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Balancing-Based Model Reduction for Fast Power Integrity Verification

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Abstract—This paper presents a novel Model Order Reduction approach as an extension of the well-known Balanced Truncation. The key contribution is a semi-analytical approach for the numerical evaluation of the Gramian matrices that are functional for optimal reduction of large-scale state-space systems. Application to product-level full-system Power Distribution Networks including integrated voltage regulation circuitry allows transient verification in more than 200× faster than SPICE.

I. INTRODUCTION

This work is motivated by the need of efficient and reliable power integrity verification via transient analysis of system-level Power Distribution Networks (PDN). Specifically, we address modern HPC or AI microprocessor systems where a bank of Fully Integrated Voltage Regulators (FIVRs) stabilize the voltage supplied to more than a hundred computing cores. Interaction of the switching circuitry of the FIVRs with the large-scale models of board, package, passive on-package inductors, capacitors, and chip models make an all-coupled full-system transient analysis very challenging.

To enable such verification framework, a number of Model Order Reduction (MOR) approaches have been presented by the Authors, with the aim of reducing the model sizes while preserving accuracy, hence reducing computational cost. Our previous approaches based either on parameterized rational macromodeling of the PDN impedance [1] or moment-matching through structured Krylov subspace projection [2], are not certified in their overall accuracy since only a-posteriori error control is possible. Therefore, one has no guarantee that the reduced model will fulfill the desired accuracy constraints.

In this paper, we propose a MOR approach based on Balanced Truncation [3], which does provide explicit and computable error bounds. Such an approach is traditionally very challenging for large-scale systems due to the required evaluation of some Gramian matrices through full linear algebra methods. We propose a new method for this evaluation, based on a semi-analytical approach enabled by an accuracy-controlled rational expansion of the system state responses. Application of the proposed method to two PDNs of real products (a mobile and an enterprise server platform) confirms both the accuracy and a major speedup with respect to full-system SPICE simulations.

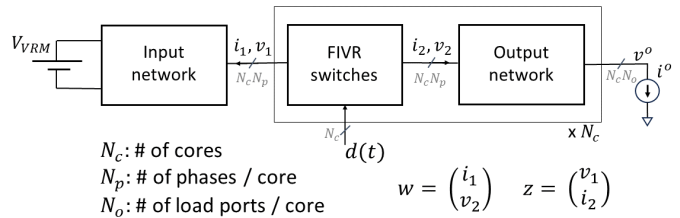


Fig. 1. Schematic representation of a FIVR-equipped power distribution network. Internal variables w, z represent currents/voltages on the primary and secondary sides of the averaged converter model (FIVR switches block).

II. PROBLEM STATEMENT AND NOTATION

The topology of power delivery networks that are used as case studies in this work is reported in Fig. 1. The array of integrated voltage regulators defines a boundary between an input subnetwork (including EM-accurate models of board and package, as well as decaps and VRM models) and an output network (including EM-accurate models of regulator inductors as well as MIM capacitances and chip core models). These two networks are passive LTI systems, for which we assume that a state-space description is available, as in [1]. Regarding the switching circuitry, we adopt an averaged converter model where the switches are modeled as ideal transformers, with time-varying conversion ratios equal to the per-core instantaneous duty cycle signals.

As depicted in Fig. 1, the system is loaded by a constant nominal VRM voltage and by load currents i^o at all output ports of all cores. Such inputs are collected in vector u . The outputs of interest are the load voltages v^o . We further identify a set of internal variables, namely the voltages and the currents on primary and secondary sides of the averaged converter models, collected in the vectors w and z . Based on these variables, the PDN system admits the following description

$$\begin{cases} \dot{x} = \mathbf{A}x + \mathbf{B}_w w + \mathbf{B}_u u \\ z = \mathbf{C}_z x + \mathbf{D}_{zw} w + \mathbf{D}_{zu} u \\ v^o = \mathbf{C}_y x + \mathbf{D}_{yw} w + \mathbf{D}_{yu} u \\ w = \mathbf{\Delta}(d)z \end{cases} \quad (1)$$

with the vector d collecting all duty cycle signals from each core. In (1), the first three equations describe the input and output networks. These are coupled with the ideal transformer constitutive relations, that can be written compactly

as $\mathbf{w} = \mathbf{\Delta}(d)\mathbf{z}$. During transient analysis, system (1) is augmented with models of the controllers that sense the output voltage and return per-core duty cycle signals in a closed-loop configuration (not shown, see [1] for details).

