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Vol. 7  No. 1  Summer 1992

TABLE OF CONTENTS

"Electromagnetic Scattering From Radially or Axially Inhomogeneous Objects"
by A.A. Kishk and M. Abouzahra ................................................................. 4

"Application of Equivalent Edge Currents to Correct the Backscattered Physical Optics
Field of Flat Plates"
by V. Stein ........................................................................................................ 24

"Application of Parallel Processing To a Surface Patch/Wire Junction EFIE Code"
by L.C. Russell and J.W. Rockway ................................................................. 48

"A Recursive Technique to Avoid Arithmetic Overflow and Underflow When Computing
Slowly Convergent Eigenfunction Type Expansions"
by G.A. Somers and B. A. Munk ................................................................. 67

"H-Oriented' and 'B-Oriented' Methods In a Problem of Nonlinear Magnetostatics: Some
Methodological Remarks"
by A. Bossavit ............................................................................................. 78

Institutional Members .................................................................................. 90

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ELECTROMAGNETIC SCATTERING FROM RADially OR AXIALLY INHOMOGENEOUS OBJECTS

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ABSTRACT

A computer program based on the method of moments approach is developed to compute electromagnetic scattering from axisymmetric objects. The object may consist of N linear isotropic homogeneous regions. These regions may be arranged axially and/or radially with the axis of symmetry. Surface integral equations (SIE) formulation, E-PMCHW, is used to formulate the problem. Other formulations can easily be incorporated in the computer code. Bistatic and monostatic Radar Cross Sections (RCS) for several benchmark geometries are computed. The computed results are verified by comparison with measured and exact calculated results. In some cases the self-consistency method is used to perform the verification. The measured and calculated data presented in this paper are expected to serve as benchmarks for other researchers in the field.¹

1 INTRODUCTION

The study of electromagnetic scattering from composite materials has become of interest to many engineering societies. For example, in biomedicine modeling of human bodies and tissues requires very complex composite objects. With the relative increase in the complexity of the objects of interest, numerical solutions become necessary to study the electromagnetic characteristics of these objects. Various numerical methods can be used to solve such problems

¹ This work was sponsored by the Department of the Air Force. The views expressed are those of the authors and do not reflect the official policy or position of the US Government.
The method of moments has been proven to be efficient in solving Surface Integral Equations (SIE). The SIE formulation is most suitable for objects made of linearly isotropic homogeneous materials. For inhomogeneous objects, other formulations may be used, such as the Volume Integral Equations (VIE) or Partial Differential Equations (PDE) formulations. However, if the inhomogeneity of the material is simple, the use of SIE may be preferred, because the matrix size will be smaller using SIE than the matrices that can be obtained from other formulations.

In this paper, the SIE formulation is used to assess the problem of electromagnetic scattering from bodies of revolution made of multihomogeneous regions. The present formulation is similar to the one given in [1]. The method of moments is used to solve the SIE. The resulting matrix system is a buildup of the basic Z and Y matrix obtained for conducting or dielectric bodies of revolution [7-8]. The computer program that has been developed to compute the bistatic or monostatic Radar Cross Sections (RCS) is tested and verified, and several examples are selected to show the accuracy of this program in predicting RCS. Only one formulation is discussed; however, the program has been written to implement easily other surface formulations, such as those reported in [9]. Also, the program is written to efficiently fill the matrix. The properties of symmetry in reference to the impedance and admittance matrices that are used to build the final matrix are implemented. All the numerical results presented in this paper are obtained by the same program (MRBOR) to show its flexibility and generality.

II DEVELOPMENT OF THE SIE

a. Statement of the Problem:

In this section, the concept of the equivalence principle is used to derive the SIE formulation for composite scatterers with N homogeneous regions. The geometry and notations for such a scatterer are given in Fig. 1. The whole space is divided into N+1 homogeneous regions with permittivities $\varepsilon_i$ and permeabilities $\mu_i$, $i = 0, 1, 2, \ldots, N$. Lossy materials are considered by allowing $\varepsilon_i, \mu_i$, $i = 1, 2, \ldots, N$ to be complex. Some homogeneous regions are considered to be perfect conductors. The region $V_i$ is surrounded by a closed surface $S_i$ and recognized by the inward normal unit vector $n$. The surface interface between regions $V_i$ and
V_j is S_{ij}, i \neq j. Thus, S_i is the set of all interface surfaces S_{ij}, where j represents all region numbers interfacing with region V_i. Note that S_{ij} is the same surface as S_{ji}; however, the normal unit vectors n_i and n_j are in opposite directions to each other on S_{ij}.

\[ V_0 \quad \varepsilon_0, \mu_0 \quad S_{02} \]
\[ \begin{array}{c}
S_{01} \\
n_0 \\
\end{array} \quad \begin{array}{c}
S_{12} \\
\varepsilon_1, \mu_1 \\
n_1 \\
S_{1i} \\
S_{1N} \\
n_N \\
\varepsilon_N, \mu_N \\
S_{0N} \end{array} \quad \begin{array}{c}
S_1 \\
S_{2i} \\
n_i \\
S_{2N} \\
\varepsilon_{i}, \mu_{i} \\
S_{3N} \\
S_{0N} \end{array} \quad \begin{array}{c}
S_{22} \\
\varepsilon_2, \mu_2 \\
n_2 \\
V_1 \\
V_2 \\
V_3 \\
S_{03} \\
\end{array} \]

Fig. 1. General geometry of an object consisting of N regions.

b. SIE Formulation:

The total fields in each homogeneous region are denoted by \( \mathbf{E}_i \) and \( \mathbf{H}_i \), \( i = 0, 1, 2, \ldots, N \) for the electric and magnetic fields, respectively. If \( V_i \) is a perfectly conducting region, the fields are equal to zero. In the free-space region \( V_0 \), the total fields (\( \mathbf{E}_0, \mathbf{H}_0 \)) are the summation of the incident and scattered fields (\( \mathbf{E}^{\text{inc}} + \mathbf{E}^s, \mathbf{H}^{\text{inc}} + \mathbf{H}^s \)). From Maxwell's equations and the equivalence principle, one can express the field in each region in terms of unknown electric- and magnetic-equivalent surface currents. In this paper, the sources of excitation are considered to be due to a plane wave in the free space region; therefore, the fields at any observation point \( r \) in the free space can be expressed as [1] (these expressions are given here for convenience)

\[ \theta(r) \mathbf{E}_0(r) = \mathbf{E}^{\text{inc}} - L^0_{S_0} \mathbf{J}_0(r') + \kappa^0_{S_0} \mathbf{M}_0(r') \]  
\[ \theta(r) \mathbf{H}_0(r) = \mathbf{H}^{\text{inc}} - \kappa^0_{S_0} \mathbf{J}_0(r') - (1/\varepsilon_0^2) L^0_{S_0} \mathbf{M}_0(r') \]  

and
\[ \theta(\mathbf{r}) \mathbf{E}_i(\mathbf{r}) = -L^i_{S_i} \mathbf{J}_i(\mathbf{r}') + \kappa^i_{S_i} \mathbf{M}_i(\mathbf{r}') \] (3)

\[ \theta(\mathbf{r}) \mathbf{H}_i(\mathbf{r}) = -\kappa^i_{S_i} \mathbf{J}_i(\mathbf{r}') - (1/\eta_i^2) L^i_{S_i} \mathbf{M}_i(\mathbf{r}') \] (4)

in the region \( V_i \), where \( i = 1, 2, \ldots, N \). Time variation of \( e^{\jmath \omega t} \) is implied and suppressed throughout. The electric- and magnetic-surface currents along the boundaries are

\[ \mathbf{J}_i = \mathbf{n}_i \times \mathbf{H}_i \]
\[ \mathbf{M}_i = -\mathbf{n}_i \times \mathbf{E}_i \quad \text{on} \quad S_i \] (5)

In these equations, \( \eta_i = \eta_0 \sqrt{\mu_{ir}/\epsilon_{ir}} \), \( \epsilon_i = \epsilon_0 \epsilon_{ir} \), \( \mu_i = \mu_0 \mu_{ir} \), \( \epsilon_{ir} \) and \( \mu_{ir} \) are the relative permittivity and permeability in the region \( V_i \), and \( \eta_0 \) is the intrinsic impedance of the free space. The operators \( L^i_{S} \) and \( \kappa^i_{S} \) are defined as

\[ L^i_{S_i} \mathbf{C}_i(\mathbf{r}') = j \omega \mu_i \int_{S_i} \left[ \mathbf{C}_i(\mathbf{r}') + (1/\omega^2 \epsilon_i \mu_i) \nabla \nabla'. \mathbf{C}_i(\mathbf{r}') \right] \Phi_i \mathrm{d}S' \] (6)

\[ \kappa^i_{S_i} \mathbf{C}_i(\mathbf{r}') = \int_{S_i} \mathbf{C}_i(\mathbf{r}') \times \nabla \Phi_i \mathrm{d}S' \] (7)

where \( \mathbf{C}_i(\mathbf{r}') \) represents the currents \( \mathbf{J}_i \) or \( \mathbf{M}_i \). For \( \mathbf{r} = \mathbf{r}' \), the operators are interpreted as Cauchy principal-value integrals. \( \Phi_i \) is the Green's function of unbounded region, which can be represented as

\[ \Phi_i(\mathbf{r} - \mathbf{r}') = e^{-jk_i|\mathbf{r} - \mathbf{r}'|/|\mathbf{r} - \mathbf{r}'|} \] (8)

where \( k_i \) is the wave number of the region \( i \), which is equal to \( \omega \sqrt{\epsilon_i \mu_i} \). In Equations (1) to (4) the value of \( \theta(\mathbf{r}) \) is constant, depending on the position of \( \mathbf{r} \) as

\[ \theta(\mathbf{r}) = \begin{cases} 1 & \text{for} \quad \mathbf{r} \in V_i \\ 1/2 & \text{for} \quad \mathbf{r} \in S_i \\ 0 & \text{elsewhere} \end{cases} \] (9)
Applying the boundary conditions on each $S_{ij}$ yields a set of coupled integral equations for the unknown electric and magnetic currents on these surfaces. On the dielectric interfaces, the tangential fields are continuous. Thus,

$$E_i|_{\text{tan}} = E_j|_{\text{tan}} \quad \text{on } S_{ij}$$  \hspace{1cm} (10)

$$n_i \times H_i = n_i \times H_j \quad \text{on } S_{ij}$$  \hspace{1cm} (11)

On the conductor interfaces, the tangential electric fields vanish, yielding

$$E_i|_{\text{tan}} = 0 \quad \text{on } S_{ij} \text{ (conductor surfaces)}$$  \hspace{1cm} (12)

Substituting Equations (1) to (4) into (10) to (12), one obtains

$$[L^i_{S_i} J_i - \kappa^i_{S_i} M_i - L^j_{S_j} J_j + \kappa^j_{S_j} M_j]|_{\text{tan}} = \begin{cases} 0 & \text{if } j = 0 \\
^\text{inc} E_i & \text{if } j \neq 0 \end{cases} \quad \text{on } S_{ij}$$  \hspace{1cm} (13)

$$n_i \times \left[\kappa^i_{S_i} J_i + \frac{1}{\eta_i^2} L^i_{S_i} M_i\right] - n_i \times \left[\kappa^j_{S_j} J_j + \frac{1}{\eta_j^2} L^j_{S_j} M_j\right] = \begin{cases} 0 & \text{if } j = 0 \\
_i \times H^{\text{inc}} & \text{if } j \neq 0 \end{cases} \quad \text{on } S_{ij}$$  \hspace{1cm} (14)

$$[L^i_{S_i} J_i - \kappa^i_{S_i} M_i]|_{\text{tan}} = \begin{cases} 0 & \text{if } j = 0 \\
^\text{inc} E_i & \text{if } j \neq 0 \end{cases} \quad \text{on } S_{ij}$$  \hspace{1cm} (15)

To enable the reader to have better understanding of Equations (13) to (15) (which yield a coupled system of integral equations), a specific example will be considered. Consider a scatterer consisting of two dielectric regions attached to a perfectly conducting body, all in free space, as shown in Fig. 2. The boundary conditions in integral forms are
Fig. 2. Example of a three regions object.

\[
\begin{bmatrix}
L_{S_0}^0 J_0 - \kappa_{S_0}^0 M_0 - L_{S_1}^1 J_1 + \kappa_{S_1}^1 M_1
\end{bmatrix}_{\text{tan}} = E_{\text{tan}}^{\text{inc}} \quad \text{on } S_{01} \tag{16}
\]

\[
n_0 \times \left[ \kappa_{S_0}^0 J_0 + \frac{1}{\eta_0^2} L_{S_0}^0 M_0 \right] - n_0 \times \left[ \kappa_{S_1}^1 J_1 + \frac{1}{\eta_1^2} L_{S_1}^1 M_1 \right] = n_0 \times H^{\text{inc}} \quad \text{on } S_{01} \tag{17}
\]

where

\[
\begin{align*}
J_0 &= (J_{01}, J_{02}, J_{03}) \\
J_1 &= (-J_{01}, J_{12}, J_{13}) \\
M_0 &= (M_{01}, M_{02}) \\
M_1 &= (-M_{01}, M_{12})
\end{align*} \tag{18}
\]
\[ [L_{S_0}^0 J_0 - \kappa_{S_0}^0 M_0 - L_{S_0}^2 J_2 + \kappa_{S_2}^2 M_2 ] \tan = E_{\tan}^{inc} \quad \text{on } S_{02} \quad (19) \]

\[ n_0 \times [\kappa_{S_0}^0 J_0 + (1/\eta_0^2) L_{S_0}^0 M_0] - n_0 \times [\kappa_{S_2}^2 J_2 + (1/\eta_2^2) L_{S_2}^2 M_2] = n_0 \times H^{inc} \quad \text{on } S_{02} \quad (20) \]

where

\[ J_2 = (-J_{02}, J_{12}, J_{23}) \]
\[ M_2 = (-M_{02}, -M_{12}) \quad (21) \]

\[ [L_{S_0}^0 J_0 - \kappa_{S_0}^0 M_0 ] \tan = E_{\tan}^{inc} \quad \text{on } S_{03} \quad (22) \]

\[ [L_{S_1}^1 J_1 - \kappa_{S_1}^1 M_1 - L_{S_2}^2 J_2 + \kappa_{S_2}^2 M_2 ] \tan = E_{\tan}^{inc} \quad \text{on } S_{12} \quad (23) \]

\[ n_1 \times [\kappa_{S_1}^1 J_1 + (1/\eta_1^2) L_{S_1}^1 M_1] - n_1 \times [\kappa_{S_2}^2 J_2 + (1/\eta_2^2) L_{S_2}^2 M_2] = n_1 \times H^{inc} \quad \text{on } S_{12} \quad (24) \]

\[ [L_{S_1}^1 J_1 - \kappa_{S_1}^1 M_1 ] \tan = E_{\tan}^{inc} \quad \text{on } S_{13} \quad (25) \]

\[ [L_{S_2}^2 J_2 - \kappa_{S_2}^2 M_2 ] \tan = E_{\tan}^{inc} \quad \text{on } S_{23} \quad (26) \]

These are nine vectorial equations in nine vectorial unknowns. The unknowns are the
electric- and magnetic-surface currents \( J_{01}, J_{02}, J_{03}, J_{12}, J_{13}, J_{23}, M_{01}, M_{02}, \) and \( M_{12}. \) The
integral equations are reduced to matrix equations using the method of moments by using
triangle testing and triangle expansion functions [8].
The above surface integral equations are applied to rotationally-symmetric bodies. The reduction of the integral equations to matrix equations involving unknown surface currents follows a well-known procedure [9]. After the necessary manipulations of the method of moments, the general matrix takes the form

\[
[T \ I_n \ I_n = [V]_n
\]

(27)

where \( T \) is a square matrix, representing a combination of impedance and admittance submatrices that are given in [9]; \( I_n \) is a column matrix for the unknown expansion coefficients of the unknown current components \( J \) and \( M \); and \( V_n \) is the excitation column matrix. Once the matrix is solved, the induced currents on all surface interfaces can be determined. Scattered far fields can be determined from the induced currents on the outer surfaces.

III RESULTS AND DISCUSSION

a. Bistatic RCS:

In this section the bistatic RCS is computed numerically. The numerical results are verified by comparison with analytical and supplemental numerical results.

Dielectric sphere: First, to verify the numerical results of objects made of more than one homogeneous region, the example of a sphere geometry is considered. The series solution for a dielectric sphere [10] having \( \epsilon_r = 4-0.5 \), \( \mu_r = 2-0.25 \), and \( \kappa a = 3 \) is obtained and compared with the numerical solution of the same object when it is divided into three homogeneous regions of the same material type. The results obtained from both solutions must be identical, because physically both cases represent the same object. The agreement between the two solutions is excellent, as shown in Fig. 3. Several spherical objects have been considered, such as the coated sphere and the multilayered sphere; however, their results have been omitted for brevity.
Fig. 3. Bistatic RCS of dielectric sphere divided into three regions of the same materials, \( \varepsilon_r=4-j0.5, \mu_r=2-j0.25, \) and \( k_s=3. \)

**Dielectric toroid:** Another geometry, of a dielectric toroid, is considered, as shown in Fig. 4a. The numerical solution is obtained, once, when the circular cross section is divided into six regions, all of which have the same type of material. This numerical solution is compared with the numerical solution of the same object when it is treated as one region. The agreement between both solutions is excellent, as shown in Fig. 4b. When each region is filled with different homogeneous materials, the computed bistatic RCS is shown in Fig. 5. Notice that the scattering level has increased in the whole bistatic range.

Fig. 4a. Toroid cross section divided into six homogeneous regions.
Fig. 4b. Bistatic RCS of the toroid when all regions have the same materials, 
\( \varepsilon_r = 4 - j0.5, \mu_r = 2 - j0.25, ka = 3, kb = 4.4. \)

Fig. 5. Bistatic RCS of the toroid with, \( \varepsilon_{r1} = 3 - j3, \varepsilon_{r2} = 2 - j2, \varepsilon_{r3} = 4 - j0.5, \varepsilon_{r4} = 6 - j6, \varepsilon_{r5} = 5 - j5, \varepsilon_{r6} = 4 - j4, \) and \( \mu_r = 2 - j0.25 \) in all regions.
**Loaded conducting bicone:** To examine the numerical solution performance of objects made of dielectric and conductors, the case of a biconical conducting object is selected. The numerical solution of this object and the conducting bicone is compared with the numerical solution of a conducting bicone surrounded by an artificial dielectric material of free-space permittivity and permeability, as shown in Fig. 6. In this example, the bicone is illuminated by a plane wave of normal incidence ($\theta = 90^\circ$). Therefore, the number of the azimuthal modes that is used to obtain the bistatic RCS is eleven, $n=0,\ldots, \pm 5$; for the axial incident cases the required modes are $\pm 1$. Figure 6 shows the excellent agreement between the two solutions. Also shown is the symmetry around $\theta = 90^\circ$. In this example, the H-plane shows a zero-back and forward-scattering, and the E-plane shows its maximum values at these directions. When the free-space part of this object is filled with homogeneous materials of $\epsilon_r = 4$, the RCS is computed as shown in Fig. 7. The sidelobes of the scattered H-plane pattern disappear, and the forward scattering for the E-plane pattern becomes higher than the back-scattering.

![Fig. 6. Bistatic RCS of a conducting bicone, $ka=3$, $\epsilon_r=1$, $\mu_r=1$.](image-url)
b. **Monostatic RCS:**

In the above examples, the bistatic RCS were considered. Next monostatic RCS are also verified numerically and experimentally.

