Surrogate Models for Linear Response

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Linear response theory is a well-established method in physics and chemistry for exploring excitations of many-body systems. In particular, the quasiparticle random-phase approximation (QRPA) provides a powerful microscopic framework by building excitations on top of the mean-field vacuum; however, its high computational cost limits model calibration and uncertainty quantification studies. Here, we present two complementary QRPA surrogate models and apply them to study response functions of finite nuclei. One is a reduced-order model that exploits the underlying QRPA structure, while the other utilizes the recently developed parametric matrix model algorithm to construct a map between the system's Hamiltonian and observables. Our benchmark applications, the calculation of the electric dipole polarizability of ¹⁸⁰Yb and the β -decay half-life of ⁸⁰Ni, show that both emulators can achieve 0.1%–1% accuracy while offering a six to seven orders of magnitude speedup compared to state-of-the-art QRPA solvers. These results demonstrate that the developed QRPA emulators are well-positioned to enable Bayesian calibration and large-scale studies of computationally expensive physics models describing the properties of many-body systems.

I. INTRODUCTION

Linear response theory provides a general framework for describing the response of a system to a weak external perturbing field [1]. It is fundamentally tied to the fluctuation-dissipation theorem by relating the groundstate thermal fluctuations to a dissipative external field. Because of this universality, linear response theory finds applications across a wide range of disciplines of natural sciences. In condensed matter physics, it underlies the study of dielectric functions and optical response of the system [2, 3]; in nuclear physics, it provides a microscopic description of collective excitations governing nuclear reactions and decay processes [4, 5]. In quantum chemistry, linear response theory forms the foundation for time-dependent methods that describe excited-state and ground-state correlations in molecular systems [6, 7], while in molecular biology, it has been used to analyze cellular responses to external perturbations [8]. Beyond these domains, it also applies to classical statistical mechanical systems [9, 10], non-equilibrium fluctuations in complex quantum systems [11], and even to modeling fluctuations in financial markets [12].

A specific formulation of linear response theory was developed by Bohm and Pines [13–15] to describe electron-electron interactions within a degenerate electron gas embedded in a uniform positive background. In this system, the total contribution from background and interactions between electrons and positive ions cancels out, leaving the electron-electron interaction as the dominant

effect. The collective motion of the electrons gives rise to plasma oscillations – quantized as plasmons. Although not explicitly named in the original work, this framework became known as the *random-phase approximation* (RPA), referring to the partial cancellation of out-of-phase density-fluctuation terms in the Fourier expansion.

Within many-body perturbation theory, the same approximation corresponds to a resummation of ring diagrams [16]. Today, the RPA is widely used in quantum chemistry and material science to study electronic excitations, and evaluate ground-state correlation energies [17–19].

The RPA equations can be derived in many ways. Within the Green's function approach [20, 21] the starting point is the two-particle Green's function, which is treated by the many-body perturbation theory. To derive the RPA equation, one assumes that the one-particle propagators are diagonal and single-particle levels are solutions of Hartree-Fock (HF) equations, and performs a re-summation of ring diagrams. Starting from the timedependent Hartree-Fock (TDHF) equation, by assuming a harmonic time-dependence of the induced density, and expanding the mean-field Hamiltonian up to linear order, one can derive the linear response equation, which has the same form as the RPA [22]. This approach can be coupled with density functional theory (DFT) in the form of time-dependent DFT (TDDFT) [23]. Lastly, one can obtain RPA equations for density-independent interactions by assuming a quasi-boson approximation [24, 25]. Formally, the RPA is related to other many-body methods, being equivalent to the coupled-cluster doubles theory [26].

The RPA was adapted to the nuclear many-body problem in Ref. [27], providing microscopic description of nuclear excitations and collective modes [4, 5]. In particular, an important extension of RPA is due to the presence of nucleonic pairing. First, the description of the

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nuclear mean field has to be extended by introducing the concept of quasiparticles, representing a linear combination of particle and hole states [28]. Subsequently, the RPA was formulated in the basis of two-quasiparticle excitations, yielding quasiparticle RPA (QRPA) [29]. Nuclear response functions and the corresponding excited states are central to a wide range of applications, from nuclear structure theory [30, 31] and astrophysics [32– 34, to the investigation of fundamental symmetries [35]. In particular, determining excitation energies and wave functions is essential for computing decay rates and transition probabilities, such as electric and magnetic transitions [36, 37], as well as charge-exchange processes such as β decay [38, 39]. In nuclear astrophysics, QRPA is used to estimate decay rates important for the synthesis of elements heavier than iron within the r-process nucleosynthesis [40].

General methods for QRPA matrix diagonalization are discussed in Refs. [24, 41]. A more computationally efficient approach, suited for deformed nuclei, is the finite-amplitude method (FAM) [42]. Although much development has been achieved in both approaches in recent years, QRPA calculations become increasingly more expensive as the number of possible two-quasiparticle (2qp) excitations grows, especially when symmetries of the nuclear ground state are internally broken.

In general, the QRPA equations can be derived within nuclear density functional theory (DFT), as it provides a robust foundation for practical implementations [43]. Nuclear energy density functionals (EDFs) — basic ingredients of DFT — can be represented by around a dozen parameters optimized to selected experimental data [44, 45]. Up to now, only a few EDFs have been calibrated by including excited-state properties obtained within the QRPA [46, 47]. Indeed, a comprehensive description of giant resonances and related observables, would require a dataset consisting of a wide range of nuclei and nuclear response functions. However, such considerations are currently computationally prohibitive – a full-fledged Bayesian model calibration [48] would require thousands to millions of model calculations, which remains beyond the reach of high-fidelity QRPA solvers.

Addressing the computational demands of modern models has motivated the development of emulators, or surrogate models. An emulator is a fast model that approximates the output of complex, computationally intensive models, referred to as Full Order Model (FOM) in the following. Using FOM evaluations, the emulators are trained to connect selected inputs (i.e., the underlying model parameters), with selected outputs (i.e. calculated observables). See Refs. [49–52] for some recent applications.