Considering a nominal duty cycle value as the operating point, the dynamics of the PDN can be approximated with a small-signal linearization as described in [2]. Hence, for the purposes of the formulation, we can focus on a generic linear system described by a state-space realization $(\mathcal{A}, \mathcal{B}, \mathcal{C}, \mathcal{D})$.

III. FORMULATION

Balanced Truncation (BT) [3] is a well known method to obtain a simplified model of reduced state-space dimension n that preserves the input-output behavior of a given full-order system. BT finds principal directions in the state space that are simultaneously most controllable and observable using a pair of special matrices, namely the controllability Gramian \mathbf{P} and the observability Gramian \mathbf{Q} . Based on these, projection matrices \mathbf{S}_r and \mathbf{S}_l can be constructed [3], [4], which are used to remove the least important states. The most appealing feature of BT is the availability of an explicit error bound on the approximation error, computed from the truncated *Hankel singular values* [5].

A. Approximate Gramians via Vector Fitting

Computation of system Gramians \mathbf{P} , \mathbf{Q} can be carried out by solving two associated Lyapunov equations. For very large-scale systems, this direct solution becomes too computationally expensive, so that alternative methods have been devised, e.g. iterative methods [6]. A simpler way of approximating \mathbf{P} and \mathbf{Q} , proposed in [7], is based on the following identities

$$\begin{aligned} \mathbf{P} &= \frac{1}{2\pi} \int_{-\infty}^{+\infty} \mathbf{X}(j\omega) \mathbf{X}^T(-j\omega) d\omega \\ \mathbf{Q} &= \frac{1}{2\pi} \int_{-\infty}^{+\infty} \mathbf{X}_d(j\omega) \mathbf{X}_d^T(-j\omega) d\omega \end{aligned} \quad (2)$$

with $\mathbf{X}(j\omega) = (j\omega\mathbb{I} - \mathcal{A})^{-1}\mathcal{B} \in \mathbb{C}^{N \times P}$ and $\mathbf{X}_d(j\omega) = (j\omega\mathbb{I} - \mathcal{A}^T)^{-1}\mathcal{C}^T$. In [7], it is suggested that these integrals can be evaluated by quadrature rules, so that \mathbf{P} is approximated as a suitably weighted sum of the snapshots $\mathbf{X}(j\omega_k)$ computed at K nodes $\{\omega_k\}_{k=1}^K$, and similarly for \mathbf{Q} . Here, we make a different use of these snapshots as we propose an alternative way of approximating the Gramians. The key observation is that, if an (approximate) pole-residue expansion

$$\mathbf{X}(s) \approx \check{\mathbf{X}}(s) = \sum_{i=1}^{\nu} \frac{\mathbf{R}_i}{s - q_i} \quad (3)$$

is available in terms of ν stable poles $\{q_i\}_{i=1}^{\nu}$, then the computation of the integral can be carried out analytically. First, we replace $\mathbf{X}(s)$ in (2) with the approximation $\check{\mathbf{X}}(s)$. Then the residue of the integrand at each left-half plane pole q_i is given by $\mathbf{R}_i \check{\mathbf{X}}^T(-q_i)$. At this point, the Residue Theorem is invoked [8, App. E.2] to express the integral along the imaginary axis in terms of these residues,

