Low-Complexity Direct and Iterative Volume Integral Equation Solvers With a Minimal-Rank $H^2$-Representation for Large-Scale Three-Dimensional Electrodynamic Analysis

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Abstract—Low-complexity iterative and direct solvers are developed for the volume integral equation (VIE) based general large-scale electrodynamic analysis. The dense VIE system matrix is first represented by a new cluster-based multilevel low-rank representation. In this representation, all the admissible blocks associated with a single cluster are grouped together and represented by a single low-rank block, whose rank is minimized based on prescribed accuracy. From such an initial representation, an efficient algorithm is developed to generate a minimal-rank $H^2$-matrix representation. This representation facilitates faster computation, and ensures the same minimal rank’s growth rate with electrical size as evaluated from singular value decomposition. For the minimal-rank representation whose rank grows with electrical size linearly for a prescribed accuracy, we develop linear-complexity $H^2$-matrix-based storage and matrix-vector multiplication, and thereby an $O(N)$ iterative VIE solver. Moreover, we develop an $O(N\log N)$ matrix inversion, and hence a fast $O(N\log N)$ direct VIE solver for large-scale electrodynamic analysis. Both theoretical analysis and numerical simulations of large-scale one-, two-, and three-dimensional structures on a single-core CPU, resulting in millions of unknowns, have demonstrated the low complexity and superior performance of the proposed VIE electrodynamic solvers.

Index Terms—Electrodynamic, electromagnetic modeling, fast direct solvers, fast solvers, linear complexity solvers, radiation, scattering, three-dimensional (3-D) structures, volume integral equations (VIE).

I. INTRODUCTION

THE volume integral equation (VIE) based methods [1]–[6] offer great flexibility in modeling both complicated geometries and inhomogeneous materials in open-region settings. From a computational perspective, integral equation (IE) methods lead to dense matrices. The size of these dense matrices, for volume based analysis, increases cubically with the size of the objects under study. Therefore, the advantages of the VIE-based analyses can be fully accentuated only if they can be performed with low computational complexity.

Existing fast VIE solvers for solving large-scale electrodynamic problems are, in general, iterative solvers. These include methods, such as fast Fourier transform [7]–[13], low-rank compression [14], [15], fast multipole method (FMM) [16]–[20], etc. The memory requirements in such methods scale at best as $O(N)$, whereas a single matrix-vector multiplication can cost as small as $O(N\log N)$ in time complexity, where $N$ is matrix size. The overall complexity for such iterative solvers is $O(N_{\text{rhs}}N_{\text{id}}N\log N)$, where $N_{\text{rhs}}$ is the number of right-hand sides, and $N_{\text{id}}$ is the number of iterations. When the number of right-hand sides under analysis and/or the number of iterations are large, iterative solvers become inefficient since an entire iteration procedure has to be repeated for each right-hand side.

There have been significant contributions in fast direct solvers [23]–[41], for the analysis of problems ranging from circuits to scattering problems. When dealing with ill-conditioned numerical problems like those arising from multiscaled problems, direct solvers have a potential to overcome the difficulty encountered by iterative solvers such as slow convergence or even failure in convergence. For VIE analysis, although $O(N)$ direct VIE solvers have been developed for full-wave general three-dimensional (3-D) circuit analysis [40], no $O(N^{\log^\alpha N})(\alpha \geq 0)$ fast solvers have yet been made possible for electronically large analysis. The main contribution of this paper is such a fast direct VIE solver whose inversion complexity is $O(N\log N)$ based on a minimal-rank $H^2$-representation of VIE operators whose rank has a linear growth rate with electrical size, in addition to a fast iterative VIE solver whose matrix-vector multiplication complexity is $O(N)$.

To achieve these low complexities, one of the key challenges is to compactly represent the dense VIE system matrix into a reduced set of parameters, despite the large and electrical-size-dependent rank in the off-diagonal blocks. This is fulfilled in this work by finding a minimal-rank representation to approximate the VIE operator based on prescribed accuracy, which does not separate sources from observers. The minimal rank required for representing the VIE operator will be analyzed in Section II. Such a minimal-rank representation can be obtained from singular value decomposition (SVD). However, a
brute-force SVD is computationally expensive. We, hence, develop an efficient algorithm to represent the original dense system matrix using its minimal rank required by accuracy, while avoiding the huge computational cost of SVD. This algorithm will be detailed in the following Section III. In Section IV, we present proposed fast VIE solvers for electromagnetic analysis, enabled by the proposed minimal rank as well as nested representation. In Section V, numerical results are presented to demonstrate the performance of the proposed iterative and direct solvers for arbitrary electromagnetic analysis. Section VI relates to our conclusions. This paper is a significant expansion of our conference paper [53] from theory, algorithm, and numerical experiments perspectives. The algorithms developed in this paper for obtaining a minimal-rank $\mathcal{H}^2$-representation of complex-valued dense matrices are purely algebraic and kernel independent. In addition to VIE, they can also be applied to other IE operators.

It is worth mentioning that the algorithm developed in this paper for obtaining a minimal-rank $\mathcal{H}^2$-matrix is very different from that in [34]. In [34], because the problem being considered is a circuit extraction problem whose electric size is small, an interpolation-based $\mathcal{H}^2$-representation is first obtained, which is then converted to a minimal-rank $\mathcal{H}^2$-matrix. Hence, the algorithm developed therein is to convert an initial $\mathcal{H}^2$-matrix whose rank is not minimal to a new $\mathcal{H}^2$ matrix, whose rank is minimal for a prescribed accuracy. In this paper, the interpolation-based $\mathcal{H}^2$-representation cannot be used since it upfront would produce a full-rank representation when handling electrically large Green’s function. Therefore, our algorithm in this paper for obtaining a minimal-rank $\mathcal{H}^2$ for an electrically large kernel is very different from the algorithm in [34], the details of which can be found from Section III. In addition, all the algorithms developed in this paper are for complex-valued numerical systems, unlike the real-valued system concerned in an electrically small analysis. Our previous direct-solver work reported in [29]–[32], [40] are all based on an interpolation-based method to obtain an $\mathcal{H}^2$-matrix, which is not amenable for handling the rank’s growth with electrical size in electrically large analysis. Therefore, the problem studied in this paper has not been addressed by our previous work.