We highlight two general approaches for building emulators: data-driven and model-driven. Data-driven emulators usually attempt to connect inputs and outputs of a model directly, as exemplified by Gaussian Processes [53] and neural networks. Emulators following a model-driven approach usually perform this task by finding a set of re-

duced coordinates for the model's latent variables (i.e., wave functions, densities), and constructing their dynamical equations by exploiting the original operators [54].

Emulators based on the reduced basis method (RBM) [55, 56] are examples that have followed the model-driven approach by identifying the reduced coordinates as amplitudes of a small tailored basis, and constructing the equations by projecting the operators into a small linear subspace. Since model-driven emulators are informed by the underlying system's equations which often include physics-driven constraints — they require appreciably less training data to achieve similar predictive accuracy compared to their data-driven counterparts. However, this structural integration also makes them more cumbersome to construct [57, 58]. These challenges have motivated developments to augment model-driven emulators to construct not only the low-dimensional latent space characterized by a set of reduced coordinates, but also the governing equations for those coordinates. This insight has enabled the creation of many successful emulators and model discovery tools in science and engineering broadly [59–64], and within nuclear science and engineering applications in particular [65-72].

Based on these developments, in this work we create and benchmark two emulators for the QRPA calculations of electric dipole polarizability and β -decay half-life $T_{1/2}$. The first emulator follows a hybrid data- and modeldriven approach by identifying effective reduced coordinates to represent the strength function, and constructing the necessary equations by mirroring the structure of the QRPA eigensystem. From the emulated strength function, the system's observables can be calculated. The second emulator adopts the Parametric Matrix Model (PMM) [67] architecture in a purely data-driven setting: its trainable parametric matrices are optimized to predict the target observables directly, bypassing the intermediate construction of the strength function. We train both emulators using the FOM evaluations, and compare their performance in reproducing observables as a function of their complexity. Emulators developed in this work are publicly available [73]. We have also developed online Yinteractive visualization tools to further illustrate key concepts of the work [74].

The paper is organized as follows. In Sec. II we present the theoretical formalism behind QRPA, while the corresponding emulators are described in Sec. III. In Sec. IV, we present the emulation results for electric dipole polarizability and β -decay half-lives. Finally, Sec. V contains the summary of the paper. The glossary of acronyms and symbols used can be found in Appendix A.

II. QUASIPARTICLE RANDOM PHASE APPROXIMATION

We now present a summary of the QRPA formalism. In this work, the ground-state configuration of the nucleus is obtained by solving the Relativistic Hartree-Bogoliubov (RHB) equations in either a spherical or axially deformed geometry, with the mean-field generated self-consistently by the underlying EDF [75]. This self-consistent solution yields the static generalized density matrix \mathcal{R}^0 . Subjected to an external field $\mathcal{F}(t)$, the system evolves according to the time-dependent generalized density:

$$\mathcal{R}(t) = \mathcal{R}^0 + \delta \mathcal{R}(t), \tag{1}$$

with the harmonic ansatz in the time domain

$$\delta \mathcal{R}(t) = \delta \mathcal{R}(\omega) e^{-i\omega t} + \delta \mathcal{R}^{\dagger}(\omega) e^{i\omega t}. \tag{2}$$

The linearization of the time-dependent RHB equations in $\delta \mathcal{R}$ leads to the QRPA eigenvalue problem. In the two-quasiparticle basis, the dimension of the QRPA matrix is given by

$$N_d = 2n_{2qp},\tag{3}$$

where n_{2qp} denotes the number of two-quasiparticle excitations allowed by selection rules [76].

The linear response equation takes the form [30]:

$$\left[\mathcal{M}(\boldsymbol{\alpha}) - \omega \mathcal{N}\right] \delta \mathcal{R}_{\boldsymbol{\alpha}}(\omega) = -\mathcal{F}(\boldsymbol{\alpha}), \tag{4}$$

where the metric \mathcal{N} is

$$\mathcal{N} = \begin{pmatrix} \mathbb{1}_{n_{2qp}} & 0\\ 0 & -\mathbb{1}_{n_{2qp}} \end{pmatrix}_{N_d \times N_d}, \tag{5}$$

and ω is the (generally complex) excitation energy, while α denotes the set of k parameters defining the interaction — typically the coupling constants of the underlying EDF. The QRPA Hermitian matrix $\mathcal{M}(\alpha)$ has the block structure [24]

$$\mathcal{M}(\boldsymbol{\alpha}) = \begin{pmatrix} A(\boldsymbol{\alpha}) & B(\boldsymbol{\alpha}) \\ B^*(\boldsymbol{\alpha}) & A^*(\boldsymbol{\alpha}) \end{pmatrix}_{N_d \times N_d}, \tag{6}$$

where A, B are complex matrices constructed from the residual two-body interaction, satisfying $A^{\dagger} = A$, and $B^T = B$. The external field vector $\mathcal{F}(\boldsymbol{\alpha}) =$ $(F^{20}(\boldsymbol{\alpha}), F^{02}(\boldsymbol{\alpha}))$ represents the external field operator in the quasiparticle basis. The QRPA amplitudes are of the form $\delta \mathcal{R}_{\boldsymbol{\alpha}}(\omega) = (x_{\boldsymbol{\alpha}}(\omega), y_{\boldsymbol{\alpha}}(\omega))$.

Once the QRPA amplitudes are computed, the strength function – a physical observable representing the response of the system to the external field– can be written as:

$$S(\boldsymbol{\alpha}; \omega) = -\frac{1}{\pi} \text{Im} \mathcal{F}(\boldsymbol{\alpha})^{\dagger} \delta \mathcal{R}_{\boldsymbol{\alpha}}(\omega). \tag{7}$$

While the QRPA amplitudes may, in principle, be obtained by the matrix inversion, the dimensionality of the quasiparticle space $n_{\rm 2qp}$ is often prohibitively large, rendering the direct approach computationally infeasible. Accordingly, alternative methods have been developed. One such method reformulates the linear response equation, Eq. (4), as a variational problem within the finite

amplitude method (FAM) [37, 42]. Other techniques exploit symmetries and structure in the residual interaction to bypass explicit inversion in the large quasiparticle basis [77, 78].

