New Numerical Techniques and Tools in SUGAR for 3D MEMS Simulation

Z. Bai*, D. Bindel†, J. Clark‡, J. Demmel†, K. S. J. Pister‡, N. Zhou†
*Department of Computer Science, University of California, Davis, USA, bai@cs.ucdavis.edu
†Department of Computer Science, University of California, Berkeley, USA
dbindel@cs.berkeley.edu, demmel@cs.berkeley.edu
‡Berkeley Sensor and Actuator Center, University of California, Berkeley, USA
jvclark@bsac.eecs.berkeley.edu, pister@eecs.berkeley.edu, nzhou@bsac.eecs.berkeley.edu

ABSTRACT

SUGAR is a modified nodal analysis package for 3D MEMS simulation that owes its heritage and its name to the SPICE family of circuit simulation. SUGAR has undergone the stage of proof-of-concept which showed that modified nodal analysis was in fact just as accurate and much faster than finite element simulation on many MEMS problems. The upcoming major release of SUGAR is version 2.0, which includes a number of new features, such as thermal expansion, linearly and rotationally accelerating frames, residual strain gradient, nonlinear stiffness and user-defined models.

In this paper, we introduce two new numerical techniques and tools to be incorporated in the future release of SUGAR, namely scaling schemes to remedy artificial ill-conditioning and Krylov subspace based reduced-order modeling techniques for efficient transient analysis of dynamical systems.

Keywords: MEMS, SUGAR, modified nodal analysis, scaling, reduced-order modeling

1 Introduction

The potential impact of simulation, verification and synthesis tools to MEMS design, fabrication and applications is hard to overstate [8], [7]. There are a large number of efforts to bring such tools to the market. SUGAR is one of such efforts and it is a modified nodal analysis package for 3D MEMS simulation that owes its heritage and its name to the SPICE family of integrated circuit simulation [4], [3]. It is an open source package. SUGAR has undergone the stage of proof-of-concept which showed that nodal analysis was in fact just as accurate and much faster than finite element simulation on many MEMS problems. The upcoming release of SUGAR is version 2.0, which includes a number of new features, such as thermal expansion, linearly and rotationally accelerating frames, residual strain gradient, nonlinear stiffness and user-defined models.

In this paper, we will introduce two new numerical techniques to be incorporated in the future release of SUGAR, namely scaling schemes to remedy artificial ill-conditioning and reduced-order modeling techniques for efficient transient analysis of dynamical systems.

The coefficients of the system matrices for the ODEs used in SUGAR vary across several orders of magnitude. This poor scaling causes the system to be artificially ill-conditioned, so that when an ODE solver and Lanczos process involve solutions to linear systems associating with these matrices they get undeservedly inaccurate results. To cope with scaling difficulties, we transform the system using diagonal matrices, or equilibrate it.

The basic idea of reduced-order modeling of a dynamical system is to replace the original system by an approximating system with much smaller state-space dimension. An accurate and effective reduced-order model can be applied for steady state analysis, transient analysis and sensitivity analysis. As a result, it can significantly reduce design time and allow for aggressive design strategies. Such a computational prototyping tool will let designers try “what-if” experiments in hours instead of days. Krylov subspace methods are emerging numerical techniques for reduced-order modeling of large scale dynamical systems. They have led to a major breakthrough in the field. In this part of work, we will demonstrate how to use Krylov subspace based reduced-order modeling techniques for transient analysis of the nonlinear ODE systems arising from the SUGAR simulation. We have achieved a factor of 60 speedup for the transient analysis of an electrostatic gap-closing actuator device.

2 Simulation Model and Case Study

Currently, SUGAR uses the following matrix representation to simulate the dynamic response of MEMS due to nonlinear, time-varying forces [3]:

\[ M \ddot{q} + D \dot{q} + K q = F(q, t) \]  (1)

where \( q \) is a state vector of length \( N \). \( M, D, \) and \( K \) are the \( N \times N \) multi-energy domain system matrices, which are analogous to the mass, damping, and stiffness of a purely mechanical system. \( F(q, t) \) is the \( N \times 1 \) excitation which is a function of state and time. We can write \( F(q, t) = B u(q, t) \), where \( B \) is an \( N \times p \) input influence array to indicate the position input excitation, and \( u(q, t) \) is the input excitation source including nonlinear electrostatic force.