II. ON THE VIE OPERATOR AND ITS RANK

A. VIE Formulation for Wave-Based Analysis

Consider an arbitrarily shaped 3-D inhomogeneous dielectric body of complex permittivity $\epsilon(r')$ occupying volume $V$, which is exposed to an incident field $E^i(r)$.

The scattered field due to the equivalent volume polarization current $J$ contributes to the total field at any point $r$ in the sense as expressed in the form of the following VIE:

$$ E'(r) = \frac{D(r)}{\epsilon(r)} - \int_V \left[ \mu_0 \omega^2 \kappa(r') D(r') + \nabla \left( \nabla' \cdot \left( \kappa(r') \frac{D(r')}{\epsilon_0} \right) \right) \right] g(r, r') dv' \quad (1) $$

where $g(r, r') = e^{-j k_0 |r-r'|/4\pi |r-r'|}$, $\omega$ being the angular frequency, $\kappa$ the contrast ratio defined as $(\epsilon(r) - \epsilon_0)/\epsilon(r)$, $D(r')$ the electric flux density, whereas $k_0$ is the free space wave number.

Tetrahedral-based discretization is used to model the arbitrarily shaped 3-D inhomogeneous dielectric scattering body and divergence conforming Schaubert Wilson Glisson (SWG) basis functions are used [1] to expand the unknown electric flux density $D(r')$. Each of the SWG basis function is defined for a face of a tetrahedron.

By expanding the unknown electric flux density $D(r')$ in terms of SWG basis functions $D_n(r')$ each with a coefficient $D_n$, and then testing the resulting equation using Galerkin method with $D_m(r)$, we obtain the following linear system of VIE:

$$ SD = E \quad (2) $$

where

$$ E_m = \int_{V_m} \tilde{E}^i \cdot \tilde{D}_m(\tilde{r}) dv $$

$$ S_{mn} = \int_{V_m} \frac{D_m(\tilde{r})}{\epsilon_n(\tilde{r})} \cdot \tilde{D}_n(\tilde{r}) dv $$

$$ - \mu_0 \omega^2 \int_{V_m} \int_{V_n} \kappa_n(r') \tilde{D}_m(\tilde{r}) \cdot \tilde{D}_n(\tilde{r}) g(r', r') dv' dv + \frac{1}{\epsilon_0} \left( \int_{V_m} \int_{V_n} (\tilde{D}_m(\tilde{r}) \cdot \hat{n}) (\nabla' \cdot \tilde{D}_n(\tilde{r}')) g(r', r') dv' ds \right. $$

$$ + \left. \int_{V_m} \int_{S_n} (\nabla' \cdot \tilde{D}_n(\tilde{r})) (\nabla' \kappa_n(\tilde{r}')) g(r', r') ds' dv' \right) $$

$$ - \int_{V_m} \int_{S_n} (\nabla' \cdot \tilde{D}_m(\tilde{r})) (\nabla' \kappa(\tilde{r})) g(r', r') ds' dv' $$

$$ = \int_{V_m} \int_{S_n} (\nabla' \cdot \tilde{D}_n(\tilde{r})) (\nabla' \kappa(\tilde{r})) g(r', r') ds' dv' \quad (3) $$

As evident, each system matrix entry involves all four possible combinations of volume and surface integrals with different terms for different observation and source locations.

B. Rank of the Electrodynamic VIE Operator

Unlike static problems, the rank of an electrodynamic IE kernel increases with electrical size for achieving a prescribed accuracy. Therefore, a fast solver built upon the low-rank property would have a higher computational complexity for solving electrically large problems as compared to electrically small problems, if no advanced algorithms are developed to effectively manage the rank’s growth with electrical size. The true indicator of the rank’s growth is SVD, since its resultant representation constitutes a minimal-rank representation of a matrix for any prescribed accuracy. The SVD does not separate
sources from observers in approximating Green’s function, and it finds a minimal-rank representation of the IE kernel as a whole. The SVD is computationally $O(N^3)$, and hence not practically feasible for studying the rank of electrically large IE operators. In view of the pivotal importance of this subject, a theoretical study has been carried out on the rank’s growth with electrical size in IEs [50]. A closed-form analytical expression of the rank of the coupling Green’s function is derived, which has the same scaling as that depicted by SVD-based rank revealing. The findings on the rank-study are summarized as follows.

1) The rank ($k$) of the off-diagonal block, irrespective of the electrical size, is far less than the size of the block, thus the off-diagonal block has a low rank representation, i.e., $k \ll N$.

2) For static and 1-D configurations of sources and observers, the rank required by a prescribed accuracy remains constant irrespective of the problem size.

3) For 2-D and 3-D configurations, the rank varies as square root of logarithm and linearly with the electrical size, respectively.

To numerically verify the findings related to the rank in the VIE setting concerned in this paper, in Fig. 1, we plot the number of singular values and hence the rank required to maintain an accuracy of $10^{-5}$ in a 1-D type VIE configuration. The size of the two dielectric rods is kept to be 1 m geometrically separated by a distance of 2 m, whereas the frequency (hence, discretization also scales accordingly) sweeps to give the electric sizes as plotted on the horizontal axis. We can observe a constant rank throughout even if the electric size grows to as large as 100 wavelengths.

In Fig. 2, we plot the rank required to maintain an accuracy of $\epsilon$ versus electrical size in a 2-D VIE configuration, as shown in Fig. 2(a). Two dielectric slabs of thickness 0.1 m, side length 1 m, and dielectric constant 2.54 are separated by 2 m (center-to-center distance). The frequency (hence, discretization, along each dimension, also scales accordingly) sweeps to give the electric sizes as plotted on the horizontal axes. To test the rank at larger electrical sizes, a mesh density of two points per wavelength is used. The rank of the matrix formed between the two slabs is obtained from adaptive cross approximation (ACA) followed by a reduced SVD with accuracy parameter $\epsilon$. The $\epsilon$ is chosen to be $10^{-4}$ and $10^{-8}$, respectively. From Fig. 2, we can observe that the rank’s growth rate is smaller than linear with electrical size. We also use a mesh density of ten points per wavelength, and test the rank using the same numerical procedure. The rank’s growth rate remains the same as can be seen from Fig. 2, which agrees with the findings of [50].