The QRPA linear response equation may also be cast as a non-Hermitian eigenvalue problem, commonly referred to as the matrix QRPA:

$$\mathcal{NM}(\boldsymbol{\alpha})\delta\mathcal{R}_{\boldsymbol{\alpha};i} = E_{\boldsymbol{\alpha};i}\delta\mathcal{R}_{\boldsymbol{\alpha};i}, \quad i = 1,\dots,N_d$$
 (8)

where $\delta \mathcal{R}_{\alpha;i} = (x_{\alpha;i}, y_{\alpha;i})$ are the N_d eigenvectors corresponding to eigenvalues $E_{\alpha;i}$. In general, the spectrum of the QRPA matrix consists of complex-conjugate pairs $\{E_{\alpha;i}, -E_{\alpha;i}^*\}$. However, if the matrix $\mathcal{M}(\alpha)$ is positive definite (this condition is often satisfied when the QRPA is constructed on top of a self-consistent RHB minimum), then the eigenvalues of (8) are real. To compute transition matrix elements between the ground state and excited states, we define the reduced transition probability:

$$\mathcal{B}_{\boldsymbol{\alpha};i} = \left| \begin{pmatrix} F^{20}(\boldsymbol{\alpha}) \\ F^{02}(\boldsymbol{\alpha}) \end{pmatrix}^{\dagger} \begin{pmatrix} x_{\boldsymbol{\alpha};i} \\ y_{\boldsymbol{\alpha};i} \end{pmatrix} \right|^{2}. \tag{9}$$

In practice, the discrete strengths $\mathcal{B}_{\alpha;i}$ are folded with a Lorentzian of width η to produce a continuous strength function:

$$S(\boldsymbol{\alpha}; \omega' + i\eta) = \frac{\eta}{2\pi} \sum_{i=1}^{N_d/2} \frac{\mathcal{B}_{\boldsymbol{\alpha};i}}{(\omega' - E_{\boldsymbol{\alpha};i})^2 + (\eta/2)^2}, \quad (10)$$

where $\omega=\omega'$. Once the strength function $S(\omega)$ is obtained, other observables may be computed. In this work, we focus on the electric dipole polarizability α_D and the β -decay half-life $T_{1/2}$, both of which are strongly sensitive to nuclear excitations. Both observables are important. The dipole polarizability correlates closely with the neutron skin thickness, offering a stringent constraint on the density dependence of the symmetry energy. The β -decay half-life, on the other hand, is essential for a realistic description of charge-exchange nuclear excitations. We now summarize how these two observables are calculated from the QRPA strength functions.

A. Dipole Polarizability α_D

Physically, the electric dipole polarizability determines the response of the nuclear system to an external electric field. This quantity is of significant interest in both nuclear structure physics and astrophysics due to its strong correlation with the symmetry energy of nuclear matter, as well as the neutron skin thickness [31, 79, 80]. It can be expressed through the QRPA strength function as [81]:

$$\alpha_D(\boldsymbol{\alpha}) = \frac{8\pi}{9} e^2 \lim_{\eta \to 0^+} \int d\omega' \frac{S(\boldsymbol{\alpha}; \omega' + i\eta)}{\omega'}.$$
 (11)

In terms of all the N_d eigenpairs, involving the discrete strengths $\mathcal{B}_{\alpha;i}$ and eigenvalues $E_{\alpha;i}$, the above equation can be rewritten as:

$$\alpha_D(\alpha) = \frac{8\pi}{9} e^2 \sum_{i=1}^{N_d/2} \frac{\mathcal{B}_{\alpha;i}}{E_{\alpha;i}}.$$
 (12)

In practical calculations of α_D in deformed nuclei with $N_d > 10^5$, contour integration techniques [36] eliminate the need to work directly with the strength function.

B. β -decay half-life $T_{1/2}$

 β -decay is a nuclear decay governed by the weak interaction. Within QRPA, we can compute the charge-exchange linear-response in the allowed Gamow-Teller (GT) channel to obtain the GT strength function $S_{GT}(\alpha;\omega)$ [38, 39, 82]. Restricting to allowed transitions, the inverse half-life is given by:

$$\frac{1}{T_{1/2}(\boldsymbol{\alpha})} = \frac{g_A^2}{K} \lim_{\eta \to 0^+} \int_{-\lambda_{np}}^{\Delta_{nH}} d\omega \, f(\Delta_{np} - \omega, Z, A) S_{GT}(\boldsymbol{\alpha}; \omega),$$
(13)

for a nucleus with Z protons and A total nucleons, where $g_A = 1.2$ is the axial-vector coupling (for which we take the vacuum value) and K = 6147 s is the constant measured in superallowed β decays [38]. The integration in (13) is performed from $\lambda_{np} = \lambda_n - \lambda_p$, where $\lambda_{n(p)}$ is the neutron(proton) chemical potential, determined by solving the underlying RHB equations, up to the neutronhydrogen atom energy difference $\Delta_{nH} = 0.782 \text{ MeV}$, effectively enclosing the Q_{β} -value window. The phasespace factor $f(E_0, Z, A)$ takes into account the kinematics of outgoing leptons with the end-point energy $E_0 = \Delta_{np} - \omega$, where the neutron-proton mass difference is $\Delta_{np} = 1.293$ MeV. The function $f(E_0, Z, A)$ accounts for Coulomb distortion of electron wave functions through the Fermi function dependence [83]. Having calculated the full QRPA spectrum, Eq. (13) can be written

$$\frac{1}{T_{1/2}(\boldsymbol{\alpha})} = \frac{g_A^2}{K} \sum_{i=1}^{N_d/2} f(\Delta_{np} - E_{\boldsymbol{\alpha};i}, Z, A) \mathcal{B}_{\boldsymbol{\alpha};i}, \qquad (14)$$

where the phase-space function imposes the condition $E_{\alpha;i} < \Delta_{nH}$.