**Half-conducting sphere:** The first example is the perfectly-conducting dielectric hemisphere. For numerical verification, the dielectric is considered as a free space. The monostatic RCS is compared with the numerical solution of a conducting hemisphere, as shown in Fig. 8. This figure indicates excellent agreement between both solutions in the whole $\theta$ range. When the free-space part is replaced by materials of $\varepsilon_r=2-j0.5$, and $\mu_r=3-j0.5$, the monostatic RCS is computed as shown in Fig. 9. The effect of the presence of dielectric materials...
Fig. 8. Monostatic RCS of a conducting hemisphere compared with the scattering from a conducting hemisphere-air hemisphere object, $ka=3$.

Fig. 9. Monostatic RCS of a conducting hemisphere-dielectric hemisphere object, $ka=3$, $\varepsilon_r=2-j0.5$, $\mu_r=3-j0.5$. 
is different from the free space. This difference is clear in the backscattering data for ranges of $\theta < 90$, which is the dielectric side. Significant reduction of the backscattering level is observed. When $\varepsilon_r = 1$, the effect of the flat surface and the sharp edges obviously contributed to the high backscattering levels.

**Round-tip cone:** In this section, new numerical results are presented and verified with measurements, which have been collected at the RCS Range of Group 95 at Lincoln Laboratory. The first geometry that is considered is the partially coated, perfectly conducting round-tip cone, shown in Fig. 10. The first case that has been considered is the one that corresponds to a homogeneous coating; in other words, the materials in both coating regions are the same. The monostatic RCS are shown in Fig. 11 for both $\theta\theta$ polarization (left) and $\phi\phi$ polarization (right), respectively. When the coating on the second region is removed, the second region is then equivalent to free-space permittivity. The monostatic RCS is computed and compared with the measurements as shown in Fig. 12 for $\theta\theta$ polarization and $\phi\phi$ polarization, respectively. Figure 13 show the monostatic RCS of both polarizations when second region is filled with materials of $\varepsilon_r = 2.60$. In the above three cases, excellent agreement is obtained between the measured data and the computed data.

![Diagram of a partially coated, perfectly conducting round-tip cone](image)

All dimensions in $\lambda_0$ (free space wavelength)

$L = 1.978,$ $b = 0.2328,$ $a = 0.0424,$ $t = 0.0847,$ $c = 0.86$

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<tr>
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<th>$\varepsilon_{r2}$</th>
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<td>2.05</td>
</tr>
<tr>
<td>2</td>
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</tr>
<tr>
<td>3</td>
<td>2.05</td>
<td>2.60</td>
</tr>
</tbody>
</table>

Fig. 10. Geometry of a partially coated, perfectly conducting round-tip cone.
Fig. 11. Computed and measured monostatic RCS of case 1 in Fig. 10.

Fig. 12. Computed and measured monostatic RCS of case 2 in Fig. 10.
Fig. 13. Computed and measured monostatic RCS of case 3 in Fig. 10.

Fig. 14. Geometry of a perfectly-conducting cylinder partially coated with two dielectric layers.
Finite cylinders: Another geometry, double layered coating of a finite conducting cylinder, is considered. Figure 14 shows the geometry of the cylinder. The values of the parameters of Fig. 14 are given in Table I for three cases. All the measurements in this section have been performed at 4GHz. Figure 15 (left) shows the comparison between the computed and measured data of the monostatic RCS for $\theta\theta$ polarization of case 1 in Table I. In this case only one layer is considered.

Table I. Parameters of Figure 14

<table>
<thead>
<tr>
<th>case</th>
<th>a(cm)</th>
<th>L(cm)</th>
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<th>$\varepsilon_{r1}$</th>
<th>$t_2$(cm)</th>
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<td>-</td>
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<tr>
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</tbody>
</table>

The measured data of $\phi\phi$ polarization are not available, however, the computed data are shown in Fig. 15 (right). The monostatic RCS of the second case in Table I of the two layered coating is shown in Fig. 16. Again, only the measurements of the $\theta\theta$ polarization are available. In the last case, the first layer is a high loss material of large permittivity and the second one is high loss of small permittivity. In the third example of Table I, the first layer is made of a lossless material of small permittivity; the second layer is made of high-loss high-permittivity materials. The monostatic RCS is shown in Fig. 17 for the $\theta\theta$ and $\phi\phi$ polarization, respectively. The last three cases represent large objects. The agreement between the measured and computed data is satisfactory. In general, the above results show accuracy of the measurements within a wide dynamic range of sensitivity.
Fig. 15. Computed and measured monostatic RCS of case 1 in Table I.

Fig. 16. Computed and measured monostatic RCS of case 2 in Table I.
IV CONCLUSION

In this paper, a computer program has been developed to compute the electromagnetic scattering from axisymmetric objects. The method of moments is used to solve the surface-integral equations formulation (E-PMCHW). A number of objects has been analyzed by this program. Each object consisted of arbitrarily arranged homogeneous regions. Both bistatic and monostatic RCS data were presented. The computed data were verified either numerically or experimentally and excellent agreement was demonstrated.

Fig. 17. Computed and measured monostatic RCS of case3 in Table I.
REFERENCES:


Application of Equivalent Edge Currents to Correct the Backscattered Physical Optics Field of Flat Plates

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Abstract
In the microwave case the physical optics (PO) method is frequently used for the analysis of complex structures which are modeled by flat plates of triangular or quadrangular shape. The study of the radar cross section (RCS) of an isolated panel, however, reveals deviations from experimental results which are due to edge diffraction effects not considered by PO. In order to correct the PO-field by an additive field term, the equivalent fringe currents (EC) of Michaeli have been used to derive the backscattering matrix of an isolated edge. By adding the matrices of the individual edges to the PO-matrix the RCS of a square flat plate with zero and finite thickness is analysed and the result is compared with measurements. The efficiency of the method is demonstrated for objects modeled by a higher number of panels and edges, namely a cylinder and a double dihedral. All computations were performed with the computer code SIG5 of the Institute.

1. Introduction
Since several years, the computer program SIG5 is applied in the Institute for Radio Frequency Technology for the prediction of the RCS of structures which are complicated in shape and large compared to the wavelength. SIG5, based on PO, is capable of analysing perfectly and imperfectly conducting structures, including double reflections. The targets are modeled by panels of triangular and quadrangular shape, see Fig. 1.1. The hidden surface problem inherent with PO is solved by an exact construction of the shadow boundary for each panel, whose size is only limited by the admissible deviation between the true surface and the model surface. SIG5 is organized in a very similar way to the computer code RECOTA, developed by the Boeing Aerospace Company, Seattle [1]. SIG5 has been successfully tested for a series of perfectly conducting basic structures such as a sphere, cylinder, cube, circular disk and a double dihedral [2-4]. Also, more complex bodies like a periscope structure have been analysed with promising results [5].

Fig. 1.1
Panel model of an airplane for the application of PO.
Despite the good results which are generally available with PO, there are special cases (certain structures, specific pattern cuts, selected polarizations) where deviations of practical significance between computational and experimental results are observed. They frequently can be explained by the feature of PO to treat the influence of an edge only as the geometrical boundary of a panel thus neglecting physical edge diffraction effects.

The choice of a theory which takes into account edge diffraction effects is influenced by the following viewpoints:

a) Since SIG5 is a very comprehensive computer code each extension should cause a minimum of changes. Therefore, theories are preferred which are able to correct the PO-solution by an additive term.

b) In computing the RCS of large and complex objects modeled by numerous panels the computer effort increases considerably. Therefore, theories which need a high computer effort are not favoured.

c) Since edges of arbitrary length, wedge angle and orientation in space occur in the target model, the theory should not present a solution for a specific panel, rather, the diffracted field of an isolated edge must be described by the specific edge parameters alone.

Bearing these points in mind, only asymptotic theories come into question, which either describe the difference between the total wedge diffracted field and the PO-field directly [6,7] or which evaluate fringe currents flowing along the edge and generating the same difference field by evaluating the radiation integral over the length of the edge [8, 9].

In this paper, the second theory is followed. Here the fringe currents given in [9] are preferred, since they are valid for arbitrary aspects of observation. On the basis of these currents, the backscattering matrix for an isolated edge is derived.

In the following section the theoretical background is discussed. In section 3, the theory is applied for the RCS-analysis of a panel. In section 4, the results for a cylinder and a double dihedral are presented. Section 5, finally, summarizes some conclusions on the basis of the proceeding analysis.

2. Theoretical Background

The results of the theories used in the computer code SIG5 are expressed by the backscattering matrix

\[
[T] = \begin{bmatrix}
t_{11} & t_{12} \\
t_{21} & t_{22}
\end{bmatrix},
\]

which relates the cartesian components of the scattered field \(\mathbf{E}_s\) to those of the incident field \(\mathbf{E}_i\):

\[
\begin{bmatrix}
E_{sx} \\
E_{sy}
\end{bmatrix} = [T] \begin{bmatrix}
E_{ix} \\
E_{iy}
\end{bmatrix}.
\]
The propagation direction of the scattered field is given by the z-axis (observer fixed coordinate system). From the complex elements \( t_{ij} \) of the scattering matrix the polarization dependent RCS is computed:

\[
\sigma_{ij} = \lim_{r \to \infty} (4\pi r^2 t_{ij} t_{ij}^*),
\]

where \( r \) is the distance between the radar observer and the test object. The RCS referred to 1 square meter and expressed in decibels yields the quantity dBsm, which is used in this paper to compare theoretical and experimental results for the selected transmitting/receiving polarizations.

The total scattering matrix \([T]\) of a complex body can be thought to be composed of a matrix \([TP^o]\) which is the sum of the scattering matrices based on PO of the individual \( N \) panels and the scattering matrix \([T_f]\) which sums up the scattering matrices of the \( M \) edges of all panels:

\[
[T] = [TP^o] + [T_f], \quad [TP^o] = \sum_{n} [TP^o_n], \quad [T_f] = \sum_{m} [T_f_m].
\]

The PO-scattering matrix of an individual panel of zero thickness and perfect conductivity is readily evaluated using the radiation integral

\[
E_s(r) = \frac{j k e^{-jkr}}{4\pi r} \int_{F_p} (s_x(s_x \mathcal{J}_p(r'))) e^{-j k \vec{s} \cdot \vec{r}'_f} df',
\]

and introducing the surface current

\[
\mathcal{J}_p(r') = \begin{cases} 2n x (e x \mathcal{P}_e) E_e / Z e^{-j k \vec{e} \cdot \vec{r}'} & \text{on the illuminated parts of the panel.} \\ 0 & \text{in the shadowed parts of the panel.} \end{cases}
\]

The geometrical parameters \( r, r', e, s, F_p \) and \( n \) are explained in Fig. 2.1, while the electric parameters are given by \( \lambda = \text{wavelength}, k = 2\pi/\lambda = \text{wave number}, Z = \text{wave impedance of the propagation medium}, \mathcal{P}_e = p_x \vec{e}_x + p_y \vec{e}_y = \text{unit polarization vector} \) and \( E_e = \text{magnitude of the incident electric field}. \)

![Fig. 2.1](image-url) Geometrical scheme for the interpretation of the radiation integral.
The evaluation of the radiation integral for the backscattered \((e = -s)\) field in the observer fixed cartesian coordinate system and the comparison with (2.2) results in the following backscattering matrix:

\[
(2.7) \quad \{T_p\} = -\frac{j}{2\pi} \frac{e^{-jkr}}{r} \int_{F_p} e^{jkz'} dx'dy' \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix}.
\]

For a panel of polygonal shape the phase integral can be solved analytically. Details of the integration are given in [4] for panels of triangular and quadrangular shape which are used in the computer program SIG5 to model the target.

Normalizing the scattering matrix to \(e^{-jkr}/r\), the matrix appears purely imaginary. Independent of the orientation of the panel, no differences between xx-polarization and yy-polarization nor any cross-polarization are predicted by PO. This, however, is only true if the panel is perfectly conducting and no double or multiple interactions can occur between pairs or a higher number of panels.

The electric and magnetic fringe currents of Michaeli [9] can be written in a very compact way using the coefficients \(D^e_f, D^e_m, D^e_{em}\) (these symbols are proposed by Knott [10]):

\[
(2.8) \quad I^e(\Phi_e, \Phi_m; \beta_e, \beta_m) = -\frac{j2E_{et}}{k2\sin^2 \beta_e} D^e_e(\Phi_e, \Phi_m; \beta_e, \beta_m) - \frac{j2H_{et}}{k\sin^2 \beta_e} D^e_{em}(\Phi_e, \Phi_m; \beta_e, \beta_m),
\]

\[
(2.9) \quad M^e(\Psi_e, \Psi_m; \beta_e, \beta_m) = \frac{j2H_{et}}{k\sin \beta_e \sin \beta_m} D^e_m(\Phi_e, \Phi_m; \beta_e, \beta_m).
\]

\(E_{et} = \vec{E}_e \) or \(H_{et} = \vec{H}_e\) is the component of the incident electric or magnetic field parallel to the edge with unit tangent vector \(\vec{t}\). The geometrical parameters \(\Phi_e, \Phi_m, \beta_e, \beta_m\) and \(n\) are explained in Fig. 2.2. The formulas for \(D^e_f, D^e_m\) and \(D^e_{em}\) are given in the appendix for the backscattering case \((\Phi_e = \Phi_m, \beta_m = \pi - \beta_e)\).

![Fig. 2.2 Wedge geometry.](image)
Before choosing the fringe currents of Michaeli, it was necessary to relate this theory to other theories, which, in principle, could be used to solve edge diffraction problems observing the viewpoints given in the introduction.

Though there are several papers [10 - 15] dealing with the relationships between asymptotic diffraction theories, it is useful to summarize the results with the aid of a few statements. Thereby the output of the different theories are given in the form of currents. For mathematical details see [15].

The fringe currents of Michaeli can be written in the following way:

\[(2.8a) \quad I^f = I^t - I^{PO}, \]
\[(2.9a) \quad M^f = M^t - M^{PO}, \]

where \(I^t, M^t\) are the equivalent electric and magnetic total currents, responsible for the total wedge diffracted field while \(I^{PO}\) and \(M^{PO}\) are the equivalent PO-components responsible for the PO-field of the wedge. The expressions derived by Michaeli are obtained by asymptotic end-point evaluation of the fringe current radiation integral over the ray coordinate measured along the diffracted ray grazing the surface of the local wedge. The resulting expressions are finite for all aspects of illumination and observation, except for the special case where the direction of observation is the continuation of a glancing incident ray coming from outside the wedge. This situation occurs only in forward scattering.

Choosing the coordinate for the radiation integral in a traditional way normal to the edge, expressions for the fringe currents are obtained [16] which display infinities for certain combinations of observation and incidence directions. These fringe currents are identical to those which can be evaluated [15] from the fringe diffracted fields of Mitzner [7]. Now adapting the fringe currents to the cone of diffracted rays one would expect that these would be identical to those which can be derived from the fringe field expressions of Ufimtsev [6] and which are used in [1]. This is the case for the electric total current, for the magnetic total and PO-current but in general not for the electric PO-current.

The total currents are identical to the filamentary currents of Keller's GTD-field [17], represented by Knott and Senior in a compact manner [18] and denoted as equivalent currents by Ryan and Peters [19] who applied the concept to compute the edge diffracted field in caustic regions. Evaluating the equivalent currents of the GTD for vertical incidence (\(\theta_e = 90^\circ\)) on a half-plane \((n = 2)\) one receives the filamentary currents of the diffracted field derived by Sommerfeld [20].

The difference in the electric PO-currents is given by a coupling term between the incident electric field, being perpendicular to the plane of incidence, and the PO-field having a component parallel to the plane of diffraction. This coupling term is not taken into account by the theory of Ufimtsev as was pointed out in [14]. Only in the case that both faces of the wedge are illuminated by the incident wave does the coupling term become identical to zero [15] and the fringe currents extracted from the theory of Ufimtsev become identical to the fringe currents of Michaeli.
Introducing now the fringe currents (2.8) and (2.9) in the radiation integral over a filamentary current with length \(L\)

\[
(2.10) \quad E_\delta^f(\mathbf{r}) = \frac{ik}{2\pi} e^{-jkr} \int_L \left( Z \mathbf{I}^f(\mathbf{r'}) (\mathbf{s} \times (\mathbf{s} \times \mathbf{e}')) + M^f(\mathbf{r'}) (\mathbf{3} \times \mathbf{e}') \right) e^{jkr' \cdot \mathbf{r}} \, dl',
\]

one arrives at the backscattering matrix

\[
(2.11) \quad [T^f] = -\frac{1}{2\pi} \frac{e^{-jkr}}{r} \int_L e^{j2k\mathbf{s} \cdot \mathbf{r'}} \, dl' \times \left[ \begin{array}{ccc}
D_e^f t_x^2 + D_m^f t_y^2 - D_m^e t_x t_y & (D_e^f - D_m^e) t_x t_y + D_m^e t_x^2 \\
(D_e^f - D_m^f) t_x t_y - D_m^e t_y^2 & D_m^f t_x^2 + D_m^e t_y^2 + D_m^e t_x t_y
\end{array} \right].
\]

The geometrical parameters are explained in Fig. 2.3.

As in the case of the PO-scattering matrix the phase integral can be solved analytically. Introducing the mid-point vector \(\mathbf{r}_M\) of the edge one obtains

\[
(2.12) \quad \int_L e^{j2k\mathbf{s} \cdot \mathbf{r'}} \, dl' = L \sin(kL_{t_z}) \frac{e^{j2kr_Mz}}{kL_{t_z}} 1 - t_z,
\]

with \(t_z = -\cos \beta_e\).

Normalizing the backscattering matrix \([T^f]\) by \(e^{-jkx/r}\) it appears purely real. In general differences between HH- and VV-polarization are predicted. Also, since \([T^f]\) is asymmetric, differences between HV- and VH-polarization are predicted which is not correct for monostatic scattering processes. This is due to corner diffraction effects which arise with an edge of finite length and which are not considered in the theory. Symmetry is observed for the special case of normal wave incidence (\(\beta_e = 90^\circ\)). Further, if the contour of a panel is a smooth curve, the matrix \([T^f]\) would be symmetric.

For the discussions in the next section, some properties of the coefficients \(D_e^f, D_m^f, D_m^e\) are needed, which are summarized in the following. Fig. 2.4 shows...
the coefficients for normal incidence ($\beta_e = 90^\circ$) and a half-plane ($n = 2$) depending on the angle $\psi_e$. In Fig. 2.5 the coefficients for a step ($n = 1.5$) are represented where $D_{em}^f$ is identical to zero. In both cases, mirror symmetries for each coefficient become relevant about $\psi_e = 180^\circ$ for the half-plane and $\psi_e = 135^\circ$ for the step. Further symmetries, now radial symmetries, occur for the half-plane between coefficient $D_e^f$ and $D_m^f$ about $\psi_e = 90^\circ$ and $\psi_e = 270^\circ$.

Fig. 2.6 shows the coefficients $D_e^f$, $D_m^f$ and $D_{em}^f$ for $\psi_e = 270^\circ$ and a half-plane now dependent of the angle $\beta_e$. One also finds symmetries, namely mirror-symmetry for $D_e^f$ and $D_m^f$ and radial-symmetry for $D_{em}^f$ about $\psi_e = 90^\circ$. These symmetries occur for all $\psi_e = \text{const}$.