In Fig. 3, we plot the rank versus electrical size in a 3-D VIE configuration shown in Fig. 3(a). Again, a mesh density of two points per wavelength is used. The rank of the matrix formed between the two cubes is obtained from ACA followed by a reduced SVD with accuracy parameter $\epsilon$. The $\epsilon$ is chosen to be $10^{-4}$, $10^{-8}$, and $10^{-12}$, respectively. From Fig. 3, it can be seen that for a lower order accuracy setting like $10^{-4}$, the rank’s growth rate is higher than linear. However, it has not saturated to its asymptotic value yet, which can be seen from the reduced growth rate when the accuracy setting is higher. One has to wait until the rank’s growth rate saturates to obtain its actual growth rate with electrical size, which is also pointed out in [50]. The converged growth rate agrees with the theoretical prediction, which is linear, as can be seen from Fig. 3. To be clearer, when accuracy setting is low, only the plane waves whose wave numbers are the closest to the working wave number are necessary. Since they are distributed on a spherical shell, their number grows quadratically with electrical size. However, when the accuracy setting is higher, the plane waves whose wave numbers are distributed within a certain distance away from the shell need to be incorporated, and this distance is inversely proportional to electrical size for a given accuracy, rendering a linear growth rate of the rank. The rank obtained
using a mesh density of ten points per wavelength is also plotted in Fig. 3. It exhibits the same growth rate with electrical size as that obtained from two points per wavelength. The above numerical results obtained from the VIE operator agree very well with the theoretical findings in [50].

Since the number of unknowns in a VIE-based analysis scales with the electric size in a cubic way, an error-bounded low-rank representation exists for VIE operators, irrespective of the electric size and object shape. Since SVD is computationally expensive, in the following section, we present an efficient algorithm for generating a minimal-rank $H^2$-representation of the VIE dense system matrix, which has the same rank’s growth rate with electrical size as that dictated by SVD.

III. PROPOSED ALGORITHM FOR GENERATING MINIMAL-RANK $H^2$-REPRESENTATION OF THE ELECTRODYNAMIC VIE OPERATOR

Starting from the original VIE operator, we develop a two-stage algorithm to obtain a minimal-rank $H^2$-representation, which paves the way to the low-complexity VIE solvers to be described Section IV. To help readers better understand the proposed two-stage algorithm, it is necessary to first review the cluster tree structure used to model an $H^2$-matrix and the $H^2$-matrix partitioning.

A. Cluster Tree and $H^2$-Matrix Partitioning

A cluster tree captures the hierarchical dependence of the entire unknowns to be solved in a given problem. To build a cluster tree, we recursively split the 3-D computational domain that is composed of the SWG basis functions for $D_n$ into two subdomains till the number of unknowns in each subdomain becomes less than or equal to the leaf size ($n_{\text{min}}$). Typical values of $n_{\text{min}}$ can be as small as 2 to as large as 100. The process results in a cluster tree with each node in the tree called a cluster, as illustrated by the left (right) tree of Fig. 4(a). The root cluster is nothing but the entire unknown set, and the clusters at the bottom leaf level correspond to the subdomains whose unknown number is no greater than leaf size. The leaf size essentially controls the depth of the tree. While doing the splitting operation, special care is taken to adaptively make sure that at each nonleaf tree level, every cluster has two children of similar size. We call this splitting as a balanced splitting. Such a splitting facilitates most efficient computational cases in the arithmetic operations to be performed with the resultant matrix.

From the cluster tree, we partition the original dense VIE matrix into multilevel admissible blocks based on a strong admissibility condition [42]. To explain the process, we can place the cluster tree in parallel with itself as illustrated in Fig. 4(a). We can call the left tree row tree, and the right one column tree, as their interaction forms a matrix. Starting from the root level, we level-by-level check whether a cluster in the row tree (denoted by $t$), and a cluster in the column tree (denoted by $s$) are admissible or not. The two clusters are said to be admissible
if they satisfy the following strong admissibility condition [42]:

$$\max \{ \text{diam}(\Omega_t), \text{diam}(\Omega_s) \} \leq \eta \text{dist}(\Omega_t, \Omega_s)$$  \hspace{1cm} (4)

where $\eta$ is a positive parameter, and $\text{diam}(\cdot)$ and $\text{dist}(\cdot, \cdot)$, respectively, denote the Euclidean diameter of the support of a cluster denoted by $\Omega$, and Euclidean distance between the supports of any two clusters. As apparent from the condition that once clusters $t$ and $s$ are admissible, they ought to be physically apart. If $t$ and $s$ are admissible, they form an admissible block at that tree level, and we do not check their children clusters. Such an admissible block is marked in green in Fig. 4(b), denoted by a green link in Fig. 4(a). If they are not admissible, we proceed to check whether their children clusters satisfy the admissibility condition or not. This procedure continues until we reach the leaf level. At the end, the original dense VIE system matrix is partitioned into multilevel admissible and inadmissible blocks, as illustrated in Fig. 4(b). All the inadmissible blocks are formed at the leaf level between leaf clusters.

B. Stage I: Cluster-Based AB$^T$-Representation

With the $\mathcal{H}^2$ cluster tree and matrix partition built, we first construct a cluster-based nondegenerate hierarchical low-rank representation. This is different from a commonly used block-based low-rank representation like that in an $\mathcal{H}$-matrix [42]. With this new approach, the admissible blocks formed by a single cluster at its tree level are grouped together to be represented by a single low-rank block. Such a representation significantly reduces the storage and time requirements for obtaining a low-rank representation of the original dense system matrix, especially in 3-D settings. This makes the solution of millions of unknowns resulting from the electrically large VIE feasible on a single-core CPU. The accuracy of the low-rank representation of the multiple admissible blocks formed by a single cluster is also better controlled since their weights in the matrix relative to each other are considered, as compared to individually building a low-rank form for each admissible block. After the initial cluster-based low-rank representation is generated, we proceed to the second stage to obtain a minimal-rank $\mathcal{H}^2$-matrix.