Furthermore, system (1) can be rewritten in the fol-
lowing multi-input and multi-output (MIMO) form
\[
\begin{cases}
M\ddot{q} + D\dot{q} + Kq = Bu(q, t) \\
y = L^T q,
\end{cases}
\]  
(2)

where \(y(t)\) is the output of the system, and \(L\) is an \(N \times m\) output influence array and is chosen to extract the components of state vector of interest. The equation (2) can be equivalently cast in the following form
\[
\begin{cases}
C\ddot{x} + Gx = \tilde{B}\tilde{u}(x, t) \\
y = \tilde{L}^T x
\end{cases}
\]  
(3)

where
\[
x = \begin{bmatrix} q \\ \dot{q} \end{bmatrix}, \quad \tilde{B} = \begin{bmatrix} B \\ 0 \end{bmatrix}, \quad \tilde{L} = \begin{bmatrix} L \\ 0 \end{bmatrix}
\]

and
\[
C = \begin{bmatrix} D & M \\ I & 0 \end{bmatrix}, \quad G = \begin{bmatrix} K & 0 \\ 0 & -I \end{bmatrix}
\]

The system (3) is similar to well-studied linear time-invariant MIMO systems (in that case, \(u(x, t) = \tilde{u}(t)\), independent of \(x\)). We can try to benefit from techniques developed for the linear MIMO systems, and exploit the idea of so-called “nonlinear dynamics using linear modes” as in [1].

We use the electrostatic gap-closing actuator shown in Fig. 1 as an example throughout this paper. A 2\(\mu\)m by 100\(\mu\)m flexure is attached to a 5\(\mu\)m by 100\(\mu\)m movable plate that extends from node b to node c. We use the parallel plate approximation to calculate the total force the plate. \(F(q, t) = Bu(q, t)\), where \(B\) only has ones in the components corresponding to the direction of force at nodes b and c,

\[
u(q, t) = -\frac{1}{2} \varepsilon_0 A \frac{v(t)^2}{\text{gap}(q)^2},
\]

\(\varepsilon_0\) is the permittivity of free space, \(\text{gap}(q)\) is the distance between the 2 plate electrodes, \(A\) is the area facing the gap, and \(v(t)\) is the voltage between the electrodes. \(M\) and \(K\) are derived using linear beam theory, while \(D\) is based on simple Couette damping, and is proportional to \(M\). All structures are fabricated in a 2\(\mu\)m polysilicon layer. While this model may not be highly physically accurate, it serves to illustrate our techniques. For more detail about the models used in the electrostatic gap-closing actuator, see [3].

3 Scaling

The nonzero coefficients of the matrix \(C\) in our example vary across 40 decimal orders of magnitude, causing \(C\) to be very poorly scaled. Consequently, \(C\) has an enormous 2-norm condition number of around 7.7 \(\times\) 10^{23}. While such a large condition number does not guarantee that a particular linear system involving \(C\) will be inaccurately solved, it does raise suspicion, and renders many of the standard error bounds ineffective. Since we solve linear systems involving \(C\) when forming the Krylov subspaces for the Lanczos procedure and at each step of the ODE solver, these large condition numbers cause concern us.

There are several standard techniques for dealing with ill-scaled problems [6]. The simplest is equilibration, which involves multiplying \(C\) on the left by a diagonal matrix which scales each row of the matrix to have unit norm. Multiplying the matrix by diagonal matrices on both the left and the right can improve the condition number even further, but undoing the column scaling subsequent to solving a linear problem with \(C\) can undo whatever benefits where conferred by the column scaling. Another technique which often remedies the effects of ill-scaling is iterative refinement, which involves computing the residual \(r = b - \hat{C}\hat{z}\) for the computed solution \(\hat{z}\), solving a linear system to find the approximate error \(C(x - \hat{z}) = r\), and then adding the approximate error to to obtain a corrected solution.