3. Analysis of the RCS of a Square Test Panel by the PO- and EC-Method

Studies similar to the ones reported here were performed by Ross [21], Sikta, Burnside, Chu and Peters [22], Balanis, Griesser and Marsland [23], Volakis and Ricoy [24], Pelosi, Tiberio, Puccini and Maci [25], Ivrissimtzis and Marhefka [26]. Ross applied the GTD up to triple diffraction to compute the RCS of rectangular flat plates. Sikta et al. used a modified equivalent current concept based on GTD and a corner diffraction analysis to study the RCS of flat plate structures such as a square flat plate, a finlike plate and a disc. Balanis et al. used GTD up to third order diffraction for principal
plane and EC-currents of Michaeli for off-principal plane backscattering of plates. The angular spectrum method along with the generalized matrix formulation were employed for the diffraction analyses of a thick perfectly conducting half-plane by Volakis and Ricoy. Pelosi et al. applied GTD to calculate the RCS of a square plate. Ivrissimtzis and Marhefka used a uniform ray approximation including higher order terms for the analysis of polyhedral structures.

The procedure presented in the previous section is not limited to the simple structure of a flat plate as will be demonstrated in the next section. However, in this section the scattering from a square test plate is investigated in great detail to estimate the efficiency and the limits of application of the chosen method.

The square test panel is analysed at a frequency of 16.66 GHz which results in a wavelength of 17.995 mm. The edge length is 91.4 mm (5.08 λ) and the thickness is 0.8 mm (0.044 λ). In order to have an independent external accuracy check for the following computations, the test panel was chosen to be equal to one of the panels investigated by Ross [21]. The panel is defined in the observer-fixed coordinate system, see Fig. 3.1. The rotations around the x, y and z-axis are defined by the angles \( \psi_x, \psi_y, \psi_z \). An incident wave with electric field vector parallel to the x-axis is designated as horizontally polarized while a field vector parallel to the y-axis describes a vertically polarized field, if the object is rotated around the y-axis for instance.

\[
\begin{align*}
\text{edge 1} & \quad \text{edge 2} & & \text{edge 3} & & \text{edge 4} \\
\text{t}_1 & & \text{t}_2 & & \text{t}_3 & & \text{t}_4
\end{align*}
\]

\[
\begin{align*}
\text{y} & & \psi_y & & \text{z} \\
\text{x} & & \text{t}_1 & & \text{t}_2 & & \text{e}_z
\end{align*}
\]

Fig. 3.1 Orientation of the panel in the observer-fixed coordinate system and numbering of the edges for a panel with zero thickness.

Fig. 3.2 resp. Fig. 3.3 shows the experimental RCS-result for horizontal resp. vertical polarization and for the principal plane (\( \psi_x=0, \psi_y=\alpha, \psi_z=0 \)). For \( \alpha = 0^\circ \) one has normal incidence, for \( \alpha = 90^\circ \) the panel is seen under grazing incidence. The PO-result for a panel with zero thickness, which is usually compared with experimental results, is shown in Fig. 3.4. The maximum RCS-value amounts to 4.33 dBsm.
Besides the smoothing of the deep nulls of the PO-result, the sidelobe peaks far from the mainlobe are significantly higher in the experiment than in the theory. At grazing incidence a deep null of $-\infty$ dB is predicted from theory. In the experiment this null occurs only for horizontal polarization, while for vertical polarization a level of about -27 dB under the main lobe peak can be observed. In view of the experimental results the theoretical results need improvement.

However, it must be emphasized, that for the experiment a panel with a thickness of 0.044 $\lambda$ was used while for the theory the panel was assumed with zero thickness. Thus far the comparison is a little unfair. If one models the flat plate with the same thickness as was used for the experiment, then it consists of six faces. Each of them can contribute to the total scattered field if it is illuminated by the incident field. The result is presented in Fig. 3.5. One can see that the deep nulls are filled up, that the peaks of the distant sidelobes are higher than in the case of zero thickness, and that, at grazing incidence, an RCS on the order of -41 dB under the peak is predicted compared to $-\infty$ dB for the zero thickness panel and the -27 dB of the experiment.
In the following the influence of the edges is considered. First the panel with zero thickness is analysed.

Edges 2 and 4 make an angle $\beta_e = 90^\circ$ with the incident ray independent from $\phi_y$, see Fig. 3.1, while $\beta_e$ of edges 1 and 3 varies with $\beta_e = 90^\circ + \phi_y$. From Eq. (2.11) and Eq. (2.12), one can expect an oscillating behaviour of the RCS of edges 1 and 3 and a monotonic function for edges 2 and 4 with respect to the rotation angle $\phi_y$. For rotation angles $0^\circ < \phi_y < 180^\circ$, edge 2 is nearer to the radar observer than edge 4, therefore, it is sometimes called the leading edge while edge 4 is the trailing edge.

First the influence of edges 1 and 3 on the RCS computed by PO is considered. The scattering matrix of edge 1 is given by

$$[T^s_1] = -\frac{1}{2\pi} \frac{e^{-jkr}}{r} L \frac{\sin(kLsin\phi_y)}{kLsin\phi_y} \begin{bmatrix} D^f_0 & D^f_{em} \\ 0 & D^f_m \end{bmatrix},$$

with arguments $\psi_e = 90^\circ$, $\beta_e = 90^\circ + \phi_y$ for $D^s_0$, $D^s_{em}$ and $D^s_m$. For edge 3 the following formula is obtained:

$$[T^s_3] = -\frac{1}{2\pi} \frac{e^{-jkr}}{r} L \frac{\sin(kLsin\phi_y)}{kLsin\phi_y} \begin{bmatrix} D^f_0 & D^f_{em} \\ 0 & D^f_m \end{bmatrix},$$

with $\psi_e = 90^\circ$, $\beta_e = 90^\circ - \phi_y$.

We can make the following observations:

1. Both scattering matrices are purely real, after normalizing by the factor $e^{-jkr}/r$.

2. The maximum value is reached for $\phi_y = 0^\circ$ (the incident ray hits the plate normally). The maximum value amounts to about -32 dBsm which is about -36 dB under the PO-value. This means that the peak value computed by the PO-method is negligibly influenced by the diffraction effects of edges 1 and 3. Since, further, the RCS decreases with increasing angle $\phi_y$ in an oscillating manner, both edges have practically no influence on the PO-result.

3. Since mirror symmetries for $D_0$ and $D_m$ hold, see Fig. 2.6, the copolar scattered fields of each edge are equal in amplitude and phase.

4. Both matrices indicate a cross-polarization term and, therefore, are asymmetric. Since, however, for the edge coefficient $D_{em}$ a radial symmetry, see Fig. 2.6, holds, the sum matrix $[T^s_{1,3}] = [T^s_1] + [T^s_3]$ is symmetric and is given by

$$[T^s_{1,3}] = \frac{1}{\pi} \frac{e^{-jkr}}{r} L \frac{\sin(kLsin\phi_y)}{kLsin\phi_y} \begin{bmatrix} D^f_0 & 0 \\ 0 & D^f_m \end{bmatrix},$$

with arguments $\psi_e = 270^\circ$, $\beta_e = 90^\circ - \phi_y$. The sum matrix predicts differences between horizontal and vertical polarization. However, this effect has practically no influence on the final result.
Fig. 3.6 shows the RCS of edge 1 and 3 for horizontal polarization and Fig. 3.7 for vertical polarization. Fig. 3.8 shows the sum RCS of edges 1 and 3 for horizontal polarization. For vertical polarization only negligible differences exist.

Contrary to edges 1 and 3, edges 2 and 4 generate contributions of significant practical interest. Bearing in mind that $D_{em}(\beta_e = 90^\circ) = 0$, one obtains for edge 2

$$ (3.4) \quad [T_2^e] = -\frac{1}{2\pi} \frac{e^{-jkr}}{r} L e^{jk\sin\psi_y} \begin{bmatrix} D_m^e & 0 \\ 0 & D_\delta^e \end{bmatrix}. $$

The arguments of the edge coefficients are $\nu_e = 270^\circ - \psi_y$, $\beta_e = 90^\circ$. For edge 4, we get for the parameters $\nu_e = 270^\circ + \psi_y$, $\beta_e = 90^\circ$

$$ (3.5) \quad [T_4^e] = -\frac{1}{2\pi} \frac{e^{-jkr}}{r} L e^{-jk\sin\psi_y} \begin{bmatrix} D_m^e & 0 \\ 0 & D_\delta^e \end{bmatrix}. $$

The following observations can be made:
1. After normalization both scattering matrices appear to be complex.
2. The RCS contributions of both matrices vary monotonically with respect to the rotation angle and are significant compared to the PO-solution.
3. The individual RCS contributions are different from each other and show differences for horizontal and vertical polarization.
4. For horizontal polarization, the contribution of edge 4 dominates, while, for vertical polarization, the contribution of edge 2 dominates.
5. Since symmetry, see Fig. 2.4, is given, the RCS of edge 2 for horizontal (vertical) polarization is identical to the RCS of edge 4 for vertical (horizontal) polarization. For the sum matrix one obtains

\[
[T_{2,4}^s] = -\frac{1}{2\pi} \frac{e^{-jkr}}{r} L \cos(\text{kasin}\varphi_y) (D_m^f - D_e^f) \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix} - j \frac{1}{2\pi} \frac{e^{-jkr}}{r} L \sin(\text{kasin}\varphi_y) (D_m^f + D_e^f) \begin{bmatrix} 1 & 0 \\ 0 & 1 \end{bmatrix},
\]

where the edge coefficients have the arguments \(\varphi_e = 270^\circ - \varphi_y, \beta_e = 90^\circ\).

The individual RCS contribution of edges 2 and 4 is represented in Fig. 3.9 and Fig. 3.10. The sum RCS of edges 2 and 4 is given in Fig. 3.11.

![Fig. 3.9](image1)
RCS of edge 2 (leading edge) for horizontal polarization and of edge 4 (trailing edge) for vertical polarization.

![Fig. 3.10](image2)
RCS of edge 2 for vertical polarization and of edge 4 for horizontal polarization.

![Fig. 3.11](image3)
RCS of edges 2 and 4 for both polarizations.
Fig. 3.12 finally shows the PO-result plus the contribution of all four edges which is identical for horizontal and vertical polarization. One can confirm excellent agreement with the experimental results for vertical polarization. It is, however, unsatisfactory that the theoretical solution doesn't indicate any polarization dependence.

Mathematically this can be explained by discussing the scattering matrix \([T_{2,4}]\). The imaginary part has identical elements for horizontal and vertical polarization. The real part has elements of equal magnitude but opposite sign, so that, with the additive correction of the purely imaginary PO-solution, the matrix behaves like \(-a-jb\) for horizontal and \(+a-jb\) for vertical polarization. This means that the final RCS result (PO-solution + EC-solution) presents identical values for horizontal and vertical polarization. This is not in agreement with measurements. Physically the effect is explained by the fact that the theory used does not take into account second and higher order diffraction effects. For horizontal polarization, the fields associated with double diffraction e.g. are 4 kL greater than the corresponding vertical polarization contribution [21]. Polarization dependent effects, however, are predicted by the theory for other diagram cuts, other panel shapes (e.g. triangle) or other wedge angles.

The latter is demonstrated for a flat plate with dimensions of the square panel, however, with the same thickness, namely 0.044 \(\lambda\), as in the experiment. This plate is modeled now by 6 panels and 12 edges \((n = 1.5)\). Without discussion if it is allowed to use the theory for close adjacent edges the result.

Fig. 3.13
RCS of the test panel with 0.044 \(\lambda\) thickness for horizontal polarization.

Fig. 3.14
RCS of the test panel with 0.044 \(\lambda\) thickness for vertical polarization.
of the computations is presented in Fig. 3.13 and Fig. 3.14. For horizontal polarization, the discrepancies between theory and experiment have become minor, but the deep null for grazing incidence again is not predicted by the theory.

If the plate is rotated around the $z$-axis by an angle of $\varphi_z = 45^\circ$ a diagonal cut can be achieved by a following rotation with $\varphi_y$. The theoretical results are presented in Figs. 3.15 and 3.16. Again no cross-polarization occurs which is in agreement with symmetrical properties. The experimental results are presented in Figs. 3.17 and 3.18. Since the RCS-values drop very quickly down to levels of about -40 dB under the mainlobe it is not very meaningful to discuss minor deviations between experimental and theoretical results.

![Graph](image1)

**Fig. 3.15**
Theoretical RCS of the test panel for a diagonal cut, horizontal polarization.

![Graph](image2)

**Fig. 3.16**
Theoretical RCS of the test panel for a diagonal cut, vertical polarization.

![Graph](image3)

**Fig. 3.17**
Experimental RCS of the test panel for a diagonal cut, horizontal polarization.

![Graph](image4)

**Fig. 3.18**
Experimental RCS of the test panel for a diagonal cut, vertical polarization.

Finally, the RCS has been computed for a cut where the rotation axis makes an angle of $\varphi_z = 30^\circ$ with the main axis ($15^\circ$ with the diagonal) of the plate. In this case no symmetry occurs and the theory predicts cross-polarization. Because of the asymmetry in the scattering matrix, the cross-polarization HV, see Fig. 3.19, is slightly different from the cross-polarization VH, see Fig. 3.20. No experiments could be carried out at this time in our institute; see however [27].

37
This section is closed with simple formulas derived from the above matrices to estimate the effect of a single edge in relation to the RCS peak value of a panel computed by PO. The panel may be of arbitrary polygonal structure with size A. The edge under consideration has the length L, is hit normally by the incident wave and is rotated around the y-axis by the angle $\phi_y$ (this is just the situation of edge 2 in Fig. 3.1). Relating the magnitude of the elements of the scattering matrix given by Eq. (3.4) to the peak value of the PO-scattering matrix, one obtains for horizontal polarization

$$(3.7) \quad r_{\text{HH}} = 20 \log \frac{L |D_m^f|}{kA},$$

and for vertical polarization

$$(3.8) \quad r_{\text{VV}} = 20 \log \frac{L |D_e^f|}{kA}.$$

The arguments of $D_m^f, D_e^f$ are $\psi_e = 270^\circ - \phi_y, \beta_e = 90^\circ$.

Using the values for $D_m^f$ and $D_e^f$ given in Fig. 2.4 for a half-plane ($n=2$) and in Fig. 2.5 for a step ($n=1.5$), the following table may be established choosing the test panel of this section as an example.

<table>
<thead>
<tr>
<th>$\phi_y$</th>
<th>$0^\circ$</th>
<th>$45^\circ$</th>
<th>$90^\circ$</th>
<th>$135^\circ$</th>
<th>$180^\circ$</th>
<th>$225^\circ$</th>
<th>$270^\circ$</th>
</tr>
</thead>
<tbody>
<tr>
<td>half-plane ($n=2$)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$r_{\text{HH}}$ [dB]</td>
<td>-36.1</td>
<td>-40.8</td>
<td>$-\infty$</td>
<td>-40.8</td>
<td>-36.1</td>
<td>-33.1</td>
<td>-30.1</td>
</tr>
<tr>
<td>$r_{\text{VV}}$ [dB]</td>
<td>-36.1</td>
<td>-33.1</td>
<td>-30.1</td>
<td>-33.1</td>
<td>-36.1</td>
<td>-40.8</td>
<td>$-\infty$</td>
</tr>
<tr>
<td>step ($n=1.5$)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$r_{\text{HH}}$ [dB]</td>
<td>-32.3</td>
<td>-36.8</td>
<td>-44.5</td>
<td>-42.8</td>
<td>-44.5</td>
<td>-36.8</td>
<td>-32.3</td>
</tr>
<tr>
<td>$r_{\text{VV}}$ [dB]</td>
<td>$-\infty$</td>
<td>-40.3</td>
<td>-34.8</td>
<td>-35.4</td>
<td>-34.8</td>
<td>-40.3</td>
<td>$-\infty$</td>
</tr>
</tbody>
</table>

Table 3.1 Level of edge diffraction effects related to the PO-peak value for the square test panel with edge length $L=5.08 \lambda$. 

Fig. 3.19 Theoretical RCS of the test panel for a $30^\circ$-cut, HV-polarization. Fig. 3.20 Theoretical RCS of the test panel for a $30^\circ$-cut, VH-polarization.
4. Application of the EC-Method for a Circular Cylinder and a Double Dihedral

4.1 Circular Cylinder

The test cylinder has a length of 5\( \lambda \) and a diameter of 1\( \lambda \). The wave length is 1128 mm. The broadside RCS amounts to 20 dBsm and the front face RCS to 9.9 dBsm. The cylinder is rotated around an axis vertically to its own axis. Theoretical results for the smooth cylinder based on PTD are published together with experimental results by Ufimtsev in [6]. The circumference of the cylinder was modeled (see Fig. 4.1) by 14 rectangular panels with dimensions 5\( \lambda \times 0.22 \lambda \). The deviation from the true cylinder surface was about \( \lambda/80 \).

Each of the front faces was modeled with 14 triangles. In the geometrical model artificial edges (\( n = 1.29 \)) arise between the rectangular panels. The natural circular edges (\( n = 1.5 \)) between the cylinder circumference and the front faces are approximated by straight lines of length 0.22\( \lambda \). Both types of edges are treated in the same way by the theory.

![Panel model of the test cylinder.](image1)

![RCS of the cylinder modeled by panels, PO-solution.](image2)

The PO-solution, insensitive to polarization, is presented in Fig. 4.2. The results of the experiment and of the PTD-theory for horizontal and vertical polarization are given in Fig. 4.3. The broadside peak of the theoretical curves, however, should not exceed 20 dBsm. The pictures of Fig. 4.4, finally present the results of the procedure outlined in this paper. For this special cut they should be identical to the results of Ufimtsev. This is the case except for the difference at broadside incidence and some deviations of minor practical interest.
Fig. 4.3 RCS of the smooth cylinder, experiment (---) and PTD-solution (—), left side: HH-polarization, right side: VV-polarization.

Fig. 4.4 RCS of the cylinder modeled by panels, PO- and EC-solution, left side: HH-polarization, right side: VV-polarization.
4.2 Double Dihedral

A double dihedral constructed on the basis of a cube with additional shado-wing surfaces, see Fig. 4.5, is rotated in an unconventional way as shown by Fig. 4.6. For $\alpha = 0^\circ$ the edges of the double dihedral make an angle of $45^\circ$ with the axis of rotation. The purpose was to generate a strong depolarized backscattered field. Previous PO-results are published for the main cut in [2] and for the diagonal cut within a restricted range of aspect angles in [4].

![Fig. 4.5](image)

Cube with additional shado-wing faces forming a double dihedral, dimensions in mm.

![Fig. 4.6](image)

**Fig. 4.6** Geometry and axis of rotation.

Figs. 4.7, 4.8 and 4.9 present the results for VH-polarization of experiment, of PO including double reflection and of PO + EC for the full range of aspect angles and a frequency of 15.5 GHz ($\lambda = 19.4$ mm). The measurements had to be arranged with great care since a wide dynamic range was needed. In addition, the exact positioning of the double dihedral according Fig. 4.6 caused major problems.

The structure of the pattern around $0^\circ$ is well represented by PO alone but the decrease is too rapid with increasing aspect angles. In addition, the spikes at $-235^\circ$, $-180^\circ$, $-125^\circ$, $-55^\circ$ and $+55^\circ$ are not predicted by PO. This, however, is the case when the EC-field is added to the PO-field. The spikes, therefore, are due to edge diffraction only.
Fig. 4.7 PO-result for the double dihedral, VH-polarization.

Fig. 4.8 Experimental result.

Fig. 4.9 PO- and EC-result.
5. Conclusion

The PO-method is frequently applied with good success to predict the RCS of large and complicated structures modeled by a collection of quadrangular and triangular panels. However, there are certain situations (specific structures, pattern cuts, polarizations) for which a correction of the PO-field by an edge diffracted field is required.

In this paper the concept of equivalent currents (EC) is applied. The fringe currents of Michaeli are used to derive the scattering matrix of an isolated edge with arbitrary length and orientation in an observer fixed coordinate system. The relationship of the method to other theories which are concerned with edge diffraction are summarized by some statements. The theory is applied for the analysis of a square flat plate, of a cylinder and of a double dihedral. All theoretical results are compared with measurements.

