A Multistage Adaptive Sampling Scheme for Passivity Characterization of Large-Scale Macromodels

Marco De Stefano, Graduate Student Member, IEEE, Stefano Grivet-Talocia, Fellow, IEEE, Torben Wendt, Graduate Student Member, IEEE, Cheng Yang, Member, IEEE, and Christian Schuster, Senior Member, IEEE

Abstract—This article proposes a hierarchical adaptive sampling scheme for passivity characterization of large-scale linear lumped macromodels. In this article, large scale is intended both in terms of dynamic order and especially number of input–output ports. Standard passivity characterization approaches based on spectral properties of associated Hamiltonian matrices are either inefficient or nonapplicable for large-scale models, due to an excessive computational cost. This article builds on existing adaptive sampling methods and proposes a hybrid multistage algorithm that is able to detect the passivity violations with limited computing resources. Results from extensive testing demonstrate a major reduction in computational requirements with respect to competing approaches.

Index Terms—Adaptive sampling, Hamiltonian matrices, macromodeling, model order reduction, multiscale optimization, passivity, scattering, shielding.

I. INTRODUCTION

MACROMODELING techniques are generally considered as key enabling factors for the efficient simulation of complex devices and systems. A macromodel approximates the input–output response of a given structure through a compact set of behavioral equations. Such equations are most commonly derived through some fitting process [1] applied to a set of sampled responses, usually in the frequency domain (see [2] for an overview).

The focus of this work is on macromodeling of passive structures, which are unable to generate energy on their own [3]–[6]. Model passivity must be, therefore, checked and enforced through suitable constraints, which are usually applied through a perturbation step applied to an initially non-passive model obtained by some unconstrained fitting [7]–[19]. Lack of model passivity can trigger instabilities in transient analyses [20] and must be therefore avoided with care.

Macromodels are intended to be compact and reduced order and, hence, small size. Nonetheless, the robustness of state-of-the-art macromodeling algorithms has enabled applications that are aggressively driving model complexity beyond current capabilities. One typical example can be full-package modeling for signal, power, or coupled signal-power integrity analysis, in which case the number of input–output ports can reach hundreds or even thousands [21]. A second application is modeling of energy-selective electromagnetic shielding structures [22] formed by conventional enclosures having openings that are closed by regular grids of nonlinear elements [23]–[25]. An efficient time-domain simulation calls for a macromodel that represents in a compact way the electromagnetic behavior of the shield. Also, in this case, the number of input–output ports can reach the order of thousands or more. A further complication is provided by the possibly high dynamic order (number of poles) of the macromodels, as required by accuracy requirements over extended frequency bands.

Recent vector fitting (VF) [1] implementations [26], [27], including parallel codes [28], [29], have been demonstrated to scale very favorably with the model size. Conversely, standard passivity enforcement schemes are more critical and less scalable, despite the advancements that have been documented over the last two decades [30]. The major bottleneck is the passivity verification step, which must detect not only whether a given initial model is passive but should also provide a precise localization of the passivity violations so that a successive perturbation stage can be set up to remove them. Complexity of this model perturbation step is less critical, so passivity enforcement is not considered here. Passivity characterization is far more demanding in terms of computational resources, both memory and runtime, and therefore more problematic. To the best of authors’ knowledge, there is no documented passivity characterization approach that can provide reliable results in a limited runtime and with limited memory occupation, for systems with input–output ports in the order of hundreds and possibly exceeding 1000.

The most prominent passivity characterization approaches can be roughly grouped in three main classes, based,
respectively, on linear matrix inequalities (LMIs) [6], direct sampling, and Hamiltonian matrix properties [16], [31]. LMIs provide the best approach from a theoretical standpoint, but applicability is limited to small-scale systems. Pure sampling approaches are very fast and provide potentially good scalability, but they are likely to miss passivity violations making the verification not reliable. Hamiltonian matrices with their spectral properties are the usual method of choice for passivity verification, for small- and medium-scale problems. Extension to large-scale systems has been attempted in [17], possibly through hybridization, preprocessing, or postprocessing with adaptive sampling methods [18]. However, the necessary Krylov subspace iterations for the spectral characterization of large and sparse Hamiltonian matrices make the entire approach infeasible when the number of ports exceeds about 100.

The above difficulties motivate the passivity verification approach that we propose in this work. We build on the adaptive sampling scheme mentioned in [18], and we try to avoid completely the use of Hamiltonian matrices and fully coupled LMI conditions, which are the main ones responsible for driving computational cost beyond what is usually affordable in design flows. The main objective that we pursue is to construct an adaptive sampling scheme that is fast and at the same time reliable. We should stress that there is no mathematical guarantee that checking local passivity conditions on a finite number of points will not miss a violation. Therefore, the main objective of this work is to make the presence of undetected passivity violations extremely unlikely. This is the reason why we propose a hybrid scheme that combines different basic sampling methods in a hierarchical way.

The proposed algorithm performs an initial coarse pole-based sampling similar to [18], intended to locate fast variations. Differently from [18], the proposed scheme detects passivity violation subbands avoiding both eigenvector tracking, which causes unnecessary overhead, and especially any Hamiltonian eigenvalue search through a dedicated multishift Arnoldi process. These two improvements provide much better scalability with model size, as demonstrated by the examples. The results of the above coarse sampling are then used to produce an adaptive frequency warping and subdivision on a possibly large number of subbands, which are then subjected to hierarchical refinement. The basic scheme mentioned in [32] is enriched with a set of passivity-driven strategies to tune the algorithm behavior to the goal of passivity verification.

The proposed scheme was tested against a large number of regression test cases, with a number of ports ranging from $P = 1$ to $P = 640$ and a dynamic order (number of states) ranging from $N = 10$ to more than $25 \times 10^3$. The results show that standard Hamiltonian-based approaches remain competitive and should be preferred for small-scale systems. Conversely, the proposed approach outperforms the Hamiltonian tests for medium- and large-scale systems, with speedup in runtime reaching and exceeding $200\times$ for the largest cases.

This article is organized as follows. Section II provides some background information and sets notation. Section III states the main problem by providing an illustrative example that highlights the main difficulties to be addressed. Section IV introduces the adaptive frequency warping scheme that forms the initial step of the proposed sampling scheme. Section V introduces the multiscale search to be applied as a second-stage adaptive refinement. Numerical results are presented in Section VI and conclusions are drawn in Section VII.

