \begin{cases} 
0, & i \neq j \\
1, & i = j
\end{cases},
\]

i.e. the values at the nodes are fitted exactly.

The choice of the shape parameter \( \alpha_\kappa \) is a crucial step in order to achieve accurate results. Large values of \( \alpha_\kappa \) lead to low accuracy. Increasingly small values of \( \alpha_\kappa \) on the other hand lead to flatter basis functions and increased accuracy [8], but also to an increasingly poorly conditioned interpolation matrix \( \mathbf{R}_0 \) [9]. At one point, \( \mathbf{R}_0 \) becomes ill-conditioned, leading to a failure of the numerical inversion for the interpolation matrix. This results in degenerate interpolation problems for the shape functions. Here, the "leave-one-out-cross-validation" algorithm (LOOCV) [10] is applied to find an optimized shape parameter for a given node distribution, i.e. the value of the shape parameter \( \alpha_\kappa \) where the numerical inversion leads to a maximized interpolation accuracy. The cost function \( CF \) of this optimization is based on the pseudo-inverse [11] \( \text{pinv}(\cdot) \) of the interpolation matrix:

\[
CF(\alpha_\kappa) = \max_k |e_k(\alpha_\kappa)|
\]

\[
e_k(\alpha_\kappa) = \sum_{j=1}^{N} \text{pinv}(\mathbf{R}_0)_{k,j} f_j^t(x_j)
\]

where \( \mathbf{R}_0 \) depends on the global shape parameter \( \alpha_\kappa \). The test function \( f_j^t(x) \) is a smooth function, defined over the whole computational domain. In this case, we chose the two-dimensional sinc-function

\[
f_j^t(x, y) = \text{sinc}(x, y) = \begin{cases} 
\frac{\sin(\pi x) \sin(\pi y)}{\pi x \pi y}, & x, y > 0 \\
1, & x, y = 0
\end{cases}
\]

The optimization algorithm is used to find the minimum of the cost function \( CF(\alpha_\kappa) \). The relation between the condition number and the cost error function for an exemplary node distribution is shown in Fig. 2. In this case, the algorithm yields a value for the shape parameter of \( \alpha_\kappa = 0.1 \).

### III. EIGENVALUE SOLVER

In [12], an eigenvalue solver for the Helmholtz problem, based on compactly supported basis functions is presented.

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**Fig. 1:** Gaussian radial basis functions are used to interpolate the field value at \( x \), based on the field values at the surrounding node locations \( x_n \). The shape parameter \( \alpha_\kappa \) is used to control the flatness of the basis functions.

**Fig. 2:** Relation between condition number of interpolation matrix and the cost error function to be minimized in LOOCV algorithm for different shape parameters \( \alpha_\kappa \).
There the eigenmodes are solved for the interpolation coefficient vector \( \mathbf{a} \) (1). A different approach is taken here, where we compute the shape functions \( \Psi(x) \) in a preprocessing step and solve for the eigenmodes, i.e. the resonance frequency and field distribution of the electric field at these frequencies. We consider a 2D transverse magnetic (TM) case, where the electric field points perpendicular to the computational domain \( E_x(x, y) \). The discretized eigenvalue problem can be expressed as:

\[
\mathcal{L}^h \mathbf{E}_z = \lambda \mathbf{E}_z \quad \text{in } \Omega \tag{11}
\]

\[
\mathbf{E}_z = 0 \quad \text{on } \partial \Omega. \tag{12}
\]

with the wave equation inside the domain and a Dirichlet condition for perfect electric conductors (PEC) on the boundary. \( \mathbf{E}_z \) is a vector with a length equal to the number of nodes and \( \lambda = \omega^2/c_0^2 \) are the eigenvalues. \( \mathcal{L}^h \) contains the Laplace operator, discretized with the RBF interpolation shape functions (6c):

\[
\mathcal{L}^h \eta = \partial_x^2 \Psi_j(x_i) + \partial_y^2 \Psi_j(x_i). \tag{13}
\]

The eigenproblem is solved with the numerical full matrix library ARPACK [13], and yields the eigenpairs \( \{ \mathbf{E}_z^i, \lambda^i \} \) with \( i = 1, \ldots, N \).