4 Reduced-Order Modeling

Krylov subspace methods are emerging numerical techniques for reduced-order modeling of large scale dynamical systems. They have led to a major breakthrough in the field; see, for example, the survey paper [5] and the references therein. The need and challenges of reduced-order modeling techniques for simulating MEMS devices are discussed in [8], [7]. In this part of our work, we will demonstrate how to use Krylov subspace based reduced-order modeling techniques for transient analysis of the nonlinear system (3).

The idea of nonlinear dynamics using linear modes is presented in [1]. A selected set of eigenvectors (modes) of the matrix \(M^{-1}K\) are used and the damping term \(D\)
is neglected. In our approach, we use the basis vectors of Krylov subspaces computed via the efficient Lanczos process directly, without the need to further compute the eigenvectors. The damping term is included in our approach.

Krylov subspace based methods for reduced order modeling have two steps: generating the bases of the underlying Krylov subspaces via a Lanczos or Arnoldi procedure, and model order reduction by coordinate transformation using the Krylov bases. We will use Lanczos process here. For simplicity, we only present the single vector Lanczos process, i.e., for the SISO (p = m = 1) system of (3). For the detail of Lanczos process including the multi-vector Lanczos process, see [2, secs 7.8, 7.9 and 7.10].

Let \( A = -(G + s_0 C)^{-1} C \) and \( R = (G + s_0 C)^{-1} \hat{B} \), where \( s_0 \) is chosen to be an expansion parameter close to the frequency of interest. The governing equations of the Lanczos process with \( A, R \) and \( \hat{L} \) can be summarized compactly in matrix form as follows:

\[
\begin{align*}
AV_n &= V_nT_n + \rho_{n+1}u_{n+1}e_n^T \\
A^TW_n &= W_nT_n + \eta_{n+1}w_{n+1}e_n^T
\end{align*}
\]

(4)

(5)

where \( T_n \) and \( T_n^\perp \) are tridiagonal matrices, and the columns of \( N \times n \) matrices \( V_n \) and \( W_n \) are called right and left Krylov vectors and span the so-called right and left Krylov subspaces

\[
\begin{align*}
K_n(A, R) &= \text{span}\{R, AR, \ldots, A^{n-1} R\} \\
K_n(A^\perp, \hat{L}) &= \text{span}\{\hat{L}, A^\perp \hat{L}, \ldots, (A^\perp)^{n-1} \hat{L}\}
\end{align*}
\]

and furthermore satisfy the biorthogonality conditions

\[
W_n^TV_n = \Delta_n, \quad W_n^Tv_{n+1} = 0, \quad V_n^Tw_{n+1} = 0.
\]

(6)

where \( \Delta_n \) is a diagonal matrix. From (4) and (6), it follows that

\[
W_n^TA V_n = \Delta_n T_n
\]

(7)

We note that for the implementation of the Lanczos process, the matrix \( A \) is involved only in the form of matrix-vector multiplication. Hence the structure and sparsity of \( A \), and correspondingly the matrices \( M, D \) and \( K \) can be exploited for memory saving and computational efficiency.

This Krylov-Lanczos process can be used as a building block for reduced-order modeling and applications for steady-steady, transient and sensitivity analysis of dynamical systems. In this paper, we will focus on how to do transient analysis of the system (3).

Again, let \( A = -(G + s_0 C)^{-1} C \) and \( R = (G + s_0 C)^{-1} \hat{B} \), then the equation (3) can be written as

\[
\begin{align*}
-\hat{A} z(t) + (I + s_0 A) z(t) &= \hat{R} z(t), \\
y(t) &= \hat{L}^T z(t),
\end{align*}
\]

where \( z(t) \) is the new state vector of length \( n \), then an approximate dynamical system is

\[
\begin{align*}
-\hat{A} z(t) + (I + s_0 A) z(t) &= \hat{R} z(t), \\
y(t) &= \hat{L}^T z(t).
\end{align*}
\]

(8)

Multiplying \( W_n^T \) from the left, we have

\[
\begin{align*}
-\bar{W}_n^T \hat{A} V_n z(t) + \bar{W}_n^T (I + s_0 A) V_n z(t) &= \bar{W}_n^T \hat{R} V_n z(t), \\
y(t) &= \hat{L}^T V_n z(t).
\end{align*}
\]