II. BACKGROUND AND NOTATION

Throughout this work, we consider a lumped linear time-invariant (LTI) model, which is known either through a pole-residue expansion or a state-space realization

$$
H(s) = \sum_{n=1}^{N} \frac{R_n}{s - p_n} + R_0 = C(sI - A)^{-1}B + D
$$

where $s$ is the Laplace variable and the transfer matrix $H(s) \in \mathbb{C}^{P \times P}$ with $P$ number of input–output ports. The dynamic order of the model is denoted as $N$, which is the size of the state-space matrix $A$. In case the residue matrices $R_n$ are full-rank, we have $N = nP$ (see [2], [33], [34] for details).

In general, $N \gg P$. We assume that the bandwidth where the model is assumed and intended to be accurate is $[0, \omega_{\text{max}}]$, and that all model parameters (poles-residues or state-space matrices) are available from some initial fitting/approximation stage. We assume that all state-space matrices are real-valued and that the model poles $p_n$ (equivalently, the eigenvalues of $A$) are strictly stable. This condition is easily enforced by widespread fitting schemes such as VF [1].

A. Passivity Verification

We will consider only models (1) in scattering representation so that $H(s)$ approximates the scattering matrix of the modeled system. This is not a limitation since the proposed approach can be extended to other representations (impedance, admittance, or hybrid) with simple modifications (see [2] for a complete theoretical framework applicable to all representations). We review below the three main available approaches for passivity verification.

1) Sampling Local Passivity Conditions: For scattering models (1) with real-valued realization and strictly stable poles, the passivity conditions can be stated as the unique constraint

$$
\sigma_{\text{max}}(H(j\omega)) \leq \gamma = 1 \quad \forall \omega \in \mathbb{R}.
$$

This condition expresses that the maximum singular value of the scattering matrix must not exceed the threshold $\gamma = 1$ at any real frequency. Under the working assumptions, (2) is a sufficient condition for $H(s)$ to be Bounded Real, hence passive [3], [5], [6].

Sampling approaches test (2) over a finite number of frequency points $\{\omega_k, \ k = 1, \ldots, K\}$. Determination of $\sigma_{\text{max}}$ at a single frequency requires $O(P^3)$ operations, and hence, the overall complexity for $K$ samples amounts to $O(KP^3)$. Since each individual frequency $\omega_k$ can be processed independently, there is no significant memory cost, even for large $P$. The main advantage of this sampling-based check is the availability of all local maxima of $\sigma_{\text{max}}$, obtained as
trivial postprocessing. These maxima are essential to setup passivity enforcement through constrained perturbation. The main disadvantage of checking (2) over $K$ finite points is the possibility to miss local maxima exceeding $\gamma = 1$ by a small amount or over a small frequency extent, due to the limited number of points that can be processed.

We remark that the (scalar) function of frequency

$$\varphi(\omega) = \sigma_{\max}[H(\omega)]$$

henceforth denoted as passivity metric is generally smooth and differentiable, with the possible exception of a finite number of points where two singular values cross. In the latter case, $\varphi(\omega)$ is guaranteed to be continuous since all singularities (poles) of $H(s)$ fall outside the imaginary axis.

2) Linear Matrix Inequalities: LMI conditions provide a fully algebraic test that does not require sampling. We recall that the scattering system (1) is passive (dissipative) if and only if $\exists P = P^T > 0$ such that

$$\begin{bmatrix} A^T P + P A + C^T C & P B + C^T D \\ B^T P + D^T C & -(I + D^T D) \end{bmatrix} \leq 0.$$  (4)

This condition, known as bounded real lemma (BRL) or Kalman–Yakubovich–Popov (KYP) lemma [6], [35], has the major disadvantage of requiring the determination of the $N \times N$ matrix $P$ related to the internal energy storage. The computational cost for checking the feasibility of (4) is $O(N^6)$, which can be reduced to $O(N^4)$ with specialized formulations [36]. This cost becomes exceedingly high for large-scale systems. One additional disadvantage of (4) is that this is a pass/fail test, which does not provide localization of passivity violations that can be used in a successive enforcement. Passivity enforcement schemes exist that use directly (4) as constraint [7], [37], but these inherit the poor scalability with large $P$ and $N$ (see [19]).

3) Hamiltonian Matrices: Spectral properties of Hamiltonian matrices provide the standard approach for checking passivity of small- and medium-scale models. Under the working assumptions, the model is (strictly) passive if and only if the Hamiltonian matrix

$$\mathcal{M} = \begin{pmatrix} A + BR^{-1}D^T C & BR^{-1}B^T \\ -C^T S^{-1} C & -A^T - C^T D R^{-1} B^T \end{pmatrix}$$  (5)

with $R = I - D^T D$ and $S = I - D D^T$ has no purely imaginary eigenvalues $\mu_k = j\omega_k$. Such eigenvalues denote the crossing points of any singular value of $H(j\omega)$ with the passivity threshold $\gamma = 1$ (see [16], [31]). Therefore, they also provide localization of passivity violations. Unfortunately, the eigenvalue spectrum of $\mathcal{M}$ is required, whose numerical determination scales as $O((2N)^3)$. Extension to large-scale and sparse (decompositions) was documented in [17] and [18], allowing some improvement for systems with moderate $P$ and possibly large $\bar{n}$. Extension to large port count $P$ appears problematic since the Krylov subspace iterations required for detection of purely imaginary eigenvalues of $\mathcal{M}$ require the repeated inversion of full $P \times P$ matrices, leading to a cost that once again becomes impractical in the large-scale case.

We remark that (5) is applicable only when state-space matrix $D$ has no singular values too close to $\gamma = 1$. In this case, it is preferable to process the extended Hamiltonian pencil $(\mathcal{M}_e, \mathcal{K})$ defined as [38]

$$\mathcal{M}_e = \begin{bmatrix} A & 0 & B & 0 \\ 0 & -A^T & 0 & -C^T \\ 0 & B^T & -I & D^T \\ C & 0 & D & -I \end{bmatrix}$$

$$\mathcal{K} = \begin{bmatrix} I & 0 & 0 \ 0 & 1 & 0 \ 0 & 0 & 0 \ 0 & 0 & 0 \end{bmatrix}$$

to find its purely imaginary generalized eigenvalues $\mu_k$. Some of the test cases that will be investigated in Section VI fall in this situation, in which the standard Hamiltonian matrix (5) is ill-defined. Throughout this work, we use (6) only when $|\sigma_{\max}[D] - 1| < 10^{-4}$ since the associated computational cost is higher than for (5).

B. Discussion

Table I summarizes the number of elementary operations for the three leading passivity characterization approaches. Only the leading terms are included, and the number of floating-point operations is reported up to a constant factor, which depends on the specific algorithm or numerical library used to implement each method.

