Abstract—The performance of micro-electro-mechanical systems depends on the interaction between electrical, mechanical, and fluidic forces. Simulating this coupled problem is made more difficult by the fact that most MEMS devices are innately three-dimensional and geometrically complicated. It is possible to simulate efficiently these devices using domain-specific solvers, provided the coupling between domains can be handled effectively. In this paper we will survey recent developments in coupled-domain simulation, and give computational comparisons between relaxation, multi-level Newton, and Newton-Iterative methods for 3-D electromechanical analysis.

I. INTRODUCTION

 Because of the specialized processing involved, the cost of prototyping even simple microsensors, microvalves, and microactuators is enormous. In order to reduce the number of prototype failures, designers of these devices make frequent use of finite-element based simulation tools. Finite-element techniques are very general, but are very inefficient when used to simulate three-dimensional geometrically-complicated micromechanical structures, where device performance is critically dependent on the interaction between electrostatic, mechanical and fluidic forces. Simulation efficiency can be substantially improved by using domain-specific solvers, provided the coupling between domains can be handled effectively. In this paper, we will survey recent developments in coupled-domain simulation, and give computational comparisons between relaxation, multi-level Newton, and Newton-Iterative methods for 3-D electromechanical analysis.

II. ELASTOSTATICS AND ELECTROSTATICS.

Micro-mechanical structures undergo large deformation when subjected to electrostatic forces. This structural deformation can be determined by solving a nonlinear force balance equation which includes the material stresses and the electrostatic pressures. In particular, standard continuum mechanical analysis [6] leads to a system of partial differential equations of the form

\[
\nabla \cdot (S \nabla u(x)) = 0 \text{ in } \Omega
\]

(1)

\[
u = 0 \text{ on } \Gamma_d
\]

(2)

\[
\nabla \cdot (S \nabla u(x)) = p \text{ on } \Gamma_h
\]

(3)

\[
\mathbf{n} \times \nabla \cdot (S \nabla u(x)) = 0 \text{ on } \Gamma_h
\]

(4)

where \( u \) is a point on the initial structure, \( u(x) \) is the displacement of that point from its initial position, \( S \) is the nonlinear algebraic operator which relates the displacement gradient to the material stresses, \( \Omega \) is the interior of the structure, \( \Gamma_d \) is that part of the structure surface whose position is fixed, \( \Gamma_h \) is the movable part of the structure surface which is acted on only by electrostatic pressure forces \( p \).

The electrostatic pressure, \( p \), on micro-mechanical conductors is related to the surface charge density, \( \sigma \), as in \( p = \frac{1}{2} \sigma \), where \( \sigma \) is the dielectric permittivity. This surface charge density can be determined by solving the integral equation

\[
\psi(x) = \int_{S_{surf}} \frac{1}{4\pi|x-x'|} da', \quad x \in S_{surf}
\]

(5)

where \( \psi(x) \) is the known conductor surface potential, \( \sigma \) is the surface charge density, \( da' \) is the incremental conductor surface area, \( x, x' \in \mathbb{R}^3 \), and \( \|x\| \) is the usual Euclidean length of \( x \) given by \( \sqrt{x_1^2 + x_2^2 + x_3^2} \).

The most commonly used approach to solving (1) are the finite-element methods [3]. Finite-element methods can also be used to solve a partial differential equation form of (5), and is a commonly used approach to solving the coupled electromechanical problem. However, for very complicated three-dimensional geometries, the electrostatic pressures can be computed much more efficiently using multipole or precorrected-FFT accelerated iterative methods applied directly to (5)[7], [8]. The difficulty addressed in this paper is finding approaches which lets one use the most efficient algorithm for each of the domains but still allows one to solve the coupled problem.

III. COUPLED METHODS

Boundary-element discretization of (5) leads to a system of equations of the form

\[
P(u + x)q = \psi = R_E(u, q) = 0
\]

(6)

where \( n \) is the number of discretization unknowns, \( q \in \mathbb{R}^n \) is a vector of surface charges, \( u + x \in \mathbb{R}^m \) is the vector of absolute surface positions, and \( P(u + x) \in \mathbb{R}^{n \times m} \) is a dense geometry dependent matrix which relates discretized surface charges to discretized surface potentials. Here, \( R_E \) is our notation for the electrostatic equilibrium equations, note that it indicates the dependence on geometry.

Finite-element discretization of (1) leads to a nonlinear system of equations of the form

\[
F_M(u) - F_E(q) = R_M(u, q) = 0
\]

(7)

where \( m \) is the number of discretization nodes, \( u \in \mathbb{R}^m \) is the vector of discretized node displacements, \( F_M(u) \in \mathbb{R}^m \) is a vector of integrated forces due to material displacement, and \( F_E \) is the vector of integrated electrostatic pressure forces. Here, \( R_M \) is our notation for the elastostatic equilibrium equations, note that it indicates the dependence on electrostatic forces.