For each cluster in the cluster tree, it can form multiple admissible blocks at its tree level, as shown by the links in Fig. 4(a). The number of such admissible blocks is bounded by a constant $C_{sp}$ [46]. The $C_{sp}$ can be as large as hundreds in a 3-D configuration. Hence, it is not efficient to handle each admissible block one by one and generate its low-rank form individually. Instead, we propose to group these admissible blocks together and generate a single low-rank representation. Although physically, these blocks can be scattered in the matrix as disconnected blocks, algorithm wise, we can put them together to form a single block. After generating the low-rank representation for this single block, we can distribute it back to the original location of each admissible block if needed. As a result, our low-rank representation has a one-to-one correspondence with each cluster in an $\mathcal{H}^2$-tree, instead of being individually constructed for each admissible block. This concept is illustrated in Fig. 5.

Consider an arbitrary cluster $t$, and its associated admissible blocks $G^{(t,s_1)}$, $G^{(t,s_2)}$, ..., and $G^{(t,s_p)}$ at its tree level. We group them into a single matrix as the following:

$$G^t = \left[ G^{(t,s_1)}, G^{(t,s_2)}, \ldots, G^{(t,s_p)} \right]$$  \hspace{1cm} (5)

where $p$ is the number of admissible blocks formed by $t$ at its tree level. We then perform an ACA (see [42, p. 69]) on this matrix and obtain a factorized form of

$$G^t \approx \tilde{A}_{\#t \times \#k} \tilde{B}_{\#s \times k} \cdot \tilde{C}_{\#k \times \#s}$$  \hspace{1cm} (6)

based on prescribed accuracy $\epsilon_A$. The computational cost of this step is simply $O(k^2(\#t + \#s))$, where $\#s = \#s_1 + \#s_2 + \cdots + \#s_p$, and $\#$ denotes the cardinality of a set. This is much smaller than the cost of a brute-force SVD, which scales cubically with block dimension. It is also smaller than a block-by-block ACA procedure, whose cost is $O(k^2(\#t) C_{sp} + k^2 \#s)$. In addition, the storage of $G^t$ is also greatly reduced from $O(k'(\#t) C_{sp} + k' \#s)$ required by a block-based ACA to $O(k'(\#t + \#s))$ units. In addition to computational efficiency, the accuracy of such a low-rank representation is also better controlled since now all admissible blocks are put together and their approximation error is controlled by $\epsilon_A$ as a whole in (6). The resulting rank is also the minimal one required to represent the entire admissible blocks formed by cluster $t$ for $\epsilon_A$ accuracy in the ACA procedure. For example, for an admissible block whose column cluster is very far from its row cluster, to achieve a 1% accuracy in representing itself may require a rank of 40. However, when being put together with other admissible blocks formed by the same cluster, this block may not contribute any additional rank as it is negligible in matrix norm. Hence, it is more efficient to group the admissible blocks together to construct a single low-rank block as the relative weight of each admissible block in the entire matrix is taken into account in this representation.

After ACA, we perform another SVD, obtaining

$$G^t \approx A_{\#t \times k} B_{\#s \times k}^T$$  \hspace{1cm} (7)

where $k$ is the further reduced rank, as compared to $k'$ in (6), based on the required accuracy $\epsilon_{acc}$. This step is performed because the rank $k'$ determined from ACA is not the minimal rank determined by accuracy [33].

Furthermore, because the initial matrix to perform SVD is a factorized low rank form obtained from ACA, the SVD can be performed as a reduced SVD (r-SVD) [42], whose computational cost is much reduced to $O(k^2(\#t + \#s))$ as compared to a brute-force SVD. As shown in [33], adding an additional
step of r-SVD after ACA is effective in further reducing the rank without sacrificing prescribed accuracy.

After the aforementioned procedure, we obtain a cluster-based $\mathbf{AB}^T$ representation of the original VIE system matrix, as illustrated in Fig. 5. The rank of this representation is minimized based on accuracy for each cluster.

C. Stage II: Minimal-Rank $\mathcal{H}^2$-Matrix Construction

With the $\mathbf{AB}^T$ form obtained for each cluster of an $\mathcal{H}^2$-tree, in this section, we show how to construct an $\mathcal{H}^2$-matrix representation out of it. Such a minimal-rank $\mathcal{H}^2$-matrix provides a compact nested structure for more efficient computation.

In an $\mathcal{H}^2$-matrix, the inadmissible blocks keep their original full-matrix representations, while admissible blocks are represented in the following factorized form:

$$
\mathbf{G}_{t,s}^{l,s} = \mathbf{V}^l \mathbf{S}^{l,s} \mathbf{V}^s^T
$$

$$
\mathbf{V}^l \in \mathbb{C}^{n_l \times k_l^t}, \quad \mathbf{S}^{l,s} \in \mathbb{C}^{k_l^t \times k_s}, \quad \mathbf{V}^s \in \mathbb{C}^{n_s \times k_s}
$$

with $\mathbf{V}^l(\mathbf{V}^s)$ called a cluster basis associated with $t$ and $s$, respectively, $\mathbf{S}^{l,s}$ is called a coupling matrix, and $k_s^{l,t}$ is the rank of $\mathbf{V}^l(\mathbf{V}^s)$. The cluster basis $\mathbf{V}^l$ is nested in an $\mathcal{H}^2$-matrix, which satisfies the following relation:

$$
\mathbf{V}^l = \begin{bmatrix} \mathbf{V}^{l_1} & \mathbf{T}^{l_1} \\ \mathbf{V}^{l_2} & \mathbf{T}^{l_2} \end{bmatrix} = \begin{bmatrix} \mathbf{V}^{l_1} & 0 \\ 0 & \mathbf{V}^{l_2} \end{bmatrix} \begin{bmatrix} \mathbf{T}^{l_1} \\ \mathbf{T}^{l_2} \end{bmatrix}
$$

(9)

where $l_1, l_2 \in \text{children}(t)$ are the two children clusters of $t$. $\mathbf{T}^{l_1}$ and $\mathbf{T}^{l_2}$ are called transfer matrices associated with a nonleaf cluster $t$, and they are used to build a relationship between $t$ and its two children.