As an example, let us consider a system with $P = 400$ ports and $\bar{n} = 50$ poles (hence, $N = 20000$). The LMI and the Hamiltonian approaches would require about 3 and 12 GB of storage, respectively, with a relative CPU cost with respect to sampling a huge number $K = 10^5$ of frequencies about $20000 \times 9 \times$, respectively. From this example and Table I, it is evident that the only framework that has the potential to scale favorably with the problem size is the sampling approach. Therefore, we concentrate our efforts on the sampling scheme, with the main objective of reducing the likelihood of missing passivity violations and classifying a nonpassive model as passive.

III. Problem Statement and Goals

Considering the passivity condition (2) and the definition of the passivity metric (3), we can state the main problem that is here addressed as: find all local maxima larger than a given threshold of a continuous univariate function of frequency over the entire frequency axis. This problem may appear simple, but various difficulties arise in practical implementation. These are discussed below with reference to the top of Fig. 1 and its enlarged view in the top of Fig. 2.

1) The passivity metric $\varphi(\omega)$ is usually characterized by multiple peaks, and it is necessary to identify all of them.
Fig. 1. Top: graphical illustration of the adaptive sampling-based passivity characterization. Bottom: rescaled passivity metric \( \theta(\zeta) \) after adaptive frequency warping.

Fig. 2. Enlarged view of Fig. 1.

in order to check whether they exceed the threshold \( \gamma \). It is imperative not to miss any local maximum.

2) Some peaks can be extremely narrow, and hence, they can be almost invisible with an inadequate sampling.

3) Some peaks may be characterized by a local maximum that is very close to the threshold; in such cases, it may be difficult to detect whether there exists a passivity violation with \( \varphi(\omega) > \gamma \) at some frequency point.

The above difficulties can only be addressed by adapting the sampling process, in order to automatically refine the resolution where \( \varphi(\omega) \) is characterized by fast variations and/or its values are close to the threshold \( \gamma \). At the same time, it is mandatory to limit as much as possible the total number \( K \) of computed samples.

These objectives are here pursued through a multistage adaptive sampling scheme. First, a preprocessing step is applied to split the frequency band into a possibly large number of subbands, whose individual extent is adaptively defined and rescaled so that the total variation of the passivity metric becomes nearly uniform (see Section IV). Second, a hierarchical tree-based refinement is applied within each individual subband to track all local maxima (see Section V). The combination of these two stages will be demonstrated to outperform competing approaches, still retaining (and in some cases even improving) the reliability of algebraic passivity verification methods.

IV. STEP 1: POLE-BASED ADAPTIVE FREQUENCY WARping

A preview of the results of this first step is available in the bottom of Fig. 1 and its enlarged view of Fig. 2. We want to identify a nonlinear (invertible) frequency transformation \( \zeta = W(\omega) \) that induces the following change of variable:

\[
\theta(\zeta) = \varphi(W^{-1}(\zeta)) \tag{7}
\]

with the main objective of “flattening” the local variations of \( \varphi(\omega) \) so that the resulting \( \theta(\zeta) \) is characterized by peaks with an approximately uniform width.

The change of variable is constructed based on a set of control points \( \Omega = \{\hat{\omega}_0, \ell = 0, \ldots, L\} \) such that

\[
0 = \hat{\omega}_0 < \hat{\omega}_1 < \cdots < \hat{\omega}_L < \hat{\omega}_{L+1} < \cdots < \hat{\omega}_\infty = \infty. \tag{8}
\]

These points are used to build a local linear map between the normalized subband \( \zeta \in [\ell, \ell + 1] \) and the corresponding subband \( \omega \in [\hat{\omega}_\ell, \hat{\omega}_{\ell+1}] \). The global change of variable \( W \) is constructed piecewise as

\[
\zeta = \ell + \frac{\omega - \hat{\omega}_\ell}{\Delta_\ell} \quad \forall \omega \in [\hat{\omega}_\ell, \hat{\omega}_{\ell+1}], \quad \ell = 0, \ldots, L - 2 \tag{9}
\]

where \( \Delta_\ell = \hat{\omega}_{\ell+1} - \hat{\omega}_\ell \) is the extent of the \( \ell \)-th subband. The last (infinite-sized) subband is handled as

\[
\zeta = \ell + \frac{\omega - \hat{\omega}_\ell}{\hat{\omega}_\infty - \hat{\omega}_\ell} \quad \forall \omega \in [\hat{\omega}_\ell, \hat{\omega}_{\ell+1}], \quad \ell = L - 1. \tag{10}
\]
that the number of local maxima in each subband is at most few units (usually only one), with a most probable situation characterized by this maximum occurring at one edge of the subinterval. This favorable situation is guaranteed by a careful selection of the control points as discussed next.

A. Choosing Control Points

The strategy for constructing the control points \( \hat{\omega}_r \) aims at reducing \( \Delta_\ell \) in those regions that are characterized by fast variations of \( \varphi(\omega) \) and conversely relaxing and enlarging \( \Delta_\ell \) where \( \varphi(\omega) \) is slowly varying or nearly constant. The main underlying assumption is that variations of the singular values are induced by variations in the transfer matrix elements, which in turn are induced by the location of each pole/residue term. Therefore, we start with the pole-based sample distribution discussed in [18], which for each pair of complex conjugate poles, \( p_n = \alpha_n \pm j\beta_n \) contributes the following candidate frequency samples:

\[
\omega_{n,r} = \beta_n + \alpha_n \tan \frac{r\pi}{2(K+1)}, \quad r = -R, \ldots, R.
\] (11)

The parameter \( R \) is used to control how many samples are due to each pole term in a partial fraction expansion of the model response. Only frequencies \( \omega_{n,r} \geq 0 \) are retained and assembled in a single set \( \Omega \), that is, then sorted in ascending order. A final scan is performed to remove samples that are closer than a predefined resolution \( \Delta\omega \), which is defined as in [18] as

\[
\Delta\omega = \frac{p_{\max}}{N\rho}
\] (12)

where \( \rho \gg 1 \) is another control parameter and \( p_{\max} = \max(\omega_{\max}, \max_n |p_n|) \).

The above-defined set \( \Omega \) was tested and proved to be inadequate for a pure sampling-based passivity check. In particular, the following conditions hold.

1) The parameters setting suggested in [18] results in a large set of subbands, improving the robustness of the procedure but reducing its overall speed. A more efficient check would call for small \( R \). This strategy would provide an adequate sample distribution in band, but the passivity metric variations for \( \omega > \omega_{\max} \) might not be sampled appropriately.

2) Variations induced by real poles are not adequately sampled by (11).

3) Poles that are very close to the imaginary axis lead to subbands with very small \( \Delta_\ell \), whose edges may be removed accidentally by (12), leading to inaccurate sampling.