This research was supported by ARPA under ONR contract DABT63-94-C-0053 and FBI contract F30602-92-1-0003, by SRC under contract SJ-558, and by grants from IBM and Digital Equipment Corporation. Permission to make digital/hard copy of all or part of this work for personal or classroom use is granted without fee provided that copies are not made or distributed for profit or commercial advantage, the copyright notice, the title of the publication and its date appear, and notice is given that copying is by permission of ACM, Inc. To copy otherwise, to republish, to post on servers or to redistribute to lists, requires prior specific permission and/or a fee.© 1997 ACM 0-89791-920-3/97/06..$3.50
The most obvious approach to solving the coupled system of (6) and (7) is to use a simple relaxation scheme [4], [5], [11], as diagrammed in Figure 1. The relaxation algorithm does not always converge, however, particularly when the electrostatic forces are large and the structure is extremely compliant [12].

A more robust approach is to use a full Newton algorithm to solve the coupled system of (6) and (7) is to use a simple relaxation scheme [4], [5], [11], as diagrammed in Figure 1. The relaxation algorithm does not always converge, however, particularly when the electrostatic forces are large and the structure is extremely compliant [12].

A coupled algorithm for self-consistent electro-mechanical analysis is extremely compliant [12]. However, particularly when the electrostatic forces are large and the structure is extremely compliant [12], the coupled system, as diagrammed in Figure 2 [1], the difficulty with (6) is to use a simple relaxation scheme [4], [5], [11], as diagrammed in Figure 1. The relaxation algorithm does not always converge, however, particularly when the electrostatic forces are large and the structure is extremely compliant [12].

Next Relaxation Step

\[
\begin{bmatrix}
\frac{\partial R_M}{\partial u} & \frac{\partial R_M}{\partial q} \\
\frac{\partial R_E}{\partial u} & \frac{\partial R_E}{\partial q}
\end{bmatrix}
\begin{bmatrix}
\Delta u^{(i)} \\
\Delta q^{(i)}
\end{bmatrix} =
\begin{bmatrix}
R_M(u^{(i)}, q^{(i)}) \\
R_E(u^{(i)}, q^{(i)})
\end{bmatrix}
\]

Fig. 2. A coupled algorithm for self-consistent electro-mechanical analysis.

It should be noted that if Krylov-subspace methods like GMRES are used to solve the system in Figure 2, then as much of the diagonal blocks as is possible should be explicitly factored and used as a preconditioner [1]. Otherwise, the GMRES algorithm will converge too slowly to be practical.

Next Newton Iteration

\[
\begin{align*}
\Delta u^{(i+1)} &= u^{(i)} + \Delta u^{(i)} \\
\Delta q^{(i+1)} &= q^{(i)} + \Delta q^{(i)}
\end{align*}
\]

Fig. 3. A multi-level Newton algorithm for self-consistent electro-mechanical analysis.

Preconditioning the linear system of Figure 2 requires modifying the individual solvers somewhat. Therefore, the coupled method is not really a “black box” approach in which different domain solvers can be swapped in and out. Consider instead that a program which solves (6) can be thought of as producing charges given geometric displacements, and we will denote this as

\[
q = R_E(u).
\]

In addition, a program which solves (7) can be thought of as producing geometric displacements given charges, and we denote that as

\[
u = R_M(q).
\]

A multilevel-Newton method, given in Figure 3, can be used to determine the solution to the coupled system. We refer to this as a multilevel Newton method because application of the transfer operators implies solving systems of equations, typically with an inner Newton’s method. Note that in the multilevel-Newton method, the block diagonals which are already identity matrices and need not be preconditioned. Also, application of \(\frac{\partial R_E}{\partial u}\) and \(\frac{\partial R_M}{\partial q}\) to a vector can be performed using finite-differences, and therefore does not require modifying the domain-specific solvers.

IV. RESULTS

Numerical results are presented for two examples: a beam over a ground plane and a comb drive structure. The performance of the relaxation, multi-level Newton and coupled algorithms is examined for both the examples. In particular, the convergence characteristics and the simulation times are compared.

A. Beam Example

The beam example considered here is 500 μm long, 50 μm wide, 14.35 μm thick and is positioned 1 μm above the ground plane. Figure 4 shows a top view of the beam example. The beam is discretized...
into 50 parabolic elements and the ground plane is discretized into 250 4-node elements. When a positive potential with reference to the ground plane is applied on the beam, the beam deflects towards the ground plane because of the electrostatic force. As the potential difference increases, the tip of the beam approaches the ground plane, and touches the ground plane for a certain bias defined as the pull-in voltage. The pull-in voltage for the beam considered here is 17.24 volts.

Figure 5 compares the peak deflection obtained from the relaxation, multi-level Newton and coupled algorithms. The results are identical verifying the accuracy of each solver. The deflection of the beam for an applied bias of 17.23 V is shown in Figure 6.