It can be seen clearly that to build an $\mathcal{H}^2$-representation, we need to find cluster basis $\mathbf{V}$ for each leaf cluster, transfer matrices $\mathbf{T}^{l_1}$ and $\mathbf{T}^{l_2}$ for each nonleaf cluster, as well as coupling matrix $\mathbf{S}$ for each admissible block. In the following, we show how to find them efficiently from the $\mathbf{AB}^T$ form obtained for each cluster at the first stage. This algorithm is a bottom-up tree-traversal procedure, where we first compute the cluster basis for the leaf clusters, then obtain the two transfer matrices of each nonleaf cluster level by level. After the cluster bases are obtained, we compute the coupling matrices.

1) Leaf Clusters: We first generate cluster bases at the leaf level. To build a nested relationship among cluster bases, for each leaf cluster, we require its cluster basis not only represent the admissible block formed by this cluster at the leaf level, but also the admissible blocks involving this cluster at all the other levels. To do so, for each leaf cluster $t$, we build the following Gram matrix:

$$
\mathbf{G}_t^l = \mathbf{G}(\mathbf{A}_t, \mathbf{B}_t^T, \mathbf{B}_t) = \mathbf{A}_t^H(\mathbf{B}_t^T \mathbf{B}_t) \mathbf{A}_t^H + \sum_{j \in \text{ancestors}} \mathbf{A}_{t,j}(\mathbf{B}_j^T \mathbf{B}_j) \mathbf{A}_{t,j}^H
$$

(10)

where $\mathbf{G}_t^l$ represents the low-rank block in $\mathbf{G}$ whose rows correspond to the unknowns in cluster $t$. This block is composed of a single $\mathbf{A}_t \mathbf{B}_t^T$ (7) that captures all the admissible blocks formed by $t$ at $t$’s level, as well as $t$-related rows in the $\mathbf{A}_t \mathbf{B}_j^T$ blocks formed by $t$’s parent clusters at nonleaf levels. Here, $j$ is the index of the parent clusters of $t$ at nonleaf levels. In (10), $\mathbf{B}_j^T \mathbf{B}_j$ is a $k \times k$ matrix, prepared a priori for each cluster and is just referenced to each of the lower level children. $\mathbf{A}_{t,j}$, in the summation term, represents the rows corresponding to cluster $t$ of the bigger matrix $\mathbf{A}_t$ at each ancestral level. $\mathbf{B}_j^T \mathbf{B}_j$ is the complex conjugate of $\mathbf{B}_j$. The $\mathbf{G}_t^l$ shown in (10) is clearly of leaf size $O(n_{\text{min}})$.

Next, we perform an accuracy controlled ($\epsilon_{\text{acc}}$) Schur or SVD decomposition to get

$$
(\mathbf{G}_t^l)^{\text{min}}_{n_{\text{min}} \times n_{\text{min}}} \overset{\epsilon_{\text{acc}}}{=} \mathbf{P} \mathbf{D} \mathbf{P}^H.
$$

The cluster basis for $t$ now can be obtained as

$$
\mathbf{V}^l = \mathbf{P}_{n_{\text{min}},k}
$$

(12)

where $k$ is determined based on $\epsilon_{\text{acc}}$ when truncating (11).

2) Nonleaf Clusters: For clusters at a nonleaf level, if we follow the same procedure as that in the leaf level, the Gram matrix size will become increasingly large when we traverse the tree from bottom to top. Since the cluster basis generated at the leaf level $l = L$ for a leaf cluster has already taken into account upper level blocks related to this leaf cluster, the Gram matrix formed at one level up $l = L - 1$ can be accurately projected onto the cluster bases formed at $l = L$ level. This will yield a small $k \times k$ matrix for which the cost of SVD is trivial. Similarly, the Gram matrix formed at $l = L - 2$ level can be accurately projected onto the cluster bases formed at $l = L - 1$ level. Hence, the nonleaf cluster bases are generated level by level from bottom to top so that the entire computation becomes efficient. This enables performing an SVD on an $O(k)$ matrix at each level $l$.

Consider an arbitrary nonleaf cluster $t$, instead of directly building its Gram matrix, as shown in (10), we project it onto its children’s cluster bases to get a $k \times k$ matrix as the following:

$$
\mathbf{G}_{t,\text{proj}}^l = \mathbf{A}_{t,\text{small}}(\mathbf{B}_t^T \mathbf{B}_t) \mathbf{A}_{t,\text{small}}^H + \sum_{j \in \text{ancestors}} \mathbf{A}_{j,\text{small}}(\mathbf{B}_j^T \mathbf{B}_j) \mathbf{A}_{j,\text{small}}^H
$$

(13)

where

$$
\mathbf{A}_{t,\text{small}} = \begin{bmatrix} \mathbf{V}^{l_1} & 0 \\ 0 & \mathbf{V}^{l_2} \end{bmatrix}^H \mathbf{A}_t
$$

(14)

and each of this multiplication costs $O(k^2(\#t))$ time requiring only $O(k^2)$ storage units. The remaining multiplications in (13) involve three $O(k)$-sized matrices requiring $O(k^3)$ operations only.

We then perform an SVD on $\mathbf{G}_{t,\text{proj}}^l$ to obtain

$$
(\mathbf{G}_{t,\text{proj}}^l)_{k \times k}^{\epsilon_{\text{acc}}} \overset{\epsilon_{\text{acc}}}{=} \mathbf{P} \mathbf{D} \mathbf{P}^H
$$

(15)
the cost of which is $O(k^3)$ only. The two transfer matrices of the nonleaf cluster $t$ now can be obtained as

$$
\begin{bmatrix}
T^{t_1}
\end{bmatrix} = \begin{bmatrix}
P_1 \\
P_2
\end{bmatrix}
$$

(16)

where $P_1$ and $P_2$ are $P$’s block rows corresponding to $t$’s two children clusters $t_1$ and $t_2$ respectively.