III. FULL AND REDUCED ORDER MODELS

In this section, we illustrate the methodology we developed to reduce the computational cost associated with QRPA calculations. The core principle behind various emulation strategies is that the parametric or time dynamics of high-dimensional objects can be approximated

through a much smaller set of coordinates or degrees of freedom. Here, we are interested in characterizing the outcomes of the QRPA calculations, namely the strength function $S(\omega)$, Eq. (10), and the two observables $\mathcal{O} \equiv [\alpha_D, T_{1/2}]$ as we vary the parameters α .

Figure 1 shows a diagrammatic comparison of how the FOM and the two emulators we developed connect the control variables α to the response variables $S(\omega)$ and \mathcal{O} . In the following, we use the "hat" symbol to represent approximations to quantities of interest. For example, $\hat{S}(\omega)$ represents a low-dimensional approximation to the original strength function $S(\omega)$. We now proceed to describe these approaches and how the two emulators are trained by using a set of FOM evaluations.

A. Full Order Model

The computational chart for the FOM is shown in the left column of Fig. 1. The FOM operates with matrices of dimensions $N_d \sim 10^4-10^5$ and connects α with the observable \mathcal{O} by first solving for the system's ground state through the RHB equations (blue box) from which the high-dimensional QRPA matrix $\mathcal{M}_{N_d \times N_d}(\alpha)$ is created.

The isovector dipole strength $S(\alpha; \omega)$ is computed with the axially-deformed FAM code of Ref. [84], based on the relativistic point-coupling EDF DD-PC1 [85] in the mean field channel, and the pairing interaction from Ref. [86]. The DD-PC1 EDF contains twelve parameters: four scalar couplings (a_S, b_S, c_S, d_S) ; three vector couplings (a_V, b_V, d_V) ; two isovector couplings (b_{TV}, d_{TV}) ; one derivative coupling δ_S , and two pairing strengths (G_p, G_n) .

Since α_D – and more generally the isovector giant dipole resonance (IVGDR) – are most sensitive to the isovector sector of the EDF [79, 80], we perform systematic calculations of the strength function by varying the two parameters: b_{TV} and d_{TV} .

All FAM QRPA calculations are carried out for a deformed nucleus ¹⁸⁰Yb, in a stretched axially-deformed harmonic oscillator basis with $N_{osc}=16$ shells. Both K=0 and K=1 projections of the dipole operator are included. The strength function is evaluated on the interval $\omega \in [0, 40]$ MeV with a step size of 0.1 MeV and a Lorentzian smearing of $\eta = 1$ MeV. The dipole polarizability is calculated by placing the semi-circular contour around $\omega = 0$ in the complex energy plane with radius $\gamma = 0.1$ MeV, and performing the integral by Simpson integration using 10 discretized points, following the prescription from Ref. [87]. Each calculation is run on 40 CPUs with 10 GB of memory per CPU on the MSU High Performance Computing Cluster. Evaluating the full strength function requires $\mathcal{O}(10^4)$ s of wall-clock time. Figure 2 displays the resulting strength distributions as b_{TV} and d_{TV} are varied.

As an example of charge-exchange response, we focus on studying the β -decay spectrum of the neutron-rich ⁸⁰Ni nucleus, as the GT strength of nuclei close to the

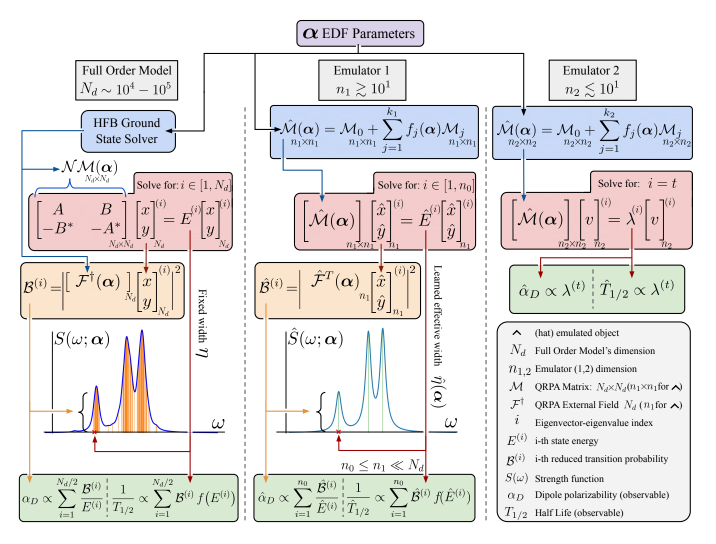


FIG. 1. Flowchart illustrating the computational algorithms for the full order model described in Sec. III A and the two emulators described in Secs. III B and III C. The three approaches aim to connect the controlling EDF parameters α (purple box at the top) with the two observables α_D and $T_{1/2}$ (green boxes at the bottom). A detailed description of the diagram is given in the text.

drip line shows considerable complexity within the Q_{β} window. Since this nucleus is spherical in its ground state, we employ the spherical matrix QRPA solver from Ref. [88], which guarantees that $N_d \approx 10^4$, allowing for direct diagonalization and obtaining the complete spectrum $\{E_{\alpha;i}, \mathcal{B}_{\alpha;i}\}$. Therefore, in this case, we directly employ Eq. (14) to calculate the half-lives. The groundstate configuration is obtained by solving the RHB equations in spherical harmonic oscillator basis with N_{osc} = 20. The strength function is sampled over two parameters, the isoscalar pairing V_0^{is} and the Landau-Migdal coupling g_0 . These correspond to time-odd terms, which need careful calibration to reproduce measured half-lives within the EDF framework [38, 39, 82]. The matrix diagonalization and subsequent calculation of half-life is performed on one node utilizing 56 cores, requiring around $\mathcal{O}(10^2)$ s. Figure 3 shows the calculated Gamow-Teller strength functions for ⁸⁰Ni across the ranges of the two parameters g_0 and V_0^{is} we explore in this work.