The square flat plate with edge length 5.08 \( \lambda \) is analysed in great detail. The results for the principal plane pattern are in excellent agreement for vertical polarization, independent of whether the plate was assumed with zero thickness or with 0.044 \( \lambda \) thickness (as used for the measurements). For horizontal polarization the agreement is unsatisfactory. In the case of zero thickness beyond that identical results for vertical and horizontal polarization are obtained. These effects are due to second order diffraction effects neglected in the EC-theory. This means that further effort is required if one is interested in the improvement of RCS calculations of an isolated plate.

The method is further applied for a cylinder with length 5 \( \lambda \) and diameter 1 \( \lambda \). The circumference was modeled by 14 rectangular panels, thus introducing artificial edges. Each of the front faces consisted of 14 triangular panels, thus modeling a circular edge by short straight edges. The RCS-results of the EC-method are in good agreement with those of the PTD-method and experiment, both applied for a smooth cylinder.

Finally the concept is used for a double dihedral which was positioned in such a way that strong depolarizations could occur during rotation. Also in this case the correction of the PO-field by the fringe current field was very efficient. The calculated RCS-values are again in good agreement with experimental results.

So one can conclude that the implementation of the presented procedure in a computer program would be efficient enough to treat edge diffraction effects with sufficient accuracy under practical viewpoints.

Acknowledgement

The author thanks Dr. Röde and Dr. K.H. Bethke for the RCS-measurements, Mr. D. Klement for the RCS-computations of the panel with finite thickness and Mrs. M. Malchow for typing the manuscript. Special thanks are expressed to Dr. A. Michaeli whom I was able to contact during the work.
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Appendix

The coefficients $D_e$, $D_m$, $D_{em}$ of the backscattering matrix are given by the theory of Michaeli [9]. For more details see also [15].

(A1) $D_v = D_{1v} - D^{co}_{1v} + D^{co}_{2v} - D^{co}_{2v}, \ v = e, m, em$

(A2) $D_{1e} = \frac{\sin \frac{\pi}{n}}{n} \frac{\sin }{n} \frac{\pi - \alpha}{n} - \cos \frac{\pi}{n} \frac{\sin }{n} \frac{\pi - \alpha}{n}$

(A3) $D_{1m} = \frac{\sin \omega_e}{n} \frac{\sin }{\sin \alpha_1} \frac{\pi - \alpha_1}{n} - \cos \frac{\pi - \alpha_1}{n} \frac{\sin }{n} \frac{\pi - \alpha_1}{n}$
\( \begin{align*} 
\text{(A4)} & \quad D_{1\text{em}} = \frac{1}{n} \cos \beta_e (\mu_1 + \cos \psi_e) \frac{\sin \frac{\pi - \alpha_1}{n}}{\sin \alpha_1} \frac{1}{\cos \frac{\pi - \alpha_1}{n} - \cos \frac{\psi_e}{n}}, \\
\text{(A5)} & \quad D_{1\text{e}}^{\text{po}} = -U(\pi - \psi_e) \frac{\sin \psi_e}{\mu_1 + \cos \psi_e}, \\
\text{(A6)} & \quad D_{1\text{m}}^{\text{po}} = D_{1\text{e}}^{\text{po}}, \\
\text{(A7)} & \quad D_{1\text{em}}^{\text{po}} = 0, \\
\text{(A8)} & \quad \mu_1 = \cos \psi_e - \frac{2}{\tan^2 \beta_e}, \\
\text{(A9)} & \quad \alpha_1 = \arccos \mu_1, \\
\text{(A10)} & \quad U(x) = \begin{cases} 
1 & \text{for } x > 0, \\
0 & \text{for } x < 0.
\end{cases}
\end{align*} \)

The coefficients \( D_{2v}^{\text{po}} \), \( D_{2v}^{\text{po}} \) result from \( D_{1v}^{\text{po}} \), \( D_{1v}^{\text{po}} \) by the following transformations:

\( \text{(A11)} \quad \psi_e \rightarrow n\pi - \psi_e, \quad \beta_e \rightarrow \pi - \beta_e. \)
Application of Parallel Processing
to a Surface Patch / Wire Junction EFIE Code

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ABSTRACT

A surface patch / wire junction EFIE method of moment algorithm, JUNCTION, developed by the University of Houston has been implemented in a transputer based parallel processing environment installed in a personal computer. This paper addresses transputer hardware and software options, the JUNCTION algorithm, techniques for parallelizing matrix analysis algorithms, and performance results. The transputer array was found to provide a flexible, low-cost, high performance desktop computing environment for method of moment analysis.

1.0 INTRODUCTION

This paper presents the application of parallel processing techniques to an existing computational electromagnetic (CEM) code. The goal was to demonstrate that high performance computing (HPC) can improve the utility and efficiency of computational techniques used in electromagnetic ship analysis [Li, 1988]. The parallel processing platform was an array of four transputers on a board inside an IBM-compatible PC. The transputers were run using the ParaSoft EXPRESS operating environment [ParaSoft, 1990]. This operating environment was chosen to allow portability of the code to other HPC platforms and minimize the number of required programming changes to the CEM code. The selected CEM code was the method of moment (MoM) algorithm JUNCTION developed by the University of Houston [Hwu et al, 1988; Wilton et al, 1988].

2.0 PARALLEL PROCESSING

The concept of parallel processing has been around for a long time but only recently was its utility generally recognized. By 1986 more than a dozen companies were either selling or in the process of building parallel processors [Tazelaar, 1988].

2.1 Transputer hardware

The hardware platform chosen was an array of four transputers on a motherboard which could be installed inside a PC. This decision was based on the low cost, ease of use, and flexibility of a transputer based system. Transputer systems provide an inexpensive entry into the world of parallel processing. For this entry level application a decision was made to purchase only four floating point transputer modules (TRAMs) each with one Megabyte of external RAM. These size 1 TRAMs are an industry standard and provide sufficient memory for most applications at a reasonable cost and in a compact package. The T800 transputer used has a peak performance of 1.5 Mflops.

2.2 Transputer Software

There are several different ways to program an array of transputers. Each method has trade-offs in portability, performance, and ease of use. The parallel language, Occam, was developed concurrently with the transputer. Parallel versions of high level languages such as Fortran, C, Pascal, and Modula 2 can be obtained. The parallel operating environment EXPRESS (ParaSoft Corporation) is available for transputer systems.
2.2.1 Occam

There are a number of advantages to using Occam for programming a transputer system. Since Occam was developed simultaneously with the transputer, it is one of the few languages designed for concurrency. Occam has the performance and efficiency of assembly language. Occam can be used as a harness to link modules written in other industry standard languages. Its other features include: well implemented timing capabilities, straightforward control of process scheduling, and a high level of structure.

The main disadvantage to using Occam is that programs written in Occam are not portable to other parallel computers. Existing codes written in other languages need to be completely recoded. Occam works only with the transputer. Other disadvantages are that Occam does not support many features of other high level languages (no recursion or dynamic memory allocation), use of it requires learning a new language, and Occam may not be supported in the future. In the trade-off space (portability versus performance versus ease of use) Occam scores high only in performance.

2.2.2 High Level Languages

The advantage of using a parallel version of a high level language such as Fortran, C, Pascal, or Modula 2 is that one can use a language with which one is already familiar. All that is required is to learn the parallel extensions. Existing sequential code can be ported over.

There are, however, a number of disadvantages to using a parallel version of a high level language. Programming an array of transputers requires the user to develop both a host program and a node program. The host program runs on the host processor and handles I/O calls and manages the other processors. The node program runs on all the other processors and does the bulk of the computations. The user is required to maintain two programs instead of the usual one program. This can make program development and maintenance somewhat complex. Performing I/O, making system calls, and measuring timing can be very difficult. Debuggers for parallel versions of high level languages are starting to appear, but they are not widely available. Debugging parallel programs is very difficult. In addition to all this, parallel versions of high level languages do not offer the performance that one can get from Occam.

2.2.3 ParaSoft EXPRESS

Parallel operating environments such as ParaSoft’s EXPRESS allow the user the convenience of using a familiar high level language while mitigating some of the disadvantages to parallel high level languages which were mentioned in the previous section. EXPRESS is based on the CUBIX programming model which was developed at CalTech. EXPRESS is available for both Fortran and C. There is no need to develop both a host and node program. Instead only one program (which runs on all processors) needs to be developed and maintained. EXPRESS does not include a compiler so one of the parallel high level language compilers mentioned above is still needed to compile and link the code. During linking the EXPRESS library is linked into the user’s source code. The EXPRESS library provides simple Fortran or C language function calls to handle I/O and system calls. EXPRESS also provides timing capabilities, a source level debugger, and a performance monitor. The number of processors being used does not have to be hardwired into the code. Instead this is specified at runtime by a switch in the run command. In addition, EXPRESS is available for a wide variety of parallel machines, not just the transputer. Currently EXPRESS is available for: the multi-headed IBM 3090 (AIX) and the multi-headed CRAY (UNICOS) mainframes; the Intel iPSC 2 and iPSC 860; the nCUBE 1 and 2; PC, MAC, and SUN hosts for transputers; and the IBM RS/6000, Silicon Graphics, and SUN workstations. All this leads to very portable source code.
The only significant disadvantage to EXPRESS is that it possibly degrades system performance to some extent. However, this disadvantage is more than offset by the advantage of having portable code. For these reasons EXPRESS by ParaSoft was chosen for the operating environment. Portability was the key feature since the ultimate goal of this effort was to run the code on a high performance computer. The language chosen was Fortran since this is the principal language of the JUNCTION method of moments code. For obvious reasons, it was desirable to make as few changes to the existing code as possible.

3.0 The JUNCTION METHOD OF MOMENTS CODE

The JUNCTION computer code invokes the method of moments to solve a coupled electric field integral equation (EFIE) for the currents induced on an arbitrary configuration of perfectly conducting bodies and wires [Wilton et al, 1989]. There are three principal advantages to the JUNCTION formulation. First, the EFIE formulation for the surfaces of bodies, in contrast to the magnetic field integral equation (MFIE), applies to open bodies. Second, JUNCTION allows voltage and load conditions to be easily specified at terminals defined on the structure. Third, the triangular patches of JUNCTION are the simplest planar surfaces which can be used to model arbitrary surfaces and boundaries, and triangular patches permit patch densities to be varied locally so as to model a rapidly varying current distribution.

The method of moments is a numerical procedure for solving field integral equations [ Harrington, 1968]. Basis functions are chosen to represent the unknown currents. Testing functions are chosen to enforce the integral equation on the surface of the conducting structure. With the choice of basis and testing functions a matrix approximating the integral equation is derived. In JUNCTION a system of \( N \) linear equations results, where \( N = N_B + N_w + N_J \).

\[
\begin{bmatrix}
[Z^{BB}] & [Z^{BW}] & [Z^{BJ}]
\end{bmatrix}
\begin{bmatrix}
[I^B]
\end{bmatrix}
= \begin{bmatrix}
[E^B]
\end{bmatrix}
\]

\[
\begin{bmatrix}
[Z^{WW}] & [Z^{WW}] & [Z^{WJ}]
\end{bmatrix}
\begin{bmatrix}
[I^W]
\end{bmatrix}
= \begin{bmatrix}
[E^W]
\end{bmatrix}
\]

\[
\begin{bmatrix}
[Z^{JJ}]
\end{bmatrix}
\begin{bmatrix}
[I^J]
\end{bmatrix}
= \begin{bmatrix}
[E^J]
\end{bmatrix}
\]

where \( Z \) is the impedance matrix, \( I \) is the current vector, and \( E \) is the excitation vector. The superscripts are \( B \) for body elements, \( W \) for wire elements, and \( J \) for junctions between wires and bodies. The analytical expressions for each of the individual submatrices are different. Solution of the linear system of equations yields the set of unknown coefficients used in the representation of the surface, wire, and junction currents. Once these currents are known, the scattered field or any other electromagnetic quantity of interest may be determined.

4.0 IMPLEMENTATION OF SEQUENTIAL PROGRAM

The first step in parallelizing an existing program is to verify the sequential version of the program. This is done by running the program on a serial computer. Once this has been successfully done, the next step is to get the program to run on a single processor on the parallel computer. This is necessary to eliminate any problems caused by compiler differences between the serial compiler and the parallel compiler and also to "check out" the hardware. Only after this has been completed can the parallelization of the program be considered.
JUNCTION was broken into four separate programs: DATGN, GEOMETRY, CURRENT, and FIELD. DATGN creates input data set files. GEOMETRY calculates the geometrical parameters. CURRENT computes the currents and impedance. FIELD computes charges, near fields, far fields, radar cross sections, and power gain. The computationally intensive programs are CURRENT and FIELD. This paper is concerned with the parallelizing of CURRENT.

From a computational standpoint CURRENT consists of filling the impedance matrix and the excitation vector, and then solving for the current vector. The current vector gives the currents on the bodies, wires and junctions. The methods used are standard matrix manipulation techniques. Once the matrix is filled, Gaussian elimination is performed using LU decomposition with partial pivoting to factor the matrix. Finally, the current vector is solved for by using backward and forward substitution. The library subroutines used are the LINPACK routines CGEFA and CGESL (Dongarra et al, 1979). CGEFA factors a general complex matrix. CGESL solves a complex matrix equation using the output from CGEFA.

5.0 PARALLEL MATRIX FACTOR AND SOLVE

5.1 Parallel LINPACK

When this effort was initiated, general parallel versions of the LINPACK subroutines were not available. This is because parallel subroutines are by their nature very machine specific and one of the goals of the LINPACK project was for the subroutines to be machine independent.

On sequential computers the number of computations required to factor a matrix into a product of triangular matrices is of the order \( n^3 \) where \( n \) is the dimension of the matrix. In comparison, the number of computations required for the triangular solution are of the order \( n^2 \). Because of this, unless multiple solutions are required, most effort has focused on optimizing the factorization algorithm.

On parallel computers good efficiency is more difficult to obtain for matrix solving compared to matrix factoring. This is because much more interprocessor communication is required during solving. Solving is inherently a finer grain process than factoring. Various parallel algorithms have been proposed. Each one has its advantages and disadvantages which are dependent on the number of processors being used, the size of the problem being solved, and the relative cost of communication and computation.

The matrix columns were distributed onto the processors using an interleaving technique known as column wrap mapping. Column wrap mapping has good load balancing properties and gives excellent performance when doing LU decomposition.

Intel Corporation provided the source code for Intel's parallel versions of DGEFA and DGESL from LINPACK. DGEFA and DGESL are double precision non-complex versions of CGEFA and CGESL, respectively. Included were two different matrix solving routines: a cyclic version and a wavefront version of DGESL. The differences between these versions are described below. These routines were all developed for Intel's iPSC parallel computer so a number of modifications were needed to implement them under EXPRESS on the transputer array. The matrix factoring routine was straightforward to parallelize and provided very high efficiencies.

The difficult problem was the solution of the lower triangular linear system

\[
Lx = b,
\]
where \( L \) is a lower triangular matrix of order \( n \), \( b \) is a known vector of dimension \( n \), and \( x \) is the unknown solution vector of dimension \( n \). The sequential version of LINPACK solves this by forward substitution using the following doubly nested loop:

\[
\begin{align*}
\text{for } j = 1 \text{ to } n \\
x_j &= b_j/L_{jj} \\
\text{for } i = j+1 \text{ to } n \\
b_i &= b_i - x_jL_{ij}
\end{align*}
\]

There are a number of ways to parallelize this problem on a distributed memory parallel computer. Several algorithms have been developed. Two of the most popular are the \textit{wavefront} algorithm and the \textit{cyclic} algorithm [Heath et al, 1988]. These algorithms assume the matrix is column wrap mapped onto the processors.

The wavefront algorithm breaks the updating of \( b \) into segments and pipelines the segments through the processors in a wavefront fashion. The \( n \)-vector \( z \) is used to accumulate the updates of \( b \) so that the components of \( b \) remain distributed among the processors. The size of the circulated segments is an adjustable parameter that controls the granularity of the algorithm and therefore affects performance. The optimal value of the segment size depends on the characteristics of the hardware.

The wavefront algorithm is written in pseudo-code as:

\[
\begin{align*}
\text{for } j \in \text{mycols} \\
\text{for } k = 1 \text{ to } \# \text{ segments} \\
&\text{receive segment} \\
&\text{if } k = 1 \text{ then} \\
&\quad x_j = (b_j - z_j)/L_{jj} \\
&\quad \text{segment} = \text{segment} - \{z_j\} \\
&\text{for } z_i \in \text{segment} \\
&\quad z_i = z_i + x_jL_{ij} \\
&\text{if } |\text{segment}| > 0 \text{ then} \\
&\quad \text{send segment to processor map}(j+1)
\end{align*}
\]

The cyclic algorithm is similar to the wavefront algorithm in that they both send a segment \( z \) between processors. However, the cyclic algorithm circulates a single segment of the fixed size \( p-1 \), where \( p \) is the number of processors. The name \textit{cyclic} comes from the segment cycling through all other processors before returning to a given processor. The updates computed by the processor while the segment is circulating elsewhere are stored in the vector \( t \). The cyclic algorithm is written in pseudo-code as:

\[
\begin{align*}
\text{for } j \in \text{mycols} \\
&\text{receive segment} \\
x_j = (b_j - z_j - t_j)L_{jj} \\
\text{segment} = \text{segment} - \{z_j\} \\
\text{for } z_i \in \text{segment} \\
&\quad z_i = z_i + t_i + x_jL_{ij} \\
&\quad z_{j+p-1} = t_{j+p-1} + x_jL_{j+p-1,i}
\end{align*}
\]
segment = segment ∪ \{z_{j+p-1}\}

send segment to processor \text{map}(j+1)

for i = j + p to n

\[ t_i = t_i + x_i L_{ij} \]

Theoretical analysis shows that the cyclic algorithm performs best on a small number of processors whereas the wavefront algorithm performs best on a large number of processors.

5.2 Implementation of parallel LINPACK under EXPRESS

A number of modifications were needed to the Intel parallel LINPACK routines in order to implement them on the transputer array operating under ParaSoft EXPRESS. These modifications fell into three categories:

1. converting double precision variables to complex variables,
2. using complex LINPACK routines instead of the equivalent double precision LINPACK routines,
3. using EXPRESS communication functions/routines instead of iPSC communication functions/routines.

5.3 Results

Performance measurements were made of both the parallel matrix factor routine and the two parallel matrix solve routines. Timings were made using one, two, three and four processors. The timing measurements were converted to the standard performance measurement parameters, speed-up and efficiency. Speed-up is the ratio of the execution time of the algorithm on a single processor to the execution time of the parallel algorithm on the \( n \) processors. Efficiency is the speed-up divided by \( n \) and expressed as percent.

Due to memory limitations, examples with more than 200 unknowns could not be run on a single transputer. Several sample problems were developed to test out the parallel factor and solve routines. The four sample problems had the following number of unknowns: 104, 116, 155, and 189.

5.3.1 Factor

Figure 5-1 shows the timing results on the parallel version of CGEFA for the four sample problems. Figure 5-2 shows speed-up and figure 5-3 shows efficiency. The important issue is that as the number of unknowns increases the parallel matrix factoring routine becomes more efficient. This is expected since small problems do not achieve good load balancing and cannot benefit greatly from parallel computation. An efficiency of over 90\% is considered very good.
Figure 5-1. Timing for parallel matrix factoring routine.

Figure 5-2. Speedup achieved for parallel matrix factoring routine.
Figure 5-3. Efficiency achieved for parallel matrix factoring routine.

5.3.2 Solve

The performance of the two parallel matrix solve routines were measured for all the example problems. The results are shown here for the largest problem, the one with 189 unknowns. Table 5-1 shows the results for the cyclic algorithm.