3) Formation of Coupling Matrices: After the factorized $AB^T$ form is generated for each cluster, for each admissible block in the $H^2$-matrix, its factorized form is readily known as $G^{(t,s)} = A_{#t \times k}B_{#s \times k}^T$. The $A_{#t \times k}$ is the same as that generated for cluster $t$, and the $B_{#s \times k}$ is simply the $t$-cluster-based $B^T$’s columns corresponding to column cluster $s$.

To obtain the coupling matrix for each admissible block, we utilize the following relationship:

$$
G^{(t,s)} = A_{#t \times k}B_{#s \times k}^T = V^T S V^s T
$$

(17)

and the fact that the proposed cluster bases are unitary. Hence, we have

$$
(S)_{k \times k} = V^T H A B^T V^s
$$

(18)

where $V^s$ is the complex conjugate of $V^s$. The total cost of computing the coupling matrix is again $O(k^2 (#t + #s))$. Since $V$ is nested, at the nonleaf level, the cluster basis is manifested as transfer matrices.

4) Transforming Complex-Valued Cluster Bases to Real-Valued Ones: The nested cluster bases constructed in the proposed algorithm are complex valued. They can be readily converted to real-valued cluster bases following the procedure described in [34, Sec. III.E]. As a result, the new cluster bases satisfy $V^T V = I$ instead of the original $V^H V = I$. This property helps make the subsequent matrix inverse more efficient, since our IE system matrix is symmetric instead of complex-conjugate symmetric. With the cluster bases updated to real-valued ones, the coupling matrices of the admissible blocks are also correspondingly updated, as shown in [34, Sec. III.E].

IV. PROPOSED LOW-COMPLEXITY ITERATIVE AND DIRECT VIE SOLVERS FOR ELECTRICALLY LARGE ANALYSIS

A. Storage and Complexity

The cluster bases are stored at the leaf level. For each nonleaf cluster, we store its two transfer matrices, each of size $k_{var} \times k_{var}$. Coupling matrices of size $k_{var} \times k_{var}$ are stored at corresponding levels to represent the off-diagonal admissible interactions between clusters. Considering the fact that the number of admissible blocks formed by a cluster at each level is bounded by sparsity constant $C_{sp}$, the total memory cost can be evaluated as

$$
\text{Memory Cost} = \sum_{l=0}^{L} O(k_{var}^2 (2^l + C_{sp} 2^l)) + 2C_{sp} O(\eta_{min}) N
$$

(19)

where the $n_{min}$-related term is associated with the storage of inadmissible blocks.

With the proposed minimal-rank $H^2$-representation, the rank ($k_{var}$) scales linearly with electrical size. Hence, for VIE, we have

$$
k_{var} = O(N^\frac{2}{3}).
$$

(20)

Substituting it into (19), we obtain

$$
\text{Memory Cost} = \sum_{l=0}^{L} O(C_{sp} 2^l (\frac{N}{2^l} \frac{2^l}{})
$$

(21)

which is linear regardless of electrical size.

B. Matrix-Vector Multiplication and Its Complexity

Multiplying minimal-rank $H^2$-based $S$ with a vector $x$ comprises of multiplying its inadmissible blocks and admissible blocks with $x$ [21], [42] [48]. In the multiplication with admissible blocks, we can fully take advantage of the $H^2$-tree structure $T_I$ and the nested cluster bases as follows.

1) For Admissible Blocks: We perform the following three steps.

1. Forward transformation: Compute $x^s := (V^s)^T x$ for all clusters $s \in T_I$.

2. Coupling-matrix multiplication: Compute $y^s := \sum_{r \in B^l} S_{l,s} x^s$ for all clusters $t \in T_I$ where $B^l$ contains all the clusters $s$ such that $(t, s)$ is an admissible block.

3. Backward transformation: Compute $y_t := \sum_{l, s \in f(T)} (V^l y^s)$.

2) For Inadmissible Blocks: For these blocks, a full matrix-vector multiplication is performed.

With the proposed minimal-rank $H^2$-representation, the total operation count for a matrix-vector multiplication is given by

$$
\text{MVM Cost} = \sum_{l=0}^{L} O(C_{sp} 2^l k_{var}^2)
$$

(22)

We can see that with a minimal-rank $H^2$-representation, even though the rank grows with electrical size, the complexity of $H^2$-based matrix-vector multiplication is kept linear for general VIE-based electrodynamic analysis.
C. Matrix Inversion and Its Complexity

The $H^2$-represented system matrix $S$ can be written as

$$ S = \begin{bmatrix} S_{11} & S_{12} \\ S_{21} & S_{22} \end{bmatrix} $$

then by Sherman–Morrison–Woodbury formula, the inverse for $S$ given by $S^{-1}$ can be recursively evaluated as

$$ S^{-1} = \left[ S_{11}^{-1} \otimes S_{12}^{-1} \otimes S_{12} \otimes F^{-1} \otimes S_{21} \otimes S_{11}^{-1} - S_{11}^{-1} \otimes S_{12} \otimes F^{-1} \right] $$

$$ -F^{-1} \otimes S_{12} \otimes S_{11}^{-1} \quad F^{-1} $$

where $F = S_{22} \oplus (-S_{21} \otimes S_{11}^{-1} \otimes S_{21})$ and $\oplus, \otimes$ are addition and multiplication defined for the $H^2$-matrix elaborated. The recursive inverse formulation can be realized by the code in Table I. It can be seen that the computation of inverse involves a full-matrix inverse at the leaf level and a number of block matrix–matrix multiplications at the nonleaf levels. The main operation in the inverse algorithm is to perform fast block matrix multiplications based on orthogonal nested cluster basis. For example, since $V^T V = I$ is satisfied for each cluster $s$, an admissible block based matrix multiplication encountered in the inverse procedure can be done based on

$$ V^T S_1 V^T \times V^T S_2 V^T = V^T S_1 (V^{sT} \times V^s) S_2 V^T $$

$$ = V^T S_1 (I) S_2 V^T $$

$$ = V^T S_1 S_2 V^T $$

where only $S_1 S_2$ needs to be computed, the cost of which is $O(k_{var}^3)$. Similarly, all the remaining 6 cases which are encountered out of the possible 27, have been presented in [33]. Each is bounded by $O(k_{var}^3)$ operations. As shown in [30], each of the $C_{sp}^2 l$ number of admissible blocks at level $l$ requires $C_{sp}$ number of block multiplications each costing $O(k_{var}^3)$ operations. So, the total cost to get an $H^2$-matrix based inverse is