The above issues 1 and 2 are here solved by introducing three separate control parameters \( R_v \) with \( v = \{c, r, h\} \) to be used, respectively, for in-band complex pole pairs, real poles, and poles with either real or imaginary part magnitude that is close to or larger than \( \omega_{\max} \). In particular, the following conditions hold.

1) \( R_c \) can be safely chosen very small, even \( R_c = 1 \).

2) \( R_r \) should be larger, with at least \( R_r \geq 2 \).

3) \( R_h \) should also be larger, with at least \( R_h \geq 3 \).

### Table II

| Mode | \( R_c \) | \( R_r \) | \( \nu \) | \( c \) | \( Q_{\max} \) | \( \kappa \) | \( d \) |
|------|------|------|------|------|------|------|------|
| soft | 10^4 | 1    | 5    | 50   | 500  | 3    | 0.5 |
| hard | \( \infty \) | 3    | 6    | 50   | 500  | 3    | 0.5 |
| final | \( \infty \) | 3    | 6    | 50   | 500  | 3    | 0.5 |

Three different settings are suggested, to emphasize speed (soft), accuracy (hard) and to perform model qualification (final).

Issue 3 is partially compensated by modifying \( \omega_n, r \), corresponding to highly resonant poles in (11) by setting

\[
\alpha_n \leftarrow c \cdot \alpha_n, \quad \text{if} \quad Q_n \approx \frac{|\beta_n|}{2|\alpha_n|} > Q_{\max}
\] (13)

where \( Q_{\max} \) is here set to 500 and \( c = 50 \). Finally, since the entire real line \( \omega \in \mathbb{R} \) needs to be sampled, we add \( \omega = \infty \) with the following \( \kappa = 1 \) logarithmically spaced samples extending \( d \) decades beyond the fitting model band:

\[
\alpha_k = \omega_{\max} \cdot 10^{d \pi v}, \quad v = 0, \ldots, \kappa
\] (14)

where \( d \) is an additional control parameter. Typical values are \( d \geq 0.5 \) and \( \kappa \geq 2 \).

In summary, frequency warping and renormalization are here achieved by a pole-based control point distribution, whose behavior can be controlled by the eight parameters summarized in Table II. The above guidelines and parameter settings have been heuristically proved to be quasi-optimal over a large number of test cases (see Section VI). It should be considered that when inserted in a passivity enforcement loop, the number of subbands can be adaptively increased as the passivity violations are iteratively removed and become more difficult to identify, simply by increasing the values of \( R_v \), as discussed and illustrated in Section V-B through an application example.

### V. Step 2: A Modified Naive multiscale Search Optimization

The second step of the proposed passivity verification algorithm applies a hierarchical adaptive sampling process to the rescaled passivity metric \( \theta(\zeta) \) within each normalized subband \( \zeta \in \mathcal{X}_v = \{\ell, \ell + 1\} \), with the main objective of identifying all local maxima that exceed the passivity threshold \( \gamma = 1 \).

Among the several adaptive algorithms offered by the literature, our starting point is the so-called naive multiscale search optimization (NMSO) algorithm [32], which is based on a tree-search divide-and-conquer strategy. Although based on adaptive tree refinement, the NMSO algorithm may suffer from the presence of many local maxima since its main objective is to find the unique global maximum in its search domain. This is the main reason why we performed the subband splitting and frequency warping in Section IV. Each subband \( \mathcal{X}_v = \{\zeta_\ell, \zeta_{\ell+1}\} \) is constructed to include a very limited number of local maxima, usually only one. Therefore, we propose a repeated and independent application of a suitably constructed NMSO scheme to each subband \( \mathcal{X}_v \) independently so that the identification of all local maxima is greatly simplified.
The discussion below focuses on $X = X_0 = [0, 1]$ without loss of generality.

The proposed algorithm builds a tree $T$ whose nodes are iteratively refined at multiple scales $h \geq 0$, in order to partition the search space $X$ at any level $h$ into a set of $M^h$ cells

$$X_{h,i} = [iM^{-h}, (i + 1)M^{-h}], \quad i = 0, \ldots, M^h - 1. \quad (15)$$

Each cell has a center point $\zeta_{h,i} = (i + 1/2)M^{-h}$, denoted as leaf or node in the following. The partition factor $M \geq 2$ defines the number of children to be expanded from each leaf of the tree when increasing refinement level. The main objective is to find all local maxima of $\theta(\zeta)$ over the search space $X$, by evaluating $\theta_{h,i} = \theta(\zeta_{h,i})$ at a minimal number of tree leaves, which are determined adaptively through hierarchical refinement of the tree. Fig. 3 shows a graphical illustration, where the entire set $L^h$ of leaves at a given maximum refinement level $h$ is represented by dots, whereas the set $L^h$ of evaluated leaves is depicted by filled dots.

### A. Adaptive Sampling Rules

The various assumptions and features of the proposed scheme are illustrated in the following, whereas a pseudocode version is available in Algorithm 1.

1) **General Structure:** The dynamic evolution of the tree is performed through successive iterations with index $\mu$. At any iteration, the set of evaluated nodes $E_\mu$ is characterized by three levels,

1) $h_{\min}$: The minimum level of any node in $E_\mu$.
2) $h_{\max}$: The maximum level of any node in $E_\mu$.
3) $h$: The level of the node being processed, also denoted as current node; $h$ is henceforth denoted as current level.

The elements of $E_\mu$ are split into two subsets

$$E_\mu = C_\mu \cup B_\mu \quad (16)$$

where $C_\mu$ collects candidates for refinement at the next iteration and $B_\mu$ forms the so-called basket, which includes those nodes that do not need to be refined since a stop condition is verified (see next).