Table 5-1. Performance results for cyclic algorithm for 189 unknowns.

<table>
<thead>
<tr>
<th>Number of Processors</th>
<th>Time, seconds</th>
<th>Speedup</th>
<th>Efficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.621</td>
<td>1.00</td>
<td>100%</td>
</tr>
<tr>
<td>2</td>
<td>0.367</td>
<td>1.69</td>
<td>84.5%</td>
</tr>
<tr>
<td>3</td>
<td>0.257</td>
<td>2.42</td>
<td>80.7%</td>
</tr>
<tr>
<td>4</td>
<td>0.207</td>
<td>3.00</td>
<td>75.0%</td>
</tr>
</tbody>
</table>

The performance results for the wavefront algorithm are complicated by the presence of the adjustable segment size parameter. This segment size parameter can range from 1 to the number of unknowns. Table 5-2 shows the optimal performance segment size and time. The optimal performance segment size is a function of the number of processors (as well as the number of unknowns). In addition, by comparing the timing results for the wavefront algorithm with the results for the cyclic algorithm it can be seen that even with the optimum segment size the performance for the wavefront algorithm is much worse than the cyclic algorithm. For these two reasons the wavefront algorithm was rejected for use on the size of problem which could be run on the four
processor transputer array. However, it must be noted that the performance results could turn out to be very different on a high performance computer with many more processors running larger problems.

Table 5-2. Performance results for wavefront algorithm for 189 unknowns.

<table>
<thead>
<tr>
<th>Number of Processors</th>
<th>Optimal segment size</th>
<th>Time, seconds</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>95</td>
<td>0.741</td>
</tr>
<tr>
<td>2</td>
<td>72</td>
<td>0.555</td>
</tr>
<tr>
<td>3</td>
<td>34</td>
<td>0.448</td>
</tr>
<tr>
<td>4</td>
<td>24</td>
<td>0.380</td>
</tr>
</tbody>
</table>

6.0 PARALLEL MATRIX FILLING

Developing a highly efficient general routine for parallel matrix filling was much more difficult in comparison to the work required to parallelize the matrix factor and solve routines. This was due to several reasons: the large variety of problem geometries being solved, shared calculations between matrix columns, memory limitations, and concurrent filling of the excitation vector.

6.1 Sequential matrix filling

In JUNCTION the columns of the matrix correspond to source points on the structure. The rows of the matrix correspond to observation points on the structure. The body columns/rows come before the wire columns/rows, but otherwise the ordering of the columns/rows depends on how the problem was specified in the input data portion of the code. The junction points are scattered within the wire portion of the matrix. Wire columns correspond to nodes on the wires whereas body columns correspond to sides of triangular patches on the surfaces. A junction is a node on a wire, hence its location is in the wire portion of the matrix.

6.2 Column mapping techniques

There are a wide variety of techniques which can be used to map the columns of the matrix onto the processors. A column wrap mapping technique was used for matrix factoring and solving as described in the previous chapter.

It quickly became apparent that column wrap mapping was not necessarily the best method to use for parallelizing the matrix filling portion of the code. Adjacent columns of the matrix are often associated with each other. For bodies adjacent columns are usually either on the same face or on an attached face. For wires adjacent columns often refer to nodes which are joined by the same segment. In JUNCTION calculations are made with respect to faces and segments, not edges and nodes. This avoids duplication of work in a serial computation. Alternative methods for mapping the columns onto the processors were needed. (Note: mapping the rows onto the processors, instead of the columns, was never considered since matrix factoring required the columns to be already mapped onto the processors).
Alternative mapping techniques were devised and evaluated. The techniques and performance results are described below in the section on structural dependencies. It was found that using a different mapping technique to fill the matrix did not compromise the output from matrix solving. The only effect was to scramble the output vector. The output vector is easily and rapidly unscrambled at the end of the matrix solve routines. Unscrambling the output vector at the end is much faster than unscrambling the matrix before the matrix factor routine, since much less interprocessor communication is required for the former.

6.3 Structural dependencies

The performance of the parallel matrix filling algorithm was found to be very dependent on the structure being analyzed. The first types of problems analyzed were for homogeneous structures. Homogeneous structures are defined as having either all body elements or all wire elements. Next, heterogeneous structures were evaluated. Heterogeneous structures are defined as having a mixture of body, wire, and junction elements.

6.3.1 Homogeneous structures

Six types of structures were considered: straight wires, cylinders, plates, cones, disks, or spheres. The performance of the parallel matrix filling routines was evaluated for homogeneous problems made up of each of these types of structures. Two column mapping techniques were compared for each structure type. The two techniques used were column wrap mapping and column block mapping. Column wrap mapping consists of interleaving the columns onto the processors. It is described in detail in Section 6.2. Column block mapping consists of distributing the columns onto the processors using contiguous blocks of columns rather than individual columns. As an example, if 4 processors were being used to fill a matrix with 100 columns, using column block mapping would mean that the first processor would fill the first 25 columns, the second processor would fill the second 25 columns, and so on.

Four sample problems were developed for each structure type. The number of unknowns ranged from 40 to 200. Each sample problem was run on both a single transputer and four transputers. Measurements were made of both matrix filling time and matrix solving time for both column wrap mapping and column block mapping.

Table 6-3 shows the measured times and calculated efficiencies for matrix filling for the different structures using wrap and block mapping.

For a single processor it is seen that an all wire homogeneous matrix takes longer to fill than an all body homogeneous matrix of the same size. On average an all wire matrix took about 38% longer to fill than the same size all body matrix. Improvements are being been made in JUNCTION to improve the speed of calculation for wires. The cylinder, plate, cone, disk and sphere matrices all took about the same time to fill a given size matrix.

When the sample problems were run on four processors, distinct structural dependencies emerged for filling times and efficiencies. It is important to note that column block mapping always performed better than column wrap mapping. Column block mapping filling efficiencies were almost always greater than 70%, whereas column wrap mapping filling efficiencies were always between 40 and 60%. It is also interesting to note that the different structures performed differently in filling efficiencies. For column wrap mapping wires performed best followed by plates, cones, spheres, disks, and, then, cylinders. For column block mapping wires again performed best followed by cylinders, plates, cones, spheres, and, then, disks. These results are due to the nature of the physical connectivity of the various structures and how their elements are distributed to the columns.
Table 6-3. Matrix filling times/efficiencies for various structures.

<table>
<thead>
<tr>
<th>Structure Type</th>
<th>Number of Unknowns</th>
<th>1 Processor TIME, sec</th>
<th>4 Processors Wrap Mapping TIME</th>
<th>EFFIC</th>
<th>TIME</th>
<th>EFFIC</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>TIME</td>
<td>EFFIC</td>
<td>TIME</td>
<td>EFFIC</td>
<td>Wrap Mapping</td>
</tr>
<tr>
<td>WIRE</td>
<td>47</td>
<td>7.38</td>
<td>55.6%</td>
<td>3.32</td>
<td>55.6%</td>
<td>1.97</td>
</tr>
<tr>
<td></td>
<td>97</td>
<td>30.78</td>
<td>54.7%</td>
<td>14.06</td>
<td>54.7%</td>
<td>8.08</td>
</tr>
<tr>
<td></td>
<td>147</td>
<td>70.51</td>
<td>55.9%</td>
<td>31.54</td>
<td>55.9%</td>
<td>17.95</td>
</tr>
<tr>
<td></td>
<td>197</td>
<td>126.06</td>
<td>55.4%</td>
<td>56.91</td>
<td>55.4%</td>
<td>32.22</td>
</tr>
<tr>
<td>CYLINDER</td>
<td>56</td>
<td>8.56</td>
<td>43.1%</td>
<td>4.97</td>
<td>43.1%</td>
<td>2.80</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>27.04</td>
<td>42.8%</td>
<td>15.78</td>
<td>42.8%</td>
<td>7.94</td>
</tr>
<tr>
<td></td>
<td>152</td>
<td>55.95</td>
<td>42.8%</td>
<td>32.69</td>
<td>42.8%</td>
<td>15.68</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>95.17</td>
<td>42.8%</td>
<td>55.62</td>
<td>42.8%</td>
<td>25.99</td>
</tr>
<tr>
<td>PLATE</td>
<td>40</td>
<td>5.12</td>
<td>48.1%</td>
<td>2.66</td>
<td>48.1%</td>
<td>1.66</td>
</tr>
<tr>
<td></td>
<td>96</td>
<td>25.11</td>
<td>49.5%</td>
<td>12.69</td>
<td>49.5%</td>
<td>7.88</td>
</tr>
<tr>
<td></td>
<td>133</td>
<td>45.7</td>
<td>44.0%</td>
<td>25.95</td>
<td>44.0%</td>
<td>14.01</td>
</tr>
<tr>
<td></td>
<td>176</td>
<td>76.87</td>
<td>49.1%</td>
<td>39.15</td>
<td>49.1%</td>
<td>21.92</td>
</tr>
<tr>
<td>CONE</td>
<td>52</td>
<td>7.15</td>
<td>43.8%</td>
<td>4.08</td>
<td>43.8%</td>
<td>2.54</td>
</tr>
<tr>
<td></td>
<td>100</td>
<td>24.09</td>
<td>43.4%</td>
<td>13.88</td>
<td>43.4%</td>
<td>7.36</td>
</tr>
<tr>
<td></td>
<td>155</td>
<td>55.85</td>
<td>47.9%</td>
<td>29.12</td>
<td>47.9%</td>
<td>16.56</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>90.52</td>
<td>48.6%</td>
<td>46.58</td>
<td>48.6%</td>
<td>25.31</td>
</tr>
<tr>
<td>DISK</td>
<td>49</td>
<td>6.60</td>
<td>43.8%</td>
<td>3.77</td>
<td>43.8%</td>
<td>2.77</td>
</tr>
<tr>
<td></td>
<td>98</td>
<td>25.19</td>
<td>46.0%</td>
<td>13.7</td>
<td>46.0%</td>
<td>10.42</td>
</tr>
<tr>
<td></td>
<td>150</td>
<td>55.65</td>
<td>46.6%</td>
<td>29.84</td>
<td>46.6%</td>
<td>20.05</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>97.01</td>
<td>45.7%</td>
<td>53.12</td>
<td>45.7%</td>
<td>34.33</td>
</tr>
<tr>
<td>SPHERE</td>
<td>60</td>
<td>9.06</td>
<td>47.0%</td>
<td>4.82</td>
<td>47.0%</td>
<td>3.23</td>
</tr>
<tr>
<td></td>
<td>90</td>
<td>19.75</td>
<td>46.4%</td>
<td>10.65</td>
<td>46.4%</td>
<td>6.85</td>
</tr>
<tr>
<td></td>
<td>168</td>
<td>66.56</td>
<td>43.8%</td>
<td>37.98</td>
<td>43.8%</td>
<td>20.27</td>
</tr>
<tr>
<td></td>
<td>198</td>
<td>91.84</td>
<td>47.8%</td>
<td>47.99</td>
<td>47.8%</td>
<td>28.96</td>
</tr>
</tbody>
</table>

of the matrix. Again, in JUNCTION calculations are made with respect to faces and segments. To avoid duplication of computation on the processors it is advantageous to have all the edge computations for a given face to be on the same processor. For a wire both nodes of a given segment should be on the same processor.

Table 6-4 shows the measured times and calculated efficiency for matrix factoring and solving for the different structures.

In matrix factoring and solving there are no structural dependencies. It should also be noted that as the number of unknowns increases, the factoring and solving efficiencies increase. For 200 unknowns factoring and solving efficiency is greater than 95% percent. Matrix filling efficiency also increases as the number of unknowns increases.
Table 6-4. Matrix factor and solve times/efficiency for various structures.

<table>
<thead>
<tr>
<th>Structure Type</th>
<th>Number of Unknowns</th>
<th>1 Processor TIME, sec</th>
<th>4 Processors TIME</th>
<th>EFFIC</th>
</tr>
</thead>
<tbody>
<tr>
<td>WIRE</td>
<td>47</td>
<td>0.66</td>
<td>0.22</td>
<td>75.0%</td>
</tr>
<tr>
<td></td>
<td>97</td>
<td>5.34</td>
<td>1.48</td>
<td>90.2%</td>
</tr>
<tr>
<td></td>
<td>147</td>
<td>18.09</td>
<td>4.81</td>
<td>94.0%</td>
</tr>
<tr>
<td></td>
<td>197</td>
<td>42.94</td>
<td>11.19</td>
<td>95.9%</td>
</tr>
<tr>
<td>CYLINDER</td>
<td>56</td>
<td>1.13</td>
<td>0.35</td>
<td>80.7%</td>
</tr>
<tr>
<td></td>
<td>104</td>
<td>6.70</td>
<td>1.81</td>
<td>92.5%</td>
</tr>
<tr>
<td></td>
<td>152</td>
<td>20.22</td>
<td>5.34</td>
<td>94.7%</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>45.36</td>
<td>11.79</td>
<td>96.2%</td>
</tr>
<tr>
<td>PLATE</td>
<td>40</td>
<td>0.42</td>
<td>0.15</td>
<td>70.0%</td>
</tr>
<tr>
<td></td>
<td>96</td>
<td>5.12</td>
<td>1.44</td>
<td>88.9%</td>
</tr>
<tr>
<td></td>
<td>133</td>
<td>13.35</td>
<td>3.58</td>
<td>93.2%</td>
</tr>
<tr>
<td></td>
<td>176</td>
<td>30.55</td>
<td>8.04</td>
<td>95.0%</td>
</tr>
<tr>
<td>CONE</td>
<td>52</td>
<td>0.92</td>
<td>0.29</td>
<td>79.3%</td>
</tr>
<tr>
<td></td>
<td>100</td>
<td>5.97</td>
<td>1.63</td>
<td>91.6%</td>
</tr>
<tr>
<td></td>
<td>155</td>
<td>21.45</td>
<td>5.63</td>
<td>95.2%</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>45.41</td>
<td>11.79</td>
<td>96.3%</td>
</tr>
<tr>
<td>DISK</td>
<td>49</td>
<td>0.78</td>
<td>0.25</td>
<td>78.0%</td>
</tr>
<tr>
<td></td>
<td>98</td>
<td>5.60</td>
<td>1.54</td>
<td>90.9%</td>
</tr>
<tr>
<td></td>
<td>150</td>
<td>19.48</td>
<td>5.11</td>
<td>95.3%</td>
</tr>
<tr>
<td></td>
<td>200</td>
<td>45.32</td>
<td>11.78</td>
<td>96.2%</td>
</tr>
<tr>
<td>SPHERE</td>
<td>60</td>
<td>1.37</td>
<td>0.41</td>
<td>83.5%</td>
</tr>
<tr>
<td></td>
<td>90</td>
<td>4.38</td>
<td>1.21</td>
<td>90.5%</td>
</tr>
<tr>
<td></td>
<td>168</td>
<td>27.10</td>
<td>7.11</td>
<td>95.3%</td>
</tr>
<tr>
<td></td>
<td>198</td>
<td>44.02</td>
<td>11.41</td>
<td>96.5%</td>
</tr>
</tbody>
</table>

6.3.2 Heterogeneous structures

The heterogeneous structures that were evaluated consisted mostly of square plates with numerous attached wires. This allowed the evaluation of matrices with various numbers of body, wire, and junction columns.

After evaluating a large number of heterogeneous problems a number of observations were made. These observations are summarized below:

- The junction columns are scattered within the wire portion of the matrix. The distribution of the junction columns depends on how the problem is initially specified. Fortunately, it is fairly straightforward to convert from junction number to matrix column number.

- A junction can connect to between one and six faces on a body. A quirk of JUNCTION is that the code will only recognize a maximum of one junction attached to a given face.
To achieve maximum efficiency the filling of a junction column should be done by the same processor which is filling the columns associated with the faces to which the junction is attached, since a large part of the calculations are in common. The problem is that the associated body columns are sometimes scattered all through the matrix.

The time to fill a matrix is a function of many factors including: relative/absolute number of wire, body and junction unknowns; number of separate wires; number of segments on each wire; locations of wires and junctions relative to a body; number of faces connected to each junction.

Developing a general method for balancing the work load and achieving optimum efficiency for a heterogeneous problem is very complicated, even for the simple problems studied involving a single plate with various attached wires. If the choice is between block and wrap mapping, block mapping gives better results. (For the problems studied, block mapping was 62 - 93% efficient, whereas wrap mapping was 41 - 52% efficient).

Based on the above observations a decision was made to implement two additional column mapping techniques. Both techniques are modifications of the standard block mapping technique.

The first technique is called random mapping. Random mapping is block mapping with one twist: instead of the junction columns staying grouped with the wire columns, the junction columns are filled by the processors which are filling the body columns of the matrix. The columns are redistributed to keep a balanced number of columns on each processor. As an example, suppose a matrix has 50 body columns, 50 wire columns, and 6 junctions, and the problem is being run on 4 processors. Under standard column block mapping the first 25 body columns would be filled by the first processor, the second 25 body columns would be filled by the second processor, the first 25 wire columns would be filled by the third processor, and the second 25 wire columns would be filled by the fourth processor. The junction columns, being wire columns also, would be distributed on the third and fourth processors. Under random mapping the six junction columns would first be distributed to the first two processors in the following manner: junction 1 column to first processor, junction 2 column to second processor, junction 3 column to first processor, junction 4 column to second processor, and so on. The first and second processors would each have 3 junction columns. Block processing would then be used to distribute the body and remaining wire columns, so that each processor would finish up with 25 total columns as before.

The second technique is called 1st order mapping. This is similar to the random mapping described above except now the junction columns are assigned to the body processors using knowledge of which processor will be calculating the majority of columns associated with the faces to which that junction is attached. This could potentially cause more junction columns to end up on one of the body processors than another, but the improvement in performance could be significant.

An analysis of matrix filling efficiency was run on four processors for the problem of a square plate with 8 wires attached to it. The problem involved 96 body unknowns, 96 wire unknowns, and 8 junctions. The four different column mapping techniques were used. The results are shown in table 6-5.

The results show that random mapping only gives a slight improvement over block mapping, but 1st order mapping gives a significant improvement over block mapping.
Table 6-5. Efficiencies of various mapping techniques.

<table>
<thead>
<tr>
<th>Mapping Technique</th>
<th>Efficiency</th>
</tr>
</thead>
<tbody>
<tr>
<td>WRAP</td>
<td>40.5%</td>
</tr>
<tr>
<td>BLOCK</td>
<td>61.8%</td>
</tr>
<tr>
<td>RANDOM</td>
<td>63.9%</td>
</tr>
<tr>
<td>1ST ORDER</td>
<td>76.9%</td>
</tr>
</tbody>
</table>

7.0 RESULTS AND CONCLUSIONS

7.1 Performance Comparisons

Comparisons were run for the micro-Vax, a single transputer, the four transputer array, and the Convex Model C-220. The Convex is a mini-supercomputer which can do some automatic vectorization.

Sample data sets were generated for both bodies and wires. Eight body data sets were generated with 104, 152, 200, 248, 296, 356, 404, and 452 unknowns. Eight wire data sets were generated with 47, 97, 147, 197, 247, 297, 347, and 397 unknowns. For each data set and each hardware platform timing measurements were made for both matrix filling and matrix factoring and solving. On the four transputer array the data sets were run using both column wrap mapping and column block mapping. On the Convex measurements were made of both CPU time and actual elapsed time. At the time the measurements were made there were 13 other users on the Convex and elapsed time averaged about 3.5 times longer than CPU time. This should be kept in mind when making comparisons between the various hardware platforms. Results are shown in tables 7-1 through 7-4.

Table 7-1. Time to fill all-body matrix, seconds.