**Inversion Cost**

$$ = \sum_{l=0}^{L} (\# \text{ of blocks at level } l) \text{(one block cost)} $$

$$ = \sum_{l=0}^{L} (C_{sp}^2 l) O(C_{sp} k_{var}^3) $$

$$ = \sum_{l=0}^{L} C_{sp}^2 l O(k_{var}^3) \cdot (23) $$

With the rank’s growth with electrical size taken into account, we obtain

$$ \text{Inversion Cost} = \sum_{l=0}^{L} O \left( \left( \frac{N}{2^l} \right)^{3} C_{sp}^2 l \right) $$

$$ = O \left( C_{sp}^2 N \sum_{l=0}^{L} l \right) $$

$$ = O \left( C_{sp}^2 O \left( N \log N \right) \right). \quad (24) $$

Thus, we see that for a VIE, with the proposed minimal-rank representation, the underlying storage and matrix-vector multiplication cost becomes linear while the inversion cost becomes as fast as $O(N \log N)$.

D. Matrix-Vector and Matrix-Matrix Multiplication for Solutions

The solution vector $D$ is then obtained by multiplying the resulting $S^{-1}$ having the same $H^2$-structure as $S$, by $E$. For multiple right-hand sides, we use $H^2$-based matrix–matrix multiplication to obtain final solutions.

V. NUMERICAL RESULTS

The numerical results from the proposed fast solvers are validated with the analytical Mie Series solution in the first multilayered dielectric sphere examples. Next, large-scale 1-D, 2-D, and 3-D dielectric structures, resulting in more than millions of unknowns, are simulated to demonstrate the accuracy controlled performance benefits that can be achieved with the proposed solvers. In all these numerical examples, $\eta = 1$ is used in (4) for building the $H^2$ cluster tree. The computer used has a single Intel Xeon E5-2690 CPU core running at 3 GHz.

A. Analytical Validation for an Eight Layered Dielectric Sphere

The numerical results from our proposed fast solvers are first validated with the analytical Mie Series. An eight layered dielectric sphere of one wavelength (free-space) radius is simulated with a permittivity profile which increases from the outermost to the center of the sphere. The outermost layer has a relative permittivity of 1.5, which increases in steps of 0.5 each layer to the innermost layer value of 5.0. The cross-sectional view of the simulated sphere is shown in Fig. 6(a). The field
of excitation is a normalized $-z$ directed plane wave polarized along the $x-$axis of standard cartesian coordinates. In Fig. 6(b), radar cross section (RCS in dB) of this eight layered sphere is plotted as a function of spherical coordinate polar angle ($\theta$ in degrees) for zero azimuth. It is evident that the numerical results from our proposed solvers (both direct and iterative solvers) show good agreement with the analytical Mie Series solution. The convergence criterion used in the BiCGStab iterative solver is $10^{-3}$.

**B. Large-Scale Dielectric Rod**

The performance of the proposed solvers is first demonstrated on a dielectric rod of relative permittivity 2.54, whose length is placed along $z$-axis, and cross section in the $x-y$ plane. It is centered at the origin. The cross section of the rod is fixed at $\lambda_0/10 \times \lambda_0/10$ whereas its length is increased from $1\lambda_0$ to $8.194\lambda_0$. This results in scaling the number of unknowns $N$ from 164 to as large as 1.31 million, with a mesh density of $\lambda_0/10$. A normalized $-y$ directed and $z-$polarized plane wave is used as excitation. Since only one dimension of the simulated structure is changing, this example essentially represents 1-D problem characteristics [50]. In Fig. 7, it is proven numerically that indeed the accuracy determined rank for all these simulated large-scale rod examples remain constant, irrespective of the electric size. We also plot the total number of iterations required for BiCGStab iterative solver for a convergence criterion of $10^{-3}$. It is clear that only five iterations are required to converge regardless of the electric size of the rod. In Fig. 8, the representation and inversion accuracy is shown as a function of the scaling of number of unknowns. It is clear that the accuracy of the inverse is well-controlled below 0.5% error for all the simulated large-scale problems.

Finally, the scaling of computational resources with number of unknowns is presented in Fig. 9. It is evident from the numerical results that, because of constant representation rank, storage requirements and solution times for iterative and direct solvers scale linearly with the number of unknowns. This proves the validity of (22) and (24). We can see that even for a dense matrix of size 1.31 million unknowns, it only takes about 40 min. to get an accuracy controlled inverse of the VIE system matrix. For such rod-like structures, our proposed fast iterative solution can be obtained in less than 4 min.
C. Large-Scale Dielectric Slab

For 2-D field variation, the performance of the proposed solvers is demonstrated on a dielectric slab of relative permittivity 2.54. The thickness of the slab is fixed at $\lambda_0/10$ whereas its length and width is increased from $8\lambda_0 \times 8\lambda_0$ to $16\lambda_0 \times 16\lambda_0$ and finally to $32\lambda_0 \times 32\lambda_0$. Such a dimension scaling results in the scaling of the number of unknowns $N$ from 89,920 to as large as 1,434,880 that is over 1.43 million, with a mesh density of $\lambda_0/10$. The geometry of the simulated structure is shown in Fig. 10, with the slab placed in the $x-z$ plane. Normalized $-y$ directed and $z-$polarized plane wave is used as excitation. Since two dimensions of the simulated structure are changing, this example represents 2-D problem characteristics [50]. In Fig. 11, it is shown numerically that for maintaining the same level of matrix representation accuracy ($\sim 0.03\%$), the rank for these simulated large-scale slab structures increases in a square-root of logarithmic trend with the electric size. The performance of the iterative solver is summarized in Fig. 12(a). A single matrix-vector multiplication time scales linearly with the number of unknowns as well as the memory consumption. In Fig. 12(b), the direct inverse time is shown as a function of $N$. It is clear that the inverse time scales linearly with the number of unknowns since the rank scales as square-root of log-linear of electrical size. In this figure, one more data point

![Fig. 9](image9.png)  
Fig. 9. Dielectric rod: scaling of computational resources with number of unknowns.