$$\mathbf{P} \approx \sum_{i=1}^{\nu} \mathbf{R}_i \check{\mathbf{X}}^T(-q_i).$$

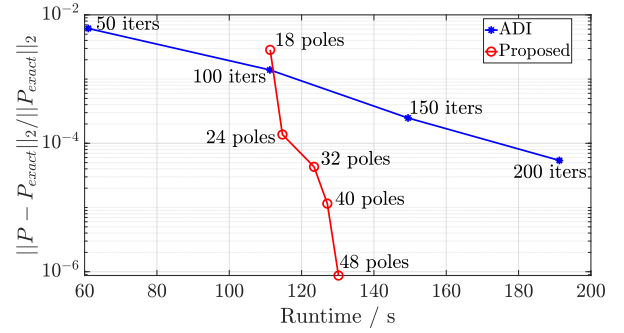


Fig. 2. Convergence of the proposed method compared to the ADI method [10]. The error norm is reported as a function of overall runtime.

The pole-residue approximation (3) is here determined using the Vector Fitting algorithm to find suitable basis poles q_i , starting from a limited number K of evaluations $\mathbf{X}(j\omega_k)$. Once the basis poles are known, the residues \mathbf{R}_i in (3) are computed by exploiting the optimality condition given in [9], which in our setting reads

$$\sum_{i=1}^{\nu} \frac{\mathbf{R}_i}{-q_j - q_i} = \mathbf{X}(-q_j), \quad j = 1, \dots, \nu,$$

where $\mathbf{X}(-q_j)$ are ν additional evaluations of $\mathbf{X}(s)$ at the mirror images of the basis poles. For each matrix entry (h, m) of \mathbf{R}_i , this condition is a $\nu \times \nu$ linear system giving the ν unknown residue entries $(\mathbf{R}_i)_{hm}$, $i = 1, \dots, \nu$. The coefficient matrix for this linear system is derived from the Cauchy matrix $\mathbf{\Phi}$ defined by $(\mathbf{\Phi})_{ij} = (-q_i - q_j)^{-1}$ as follows

$$(\mathbf{R}_1 \quad \dots \quad \mathbf{R}_\nu) (\mathbf{\Phi} \otimes \mathbb{I}_P) = (\mathbf{X}(-q_1) \quad \dots \quad \mathbf{X}(-q_\nu))$$

IV. NUMERICAL RESULTS

A. Validation of MOR via approximate Gramians

We assess the proposed algorithm for Gramian computation via VF. Results are presented for the input network of the PDN of an Intel-based enterprise server (see Fig. 1). The starting point is a passive LTI network with 181 ports and dynamic order 6170. The rational approximation (3) is determined using $K = 70$ log-spaced frequency values in $[0, 10]$ GHz. Figure 2 shows that, as ν increases, the error between the approximate Gramian \mathbf{P} and the exact solution computed via Lyapunov equation (direct method) decreases. Moreover, proposed method takes a shorter time to produce a more accurate solution with respect to ADI. Figure 3 (bottom panel) compares two selected responses of the full-size system and the reduced-order model. The latter is basically indistinguishable from the reference even if the number of states was reduced to 700, as confirmed by the Hankel singular values (top panel), which provide the explicit bound on the model approximation.

B. Modeling a full-system Power Distribution Network

We now consider a complete Power Distribution Network including multi-phase FIVRs ($N_p = 3$) for $N_c = 60$ cores,