Algorithm 1 Find $\theta_{\max} = \arg \max \theta(\zeta)$ for $\zeta \in [0, 1]$

**Require:** $h_0, M, \delta_\zeta, \delta\theta, \delta\eta, \epsilon, \eta, n^\epsilon, \delta n^\epsilon$

1. Initialize $\mu = 0$, $h \leftarrow h_0$, $B_0 = \emptyset$
2. Initialize $C_0 = \{\zeta_{h,i}, i = 0, \ldots, M^h - 1\}$, $K = M^h$
3. Initialize $\theta_{\max} = -\infty$
4. while $C_\mu \neq \emptyset$ do
5. Choose current point $\zeta_{h,i}$, via (20)
6. Expand current point via (19)
7. $K \leftarrow K + M - 1$
8. Update $\theta_{\max} \leftarrow \max\{\theta_{\max}, \zeta_{h+1,i}, j \in J_{h,i}\}$
9. if $K > n^\epsilon$ then
9.1. if $U_1$ AND (12 OR $U_2$) then
9.2. update budget via (34)
10. else
11. go to 27
15. end if
16. end if
17. if $S_1 \text{ OR } S_2 \text{ OR } S_3$ then
18. insert new points in basket via (29)
19. $h \leftarrow h_{\min}$
20. reset $\epsilon$
21. else
22. flag new points for refinement via (27)
23. $h \leftarrow h + 1$
24. end if
25. $\mu \leftarrow \mu + 1$
26. end while
27. return samples $E_\mu = C_\mu \cup B_\mu$ and maximum $\theta_{\max}$

2) **Constraints:** We aim at minimizing the number of function evaluations. Therefore, we allow a total initial budget $n^\epsilon$ of function evaluations. Another constraint on which we build our implementation is to enforce $M$ to be odd.

3) **Initialization:** The algorithm is initialized for $\mu = 0$ by choosing an initial refinement level $h_0 \geq 0$ and evaluating the passivity metric $\theta$ (our target function) at the corresponding nodes $\{\zeta_{h,i}, i = 0, \ldots, M^h - 1\}$. Such samples are inserted into $E_0 = C_0$, with the initial basket $B_0 = \emptyset$.

4) **Expansion (Refinement):** The expansion process splits a given cell $X_{h,i}$ into its $M$ children at level $h + 1$, denoted as

$$X_{h+1,i}, \quad j \in J_{h,i} = \{Mi, \ldots, M(i + 1) - 1\}. \quad (17)$$

The set $J_{h,i}$ collects the indices of the nodes within cone of influence of node $\zeta_{h,i}$ under refinement by one level. We remark that using an odd partition factor, $M$ guarantees that the midpoint of the central children cell coincides with that of the expanded leaf

$$\zeta_{h+1,M+1} = \zeta_{h,i} \quad (18)$$

which does not need to be reevaluated but simply relabeled (see Fig. 3). Note also that elimination of node $\zeta_{h,i}$ upon its refinement may trigger an update of $h_{\min}$ as a consequence of this relabeling. Refinement of one leaf $\zeta_{h,i}$ can thus be expressed as

$$E_{\mu+1} = (E_\mu - \{\zeta_{h,i}\}) \cup \{\zeta_{h+1,i} : j \in J_{h,i}\}. \quad (19)$$
5) Choosing Leaves for Refinement: One key operation is the selection of best node for refinement among a set of candidates in $C_h$. This is simply the leaf corresponding to the largest value of the passivity metric

$$
\zeta_{h,i} = \arg \max \{\theta(\zeta_{h,i}) : \zeta_{h,i} \in C_h\}
$$

selected among all leaves at the current level $h$. This choice is motivated by the main objective of finding local maxima.

6) Stopping Conditions, Classification, and Restarts: The proposed implementation checks the nodes generated upon refinement (19) to determine whether they need to be further refined. This is achieved through various stopping conditions.

1) Condition $S_1$: A target resolution has been reached. Considering current level $h$ being refined into nodes at levels $h + 1$, this condition is expressed as

$$
M^{-h-1} < \delta\zeta
$$

where $\delta\zeta$ is a control parameter.

2) Condition $S_2$: The largest variation among adjacent refined nodes is within a prescribed tolerance. Denoting

$$
\Delta_{h,i} = \max \{|\theta_{h+1,j+1} - \theta_{h+1,j}| : j = Mi, ..., M(i+1) - 2\}
$$

condition $S_2$ can be expressed as

$$
\Delta_{h,i} < \delta\theta
$$

where $\delta\theta$ is a control parameter.

3) Condition $S_3$: The largest variation among adjacent refined nodes is less than their distance from the passivity threshold

$$
\Delta_{h,i} < |\hat{\theta}_{h,i} - \gamma|
$$

where

$$
\hat{\theta}_{h,i} = \max \{\theta(\zeta_{h+1,j}) : j \in J_{h,i}\}.
$$

This latter condition is only considered when a minimum resolution has been achieved

$$
M^{-h-1} < \delta\eta
$$

where $\delta\eta$ is a control parameter. After a refinement process, if all conditions $S_1$, $S_2$, and $S_3$ are false, all nodes $\zeta_{h+1,j}$ obtained from refinement are flagged as potentially critical and will be candidates for refinement at the next iteration so that

$$
C_{h+1} = (C_h - \{\zeta_{h,i}\}) \cup \{\zeta_{h+1,j} : j \in J_{h,i}\}.
$$

At the same time, current level is increased as

$$
h \leftarrow h + 1.
$$

Otherwise, the new nodes $\zeta_{h+1,j}$ from refinement are flagged as noncritical and are inserted in the basket

$$
B_{h+1} = (B_h - \{\zeta_{h,i}\}) \cup \{\zeta_{h+1,j} : j \in J_{h,i}\}.
$$

These nodes will not be further refined. The latter condition will trigger a restart by resetting current level to

$$
h \leftarrow h_{\min}.
$$

This condition enhances the exploration capabilities of the search.

7) Budget Updates: In principle, when the budget of function evaluations is reached upon a refinement step, the algorithm should stop. However, it may be possible that some critical regions to be explored or refined are still present. Therefore, a budget update (increase) is triggered by the three conditions given in the following.

1) Condition $U_1$: All nodes from the last refinement are detected as passive

$$
\hat{\theta}_{h,i} < \gamma.
$$

2) Condition $U_2$: At least one node from the last refinement is closer to the passivity threshold than a prescribed relative threshold $\varepsilon$

$$
\frac{\gamma - \hat{\theta}_{h,i}}{\hat{\theta}_{h,i}} < \varepsilon.
$$

3) Condition $U_3$: The largest variation among adjacent refined nodes exceeds their smallest distance from the passivity threshold

$$
|\gamma - \hat{\theta}_{h,i}| < \Delta_{h,i}.
$$

As for condition $S_3$, this latter condition is only checked when a minimum resolution has been according to (26).

When $U_1$ holds and one among $U_2$ and $U_3$ is verified, budget update is achieved by setting

$$
n^\varepsilon \leftarrow n^\varepsilon + \delta n^\varepsilon
$$

where $\delta n^\varepsilon$ is a control parameter that can even be iteration-dependent. In order to preserve the effectiveness of condition $U_2$ when the tree level $h$ increases, any budget update (34) triggers a reduction of $\varepsilon$ by a factor $\varphi < 1$ as $\varepsilon \leftarrow \varphi \varepsilon$. The original value of $\varepsilon$ is restored upon restarts.