Using one of the previously described QRPA solvers, a spectral decomposition numerically yields all eigenpairs of the QRPA matrix, namely the N_d eigenstates $[x,y]^{(i)}$ and eigenergies $E^{(i)}$ (red box). These eigenpairs are then used to construct the strength function $S(\omega)$ as a sum of N_d Lorentzians, see Eq. (10). Each Lorentzian has a fixed width η , it is centered at energy $E^{(i)}$, and has an amplitude $\mathcal{B}^{(i)}$ obtained by calculating the overlap of the respective eigenstate $[x,y]^{(i)}$ with the external field $\mathcal{F}(\alpha)$ (orange box) for $i \in [1,N_d]$. The observables α_D and $T_{1/2}$ are then calculated as explicit functions of the amplitudes $\mathcal{B}^{(i)}$ through Eqs. (12) and (14).

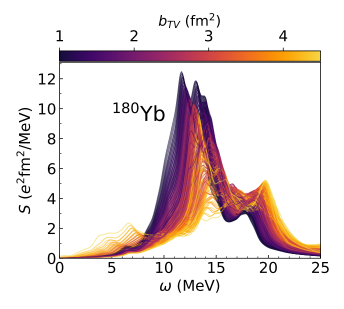


FIG. 2. The total isovector electric dipole strength function in $^{180}{\rm Yb}$ calculated with the FAM QRPA. Starting from the DD-PC1 EDF parameterization, each line corresponds to an individual QRPA calculation at fixed parameters b_{TV} and d_{TV} ; each curve is colored according to the value of b_{TV} . A total of 225 calculations were performed on a regular grid in the parameter space spanning $b_{TV} \in [1.0, 4.5]~{\rm fm}^2$ and $d_{TV} \in [0.1, 2.0]$. (See Fig. 5(a) for a graphical representation of the parameter distribution, and Fig. 6(a) for the response of α_D to changes in b_{TV} and d_{TV} .)

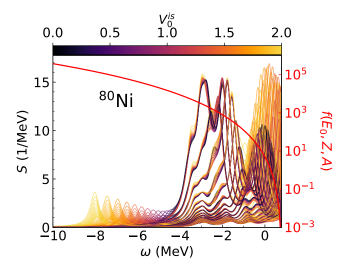


FIG. 3. The Gamow-Teller strength function in 80 Ni, within the Q_{β} -window, calculated with the linear response QRPA using DD-PC1 EDF parameterization at fixed parameters g_0 and V_0^{is} ; each curve is colored according to its value of V_0^{is} . The red solid line indicates the the phase-space factor $f(E_0, Z, A)$ at $E_0 = \Delta_{np} - \omega$. A total of 225 calculations were performed on a regular grid in the parameter space spanning $V_0^{is} \in [0.0, 2.0]$ and $V_0^{is} \in [0.0, 2.0]$ and $V_0^{is} \in [0.0, 2.0]$ are representation of the parameter distribution, and Fig. 6(b) for the response of V_0^{is} to changes in V_0^{is} .

B. Emulator 1

The computational chart for the first emulator (EM1) is shown in the middle column of Fig. 1. EM1 operates with matrices of dimension $n_1 \gtrsim 10^1$ and connects the underlying parameters α with the observables \mathcal{O} by first reproducing the strength function $S(\omega)$ as an intermediate high-dimensional latent step of the calculations. To achieve that, we observe that although $S(\omega)$ is indeed a high dimensional object comprised of the superposition of N_d Lorentzians, it can be well approximated by a much smaller number of $n_0 \ll N_d$ Lorentzians (see the schematic strength functions $S(\omega)$ and $\hat{S}(\omega)$ shown in the bottom part of columns one and two of Fig. 1, respectively). This approximation $\hat{S}(\omega) \approx S(\omega)$ effectively compresses the original high dimension, providing a much smaller set of reduced coordinates — the n_0 locations, n_0 amplitudes, and effective width $\hat{\eta}$ — that need to be calculated for each parameter evaluation.

To compute the values of the identified reduced coordinates, EM1 mirrors the structure of the FOM, except that the reduced matrix $\hat{\mathcal{M}}(\alpha)$ is not derived from solving the RHB equations. Instead, akin to data-driven operator learning methods that fit low-dimensional representations of operators directly from FOM data [61, 62, 89], and following the PMM structure [67], we represent this matrix as a linear combination of rank- n_1 matrices \mathcal{M}_i :

$$\hat{\mathcal{M}}(\boldsymbol{\alpha}) = \mathcal{M}_0 + \sum_{j=1}^{k_1} f_j(\boldsymbol{\alpha}) \mathcal{M}_j, \tag{15}$$

with k_1 chosen functions f_j that characterize the parametric dependence on α . In this work, we choose \mathcal{M}_0 as a real-valued diagonal matrix and \mathcal{M}_j to be real-valued symmetric matrices. A subset $n_0 \leq n_1$ of the obtained eigenvectors $[\hat{x}, \hat{y}]^{(i)}$ is used to construct the approximate reduced transitions $\hat{\mathcal{B}}^{(i)}$ through dot products with the approximate fixed external field $\hat{\mathcal{F}}$ learned from FOM evaluations:

$$\hat{\mathcal{B}}^{(i)} = \left| \hat{\mathcal{F}}^T(\alpha) \cdot [\hat{x}, \hat{y}]^{(i)} \right|^2, \quad i \in [1, n_0].$$
 (16)

These approximate transition probabilities $\hat{\mathcal{B}}^{(i)}$ form the amplitudes of the n_0 Lorentzians that are centered at the obtained eigenenergies $\hat{E}^{(i)}$ and used to construct the approximate strength function:

$$\hat{S}(\omega; \boldsymbol{\alpha}) = \frac{\hat{\eta}}{2\pi} \sum_{i=1}^{n_0} \frac{\hat{\mathcal{B}}^{(i)}}{(\omega - \hat{E}^{(i)})^2 + (\hat{\eta}/2)^2}.$$
 (17)

The effective Lorentzian width $\hat{\eta}(\alpha)$ is learned from FOM evaluations and changes with the controlling parameters α to compensate for the fact that the strength function is now built from a greatly reduced number of Lorentzians $n_0 \ll N_d$. The observables $\hat{\alpha}_D$ and $\hat{T}_{1/2}$ are then obtained from $\hat{\mathcal{B}}$ via Eqs. (12) and (14), respectively.