<table>
<thead>
<tr>
<th>Number of Unknowns</th>
<th>COMPUTER PLATFORM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Micro-Vax</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>104</td>
<td>73.7</td>
</tr>
<tr>
<td>152</td>
<td>151.7</td>
</tr>
<tr>
<td>200</td>
<td>257.5</td>
</tr>
<tr>
<td>248</td>
<td>394.0</td>
</tr>
<tr>
<td>296</td>
<td>555.9</td>
</tr>
<tr>
<td>356</td>
<td>801.6</td>
</tr>
<tr>
<td>404</td>
<td>1027.8</td>
</tr>
<tr>
<td>452</td>
<td>1282.3</td>
</tr>
</tbody>
</table>
Table 7-2. Time to factor & solve all-body matrix, seconds.

<table>
<thead>
<tr>
<th>Number of Unknowns</th>
<th>COMPUTER PLATFORM</th>
<th>4 Xputer</th>
<th>CONVEX</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Micro-Vax</td>
<td>1 Xputer</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wrap Map</td>
<td>Block Map</td>
</tr>
<tr>
<td>104</td>
<td>15.1</td>
<td>6.7</td>
<td>1.8</td>
</tr>
<tr>
<td>152</td>
<td>45.5</td>
<td>20.2</td>
<td>5.3</td>
</tr>
<tr>
<td>200</td>
<td>101.8</td>
<td>42.7</td>
<td>11.7</td>
</tr>
<tr>
<td>248</td>
<td>191.9</td>
<td></td>
<td>21.9</td>
</tr>
<tr>
<td>296</td>
<td>323.7</td>
<td></td>
<td>36.9</td>
</tr>
<tr>
<td>356</td>
<td>559.9</td>
<td></td>
<td>63.6</td>
</tr>
<tr>
<td>404</td>
<td>814.3</td>
<td></td>
<td>92.5</td>
</tr>
<tr>
<td>452</td>
<td>1137.2</td>
<td></td>
<td>129.0</td>
</tr>
</tbody>
</table>

Table 7-3. Time to fill all-wire matrix, seconds.

<table>
<thead>
<tr>
<th>Number of Unknowns</th>
<th>COMPUTER PLATFORM</th>
<th>4 Xputer</th>
<th>CONVEX</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Micro-Vax</td>
<td>1 Xputer</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wrap Map</td>
<td>Block Map</td>
</tr>
<tr>
<td>47</td>
<td>19.1</td>
<td>7.3</td>
<td>3.3</td>
</tr>
<tr>
<td>97</td>
<td>78.9</td>
<td>30.6</td>
<td>14.0</td>
</tr>
<tr>
<td>147</td>
<td>180.5</td>
<td>70.1</td>
<td>31.4</td>
</tr>
<tr>
<td>197</td>
<td>322.4</td>
<td>125.4</td>
<td>56.7</td>
</tr>
<tr>
<td>247</td>
<td>505.3</td>
<td></td>
<td>88.0</td>
</tr>
<tr>
<td>297</td>
<td>729.3</td>
<td></td>
<td>127.8</td>
</tr>
<tr>
<td>347</td>
<td>993.7</td>
<td></td>
<td>173.1</td>
</tr>
<tr>
<td>397</td>
<td>1302.1</td>
<td></td>
<td>227.8</td>
</tr>
</tbody>
</table>

The results show that the micro-Vax fills a wire matrix 32% slower than it fills a body matrix of the same size. Using column wrap mapping the four transputers fill a wire matrix 8% slower than they do a body matrix. In contrast, the Convex fills a wire matrix 15% faster than it does a body matrix! This may be due to the Convex's automatic vectorizing abilities. For comparison, a single transputer fills a wire matrix 36% slower than it fills a body matrix. The 8% figure for the four transputers can be accounted for by the fact that the wire filling efficiency (55%) is about 28% greater than the body filling efficiency (40%) for these problems and 28+8=36. The conclusion here is that for the JUNCTION code, sequentially speaking, wire filling takes about one third longer than body filling.

Some of the comparison data is displayed graphically in figures 7-1 through 7-3. The data is for an all-body matrix. The figures show comparisons between the four transputer array, the Convex (CPU time), and projected results for a sixteen transputer array. A four transputer array with 1 Megabyte of RAM per node can solve problems with up to 450 unknowns. With a sixteen transputer array problems with more than 900 unknowns could be solved.
Table 7-4. Time to factor & solve all-wire matrix, seconds.

<table>
<thead>
<tr>
<th>Number of Unknowns</th>
<th>COMPUTER PLATFORM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Micro-Vax</td>
</tr>
<tr>
<td></td>
<td>Wrap Map</td>
</tr>
<tr>
<td>47</td>
<td>1.6</td>
</tr>
<tr>
<td>97</td>
<td>12.3</td>
</tr>
<tr>
<td>147</td>
<td>41.2</td>
</tr>
<tr>
<td>197</td>
<td>97.4</td>
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<tr>
<td>247</td>
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<tr>
<td>297</td>
<td>327.8</td>
</tr>
<tr>
<td>347</td>
<td>520.2</td>
</tr>
<tr>
<td>397</td>
<td>776.0</td>
</tr>
</tbody>
</table>

The plots show that a sixteen transputer array would perform better than the Convex for filling the matrix and almost as good as the Convex for factoring and solving the matrix (Note: a sixteen transputer array, including the motherboard, would cost about $15K). However, no attempt was made to improve the vectorization of JUNCTION for a more efficient computation on the Convex. It is envisioned that considerable improvement could be made in the Convex times for factoring and solving of the matrix.

Figure 7-1. Matrix filling time comparisons, block mapping.
Figure 7-2. Matrix factoring and solving time comparisons.

Figure 7-3. Total computation time comparisons, block mapping.
7.2 Observations

In the course of this effort a number of observations were made regarding transputers, ParaSoft EXPRESS, and parallel processing in general:

- **A transputer array is an inexpensive, flexible parallel processing platform.** An array of four transputer modules, each with 1 Megabyte of RAM, plus the PC motherboard cost less than $5K. The transputer array is very flexible since the number of processors can be easily expanded (just plug more modules in the sockets) and any Fortran or C code which runs on the PC can be modified to run on the transputer array.

- **ParaSoft EXPRESS has been of great utility.** Without EXPRESS it would have been very difficult to make the progress that was made in parallelizing the code. In addition, major modifications to the existing code would have been required, if not a complete rewriting of the code. EXPRESS will allow the porting of the code to another computer platform to be much more direct.

- **Running an existing program on one transputer is very straightforward using EXPRESS.** By adding just a few lines of code, then recompiling using the parallel compiler, and linking the object files to EXPRESS, an executable module which runs on a single transputer can be created for virtually any Fortran program which runs on the PC.

- **The effort required to parallelize a code is algorithm dependent.** Some algorithms can be parallelized just by adding a couple of lines of code. Other algorithms require extensive restructuring. Still other algorithms are not suited for implementation on a parallel computer and must be left as sequential code. It is not always obvious beforehand which algorithms will be efficient to parallelize and which won't.

- **The optimal parallel code can be dependent on the number and type of processors as well as the size of the problem being solved.** This was shown to be the case with the matrix solve algorithms. Two algorithms were tested: a cyclic implementation and a wavefront implementation. Although the cyclic implementation outperformed the wavefront implementation for the 4-transputer array, the literature suggests that the wavefront implementation will be the best performer for a massively parallel computing platform solving larger problems.

8.0 THE FUTURE

The state-of-the-art technology in the area of parallel processing changes monthly both for hardware platforms and software tools. The transputers used for this effort will soon be outdated, replaced by faster and more flexible processing units [Pountain, 1990]. New features are continuously being added to EXPRESS to make it more powerful and adaptable.

8.1 Future Hardware

The big news in the world of transputers is the development of the T9000. This next generation processor is being developed by the same company, INMOS, which did the original pioneering work on the development of the transputer [INMOS, 1988]. According to the manufacturer the key features of the T9000 are "a high performance pipelined superscalar processor and major support for multiprocessing applications. Peak performance will be more than 150 MIPS and 20 MFLOPS, representing a major advance in parallel computing and high speed communications...The design goals for T9000 were to enhance the transputer's position as the premier multiprocessing micro-
processor, and to establish a new standard in single processor performance, while maintaining compatibility with existing transputer products." Of course, there will be some delay between the release of the T9000 and the development of compatible motherboards and software.

8.2 Future Software

A great deal of effort is going into developing programming software for parallel computing environments. This software includes parallel compilers, debuggers, performance analysis tools, visualization tools, dynamic load balancing tools, and automatic parallelization tools. Most of the presently existing software in these areas are very crude.

ParaSoft is in the process of improving EXPRESS. New tools which are being added include VTOOL, ASPAR, and EXDIST. VTOOL will allow memory access visualization. ASPAR provides automatic parallelization of sequential C programs. EXDIST will provide dynamic load balancing. ParaSoft's ultimate goal is to run EXPRESS within a heterogeneous parallel processing environment - known as a "meta" computer. A "meta" computer is made up of a number of architecturally different computers which are networked together and perform as a single entity.

REFERENCES


A RECURSIVE TECHNIQUE TO AVOID ARITHMETIC OVERFLOW AND UNDERFLOW WHEN COMPUTING SLOWLY CONVERGENT EIGENFUNCTION TYPE EXPANSIONS

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Abstract

Eigenfunction expansions for fields scattered by large structures are generally very slowly convergent. The summation often consists of two factors where one factor approaches zero and the other factor grows in magnitude without bound as the summation index increases. Each term of the expansion is bounded; however, due to the extreme magnitude of the individual factors, computational overflow and underflow errors can limit the number of terms that can be computed in the summation thereby forcing the summation to be terminated before it has converged. In this paper an exact technique that circumvents these problems is presented. An auxiliary function is introduced which is proportional to the original factor with its asymptotic behavior factored out. When these auxiliary functions are introduced into the summation, we are left with the task of numerically summing products of well behaved factors. A recursion relationship is developed for computing this auxiliary function.

1 Introduction

When solving for the fields scattered by canonical geometries, the exact solution is often available in eigenfunction form. The eigenfunction form is a viable representation for the fields providing that some characteristic dimension of the structure is "small" with respect to the wavelength, otherwise, the eigenfunction expansion is very slowly convergent and additionally can exhibit the following computational difficulty. These pathological eigenfunction expansions are in the form of infinite summations of products and quotients. Each term of the summation is well-behaved, however, the magnitude of the individual factors and/or divisors become either too small or too large to handle on the computer resulting in overflow/overflow errors. Ideally, one should find an alternate representation of the series that is more quickly convergent using a technique such as Watson's transformation [Tyras, 1969]. For many geometries the topology of the characteristic plane may be too complicated to perform the necessary function theoretic manipulations. Therefore, one may be forced to sum the slowly convergent series.
The overflow/underflow problem can be alleviated by implementing an auxiliary function that is proportional to the original pathological function with the asymptotic behavior factored out. These auxiliary functions can be calculated "exactly" from new recursion relationships which are derived from the recursion relationship of the original function. In Section 2, the development of these auxiliary functions and the corresponding recursive techniques will be presented. Section 3 contains an example of the plane wave scattering by a circular cylinder using the techniques developed in this paper and Section 4 contains some concluding remarks. Throughout this paper an \( e^{j\omega t} \) time dependence is assumed and suppressed.

2 Analytic Formulation

Symbolically, an eigenfunction expansion often has the form

\[
\phi = \sum_n C_n S_n(x) L_n(x),
\]

where \( C_n \) is a well-behaved constant, \( S_n(x) \) is a factor that becomes increasingly small as \( n \) grows,

\[
S_n(x) \xrightarrow{n \to \infty} 0,
\]

and \( L_n(x) \) is a factor that grows without bound with increasing \( n \),

\[
L_n(x) \xrightarrow{n \to \infty} \infty,
\]

such that the product, \( S_n(x)L_n(x) \), is bounded and the sum \( \sum_n C_n S_n(x)L_n(x) \) is convergent.

2.1 Computation by Recursion

Recursion relationships are very convenient, and are a common way to calculate the \( S_n(x) \) and \( L_n(x) \) functions. Typically, the functions that arise in eigenfunction solutions to electromagnetic problems satisfy a three term recursion relationship which can be expressed as follows:

\[
A(n, x) y_n(x) + B(n, x) y_{n+1}(x) + C(n, x) y_{n-1}(x) = 0.
\]

For a fixed value of \( x \), the recursion relation can be treated as a difference equation in \( n \). A three term difference equation has 2 independent solutions [Press, et al., 1988], i.e:

\[
y_n(x) = \{P_n(x), Q_n(x)\}.
\]

So the general solution to the recursion relation is:

\[
y_n(x) = \alpha P_n(x) + \beta Q_n(x)
\]
where $\alpha$ and $\beta$ are constants that need to be determined by physical considerations.

Miller's algorithm [Press, et al., 1988] is a commonly used technique to generate the sequence $\{P_n(x)\}$ or $\{Q_n(x)\}$. The algorithm begins by arbitrarily choosing two successive values to substitute into the recursion relationship which is used to generate the entire sequence. For example, let

$$y_0(x) = C_0(x) \quad \text{and} \quad y_1(x) = C_1(x),$$

which when substituted into Equation (6) yields:

$$C_0(x) = \alpha P_0(x) + \beta Q_0(x),$$

$$C_1(x) = \alpha P_1(x) + \beta Q_1(x).$$

Since $P_n(x)$ and $Q_n(x)$ are known to be independent, then by Equations (8)–(9) the values of $\alpha$ and $\beta$ are determined uniquely, hence $y_n(x)$ is well-defined. Note that $y_n(x)$ contains components of both solutions, $\{P_n(x), Q_n(x)\}$, providing that the choice of $C_0(x)$ and $C_1(x)$ are not proportional to either $P_0(x)$ and $P_1(x)$, respectively or $Q_0(x)$ and $Q_1(x)$, respectively.

\[
\begin{bmatrix}
\alpha \\
\beta
\end{bmatrix} = \begin{bmatrix} P_0 & Q_0 \\
P_1 & Q_1 \end{bmatrix}^{-1} \begin{bmatrix} C_0 \\
C_1 \end{bmatrix}, \quad \begin{vmatrix} P_0 & Q_0 \\
P_1 & Q_1 \end{vmatrix} \neq 0.
\]

At this point it is necessary to examine the stability of the desired solution. Stability refers to the relative rate of growth of the magnitude of the desired solution relative to the non-desired solution. Let’s examine the common circumstance where $|P_n(x)|$ increases as $n$ increases and $|Q_n(x)|$ decreases as $n$ increases as shown in Figure 1. It is clear that since our solution, $y_n(x)$, contains components of both $P_n(x)$ and $Q_n(x)$, then if we consider larger
and larger values of \( n \), \( Q_n(x) \) will be less and less significant compared to \( P_n(x) \), therefore:

\[
y_n(x) \sim \alpha P_n(x); \quad n \to \infty.
\]  

(11)

Under these circumstances, \( P_n(x) \) is said to be stable recursing up (in \( n \)). Similarly we could have arbitrarily chosen values for \( y_n(x) \) for two large successive values of \( n \), and then recursed down thereby recovering the solution for \( Q_n(x) \). In this case \( Q_n(x) \) is said to be stable recursing down. Mathematically,

\[
y_n(x) \sim \beta Q_n(x); \quad n \to -\infty.
\]  

(12)

This process generates a sequence proportional to the desired \( P_n(x) \) and \( Q_n(x) \) sequences. \( P_n(x) \) or \( Q_n(x) \) can be recovered by multiplying the entire sequence \( \{\alpha P_n(x)\} \) or \( \{\beta Q_n(x)\} \) by \( \frac{1}{\alpha} \) or \( \frac{1}{\beta} \), respectively. \( \alpha \) or \( \beta \) can be determined by calculating one value of \( \{P_n(x)\} \) or \( \{Q_n(x)\} \) and comparing it to the corresponding value of \( \{\alpha P_n(x)\} \) or \( \{\beta Q_n(x)\} \) which was calculated by recursion. An alternative is to use a normalization relationship of the form:

\[
\sum_n \gamma_n P_n(x) = 1 \quad \text{or} \quad \sum_n \delta_n Q_n(x) = 1.
\]  

(13)

In the introduction of *Abramowitz and Stegun* [1972, p. XIII], there is a listing of many functions and their direction of recursive stability, and in the various chapters corresponding to the functions of interest, normalization relations of the form in Equation (13) can be found.

An alternative to Miller's algorithm can be applied if two successive values of the solution are known. If the desired solution contains only one component of the two independent solutions \( \{P_n(x), Q_n(x)\} \), then it is necessary to recurse in the direction of recursive stability. Ideally, the direction of recursion should not matter, however, due to round off error in the computer, the undesired solution will be present and eventually grow to a significant value relative to the desired component of the solution.

As mentioned previously, the magnitude of the individual functions which we need to calculate can be either too large or too small for the computer to handle. This is why we introduce the auxiliary functions in Section 2.2.

### 2.2 The Auxiliary Functions

Since \( S_n(x) \) and \( L_n(x) \) are computed separately, before the sum converges individually they can become either too small or too large to calculate (due to computer limitations). To remove this upper bound limitation on the index of the summation, \( n \), we first note the asymptotic behavior of \( S_n(x) \) and \( L_n(x) \):

\[
S_n(x) \propto \sigma_n(x), \quad n \to \infty.
\]  

(14)

and,

\[
L_n(x) \propto \Lambda_n(x), \quad n \to \infty.
\]  

(15)
Introduce the auxiliary functions $A^S_n(x)$ and $A^L_n(x)$ which are defined for an appropriate interval over $n$ by the following expressions:

\[ S_n(x) = A^S_n(x) \sigma_n(x), \quad (16) \]

and,

\[ L_n(x) = A^L_n(x) \Lambda_n(x). \quad (17) \]

Since the auxiliary functions are equal to the original pathological functions with the asymptotic behavior factored out, the auxiliary functions remain well-behaved for all $n$ and are therefore computationally preferable over the original $S_n(x)$ and $L_n(x)$ functions. These expressions (Equations (16) and (17)) for $S_n(x)$ and $L_n(x)$ can be substituted into the eigenfunction expansion, Equation (1), yielding:

\[ \phi = \sum_n C_n A^S_n(x) \sigma_n(x) A^L_n(x) \Lambda_n(x), \quad (18) \]

or,

\[ \phi = \sum_n C'_n(x) A^S_n(x) A^L_n(x), \quad (19) \]

where,

\[ C'_n(x) = C_n \sigma_n(x) \Lambda_n(x). \quad (20) \]

The expression for $C'_n(x)$ can usually be significantly simplified which eliminates the necessity to compute the extremely small and extremely large values for $\sigma_n(x)$ and $\Lambda_n(x)$, respectively as $n \to \infty$.

The three factors in each term of Equation (19) are well-behaved for large values of $n$ which makes this procedure convenient for computing a slowly convergent eigenfunction expansion.

It is common practice to extract a factor from functions that grow without bound. The main contribution of this paper is the computation procedure for the auxiliary functions which is presented in the next section.

2.2.1 Computation of the Auxiliary Functions by Recursion

The direction of recursive stability of an auxiliary function is the same as the function from which it was derived, therefore Miller's algorithm can also be applied to auxiliary functions. We begin by assuming that the recursion relationship for the original function (Equation (4)) is known. If we are trying to calculate $A^L_n(x)$, the auxiliary function for $L_n(x)$, then we simply substitute Equation (17) into the recursion relationship, Equation (4).

\[ A_n(x) \Lambda_n(x) A^L_n(x) + B_n(x) \Lambda_{n+1}(x) A^L_{n+1}(x) + C_n(x) \Lambda_{n-1}(x) A^L_{n-1}(x) = 0 \quad (21) \]

In this form, the recursion relationship would experience the same overflow and underflow difficulties as the original function (due to the asymptotic factors $\Lambda_n(x)$, $\Lambda_{n+1}(x)$ and

71
Incident Plane Wave

Figure 2: Plane wave incident upon PEC circular cylinder.