8) Basket Reuse: The above-listed strategies prevent reusing and further refining the leaves in the basket $B_h$, which is appropriate when the execution speed is to be privileged with respect to accuracy. Nevertheless, the procedure can be easily modified to include the elements of $B_h$ in the set of candidates for future expansions $C_{h+1}$, as in the original NMSO scheme [32]. This basket reuse strategy should be used for more aggressive sampling, such as required for model qualification at the end of passivity enforcement loops (see next).

B. Implementation: Choosing Algorithm Parameters

The proposed two-step adaptive sampling algorithm is controlled by two sets of parameters for step-1 and step-2. These parameters are collected in Tables II and III, respectively. Both tables report our suggestion for three possible execution modes. Settings labeled as soft are adequate for passivity checks that are executed at the beginning of a passivity enforcement loop, where only the largest passivity violations are of interest and an aggressive accuracy is not necessary. Conversely, settings labeled as hard are adequate when results are expected to be very accurate, and passivity violations are very small, such as at the last iteration(s) of passivity enforcement loops. Settings labeled as final are adequate for final model qualification.
TABLE III

CONTROL PARAMETERS OF PROPOSED STEP-2: ADAPTIVE REFINEMENT

| Step 2: Modified NSMO | Mode | $\delta \xi$, $\delta \theta$ | $\delta \eta$ | $\varepsilon$ | $\phi$ | $n^*$ |
|-----------------------|------|------------------|---------------|-----------|------|-------|
| soft                  | 5    | $10^{-6}$        | $10^{-3}$     | 0.1       | 7, 10, 20, \ldots, 100 |
| hard                  | 5    | $10^{-6}$        | $10^{-3}$     | 0.1       | 10, 20, \ldots, 100  |
| final                 | 3    | $10^{-6}$        | $10^{-3}$     | 0.1       | 50, 100, \ldots, 250  |

Three different settings are suggested, to emphasize speed (soft), accuracy (hard) and to perform model qualification (final).

The differences between the two soft and final settings in algorithm performance are illustrated through an example. Fig. 4 compares the results obtained by both settings on a representative test case, precisely #444 with reference to the complete benchmark database analyzed in Section VI-A. The structure has $P = 92$ ports, with a model having $\bar{n} = 16$ pole-residue terms and a corresponding state-space matrix size $N = 1472$.

The proposed adaptive sampling check in soft mode resulted in $L = 30$ control points and $K = 2255$ evaluated frequency samples, 28 of which denoting passivity violations with $\phi(\omega) > \gamma$. Algorithm in hard mode requested $K = 4245$ samples with $L = 55$ and 47 violation points, whereas final mode led to $K = 9827$ samples evaluation, with $L = 55$ and 271 violation points. It should be noted that the above detected violation points can be at the edges of each subband $\lambda_i$, in which case they should be dropped since not corresponding to actual local maxima over the entire frequency axis. After suitable postprocessing, the actual number of detected local maxima largest than the passivity threshold resulted 7, 14, and 21 after soft, hard, and final mode checks, respectively. These results are consistent with the asymptotic outcome of the algorithm used in soft mode, imposing $n^* = \infty$. Fig. 4 and its enlarged view of Fig. 5 compare the results of soft and final mode, showing that the latter provides a more accurate detection of even very small violations over narrow frequency bands.

**VI. Numerical Results**

We document the performance of the proposed algorithm on a large set of test cases. Section VI-A presents the results of a systematic testing campaign on a database of interconnect macromodels, while Sections VI-B and VI-C concentrate on two representative large-scale application examples. All documented numerical results have been obtained on a Workstation based on Core i9-7900X CPU running at 3.3 GHz with 64-GB RAM.

**A. Consistency and Performance**

The proposed passivity check algorithm was applied to a large set of benchmarks, including 447 models of different sizes and complexities (details in the bottom of Fig. 7). A total of 243 independent models form a database constructed by applying a standard VF scheme as available in the commercial software IdEM [39]. Only 37 of these models were initially passive, whereas the remaining 206 (nonpassive) models showed in some cases very significant (and not challenging to be detected, even with simplistic algorithms) passivity violations. For this reason, additional examples were obtained by applying few passivity enforcement iterations on 204 models from the first set. Since the passivity metric changes completely through passivity enforcement iterations, these latter models can be considered as independent. Moreover, since passivity enforcement iteratively removes passivity
The proposed algorithm agreed in the 99.8% of the cases with the Hamiltonian check. The proposed algorithm gave only one different result with respect to the Hamiltonian check, providing a false negative result. It turns out that this model was actually not passive (with a very tight passivity violation region, and $|\sigma_1 - 1| \approx 1 \times 10^{-10}$), but imaginary Hamiltonian eigenvalues were not detected. Fig. 6 shows a graphical illustration. We conclude from this systematic testing campaign that the proposed scheme may be able to detect very small passivity violations also in the rare and extreme cases when the Hamiltonian algebraic test fails due to ill-conditioning.

The time performances of the two passivity check strategies are presented at the top of Fig. 7. In all cases, a MATLAB-based single-thread prototypal code was used, using a full Hamiltonian eigensolver. We see that the Hamiltonian check remains the method of choice for small-scale models (the number of ports and poles is reported at the bottom for comparison). For small-scale models, both the Hamiltonian and the proposed check produce their results in a fraction of a second. When it comes to large-scale models, the situation is drastically different. The proposed scheme has a runtime that for all investigated cases is below 10 s, which can be as much as $50 \times$ faster than the Hamiltonian check.

The timing results of Fig. 7 show that the speedup of the proposed adaptive sampling scheme with respect to the Hamiltonian tests is problem-dependent, even for models characterized by the same level of complexity. This is in fact expected based on the following considerations.

1) Some of the models required the extended Hamiltonian pencil (6), which is slower but numerically more accurate and robust than the regular Hamiltonian matrix (5) when $|D|_2 \approx 1$.

2) In presence of singular value maxima that are very close to the threshold $\gamma$, our proposed NMSO scheme keeps on refining samples until the precision on the local maxima is sufficient to discriminate possible passivity violations, while the Hamiltonian test detects the only threshold crossings.

Focusing on the latter scenario 2, it may be the case that local maxima (if any) fall below the passivity threshold. In this situation, adaptive sampling will not detect any violation, and there will be no imaginary Hamiltonian eigenvalues. Conversely, when two singular value crossings are very close so that a local maximum occurs just above the threshold, as shown in Fig. 6, the corresponding imaginary Hamiltonian eigenvalues are almost collapsed, the Hamiltonian matrix is defective, and the associated eigenproblem is ill-conditioned. In both situations, the proposed adaptive sampling scheme needs to continue refining samples until precision in the local maximum estimation is sufficient. This may lead to some small computing overhead. The execution speed of the Hamiltonian test will not be affected, but it may happen that the corresponding results are not reliable, as shown in Fig. 6.