IV. ADAPTIVE ALGORITHM

The refinement algorithm in [14] for frequency-domain source problems is adapted to the present eigenvalue problem. A residual error is calculated on a set of test nodes \( y_j \) (Fig. 3), generated though a Delaunay tessellation of the initial set of collocation nodes. The test nodes are placed on the edge centers of the Delaunay cells, in between two existing collocation nodes. The residual error function is defined as

\[
\eta(y_j) = \left\{ \begin{array}{ll}
\nu_1 d^2_y |\lambda^i \mathbf{E}_z^i(y_j) - \Delta \mathbf{E}_z(y_j)| & \text{(internal node)} \\
+ \nu_2 d^2_y |\mathbf{E}_z^i(y_j)| & \text{(boundary node)}
\end{array} \right.
\]

for a predefined eigenpair \( \{ \mathbf{E}_z^i, \lambda^i \} \). The weighting factors \( \nu_1, \nu_2 \) are used to set the importance of internal and boundary nodes. Numerical experiments showed that \( \nu_1 = 1, \nu_2 = 1000 \) leads to a good balance between the internal and boundary nodes. The residual error diminishes for very dense node distributions where the local Delaunay edge length \( d_{xy} \) becomes very small.

Test node locations \( y_j \), where the residual error is larger than a predefined threshold

\[
\eta(y_j) \geq \beta \max_n \eta(y_n), \beta \in (0, 1) \tag{15}
\]

are subsequently added to the set of collocation nodes. A value of \( \beta = 0.2 \) proved to be a good value, i.e. all nodes with an error larger than 20\% of the maximum error are added to the original set. This value is a trade-off between making the simulation less accurate by adding unnecessary nodes, and adding too few nodes, making it impossible to properly refine certain regions of the computational domain. In the next iteration step, the LOOCV algorithm is applied on the new collocation nodes to obtain an optimized shape parameter \( \alpha_e \), and the eigenvalue problem is solved again. This procedure is repeated until a required accuracy is achieved.

V. NUMERICAL EXPERIMENT

We calculate the resonance frequencies and eigenmodes of a cylindrical unit disc with perfectly conducting boundary conditions. The convergence rate is investigated. The RBF methods is initiated with a radially symmetric node distribution (Fig. 4). The proposed refinement algorithm is compared with uniform refinement, where in each step, the node distance is uniformly reduced, thus increasing the number of degrees of freedom. These approaches are compared to a uniformly refined second-order finite-difference method on a regular rectangular grid.

Fig. 5 shows the node distribution after six iterations for a TM\( \omega \) mode. The adaptive algorithm adds only nodes in the locations where necessary. Remarkably in the region near boundaries, there is a significantly higher node density than in other regions. This conforms well with the findings of [15], where it was numerically shown that higher accuracy is achieved when additional nodes are placed near the boundary.

The accuracy is studied for the relative eigenvalue error

\[
\xi_{rel} = \frac{|\lambda^i - \omega_{th}^2/c_0^2|}{\omega_{th}^2/c_0^2} \tag{16}
\]

where \( \lambda_i \) is the obtained numerical eigenvalue, and \( \omega_{th} \) is the theoretical analytical resonance frequency calculated as [16]

\[
\omega_{th} = \frac{\omega_{nm}}{\rho} \tag{17}
\]
used finite-differences or low-order finite element methods. For quite coarse discretizations, highly precise simulation results can be achieved. The node refinement algorithm automatically places collocation nodes at locations where the contribution is relevant. Thus, a node distribution refined with the presented method increases the convergence rate. This refinement algorithm could be expanded using other error indicators, such as the information of the gradients, to further increase the efficiency of the adaptive refinement.

Meshless methods are a promising approach for solving electromagnetic problems due to the increased accuracy, ability to modeling complex geometries and the ease of handling a simple node distribution.

VI. CONCLUSION

A meshless eigenvalue solver with adaptive refinement for a radial basis functions collocation method has been presented. In combination with an optimized shape parameter, this method yields excellent accuracy, superior to the commonly

REFERENCES