$\Lambda_{n-1}(x))$. It can, and must, be analytically simplified at this stage to avoid taking differences of very large numbers which is a computational faux pas. Section 3 illustrates this procedure by presenting an example which calculates the plane wave scattering by a circular cylinder.

3 Example: Scattering by a PEC Circular Cylinder

In this section we are presenting an example that applies the general technique outlined in this paper to the specific problem of determining the total (incident + scattered) $z$-directed electric field when a TM$_z$ plane wave is incident upon a perfect electric conducting (PEC) circular cylinder which has its axis along the $z$-axis. The incident field is given by:

$$E_z = E_0 e^{-j k x} = E_0 e^{-j k p \cos \phi} = E_0 \sum_{n=-\infty}^{\infty} j^{-n} J_n(k \rho) e^{j n \phi}, \quad (22)$$

where $E_0$ is the complex amplitude and the coordinates $x$, $\rho$ and $\phi$ are shown in Figure 2. The total $z$-directed field is given by [Harrington, 1961]:

$$E_z = E_0 \sum_{n=-\infty}^{\infty} j^{-n} \left[ J_n(k \rho) - \frac{J_n(ka)}{H_n^{(2)}(ka)} H_n^{(2)}(k \rho) \right] e^{j n \phi} \quad (23)$$

It is possible to convert this series into a more quickly converging representation, however, this will not be done since the goal of this section is to illustrate the recursive technique outlined in this paper by a simple example. An additional numerical difficulty which will not be addressed in detail here, with the form of Equation (23) is that when computing the total fields near the cylinder, $\rho \approx a$, there can be a loss of significant digits. This may be overcome by computing the cross product directly by means of recursion relations.
Note that this problem does not occur when computing the scattered fields alone.

Using the relationship that \( Z_n \), any integer order Bessel function \( (J_0, J_1, J_2), \), satisfies:

\[
Z_{-n} = (-1)^n Z_n, \tag{24}
\]
we can express the total fields as:

\[
E_z = E_0 \sum_{n=0}^{\infty} \epsilon_n j^{-n} \left[ J_n(k\rho) - \frac{J_n(ka)}{H_n^{(2)}(ka)} H_n^{(2)}(k\rho) \right] \cos(n\phi), \tag{25}
\]
where,

\[
\epsilon_n = \begin{cases} 1 & n = 0, \\ 2 & n \neq 0. \end{cases} \tag{26}
\]

Notice that the second term in the brackets exhibits the behavior that is discussed in this paper, namely that each individual factor grows without bound \( (H_n^{(2)}), \) or approaches zero \( (J_n), \) as \( n \) increases. The asymptotic behavior of the Bessel functions of the first and second kind are given by [Abramowitz and Stegun, 1972, p. 365]:

\[
J_n(x) \sim \frac{1}{\sqrt{2\pi n}} \left( \frac{ex}{2n} \right)^n = \sigma_n(x), \quad n \to \infty, \tag{27}
\]

and,

\[
Y_n(x) \sim -\sqrt{\frac{2}{\pi n}} \left( \frac{ex}{2n} \right)^{-n}, \quad n \to \infty, \tag{28}
\]

and since,

\[
H_n^{(2)}(x) = J_n(x) - j Y_n(x). \tag{29}
\]

Then,

\[
H_n^{(2)}(x) \sim j \sqrt{\frac{2}{\pi n}} \left( \frac{ex}{2n} \right)^{-n} = \Lambda_n(x), \quad n \to \infty. \tag{30}
\]

So then the auxiliary functions are defined by:

\[
J_n(x) = \frac{1}{\sqrt{2\pi n}} \left( \frac{ex}{2n} \right)^n A_n^J(x), \quad n = 1, 2, 3, \ldots, \infty, \tag{31}
\]
and,

\[
H_n^{(2)}(x) = j \sqrt{\frac{2}{\pi n}} \left( \frac{ex}{2n} \right)^{-n} A_n^{H^{(2)}}(x), \quad n = 1, 2, 3, \ldots, \infty. \tag{32}
\]
All Bessel functions satisfy the same recursion relationship [Abramowitz and Stegun, 1972, p. 365]:

\[ Z_{n-1}(x) + Z_{n+1}(x) = \frac{2n}{x} Z_n(x). \] (33)

By comparing the asymptotic forms of \( J_n(x) \) and \( Y_n(x) \), \( J_n(x) \) is downward stable and \( Y_n(x) \) is upward stable [Abramowitz and Stegun, 1972, p. XIII]. Since the Hankel function consists of both \( J_n(x) \) and \( Y_n(x) \) and they differ in their direction of recursive stability, it is necessary to decompose the Hankel function to determine \( J_n(x) \) and \( Y_n(x) \) separately.

The definitions for the auxiliary functions (Equations (31) and (32)) are indeterminate for \( n = 0 \). For this reason, we will extract the \( n = 0 \) term of Equation (25) and begin the sum from \( n = 1 \).

\[
E_x = E_0 \left[ J_0(k\rho) - \frac{J_0(ka)}{H_0^{(2)}(ka)} H_0^{(2)}(k\rho) \right] + 2E_0 \sum_{n=1}^{\infty} j^{-n} \left( \frac{ek\rho}{2n} \right)^n A_n^J(k\rho) - \left( \frac{ek\rho}{2n\rho} \right)^n A_n^J(ka) A_n^H(k\rho) \cos(n\phi). \] (34)

**Calculation of \( A_n^J(x) \)**

The recursion relation which defines \( A_n^J(x) \) is found by substituting Equation (31) into Equation (33). After algebraic simplification, the following form of the recursion relation is obtained:

\[
A_n^J(x) = e \left( \frac{n}{n+1} \right)^{n+1/2} A_{n+1}^J(x) - \left( \frac{1}{2} \frac{ex}{n+2} \right)^n \left( \frac{n}{n+2} \right)^{n+1/2} A_{n+2}^J(x) \] (35)

where \( e \) is the base of the natural logarithm. This is in a form suitable for downward recursion.

We will apply Miller’s algorithm to determine a sequence denoted by \( \tilde{A}_n^J(x) \) which is proportional to the desired \( A_n^J(x) \) sequence. We choose \( N \) to be larger than the maximum number of terms expected to be summed by \( M \). Also, let:

\[ \tilde{A}_0^J(x) = 1, \] (36)

and,

\[ \tilde{A}_{N+1}^J(x) = 0. \] (37)

Use the recursion relationship (Equation (35)) to determine \( \{ \tilde{A}_n^J(x) \} \) for \( n = 1, 2, 3, \ldots, N-1 \). \( \{ \tilde{A}_n^J(x) \} \) is a sequence proportional to the desired sequence \( \{ A_n^J(x) \} \). There are many ways to normalize the sequence \( \{ \tilde{A}_n^J(x) \} \). Here we will use Equation (31) to determine \( A_1^J(x) \) which will then be compared to \( \tilde{A}_1^J(x) \) to determine the constant of proportionality, \( \beta \).

\[
\beta = \frac{ex}{2\sqrt{2\pi}} \frac{\tilde{A}_1^J(x)}{J_1(x)}, \] (38)
So then,
\[
\{A_n^2(x)\} = \frac{1}{\beta} \{\tilde{A}_n^2(x)\}; \quad n = 1, 2, 3, \ldots N - M. \tag{39}
\]

The following table was constructed, using these procedures, to emphasize the favorable behavior of the magnitude of the auxiliary functions compared to the magnitude of the Bessel functions. We are showing two arguments of the auxiliary function for a wide range of orders. This table illustrates the difficulty in computing the Bessel functions directly. The triple asterisk indicates that this term is larger than \(10^{308}\), the largest number our computer can handle.

<table>
<thead>
<tr>
<th>n</th>
<th>(J_n(1))</th>
<th>(A_n^2(1))</th>
<th>(J_n(5))</th>
<th>(A_n^2(5))</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(4.4005 \times 10^{-1})</td>
<td>0.81157</td>
<td>(-3.2758 \times 10^{-1})</td>
<td>-0.12083</td>
</tr>
<tr>
<td>2</td>
<td>(1.1490 \times 10^{-1})</td>
<td>0.88200</td>
<td>(4.6565 \times 10^{-2})</td>
<td>0.014297</td>
</tr>
<tr>
<td>5</td>
<td>(2.4976 \times 10^{-4})</td>
<td>0.94323</td>
<td>(2.6114 \times 10^{-1})</td>
<td>0.31559</td>
</tr>
<tr>
<td>20</td>
<td>(3.8735 \times 10^{-25})</td>
<td>0.98405</td>
<td>(2.7703 \times 10^{-11})</td>
<td>0.73798</td>
</tr>
<tr>
<td>50</td>
<td>(2.9060 \times 10^{-78})</td>
<td>0.99345</td>
<td>(2.2942 \times 10^{-45})</td>
<td>0.88306</td>
</tr>
<tr>
<td>100</td>
<td>(8.4318 \times 10^{-189})</td>
<td>0.99670</td>
<td>(6.2678 \times 10^{-119})</td>
<td>0.93919</td>
</tr>
<tr>
<td>200</td>
<td>***</td>
<td>0.99834</td>
<td>4.7600 (10^{-296})</td>
<td>0.96898</td>
</tr>
<tr>
<td>500</td>
<td>***</td>
<td>0.99927</td>
<td>***</td>
<td>0.98737</td>
</tr>
<tr>
<td>1000</td>
<td>***</td>
<td>0.99965</td>
<td>***</td>
<td>0.99367</td>
</tr>
</tbody>
</table>

**Calculation of \(A_n^{H(2)}(x)\)**

As mentioned previously, the Hankel function, \(H_n^{(2)}(x)\), consists of Bessel functions of the first and second kind, \(J_n(x)\) and \(Y_n(x)\), respectively. \(J_n(x)\) and \(Y_n(x)\) differ in their direction of recursive stability, therefore, they must be computed separately. We note that as a consequence of Equations (29) and (32):

\[
J_n(x) = -\sqrt{\frac{2}{\pi n}} \left(\frac{e x}{2 n}\right)^{-n} \text{Im}\{A_n^{H(2)}(x)\} \tag{40}
\]

and,

\[
Y_n(x) = -\sqrt{\frac{2}{\pi n}} \left(\frac{e x}{2 n}\right)^{-n} \text{Re}\{A_n^{H(2)}(x)\}. \tag{41}
\]

From these relationships, we determine that \(\text{Im}\{A_n^{H(2)}(x)\}\) is stable recursing down and that \(\text{Re}\{A_n^{H(2)}(x)\}\) is stable recursing up. Substituting Equation (40) into Equation (33) yields the following recursion relation in a suitable form for downward recursion.

\[
\text{Im}\{A_n^{H(2)}(x)\} = \frac{1}{e} \left(\frac{2 n}{x}\right)^2 \left(\frac{n + 1}{n}\right)^{n + \frac{3}{2}} \text{Im}\{A_{n+1}^{H(2)}(x)\} \nonumber \\
- \left(\frac{2 n}{e x}\right)^2 \left(\frac{n + 2}{n}\right)^{n + \frac{3}{2}} \text{Im}\{A_{n+2}^{H(2)}(x)\} \tag{42}
\]
We can use this recursion relation along with Equation (40) with \( n = 1 \) (for normalization), to apply Miller's algorithm.

\[
\text{Im}\{A_1^{H(2)}(x)\} = -\sqrt{\frac{\pi}{2}} \frac{e^x}{2} J_1(x) \tag{43}
\]

The calculation of \( \text{Re}\{A_n^{H(2)}(x)\} \) requires two starting values and the recursion relationship since it is upward stable. The two starting values are obtained by substituting \( n = 1 \) and \( n = 2 \) into Equation (41).

\[
\text{Re}\{A_1^{H(2)}(x)\} = -\sqrt{\frac{\pi}{2}} \frac{e^x}{2} Y_1(x), \tag{44}
\]

and,

\[
\text{Re}\{A_2^{H(2)}(x)\} = -\sqrt{\pi} \left(\frac{e^x}{4}\right)^2 Y_2(x). \tag{45}
\]

The recursion relation is obtained by substituting Equation (41) into Equation (33) yielding the following recursion relationship for \( \text{Re}\{A_n^{H(2)}(x)\} \) in a suitable form for upward recursion.

\[
\text{Re}\{A_n^{H(2)}(x)\} = e^{\left(\frac{n-1}{n}\right)} \text{Re}\{A_{n-1}^{H(2)}(x)\} - \frac{\frac{1}{2} e^x}{(n-2)^2} \left(\frac{n-2}{n}\right)^{n-1/2} \text{Re}\{A_{n-2}^{H(2)}(x)\} \tag{46}
\]

The following table is presented for a comparison of the Hankel function of the second kind with its auxiliary function. We are showing two arguments of the auxiliary function for a wide range of orders.

<table>
<thead>
<tr>
<th>n</th>
<th>( H_n^{(2)}(1) )</th>
<th>( A_n^{H(2)}(1) )</th>
<th>( H_n^{(2)}(5) )</th>
<th>( A_n^{H(2)}(5) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(4.4005 + j7.8121) \times 10^{-1}</td>
<td>1.3307 - j0.74960</td>
<td>(-3.2758 - j1.4786) \times 10^{-1}</td>
<td>-1.2594 + j2.7900</td>
</tr>
<tr>
<td>2</td>
<td>(0.11490 + j1.6507) \times 10^{0}</td>
<td>1.3511 - j0.90940</td>
<td>(0.46565 - j3.6766) \times 10^{-1}</td>
<td>-7.5237 - j0.9529</td>
</tr>
<tr>
<td>5</td>
<td>(0.0000 + j2.6041) \times 10^{2}</td>
<td>1.0831 + j0.0000</td>
<td>(2.6114 + j4.5369) \times 10^{-1}</td>
<td>5.8970 - j3.3942</td>
</tr>
<tr>
<td>20</td>
<td>(0.0000 + j4.1140) \times 10^{22}</td>
<td>1.0175 + j0.0000</td>
<td>(0.0000 + j5.9340) \times 10^{8}</td>
<td>1.3996 + j0.0000</td>
</tr>
<tr>
<td>50</td>
<td>(0.0000 + j2.1911) \times 10^{77}</td>
<td>1.0068 + j0.0000</td>
<td>(0.0000 + j2.7888) \times 10^{42}</td>
<td>1.1381 + j0.0000</td>
</tr>
<tr>
<td>100</td>
<td>(0.0000 + j3.7753) \times 10^{185}</td>
<td>1.0034 + j0.0000</td>
<td>(0.0000 + j5.0849) \times 10^{115}</td>
<td>1.0661 + j0.0000</td>
</tr>
<tr>
<td>200</td>
<td>***</td>
<td>1.0017 + j0.0000</td>
<td>(0.0000 + j3.3446) \times 10^{292}</td>
<td>1.0323 + j0.0000</td>
</tr>
<tr>
<td>500</td>
<td>***</td>
<td>1.0007 + j0.0000</td>
<td>***</td>
<td>1.0128 + j0.0000</td>
</tr>
<tr>
<td>1000</td>
<td>***</td>
<td>1.0003 + j0.0000</td>
<td>***</td>
<td>1.0064 + j0.0000</td>
</tr>
</tbody>
</table>

76
4 Conclusion

In this paper we introduce a technique which circumvents the need to compute very large or very small special functions that commonly appear in eigenfunction expansions. This was accomplished by introducing a set of well-behaved auxiliary functions which are equal to the original special function with its asymptotic behavior factored out. When the eigenfunction expansion is expressed in terms of these auxiliary functions, the summation can be simplified resulting in well-behaved factors in the sum. The auxiliary functions can be computed via modified recursion formulas.

This procedure is formally exact since we are not making any approximations—only substitutions. Typically, when calculating summations of the type addressed here without the use of the auxiliary functions, at some point in the summation (which needs to be determined), an asymptotic form is substituted into the expression. The procedure described herein avoids the need to switch functional representations thereby eliminating the need to determine the value of the index to implement the asymptotic representation.

In this paper we have restricted our development of the auxiliary functions to the extraction of the asymptotic form of the function. The procedure is not limited to this asymptotic extraction. Any functional form that is convenient for the formulation of the problem at hand can be extracted in a similar manner.

References


'H-ORIENTED' AND 'B-ORIENTED' METHODS
IN A PROBLEM OF NONLINEAR MAGNETOSTATICS:
SOME METHODOLOGICAL REMARKS

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ABSTRACT

Problems which depend on a small parameter in their formulation can often be studied by a perturbation approach. Whether the perturbation is "regular" or "singular" is important in many respects. In magnetostatics, due to some inherent duality, both kinds of perturbation may happen, depending on the chosen formulation ("b-oriented" vs. "h-oriented" methods). Singularly perturbed problems are numerically more difficult than regularly perturbed ones. We suggest that this might explain why, as some recent numerical observations seem to suggest, b-oriented methods should give better accuracy in a specific class of nonlinear magnetostatic problems at high permeability.

INTRODUCTION

This paper is a case study, which butts on a methodological question of such generality that it seems almost preposterous to address it in written form: how should the particularities of a physical problem (here, the presence of "small parameters") guide the modeler in the selection of a numerical method? This is a basic subject, one about which everybody has definite opinions (and even, sometimes, strong feelings), but also an elusive one, difficult to treat in a comprehensive way. (This is why there are relatively so few books or paper collections devoted to mathematical modelling, like e.g., [1, 2, 3, 6, 7, 9, 11, 13, 15, 22]. Why most of them seem so remote from the kind of mathematical modelling we do is a puzzling question.)

One might say that, after all, this is quite normal. Why should "tricks of the trade" be honored with formal dissertations? But a moment of reflection will show that some of the most powerful of such tricks deserve to be thus treated, that they are, and that it is eventually beneficial for all those concerned. Take Fourier analysis, for instance. On one level, it is an efficient dimensionality-reduction device, which helps replace the numerical solution of a fully three-dimensional problem with a series of simpler ones, in one or two dimensions. On another level, it is the practical and usable by-product of a majestic mathematical theory: harmonic analysis [12, 17]. Is this a coincidence? Probably not. The development of the theory was stirred by the efficiency of what was at the time a modelling trick, the use of trigonometric series in the study of heat transfer, and it has paid back largely, as far as mathematical modelling is concerned. This is the classical example of a practical modelling tool—a trade trick—backed by a strong mathematical theory, and of the dialectics of their historical development. Other examples will come to
mind (the "least squares" trick in relation with control and identification theory, some aspects of Geometrical Diffraction Theory, some statistical tests . . .).

The small parameter issue (an instance of which will be presented in a moment) might deserve such a status. Here there is, no doubt, a trick: use dimensional analysis to expose "small parameters", identify the corresponding terms in the equations, drop them. There is also a grandiose theory, or rather, theories: asymptotic analysis, perturbation theory. But the theory appears incomplete, it lacks unity, and the trick itself is far from being fail-safe: when your equation is $-\varepsilon u'' + u = f$, or $-u'' + \varepsilon u = f$, should you drop the $\varepsilon$-term, or not? (Think of $\varepsilon$ as related, for instance, with the skin-depth.) The answer to this particular question is known, but lots of similar ones are still to be addressed. Here is a rich field of study, full of prospects for both intellectually stimulating mathematical theories and usable practical recipes.