Finally, we show through a toy example that the requirement of local refinements is unavoidable for all subbands. This may cause a small overhead due to the presence of computed samples that are not functional to detect passivity violations, in those cases where there are no sharp variations or local maxima. Fig. 8 shows the maximum singular value of a system with two sets of closely spaced poles $p_n^1 = \alpha_n^1 \pm j\omega_n^1$ and

\[
\begin{align*}
|\sigma_1 - 1| & \approx 1 \times 10^{-10}, \\
D & \approx 1, \\
|D|_2 & \approx 1.
\end{align*}
\]
Fig. 7. Time performances and models complexity for all the investigated 447 test cases.

\[ p_n^2 = \omega_n^2 \pm j\delta \omega, \]  
where \( \omega_n = \omega_0 + n \cdot \delta \omega \) and \( \delta \omega \ll \omega_0 \). The local variations around these poles may have local maxima or not, depending on the residues. A reduced number of samples based only on control points or on a coarse sampling (dots) is not sufficient to detect the presence of sharp local maxima: such an implementation would not distinguish between the two trajectories centered at \( \omega_0 \) and \( \omega_0 + \delta \omega \). This is the main reason why the proposed scheme places additional samples also where a first sight would deem these samples unnecessary (see Fig. 1 beyond \( 10^{10} \) GHz). Local refinement needs to guarantee that sharp peaks as in Fig. 8 (right) are not missed.

B. Large-Scale Multiport Shielding Enclosure

We compare here the performance of the proposed passivity characterization to various implementations of Hamiltonian-based tests available in the state-of-the-art commercial package IdEM [39]. The structure that is selected for this comparison is a large-scale model of a shielding enclosure, as shown in Fig. 9. The enclosure is part of an energy-selective shield, obtained by loading the \( P = 20 \times 20 = 400 \) ports located throughout the opening with nonlinear elements (back-to-back diode pairs). For an overview of energy-selective shielding, see [22], in particular [23]–[25].

The unloaded structure was initially characterized through a method-of-moments (MoM) frequency-domain code [40], obtaining a set of \( P \times P \) scattering responses, each with a total of 500 frequency samples over the frequency band [0.25, 990] MHz. These samples were subjected to VF to obtain a rational model (\( n = 26 \) poles), leading to a state-space realization of size \( N = 10400 \). A few selected port responses of the model are compared to field solver data in Fig. 10, demonstrating good model accuracy.
The model was processed by four different passivity characterization schemes given in the following.

1) **Full Hamiltonian Check**: As implemented in [39], extracting all imaginary eigenvalues of the full Hamiltonian matrix (5).

2) **Sparse Hamiltonian Check**: As implemented in [39], based on the Hamiltonian eigensolver exploiting the iterative multishift Krylov subspace iterations mentioned in [17].

3) **Adaptive Hamiltonian Check**: As implemented in [39], based on the hybrid sampling-sparse Hamiltonian eigensolver documented in [18].

4) Proposed two-stage adaptive sampling check.

The Hamiltonian eigensolvers were available in two execution modes using $T = 1$ and $T = 8$ parallel threads, respectively. The proposed approach was implemented as a serial code, executed in a MATLAB [41] environment both enabling and disabling multithread capabilities (at low level for SVD computations). The results are collected in Table V, which confirms a major speedup of the proposed approach with respect to Hamiltonian characterizations. All methods provided consistent passivity characterization results in terms of a number of violation bands. Fig. 11 shows the singular value trajectories computed by the proposed adaptive sampling scheme, highlighting the local maxima exceeding the passivity threshold.

### C. Via Array

We analyze here a via array in an eight-layer printed circuit board (PCB) structure. The array consists of $20 \times 20$ vias with a 4:1 signal to ground ratio, resulting in total of $P = 640$ electrical ports (1–320 in the top layer and 321–640 in the bottom layer). A fast frequency-domain solver [42] was applied to extract the $P \times P$ scattering responses up to 30 GHz, defining circular ports between via pad and antipad at top and bottom layers. These 150 raw samples were in turn used to generate a large-scale macromodel through VF using the IdEM software [39], resulting in $\bar{n} = 40$ pole-residue terms and a total number of states $N = 25,600$.

The model was processed by the proposed passivity characterization algorithm, as well as to all Hamiltonian checks available in IdEM. The timing results are reported in Table VI, whereas Fig. 12 shows a comparison between the selected model responses and the corresponding field solver samples. Fig. 13 shows the largest singular value samples returned by the proposed adaptive sampling scheme, highlighting all local maxima exceeding the passivity threshold. Execution in both hard and final mode identified all 11 local maxima, whereas the fast soft mode found ten maxima. Also, for this example, the execution time provided by the proposed passivity check algorithm is drastically reduced with respect to Hamiltonian checks, at least for soft and hard execution modes, with no loss of accuracy in the characterization of passivity violations. Full Hamiltonian checks could not be performed due to excessive storage requirements.
D. Discussion

The above numerical results show that the proposed algorithm appears to be the only viable and scalable solution for passivity characterization of large-sized models. The following considerations apply.

1) The number of subbands produced by the step-1 adaptive frequency warping grows with the number \( n \) of model poles and can be very large. This is not a limitation since the independent processing of those subbands lends itself to a naive data-based parallelization, which is thus expected to provide even further speedup with respect to the already highly efficient implementation that is here documented. Code parallelization is left to future investigations.

2) It may be the case that multiple passivity violation maxima in the same subband may be missed if the stopping thresholds are excessively loose or the allocated budget of evaluations is too small.

3) The speed of the proposed scheme is affected by the type of passivity violations. In case of large violations, these are detected early and further refinements are not necessary, as in the case reported in Fig. 11. In fact, since the check is designed to set up constraints for a subsequent passivity enforcement, once a significant number of constraints are available, it is not necessary to further refine the search. Instead, when the violations (if any) are smaller and the passivity metric shows many local maxima close to the threshold, a larger number of refinement iterations are potentially needed, as shown in Fig. 13. Therefore, the performance of the proposed adaptive scheme can only be assessed on a statistical basis since problem-dependent.

4) In case a very accurate test is needed, asymptotic convergence to all local maxima can be guaranteed by releasing budget constraints and enabling also basket reuse for refinement, building on the provable asymptotic consistency of the NMSO algorithm class [32]. We remark that this strategy was never required in our extensive tests since the proposed implementation never identified nonpassive models as passive.