The performance of the relaxation, multi-level Newton and coupled algorithms for the beam example is summarized in Table I. Observe that the multi-level Newton and coupled algorithms take fewer iterations and are much faster compared to the relaxation algorithm for tightly coupled cases. Figure 7 and Figure 8 compare the convergence of the relaxation, multi-level Newton and coupled algorithms for the beam and ground plane example. Note that closer to pull-in the relaxation algorithm converges slowly, but the multi-level Newton and coupled algorithms converge rapidly. The slow convergence of the relaxation algorithm, near pull-in, is due to the increased coupling between elastostatic and electrostatic systems. As the multi-level

<table>
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<th>Bias</th>
<th># Iterations</th>
<th>CPU(sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Alg I</td>
<td>Alg II</td>
</tr>
<tr>
<td>2.0</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>4.0</td>
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</tr>
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</tr>
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<td>3</td>
</tr>
<tr>
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<td>3</td>
</tr>
<tr>
<td>12.0</td>
<td>10</td>
<td>3</td>
</tr>
<tr>
<td>14.0</td>
<td>13</td>
<td>4</td>
</tr>
<tr>
<td>16.0</td>
<td>20</td>
<td>4</td>
</tr>
<tr>
<td>17.0</td>
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<td>5</td>
</tr>
<tr>
<td>17.23</td>
<td>200</td>
<td>7</td>
</tr>
</tbody>
</table>

**B. Comb Drive Example**

The comb example consists of a deformable comb structure, a drive structure and a ground plane. As shown in Figure 9, the F-shaped finger structure is the comb, the E-shaped finger structure is the drive, and the rectangular shaped structure is the ground plane. The comb is discretized into 172 parabolic elements, the drive is discretized into 144 linear bricks and the ground plane is discretized into 2688 4-node elements. When a positive potential is applied on the drive structure, and zero potential on the comb and the ground plane, the comb structure deforms out of plane. The deformation of the comb structure for an applied bias of 85 volts is shown in Figure 10. Note that only the comb structure deforms and the drive and the ground plane do not move.

A comparison of the relaxation, multi-level Newton and coupled algorithms for the comb example is summarized in Table II. At low


Fig. 7. Convergence of relaxation, multi-level Newton and coupled algorithms for a beam and ground plane structure for an applied bias of 17.20 volts

Fig. 8. Convergence of relaxation, multi-level Newton and coupled algorithms for a beam and ground plane structure for an applied bias is 17.23 volts

Fig. 9. Comb drive example

Fig. 10. Deformation of the comb (not to scale) for an applied bias of 85 volts.

TABLE II

<table>
<thead>
<tr>
<th>Bias</th>
<th># Iterations</th>
<th>CPU(sec)</th>
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<tbody>
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<td>Alg I</td>
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<tr>
<td>75.0</td>
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</tr>
<tr>
<td>80.0</td>
<td>142</td>
<td>3</td>
</tr>
<tr>
<td>85.0</td>
<td>*</td>
<td>3</td>
</tr>
</tbody>
</table>

In this paper we presented three algorithms for coupled electro-mechanical analysis. The relaxation technique is a black-box approach but does not converge very well when the electrostatic forces are large or when the structure is very compliant. The full-Newton technique exhibits excellent convergence behavior but it is not a black-box technique. The multilevel-Newton method is a black-box technique and exhibits excellent convergence behavior. For microdevices involving two or more coupled energy domains, the multilevel-Newton technique appears to be the most attractive. CPU results indicate that both voltages, the deflection of the comb is small, the coupling between the electrical and mechanical systems is weak and the relaxation algorithm works very well. At low voltages, both multi-level Newton and coupled algorithms take fewer iterations compared to the relaxation algorithm but the simulation time for both multi-level Newton and coupled algorithms is a little longer. For higher voltages, the multi-level Newton and coupled algorithms converge much faster compared to the relaxation algorithm. For a bias of 80 volts, the multi-level Newton algorithm is about 7.7 times faster and the coupled algorithm is about 5 times faster compared to the relaxation algorithm. The convergence of the relaxation, multi-level Newton and coupled algorithms at 80 V bias is shown in Figure 11. For an application of 85 V on the drive, the relaxation algorithm fails to converge, while the multi-level Newton and coupled algorithms converge very rapidly and take 3 and 10 iterations, respectively. This is illustrated in Figure 12.

V. CONCLUSION

In this paper we presented three algorithms for coupled electro-mechanical analysis. The relaxation technique is a black-box approach but does not converge very well when the electrostatic forces are large or when the structure is very compliant. The full-Newton technique exhibits excellent convergence behavior but it is not a black-box technique. The multilevel-Newton method is a black-box technique and exhibits excellent convergence behavior. For microdevices involving two or more coupled energy domains, the multilevel-Newton technique appears to be the most attractive. CPU results indicate that both
Fig. 11. Comparison of convergence of relaxation, multi-level Newton and coupled algorithms for a comb example at an applied bias of 80 V.

Fig. 12. Comparison of convergence of relaxation, multi-level Newton and coupled algorithms for a comb example at an applied bias of 85 V.

full-Newton and multilevel-Newton are faster techniques compared to relaxation. From our preliminary studies, it is hard to conclude which one of the full-Newton and the multilevel-Newton techniques is a faster approach for microelectromechanical CAD. Several improvements to the full-Newton method are possible and are currently being investigated. A comparison of the three methods with improvements to the full-Newton method should provide better insight in indentifying the most efficient approach.

REFERENCES