![Fig. 10](image10.png)  
Fig. 10. Simulated dielectric slab.

![Fig. 11](image11.png)  
Fig. 11. Dielectric slab: rank and $H^2$-representation error versus electrical size.

![Fig. 12](image12.png)  
Fig. 12. Solver performance for large-scale slab structures. (a) Memory and MVM time as a function of $N$. (b) Inverse time as a function of $N$. 


TABLE II

| N      | 89 920 | 359 040 | 876 000 | 1 434 880 |
|--------|--------|---------|---------|-----------|
| $C_{sp}$ | 55     | 53      | 57      | 57        |

$C_{sp}$ as a function of $N$ for the dielectric slab

having $N = 876 000$ for a $25\lambda_0$-size slab is added to confirm the complexity. The $C_{sp}$ is given in Table II, which is shown to have saturated to the constant range. We have also assessed the inverse accuracy by evaluating $||I - SS^{-1}||$ for matrix size whose memory cost is feasible on our computer platforms. This error is shown to be $3.9e^{-2}$ and $7.49e^{-2}$ for the $8\lambda_0 \times 8\lambda_0$ ($N = 89 920$), and $16\lambda_0 \times 16\lambda_0$ ($N = 359 040$) slab, respectively.

D. Large-Scale Array of Dielectric Cubes

A large-scale array of dielectric cubes having $\varepsilon_r = 2.54$, illuminated by a plane wave ($\vec{E} = E_0 e^{jkyz}$), is simulated to demonstrate the performance of the proposed fast VIE solvers for pure 3-D field variation. The dimension of each cube is $0.3\lambda_0 \times 0.3\lambda_0 \times 0.3\lambda_0$ and the distance between neighboring cubes is fixed at $0.3\lambda_0$. The mesh density is $0.1\lambda_0$. The number of simulation unknowns $N$ are scaled from $3 024$ to $1 037 232$ million by increasing the array size from $2 \times 2 \times 2$, $4 \times 4 \times 4$, $8 \times 8 \times 8$ to $14 \times 14 \times 14$. The geometry of the simulated structure is shown in Fig. 13. Before presenting the solvers performance, it is worth pointing out that the theoretical bounds presented in Section V inherently assume that the sparsity constant, $C_{sp}$, has saturated for all tree levels. For cubic growth of unknowns for 3-D problems, as presented here, such saturation is attained in the order of millions of unknowns, as shown in Table III. It is thus, important to analyze the performances for iterative and direct solvers as $(\text{Memory or MVM cost})/C_{sp}$ and $(\text{Inverse time})/C_{sp}^2$, respectively. In Fig. 14, we plot the rank for maintaining a matrix representation accuracy ($<0.8\%$). The rank’s growth rate is shown to be no greater than linear with the electric size in this example. The matrix representation error is also shown to be at the same order for the last two unknown numbers. The error for the first two $N$ cases is small because of smaller $N$, and hence the admissible partition has not saturated and many computations are full-matrix based. The performance of the iterative solver follows the same trend set by rank representation and is presented in Fig. 15(a). A single matrix-vector multiplication time as well as storage scales linearly with the number of unknowns. In Fig. 15(b), the direct inverse time divided by sparsity constant square, is plotted to show that indeed the inversion time complexity scales almost as $O(N^2)$, which agrees with our theoretical complexity analysis. The inverse error measured by $||I - SS^{-1}||$ is shown to be $9.03e-3$, $1.73e-2$, and $3.03e-2$, respectively, for the array size of $2 \times 2 \times 2$, $4 \times 4 \times 4$, and $8 \times 8 \times 8$.

E. Comparison Between the Proposed New $H^2$-Construction Algorithm and Existing One

We have also compared the performance of the new cluster-based low-rank algorithm described in Section III with the conventional one that is admissible-block based. Take a slab simulated in Section V-C whose side length is 80 wavelengths as an example, the new algorithm takes 153.77 s and 3.083 GB memory to finish generating the low-rank representation of the whole matrix; and another 18.72 s to obtain a minimal-rank $H^2$ representation, whereas the conventional one costs 790.17 s and 5.162 GB memory in the first stage, and 48.98 s in the second stage. Table IV has a detailed comparison between the two methods in both CPU time and memory consumption. As another example, for the $4 \times 4 \times 4$ cube array simulated in Section V-D, the computational resources cost by the new algorithm are listed in Table V in comparison with those of the old.
algorithm. Again, the new method has a better performance in both CPU time and memory usage.

VI. CONCLUSION

Low-complexity iterative and direct solvers are developed for the VIE-based general large-scale electrodynamic analysis. The large-scale system matrix is represented by an $H^2$-structure. To efficiently obtain a minimal-rank $H^2$-representation, we first generate a rank-minimized $AB^T$ form for each cluster in the $H^2$-tree. Based on such a cluster-based low-rank form, we extract nested cluster bases and coupling matrices to obtain an efficient $H^2$-matrix representation with its rank minimized by accuracy. This algorithm can also be applied to other initial representations of the IE operators such as an FMM-based representation to obtain a minimal-rank $H^2$-matrix.

Analytical expressions of complexities for storage, matrix-vector-multiplication and matrix inversion are derived for general 3-D VIE-based electrodynamic analysis. For rank that grows with electrical size linearly, $O(N)$ iterative and $O(N \log N)$ direct inverse solvers are achieved with the proposed representation of VIE operators. Numerical simulations for large-scale 1-D, 2-D, and 3-D structures, resulting in millions of unknowns, demonstrate the efficiency and complexity of the proposed VIE electrodynamic solvers. The algorithms developed in this work are kernel-independent, and hence applicable to other IE operators as well.

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