Yet, case studies may prove more effective than comprehensive studies of large scope, at this stage. One such study comes from the experience of a small electromagnetism community (the "TEAM workshop", [22, 23]) whose emphasis is on low-frequency and static situations. The test-case, described below in general terms, is a non-linear magnetostatics problem on which the same observation has been made by several independent investigators: numerical methods based on the use of the vector potential tend to perform better than those using a scalar potential. This calls for an explanation, which is the theme of the present paper. In short, there are two "small parameters" in this problem, the ratio of the airgap to some characteristic length, and the ratio of permeabilities in the air and in the iron. Their interplay results in the possibility of formulating the problem as a perturbation with respect to some similar, much simpler problem. This is a standard idea. What is new here is that, due to some symmetry inherent in Maxwell equations, there are two ways to do this, based respectively on the use of the scalar and of the vector potential, and that the nature of the perturbation is not the same for both: it's "regular" perturbation in the former case, "singular" perturbation in the latter. An attempt is made to link these facts with the numerical observations.

Similar situations where two small parameters are "competing", each one driving the problem toward different limits, seem to be common to other areas of electromagnetics, for example, skin effect, current flow around a crack, and even (for higher frequencies) surface impedance.

'H-ORIENTED' AND 'B-ORIENTED' METHODS IN MAGNETOSTATICS

We shall consider the following general situation (Fig. 1): a coil $C$ (sustaining a given, permanent, current-density $j^f$), and a ferromagnetic piece $M$. The whole space is $E$, and $A = E - C - M$ is the air-region. A non-linear behavior law $b = \Gamma(h)$ is given.
hysteresis). For the present discussion, we assume a $\Gamma$ such that vectors $b(x)$ and $h(x)$ are collinear at all points $x$ in $M$, and that $b(x)$ vanishes if $h(x)$ does. One is requested to compute the fields $b$ and $h$. The relevant equations are

$$\text{(1)} \quad \text{curl } h = j^s \text{ in } E, \quad b = \Gamma(h), \quad \text{div } b = 0 \text{ in } E.$$  

\textbf{Figure 1.} A typical problem (left). Definition of surface $\Sigma$ and domain $\overline{M}$, to be used in the sequel (right).

We are only interested here in comparing the accuracy achievable with various methods, not in things like the computing time, the number of steps in iterative procedures, etc. So we may consider a linear problem with the same solution as (1), namely

$$\text{(2)} \quad \text{curl } h = j^s \text{ in } E, \quad b = \mu \cdot h, \quad \text{div } b = 0 \text{ in } E,$$

where $\mu$ is the function $x \rightarrow lb(x)/lh(x)$, as computed from the actual solution $\{b, h\}$. Whatever can be said about the merits of various methods as regards (2) will thus be relevant to (1).

The ratio $\mu(x)/\mu_0$ is a feature of the solution (and as such, it depends on the imposed current $j^s$). It is usually high (about 2000, typically), but tends to decrease when the driving current $j^s$ is increased (saturation phenomenon). Let us call $\varepsilon$ the average of $\mu(x)/\mu(x)$ over $M$. We define, for all $x$ in $M$, $\mu(x) = \varepsilon \mu(x)$. Note that $\mu_1$ is close to $\mu_0$, except perhaps in saturated zones. This $\varepsilon$ is one of the small parameters alluded to in the Introduction. It will play an important role in what follows. The other small parameter is
the ratio $d/L$ of the gap-width $d$ to the average length $L$ of the magnetic circuit $M$. We shall assume that $d/L$, though small, is still large with respect to $\varepsilon$: thus, the reluctance of the whole device is mainly due to the air gap. This is the assumption used in the TEAM workshop study, and is the rule in problems of this kind. (But the assumption is a crucial one: if the reluctance was mainly due to the iron core, this would modify our conclusions.)

Although the discussion will be limited to linear problems, we are not losing anything significant in generality, because one can always tackle (1) by iterating on $\mu$, Newton-Raphson style, a linear system being solved at each step. In fact, almost all methods in actual use in nonlinear magnetostatics seem to rely on this approach. They fall into two main families: 'b-oriented' methods, which yield $b$ (for instance by computing the magnetic vector potential), and 'h-oriented' methods, which aim at $h$ (for instance by computing the magnetic scalar potential). For the sake of definiteness, let us formalize this:

**Definition 1.** A vector field $h$ [resp. $b$] is said "curl-conformal" [resp. "div-conformal"] if its tangential [resp. normal] part is continuous across all surfaces in its domain of definition.

**Definition 2.** A method will qualify as "h-oriented" [resp. as "b-oriented"] if it computes $h$ [resp. $b$] in such a way that its tangential [resp. normal] continuity is exactly enforced, i.e., if it yields a curl-conformal $h$ [resp. a div-conformal $b$].

Let us develop an example of b-oriented method. A tetrahedral mesh is first designed, large enough to cover $M$, $C$, and enough of the air-region to justify neglecting what happens outside the meshed volume. Let $\mathcal{N}$ be the set of nodes. Let $\lambda_n$ be the "hat-function" associated with node $n$ (i.e., the unique continuous, piecewise affine, function equal to 1 at node $n$ and to 0 at all other nodes). Call $\mathbf{IP}^1$ the set of all vector fields of the form

$$a = \sum_{n \in \mathcal{N}} a_n \lambda_n,$$

where the degrees of freedom (DoF) $a_n$ are vectors, one per node. Now look for $a$ in $\mathbf{IP}^1$ such that

$$\int_E \mu^{-1} \text{curl } a \cdot \text{curl } a' = \int_E j^\phi \cdot a' \quad \forall a' \in \mathbf{IP}^1$$

(the so-called "weak form" of the equation $\text{curl}(\mu^{-1} \text{curl } a) = j^\phi$). This is a linear system in terms of the $3 \times N$ components of the $a_n$s ($N$, the number of nodes in the mesh). There exists a solution (because $\text{div } j^\phi = 0$), perhaps not unique, but anyway, all solutions will have the same curl. (Numerical difficulties that one may encounter in solving (4) are not our concern.) This is a b-oriented method, because it gives $b = \text{curl } a$, thus enforcing the
essential requirement about the continuity of the normal component of \( b \) across all surfaces, including the faces of the mesh. The field \( h = \mu^{-1} \text{curl} \ a \) fails to have the symmetric property of continuity of its tangential part through surfaces. This is so because the relation \( \text{curl} \ h = j^\parallel \) is only weakly enforced by (4).

On the other hand, the well-known "single magnetic potential" method is \( h \)-oriented. Its principle is to look for the field \( h \) in the form \( h = \text{grad} \ \phi + h^\parallel \), where \( h^\parallel \) is a field that satisfies \( \text{rot} \ h^\parallel = j^\parallel \) and \( \text{div} \ h^\parallel = 0 \). (Such a field can be constructed from \( j^\parallel \) by Biot-Savart integration.) The unknown in this problem is thus the potential \( \phi \). According to the general finite element approach, one thus has to find a function \( \phi \) of the form

\[
\phi = \sum_{n \in X} \varphi_n \lambda_n
\]

such that

\[
\int_E \mu \left( \text{grad} \ \phi + h^\parallel \right) \cdot \text{grad} \ \phi' = 0
\]

for all test-functions \( \phi' \) themselves of the form (5). (This is a way to enforce the condition \( \text{div}(\mu h) = 0 \), in weak form.) Here the \( \varphi_n \)s are scalar degrees of freedom. This time \( h \) has the required tangential continuity, but \( b = \mu h \) fails to have normal continuity, because \( \text{div} \ b = 0 \) is only weakly enforced: the method is "\( h \)-oriented".

**Remark.** Note that \( \phi \) is single-valued here. There are variants in which a similar, but possibly multivalued magnetic potential is used [19], or more than one potential [18]. Clearly, such methods also are \( h \)-oriented.

Not all methods fall in one of the previous categories. A method can have both orientations. But in that case the behaviour law \( b = \mu h \) will fail to be exactly satisfied, in general. (This point is developed in [5].)

Now the scene is set. We shall argue as follows: "\( b \)-oriented" methods should in general work better, because the smallness of the ratio \( \mu_b/\mu \) results in a regularly perturbed problem with such formulations, whereas "\( h \)-oriented" methods suffer from singular behavior with respect to this same parameter. Therefore, they are more sensitive to numerical error.

**SINGULAR VS. REGULAR PERTURBATIONS**

Let us first recall the difference between regular and singular perturbation, with the help of a simple example. Suppose we have to solve the two-points boundary-value problem
for a numerically small value of $\varepsilon$, and a given positive function $k$. Note that this is equivalent to minimizing the functional $F_\varepsilon(v) = \int_{[0, 1]} (|v|^2 + \varepsilon k |v|^2 - 2 f v)$ among all those in the set $V = \{v \in C^0([0, 1]) : \int_{[0, 1]} |v|^2 < \infty, v(1) = 0\}$, which can easily be done by numerical methods. But instead, in order to take advantage of $\varepsilon$'s smallness, one would rather make use of the "limit-problem", corresponding to $\varepsilon = 0$, whose solution $u_0$ is obtained directly, by quadratures, without having to solve a linear system. The idea is to expand $u$ as $u_0 + \varepsilon u_1 + \ldots$, and to throw this into (7); again, $u_1$ can be obtained by straightforward integration, the same way $u_0$ was. And so on. Problem (7) is a perturbation of a simpler one (namely, to find $u$ such that $-u'' = f$, $u'(0) = 0$, $u(1) = 0$), and therefore can be solved by a cascade of similar problems. Note that $u_0$ still achieves the minimum of some functional (namely, $F_0(v)$) over the same set $V$. This is what makes the perturbation "regular": the limit solution is solution to the limit problem, as obtained by setting $\varepsilon = 0$ in either (7) or its equivalent variational formulation, "minimize $F_\varepsilon(v)$ over $V$". Note also that $u_\varepsilon$ and its first derivative both converge (in the sense of quadratic means) to $u_0$ and $u'_0$ respectively.

In contrast, "singular perturbation" occurs with

(8) $-\varepsilon u'' + k(x) u = f$, $u'(0) = 0$, $u(1) = 0$.

If one wants to solve this numerically, the smallness of $\varepsilon$ imposes a small discretization step, which is cumbersome, so making use of the limit problem seems again a good idea. But now two things go wrong. First, the would-be "limit-problem", i.e., (7) with $\varepsilon = 0$, or its variational version, "minimize $\int_{[0, 1]} (k |v|^2 - 2 f v)$ over $V$", has no solution. Next, when one relaxes its requirements by dropping the boundary conditions, the solution $u_0$ of the relaxed problem, "minimize over $L^2([0, 1])$", which is $u_0 = f$, is not the limit of $u_\varepsilon$ in the above sense: only $u_\varepsilon$ converges towards $u_0$ in $L^2$, whereas $u'_\varepsilon$ does not converge to $u'_0$ (Fig. 2).

Singular perturbation is thus characterized by the fact that, when passing to the limit, the solution "escapes" from the set where it naturally lives, and while it still does converge to something, it is in a weaker sense, and to an object which belongs to a larger universe. It does not mean that the (non-acceptable) "limit solution" $u_0$ is useless as a stepping stone to obtaining the true solution, $u_\varepsilon$, of (8). Indeed, $u_0$ is a term in some asymptotic expansion of $u_\varepsilon$, that can be derived via a moderately involved process called "matching asymptotic expansions". (Some references to these things are [10, 14, 16, 20, 21].) Quite often, a lot of numerical work can be avoided by such an asymptotic approach to the solution. This is so because, from the numerical point of view, singularly perturbed problems are tougher than regularly perturbed ones. In the case of (8) vs. (7), a glance at Fig. 2 shows why: the
step-size has to be very small, at least in the "boundary layers" near the ends of the interval.

Figure 2. Solution of (7) for $\varepsilon$ small ($k = 1$).

PROBLEM (2) AS A PERTUBED PROBLEM

For reference, let us write weak formulations analogous to (4) and (6), as they stand before any commitment to particular finite elements has been made. Call $A$ (resp. $\Phi$) the set of all square-integrable vector fields (resp. functions), with a square integrable curl (resp. gradient), over the whole space. The weak formulations are, for the (b-oriented) "a-method": find $a_e \in A$ such that

$$\int_E \mu^{-1} \text{curl } a_e \cdot \text{curl } a' = \int_E j^g \cdot a' \quad \forall \ a' \in A,$$

(where can be rewritten as

$$\int_{E^-M} \mu_0^{-1} \text{curl } a_e \cdot \text{curl } a' + \varepsilon \int_M \mu_1^{-1} \text{curl } a_e \cdot \text{curl } a' = \int_E j^g \cdot a' \quad \forall \ a' \in A),$$

and for the (h-oriented) $\varphi$-method: find $\varphi_e \in \Phi$ such that

$$\int_E \mu (\text{grad } \varphi_e + h^g) \cdot \text{grad } \varphi' = 0 \quad \forall \ \varphi' \in \Phi,$$

(where can be rewritten as

$$\int_{E^-M} \mu_0 (\text{grad } \varphi_e + h^g) \cdot \text{grad } \varphi' + \varepsilon^{-1} \int_M \mu_1 (\text{grad } \varphi_e + h^g) \cdot \text{grad } \varphi'$$

$$= 0 \quad \forall \ \varphi' \in \Phi).$$
Remark. one should compare (9) with the following:

$$\int \partial_x u \partial_x u' + \varepsilon \int k u u' = \int f u' \quad \forall \ u' \in U,$$

which is equation (7) in weak formulation. The analogy we rely on is due to the presence of the small parameter $\varepsilon$ in front of the second integral in both cases.

Let us now see in which way our problem (2), when formulated as (9) or as (10), can be considered a "perturbation" of some simpler one. A difficulty is that there are two small parameters in the picture: the above ratio $\varepsilon \approx \mu_0/\mu$, and the ratio $d/L$ of the gap-width to the circuit-length. The analysis in terms of two such "competing" small parameters is quite interesting; it leads—depending on which parameter dominates—to very different limit models (called "significant degenerations" in [10]), which do have meaningful interpretations within the present context. (See an application to skin-effect, and to the concept of "surface impedance" [8], in [4].) There is no space here to justify our main assumption that $\varepsilon$, not $d/L$, is the dominant small parameter in the present case: this would call for some lengthy dimensional analysis whose principles, anyway, are familiar to all readers. We cannot either dwell on the Taylor expansions in terms of $\varepsilon$, and similar techniques, by which models (11) and (12) below are derived: although these are familiar and elementary mathematical tools, their application to the present case requires a heavy apparatus of functional analysis, that seems out of place. So we shall proceed more dogmatically: first, some notation, next a terse statement of the results, then some a posteriori justification.

Let us call $\bar{M}$ the union of $M$ (cf. Fig. 1) and the gap, and $\Sigma$ a surface congruent to the pole surfaces, located in the middle of the gap. Enlarge space $\Phi$ by accepting into it functions which are allowed to be discontinuous across $\Sigma$. Call $\overline{\Phi}$ the bigger space so obtained.

Now, the limit problems. The one in terms of the vector potential $a$ splits into two successive problems: find $a_0 \in A$ such that both conditions

$$| \int_{E - \bar{M}} \mu_0^{-1} \text{curl} \ a_0 \cdot \text{curl} \ a' = \int_{E} j^s \cdot a' \quad \forall \ a' \in A,$$

$$| \text{curl}(\mu_1^{-1} \text{curl} \ a_0) = 0 \quad \text{in} \ \bar{M},$$

hold. The one in $\varphi$ is find $\varphi_0 \in \overline{\Phi}$ such that both conditions

$$| \int_{E - \bar{M}} \mu_0 (\text{grad} \ \varphi_0 + h^s) \cdot \text{grad} \ \varphi' = 0 \quad \forall \ \varphi' \in \overline{\Phi},$$

$$| \text{grad} \ \varphi_0 + h^s = 0 \quad \text{in} \ \bar{M} - \Sigma,$$

85
Why, on physical grounds? Imagine the air gap is shrunk to \( \Sigma \), while the ratio \( \epsilon = \mu_0/\mu_1 \) tends to zero. Then \( h \) tends to zero in the core, hence the second line of (12), and (since its circulation is held constant) tends to infinity in the air gap, hence the eventual discontinuity of the magnetic potential. As for \( b \), one has \( \text{div } b = 0 \) and \( \text{curl}(\mu_0^{-1} b) = j^* \) out of the core, and its flux lines impinge orthogonally to the surface of the core in the case of an infinite permeability: indeed, the first line of (11) implies all that. The second line of (11) describes what happens inside the core, and (since the tangential values of \( a \) are now known, from solving (11), first line) constitutes a well-posed problem. (All this, though it may not be obvious, relies on the above observation that \( d/L \) is large with respect to \( \epsilon \).)

Remark. Neither (11) or (12) are practical ways to solve the problem, if only because of the non-linearity: \( \mu_1 \), of course, is not known in advance, so no useful information on the field in the core could be obtained that way. (The stray-field thus obtained, however, might be reasonably accurate.) We do not advocate the actual use of limit models in the specific case of Pb. 13. We just expect to get insight—about how difficult it will be to solve the true problem—from an examination of the limit one.

Now it should be clear that the limit process from (9) to (11) is regular, while the one from (10) to (12) is singular: for in the latter, there is this tendency of \( \varphi_\epsilon \) to "escape from" space \( \Phi \) (towards the larger one \( \mathcal{D} \)), while nothing similar happens to \( a_\epsilon \), which stays in \( A \).

The difference can be seen more concretely from what happens in the air at the boundary of \( \Sigma \): a singularity of \( \varphi \) (actually, a jump-discontinuity), whereas \( a \) is regular (Fig. 3). Figure 3 is two-dimensional for convenience, but what is described here is a feature of the three-dimensional situation: the magnetic potential changes very rapidly across the airgap (Fig. 3, right). So, if one tries to model the situation by treating the airgap as a surface, across which \( \varphi \) can be discontinuous—which is precisely model (12)—the computed \( \varphi \) will look as on the left part of the figure, with an obvious singularity. Our point is that, although this limit model is not the one which is actually solved for, the latter is sufficiently closed to it for the singular character of the limit case to be felt, in various ways, when actually solving for \( \varphi \).

The nature of the finite elements has not been a factor in this discussion. So we may conclude that regularity (resp. singularity) pertains to the whole family of \( a \)-methods (resp. \( \varphi \)-methods). Singularly perturbed problems are notoriously more difficult to solve numerically than regularly perturbed ones. Hence the tentative conclusion: \( b \)-oriented
methods as a whole should be considered with a favorable bias, for this particular problem. Of course, this should not be construed as a seal of approval for any particular b-oriented methods.

Figure 3. Flux lines in the vertical symmetry plane, near the edge of the air gap, showing the singularity of the field: left, as predicted by the limit models, right, as it really is. The situation is nearly two-dimensional. In that case, \( a \) is a scalar, whose isolines are precisely the flux-lines shown. Dotted lines are isovalues of \( \phi \), the magnetic potential. Clearly, \( a \) is continuous at point \( P \), but \( \phi \) is not, in the limit model.

CONCLUSION

Let us summarize the main line of our argumentation. The problem which is actually solved (the one with a small but non-zero \( \varepsilon \)) is closed to a singular limit when h-oriented methods are used, and to a regular limit when b-oriented methods are used. So in the critical region (near the edges of the poles), h-oriented methods are likely to behave in a nastier way than b-oriented ones. For instance, they may require a finer mesh if the same precision is to be achieved. But meshes, as a rule, are not what would be desirable, they are what computing resources allow. So we may expect, on the average, better precision from b-oriented methods in reported numerical experiments. This seems indeed to be the present trend, as regards this particular problem.

We have seen how an analysis of the nature of the perturbation (regular or singular) in a small parameter problem can be used to predict the relative success of a particular
family of numerical methods with respect to another. This was, at best, heuristics. Can the reasoning eventually be refined into one that would yield precise, provable, statements? Perhaps, but I tend to think efforts in this direction would be misdirected. In the present state of the art, it seems better to view the above explanation as a route towards a conjecture, to be confirmed or rejected on the basis of numerical experiments: that, for these problems of non-linear magnetostatics with high relative permeability, "b-oriented methods work better".

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