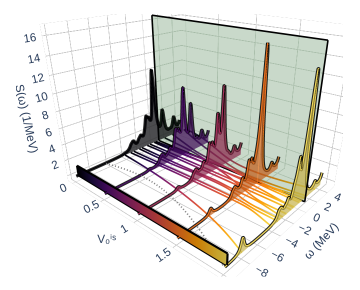


FIG. 4. Evolution of the Gamow-Teller strength function in 80 Ni as a function of V_0^{is} at $g_0=0.5$. The color coding follows the pattern of Fig. 3. The FOM strength functions $S(\omega)$ at five values of $V_0^{is}=[0.0,0.5,1.0,1.5,2.0]$ are shown. The tracks in the horizontal plane represent the subset of n_0 eigenvalues of matrix $\hat{\mathcal{M}}$. These eigenvalues represent the locations of the approximate energies \hat{E}_i that are used to construct the emulated strength function in Eq. (17). The values of $S(\hat{E}_i)$ are shown as vertical curves. The dotted line shows the lowest eigenvalue that is not retained in emulation; its behavior influences the overall dynamics. The vertical green plane shows the cutoff at $\Delta_{nH}=0.782$ MeV, representing the limit of the Q_{β} -window. For an interactive 3D visualization of the figure see Ref. [74].

EM1 is effectively constructing a low-dimensional algorithm that follows the original FOM structure as close as possible, with the numerical value of its hyperparameters — the matrix elements in Eq. (15) and the parameter dependence of $\hat{\eta}(\alpha)$ — being tuned by comparing to the FOM evaluations. The specific form of ansatz in Eq. (15) depends on the physics of the problem. For instance, in the case of β -decay emulations, it is known that the external field vector does not depend on g_0 and V_0^{is} parameters, and therefore we choose $\mathcal{F}(\alpha) = \mathcal{F}_0$. Furthermore, since there is no direct relationship between V_0^{is} and g_0 , we assume the following form for the matrix for calculations involving ⁸⁰Ni:

$$\hat{\mathcal{M}}(g_0, V_0^{is}) = \mathcal{M}_0 + (g_0 - \bar{g}_0)\mathcal{M}_1 + (V_0^{is} - \bar{V}_0^{is})\mathcal{M}_2, (18)$$

where we define:

$$(\bar{g}_0, \bar{V}_0^{is}) \equiv \alpha_C \tag{19}$$

as the central parameters on the training dataset. On the other hand, for the dipole polarizability α_D emulation, selected parameters actually impact the external field matrix $\mathcal{F}(\alpha)$. Therefore, for calculations involving ¹⁸⁰Yb, we choose:

$$\hat{\mathcal{F}}(b_{TV}, d_{TV}) = \mathcal{F}_0 + (b_{TV} - \bar{b}_{TV})\mathcal{F}_1 + (d_{TV} - \bar{d}_{TV})\mathcal{F}_2,$$
(20)

where the expansion is done around the central parameters α_C . Furthermore, in DD-PC1, the b_{TV} and d_{TV} parameters appear in a combination $b_{TV}e^{-d_{TV}x}$, where $x = \rho/\rho_{sat}$ is the ratio between vector density and the saturation density of symmetric nuclear matter [85]. Therefore, for α_D calculations, we choose $\hat{\mathcal{M}}$ as:

$$\hat{\mathcal{M}}(b_{TV}, d_{TV}) = \mathcal{M}_0 + (b_{TV} - \bar{b}_{TV})\mathcal{M}_1 + (d_{TV} - \bar{d}_{TV})\mathcal{M}_2 + (b_{TV} - \bar{b}_{TV})e^{-(d_{TV} - \bar{d}_{TV})x_1}\mathcal{M}_3,$$
(21)

where x_1 is an additional parameter to be learned from the data.

Figure 4 shows as an example the dynamical evolution of the eigenvalues (approximate energies $\hat{E}^{(i)}$) of the matrix $\hat{\mathcal{M}}(g_0, V_0^{is})$ when calibrated to reproduce the Gamow-Teller strength function of ⁸⁰Ni. For the fixed value of $\bar{g}_0 = 0.5$, the eigenvalues only depend on V_0^{is} through the diagonal matrix \mathcal{M}_0 and the symmetric matrix \mathcal{M}_2 .

The final step to build EM1 is to model the effective width $\hat{\eta}(\alpha)$. Knowing that $\hat{\eta}(\alpha)$ is positive, we decided to use the following ansatz:

$$\hat{\eta}(\boldsymbol{\alpha}) = \sqrt{\eta_0^2 + \left[\sum_{j=1}^{k_0} \eta_j(\alpha_j - \bar{\alpha}_j) + \eta_{k_0+1}\right]^2}, \quad (22)$$

where k_0 is the number of varied EDF parameters, and η_i for $j \in [0, k_0 + 1]$ are tunable parameters.

The total number of trainable parameters for EM1 for the dipole polarizability α_D calculations, with $k_0 = 2$ varied parameters $\boldsymbol{\alpha}$, $k_1 = 3$ parameter functions $f_j(\boldsymbol{\alpha})$ for the matrix parametric dependence in Eq. (15), and the selected functional form for the approximated field $\hat{\mathcal{F}}(\boldsymbol{\alpha})$ in Eq. (20), is then given by:

$$N_{param}^{\text{Em1}} = \frac{3}{2}n_1^2 + \frac{11}{2}n_1 + 5. \tag{23}$$

As an example, for a selected matrix size of $n_1 = 10$ we will have $N_{param}^{\rm Em1} = 210$ tunable parameters. For the $T_{1/2}$ emulator, and the matrix form (18), the number of tunable parameters is given by

$$N_{param}^{\rm Em1} = n_1^2 + 3n_1 + 5, (24)$$

corresponding to 135 for $n_1 = 10$.