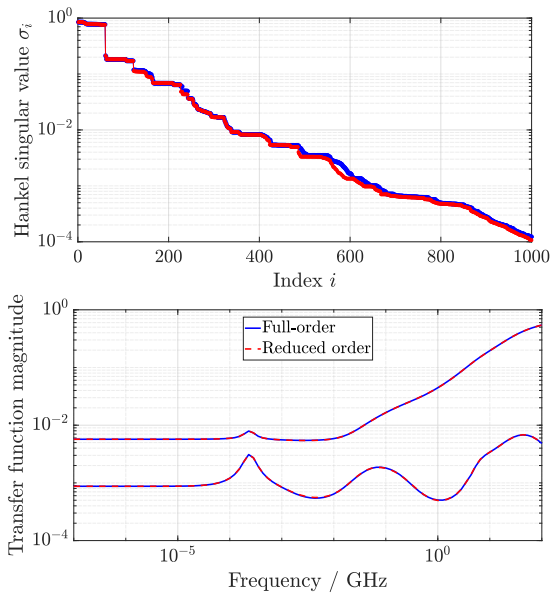


Fig. 3. The top panel shows the (normalized) system singular values of the input network computed with exact Gramians (blue) and approximate ones (red). The bottom panel shows two matrix entries of the input network transfer function computed using a reduced model of order 700, much lower than the initial 6170 states.

extracted from an Intel-based enterprise server platform. Each core has $N_o = 57$ outputs, leading to a total number $N_c N_o = 3420$ of output voltages to be stabilized. For this system, the proposed approach was applied to compute Gramians of the locally linearized approximation around the nominal duty cycle. The resulting bases were used in a structured projection with passivity preservation similarly to [2]. Figure 4 shows the results of a transient analysis where excitation currents range between 0 and 20 A/core. All cores are persistently excited starting from $t = 0.1 \mu\text{s}$ with diversified current profiles. For this test case, we applied MOR to reduce the dynamic order from more than $5 \cdot 10^4$ down to 400. Transient simulation of the reduced system takes only 66 s, corresponding to a speedup of more than $15 \times$ with respect to the full-order simulation (1038 s), both carried out through a simple MATLAB solver based on the Backward Euler method. The maximum error among all output voltages is lower than 2 mV at all times.

In order to enable a comparison with less efficient reference approaches, a simplified version of the same structure was realized by including only 16 cores. The results of this comparison are reported in Table I, which confirms that proposed method provides clear improvements with respect to [2] in terms of accuracy for a given order, and with respect to [1] in both accuracy and efficiency.

V. CONCLUSIONS

This paper presented a method derived from balanced truncation that is applicable to large-scale systems, with a particular focus on acceleration of power integrity verification analyses for multi-core microprocessors. Besides proposing a novel procedure to compute approximate Gramian matrices

TABLE I
COMPARATIVE NUMERICAL ANALYSIS BASED ON 16-CORES BENCHMARK

Method	Order	Max. error	Runtime
Full order (HSPICE)	-	-	1410 s
Full order (MATLAB)	18074	-	139 s
Approx. balancing (this paper)	250	0.9 mV	6 s
Moment-matching as in [2]	250	1.6 mV	6 s
Parametric fitting as in [1]	-	24 mV	181 s

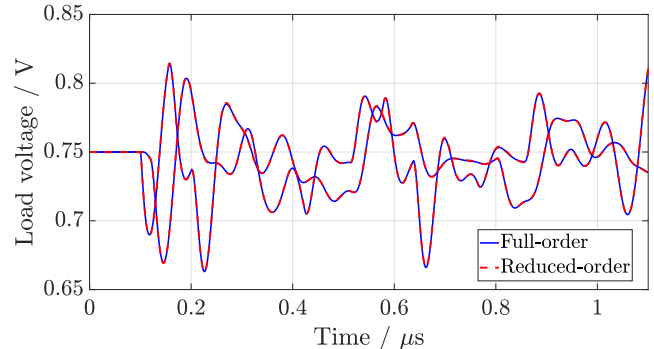


Fig. 4. Transient analysis comparing load voltage waveforms at two particular load ports obtained from the full-order model and the reduced-order model of the PDN of a 60-core Intel-based enterprise server.

using rational fitting, the presented approach improves on existing work [2] by providing higher accuracy and guidance in choosing the reduced model order through the concept of system singular value, inherited from balanced truncation. Major speedup factors in transient simulation are observed.

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