VII. Conclusion

This work introduced a two-stage hierarchical sampling-based passivity characterization algorithm for large-scale lumped macromodels. The proposed scheme was tested on about 450 test cases of increasing complexity and reaching \( P = 640 \) ports and more than \( 25 \times 10^5 \) states. The results show that the proposed approach outperforms Hamiltonian tests for medium- and large-scale systems, with speedup in runtime exceeding \( 200 \times \) in some cases.

Due to the multistage hierarchical implementation, our proposed algorithm was always in agreement with Hamiltonian checks. The only exception was a single case that was (correctly) detected as nonpassive by the proposed method, although the Hamiltonian check could not identify any passivity violation. Therefore, we conclude that the proposed scheme is at least as reliable as state-of-the-art approaches, with much more favorable scalability with model size.

Significant research work is still needed for efficient extraction and handling of large-scale macromodels. Fully coupled models with many ports and large dynamical orders are in fact not effective when synthesized as equivalent circuits and used in system-level simulations, due to the very large number of model parameters. \textit{Ad hoc} model representations based on sparsification or compression appear to provide a better solution for speeding up numerical simulations, possibly combined with specialized solvers that are aware of the model structure. We are actively investigating in this direction, and the results will be documented in a forthcoming report.

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Marco De Stefano (Graduate Student Member, IEEE) received the M.Sc. degree in mechatronic engineering from Politecnico di Torino, Turin, Italy, in 2018, where he is currently pursuing the Ph.D. degree in electrical, electronics and communication engineering.

His research interests include model-order reduction, with emphasis on parameterized macromodeling, and fast simulation methods for signal and power integrity.

Mr. De Stefano was a co-recipient of the 2018 Best Paper Award of the IEEE International Symposium on Electromagnetic Compatibility and the 2019 Best Student Paper of the IEEE Workshop on Signal and Power Integrity.

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Stefano Grivet-Talocia (Fellow, IEEE) received the Laurea and Ph.D. degrees in electronic engineering from the Politecnico di Torino, Turin, Italy, in 1994 and 1998, respectively.

From 1994 to 1996, he was with the NASA/Goddard Space Flight Center, Greenbelt, MD, USA. He is currently a Full Professor of electrical engineering with the Politecnico di Torino. He co-founded the academic spinoff company IdemWorks, Turin, in 2007, where he was the President until its acquisition by CST in 2016. He has authored over 180 journal and conference papers. His current research interests include passive macromodeling of lumped and distributed interconnect structures, model-order reduction, modeling and simulation of fields, circuits, and their interaction, wavelets, time–frequency transforms, and their applications.

Dr. Grivet-Talocia was a co-recipient of the 2007 Best Paper Award of the IEEE TRANSACTIONS ON ADVANCED PACKAGING. He received the IBM Shared University Research Award in 2007, 2008, and 2009. He was the General Chair of the 20th and 21st IEEE Workshops on Signal and Power Integrity (SPI2016 and SPI2017). He was an Associate Editor of the IEEE TRANSACTIONS ON ELECTROMAGNETIC COMPATIBILITY from 1999 to 2001. He is also serving as an Associate Editor for the IEEE TRANSACTIONS ON COMPONENTS, PACKAGING AND MANUFACTURING TECHNOLOGY.

Torben Wendt (Graduate Student Member, IEEE) received the B.S. and M.S. degrees in electrical engineering from the Technical University Hamburg (TUHH), Hamburg, Germany, in 2014 and 2017, respectively, where he is currently pursuing the Ph.D. degree with the Institute of Electromagnetic Theory.

He was a Visiting Researcher with the Politecnico di Torino, Turin, Italy, from 2018 to 2019. His current research interests include the modeling and simulation of nonlinear field circuit coupling in electromagnetic compatibility (EMC) applications.

Cheng Yang (Member, IEEE) received the B.S. degree in electronic science and technology from Wuhan University (WHU), Wuhan, China, in 2009, and the M.S. and Ph.D. degrees in electromagnetic field and microwave technology from the National University of Defense Technology (NUDT), Changsha, China, in 2012 and 2016, respectively.

Since 2019, he has been a Senior Engineer with the Institute of Electromagnetic Theory, Technische Universität Hamburg (TUHH), Hamburg, Germany. From 2013 to 2015, he was funded by the Chinese Scholarship Council (CSC) as a joint-Ph.D. student at TUHH. From 2017 to 2019, he was a Faculty Member with the State Key Laboratory of Millimeter Wave, Southeast University (SEU), Nanjing, China. His current research interests include computational electromagnetics, microwave measurement techniques, and design and analysis of nonlinearly loaded electromagnetic structures.

Christian Schuster (Senior Member, IEEE) received the Diploma degree in physics from the University of Konstanz, Konstanz, Germany, in 1996, and the Ph.D. degree in electrical engineering from the Swiss Federal Institute of Technology (ETH), Zürich, Switzerland, in 2000.

He was with the IBM T. J. Watson Research Center, Yorktown Heights, NY, USA, where he was involved in high-speed optoelectronic package and backplane interconnect modeling and signal integrity design for new server generations. Since 2006, he has been a Full Professor and the Head of the Institute of Electromagnetic Theory, Hamburg University of Technology (TUHH), Hamburg, Germany. He is currently an Adjunct Associate Professor with the School of Electrical and Computer Engineering, Georgia Institute of Technology, Atlanta, GA, USA. His current research interests include signal and power integrity of digital systems, multiport measurement and calibration techniques, and development of electromagnetic simulation methods for communication electronics.

Dr. Schuster is a member of the Board of Directors of the IEEE Electromagnetic Compatibility Society. He is also a member of the German Physical Society (DPG) and several technical program committees of international conferences on signal and power integrity, and electromagnetic compatibility. He received the IEEE TRANSACTIONS ON ELECTROMAGNETIC COMPATIBILITY Best Paper Awards in 2001 and 2015; the IEEE TRANSACTIONS ON COMPONENTS, PACKAGING, AND MANUFACTURING TECHNOLOGY Best Paper Awards in 2012 and 2016; the IEC DesignCon Paper Awards in 2005, 2006, 2010, 2017, and 2018; three IBM Research Division Awards from 2003 to 2005; the IBM Faculty Awards in 2009 and 2010; and the Sustained Service to the EMC Society Award in 2019. He served as the Chair for the German IEEE EMC Chapter from 2016 to 2019. He is also an Associate Editor of the IEEE TRANSACTIONS ON ELECTROMAGNETIC COMPATIBILITY. He served as a Distinguished Lecturer for the IEEE EMC Society from 2012 to 2013.