Principles behind EM1 can be traced back to the random-matrix theory, first introduced in nuclear physics to describe excitations in complex systems, where distances between neighboring energy levels are small [90]. In particular, it was applied in Ref. [91] to streamline

computation of the coupling between complex configurations in RPA to explain the spreading widths of giant resonances. Both matrix elements of effective couplings as well as RPA eigenmodes can be formulated within random-matrix theory, by replacing them with random variables. Although such an approach seems promising in the case of giant resonances, where the density of the states is very large, it does not apply to the case where the structure of individual low-lying states are important. On the other hand, the ideas of EM1 — with matrix elements inferred from data rather than assumed to be random numbers — can apply to both.

C. Emulator 2

TABLE I. Training and testing ranges (in square brackets) for each parameter, and the number $N_{\rm FOM}$ of FOM evaluations used in the two cases studied. The FOM evaluations were distributed on a uniform regular grid in the parameter spaces, with the training points selected inside a rectangle region and the testing points outside to benchmark extrapolation (see Fig. 5).

Case	α	Train Range	$\begin{array}{c} {\rm Train} \\ N_{\rm FOM} \end{array}$	Test Range	$\operatorname*{Test}_{N_{\mathrm{FOM}}}$
$^{180}{\rm Yb}~\alpha_D$	$\begin{vmatrix} d_{\rm TV} \\ b_{\rm TV} ({\rm fm}^2) \end{vmatrix}$	$[0.5, 1.75] \\ [1.5, 4.0]$	121	$[0.1, 2.0] \\ [1.0, 4.5]$	104
80 Ni $T_{1/2}$	$egin{pmatrix} V_0^{is} \ g_0 \end{pmatrix}$	[0.29, 1.71] [0.14, 0.86]	121	$\begin{bmatrix} 0.0, \ 2.0 \\ 0.0, \ 1.0 \end{bmatrix}$	104

The flowchart for the second emulator (EM2) is shown in the right column of Fig. 1. EM2 operates with matrices of dimension $n_2 \lesssim 10^1$ and bypasses building any intermediate latent space involving the strength function $S(\alpha;\omega)$. Instead, it aims to directly connect the controlling parameters α with the observables \mathcal{O} . To achieve that, EM2 uses a fully data-driven approach through PMMs [67] to build a parametric matrix from data with the same structure as Eq. (15) with rank- n_2 matrix and a total of k_2 functions of the controlling parameters $f_i(\alpha)$. The eigenvalues of this matrix are obtained numerically and a single selected target eigenvalue $\lambda^{(t)}$ is used to yield approximated observables. In our particular implementation, we select the innermost eigenvalue, as it shows most variability with the evolution of model parameters. The form of the matrix is selected as in Eq. (21) and (18), for α_D and $T_{1/2}$, respectively. For instance, for α_D emulations, with $k_0 = 2$ varied parameters α , $k_2 = 3$ parameter functions $f_i(\alpha)$ for the matrix parametric dependence in Eq. (15), the total number of tunable parameters is given by:

$$N_{param}^{\text{EM2}} = \frac{3}{2}n_2^2 + \frac{5}{2}n_2 + 1. \tag{25}$$

For $n_2=5$, one gets $N_{param}^{\rm EM2}=51$ tunable parameters. For $T_{1/2}$ emulations, there are only $k_2=2$ parametric

functions $f_j(\alpha)$ in Eq. (15), and no term x_1 , such that the total number of tunable parameters is

$$N_{param}^{\rm EM2} = n_2^2 + 2n_2 + 1. {(26)}$$

For $n_2 = 5$, the number of tunable parameters is 36.

D. Calibration

To accurately reproduce the observables, an emulator must be calibrated, which accounts for finding optimal numerical values for its hyperparameters. These hyperparameters are the elements of all the matrices involved in Eq. (15), any free hyperparameters used to construct the functions $f_j(\alpha)$, and in the case of EM1 the hyperparameters used to build the effective width $\hat{\eta}(\alpha)$ and external field $\hat{\mathcal{F}}(\alpha)$.

We treat the learning procedure in a traditional machine learning optimization approach and build appropriate cost functions using a set of FOM evaluations for $m \in [1, N_{\text{FOM}}]$ parameters $\boldsymbol{\alpha}^{(m)}$ sampled from a chosen region in the parameter space. For the strength function, which plays a critical role in EM1, we build the cost function:

$$\chi_S^2 = \frac{1}{w_S} \sum_{m=1}^{N_{\text{FOM}}} \frac{\int d\omega \left| S(\omega; \boldsymbol{\alpha}^{(m)}) - \hat{S}(\omega; \boldsymbol{\alpha}^{(m)}) \right|^2}{\int d\omega \left| S(\omega; \boldsymbol{\alpha}^{(m)}) \right|^2}, \quad (27)$$

where w_S is a chosen weight to scale the cost function, $S(\omega; \boldsymbol{\alpha}^{(m)})$ is the FOM strength function, and $\hat{S}(\omega; \boldsymbol{\alpha}^{(m)})$ is calculated with EM1.

For the observables we build the cost function:

$$\chi_{\mathcal{O}}^2 = \frac{1}{w_{\mathcal{O}}} \sum_{m=1}^{N_{\text{FOM}}} \left[\mathcal{O}(\boldsymbol{\alpha}^{(m)}) - \hat{\mathcal{O}}(\boldsymbol{\alpha}^{(m)}) \right]^2, \quad (28)$$

where $w_{\mathcal{O}}$ is a chosen weight to make the cost function dimensionless, \mathcal{O} and $\hat{\mathcal{O}}$ are either α_D or $\ln T_{1/2}$ (as half-lives span orders of magnitude). These cost functions are then minimized with respect to the emulator's hyperparameters. For the case of EM1, both cost functions are combined, $\chi_S^2 + \chi_{\mathcal{O}}^2$, with the weights w_S and $w_{\mathcal{O}}$ determining the relative importance between them. For both α_D and $T_{1/2}$ emulators we choose $w_S = 1$. On the other hand, for the observables, in the case of the α_D emulator we select $w_{\alpha_D} = 0.5 \text{ fm}^{-6}$, while for $T_{1/2}$ emulator, $w_{T_{1/2}} = 1$. For EM2, the hyperparameters are only adjusted to minimize the corresponding observable cost function.