Therefore, application of (7), yields an \( n \)-th reduced-order model

\[
\begin{align*}
-\Delta_n T_n \bar{z}(t) + (\Delta_n + s_0 \Delta_n T_n) \bar{z}(t) &= B_n \bar{u}(V_n z(t), \\
y(t) &= \hat{L}_n^T \bar{z}(t).
\end{align*}
\]

where \( B_n = \bar{W}_n^T \hat{R} \) and \( \hat{L}_n = V_n^T \hat{L} \).

As a result, instead of solving the original system (3) of dimension \( N \) for \( y(t) \), one can solve the reduced-order model (8) of dimension \( n \) for \( \bar{y}(t) \) as an approximation of \( y(t) \). The strength of this approach derives from the fact that \( n \) is much less than \( N \) in many cases. The constructed Krylov subspaces contain the necessary information for the dominant modes to capture the dynamic response of the system.

5 Results and Discussion

For the case study of the gap-closing actuator described in section 2, a piecewise linear voltage function \( v(t) \) is applied across the gap. The voltage \( v(t) \) ramps from 5V at \( t = 10 \mu \text{sec} \) to 12V at \( t = 500 \mu \text{sec} \), and then drops to 0V. The displacement component of node c in the direction of force is observed. The initial voltage step causes the device to oscillate. As the voltage increases at a linear rate, the gap decreases at a nonlinear rate due to the electrostatic force increasing proportionally to \( 1/\text{gap}(q)^2 \). This force also causes the period of oscillation to increase. Once the voltage is removed, the actuator exponentially decays back to equilibrium due to viscous air damping. These phenomena are captured in numerical simulation. Fig. 2(a) shows the displacement versus time. Fig. 3 shows the decays back to equilibrium. In Fig. 2(a), the displacement curves of the original system and the reduced-order models are overlapped. The accuracy of the reduced-order model can be seen in Fig. 2(b). There relative errors \( |y(t) - \bar{y}(t)|/|y(t)| \) are plotted for the 2nd and 6th order of the reduced-order model. The order of the original model is \( N = 30 \).

The solution of the full system over 1 ms took 28533 seconds. However, the construction of the 2nd and 6th
order system and the solution of the reduced-order systems took 2 seconds and 428 seconds respectively. Therefore, with the satisfactory accuracy of the 6th order system, we have achieved a factor of 60 speedup for the transient analysis of the electrostatic gap-closing actuator.

Although $M$, $D$, and $K$ are all symmetric positive definite, they are ill-scaling and ill-conditioned. Let us examine the matrix $C$ in detail. The condition number of the matrix $C$ is $7.7 \cdot 10^{23}$. Its largest singular value is $1$ and the smallest singular value is $1.3 \cdot 10^{-24}$. With row equilibration, the condition number is reduced to $8.2 \cdot 10^{9}$ and the largest and smallest singular values are $2.3$ and $2.8 \cdot 10^{-10}$, respectively. Furthermore, with both row and column equilibrations, we have a nearly perfect conditioned matrix, with condition number 48, and the largest and smallest singular values are $2.1$ and $4.4 \cdot 10^{-2}$, respectively. The issue concerning the ill-scaling and artificial ill-conditioning are pointed out. The impact of scaling and equilibration to the reduced-order modeling methods and ODE solvers are still under investigation.

All numerical simulations were run under Matlab on a SUN 440MHz Ultra 10 workstation. We used the ODE solver ode15s for the results reported in this paper.

In the future, we plan to develop a reduced-order modeling technique for a more general case where the right-hand side of equation (1) is a function of $q$, $\dot{q}$, and $t$:

$$M \ddot{q} + D \dot{q} + Kq = F(q, \dot{q}, t).$$

Here, the function $F(q, \dot{q}, t)$ may also include the variations in $M$, $D$, and $K$ matrices that depend on state and time.

Figure 3: displacement of node c vs. time, back to equilibrium, using the 6th reduced-order model

REFERENCES