Since EM1 is aiming to reproduce both the high-dimensional strength function S as an intermediate step, as well as the associated observable, each FOM evaluation is information-rich for its training. We could interpret that for an approximate strength function \hat{S} that is being created by n_0 Lorentzians, each FOM calculations provides the optimal values of n_0 locations and n_0 amplitude peaks, for a total of $2n_0 + 1$ units of data

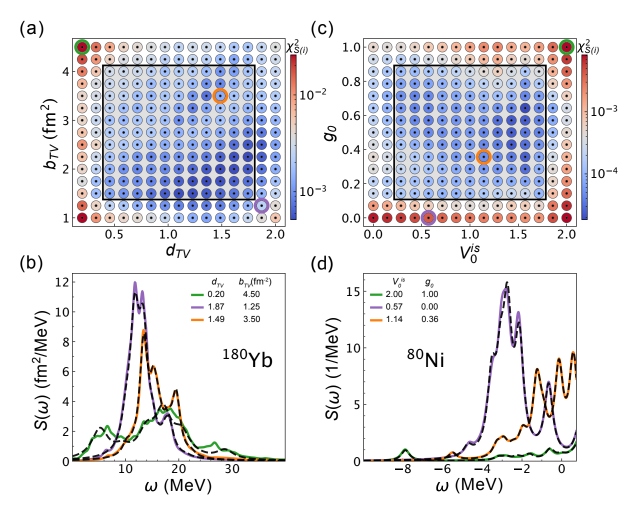


FIG. 5. Performance of EM1 in reproducing isovector dipole strength in ¹⁸⁰Yb (left panels), and the Gamow-Teller strength in ⁸⁰Ni (right panels). The emulator dimension was chosen as $n_1 = 13 (n_0 = 8)$ for the isovector dipole strength, and $n_1 = 13 (n_0 = 12)$ for the Gamow-Teller strength. The top panels show the relative error $\chi^2_{S(i)}(29)$ across the explored parameter range shown in Table I. In both cases, 121 points inside the black rectangle were used for training, while external 104 points form the test set. Bottom panels show three strength functions as calculated in QRPA (solid lines) and EM1 (dashed lines) for the selected values of parameters (shown in panels (a) and (c) as thick circles matching the colors of $S(\omega)$).

taking into account the associated value of the observable. This allows us to train EM1 with far less FOM evaluations than the total number of hyperparameters expressed in Eqs. (23) and (24). In contrast, since EM2 is only calibrated using the end value of the observable, each FOM evaluation produces only a single number for the cost function. In principle, this would suggest that one needs at least $N_{param}^{\rm Em2}$ FOM evaluations to completely fit all the hyperparameters of EM2 to obtain good performance. Nevertheless, we have observed in this and previous works [65, 66] that a good performance can be achieved with significantly lower number of evaluations than those required by these arguments. A possible reason lies in only a small subset of the eigenvalues of the parametric matrix meaningfully contributing to the response of the observable to parameter changes. While the targeted eigenvalue $\lambda^{(t)}$, and those in its vicinity are well constrained by FOM data, those in more distant

parts of the spectrum are not, yet they do not affect the emulator's performance.

Finally, in terms of the parameter initialization for the tuning, we note that the functions involved in expressions (18), (20), and (21) are expanded in terms of the difference between the parameters α and a central value α_C (19). This implies that the parametric functions involved are zero in the central parameters $f_i(\alpha_C) = 0$ for all cases, allowing for the initialization of the hyperparameter tuning to be done in an efficient way. For EM1, the parameters are initialized by fitting the energies and transition probabilities to reproduce the strength function of the central point α_C , which directly determines n_0 out of the n_1 matrix elements of \mathcal{M}_0 (energies of the peaks), as well as the n_0 of the n_1 components of the vector \mathcal{F}_0 (height of the peaks). The remaining hyperparameters are initialized with small random values sampled from a uniform distribution. We have found that

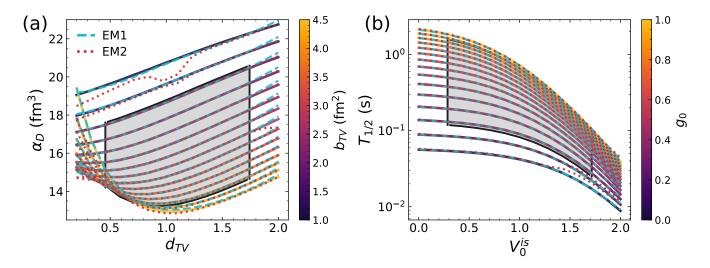


FIG. 6. Performance of EM1 with dimension $n_1=13$ (cyan dashed line) and EM2 with dimension $n_2=9$ (red dotted line), in reproducing (a) α_D of ¹⁸⁰Yb, and (b) $T_{1/2}$ of ⁸⁰Ni. The FOM calculations are shown by solid colored lines, with a color coding that matches that of Figs. 2 and 3. The calculations pertaining to the training set are shown by a gray area enclosed with a black boundary. Since α_D twists in a non-linear way near $d_{TV}\sim 0.2-0.7$, some curves at low values of b_{TV} cross into the training region even though these points were not used for training.

this initialization for EM1 achieves faster and more reliable convergence than if all hyperparameters were chosen at random. In contrast, hyperparameters of EM2 are initialized randomly in a small window around zero. To minimize the uncertainty due to optimizer finding different local minima based on random initialization, we train both types of emulators 10 times, sampling each time a different parameter set within a small window around zero. Subsequently, the training run which minimizes the overall training cost is selected.

The emulators based on both algorithms described in this section are implemented as Python scripts in tensorflow framework [92], and are publicly available in the repository [73].

IV. RESULTS

To study the performance of both emulators, we divide the parameter spaces into training and testing regions. This enables us to study the extrapolation ability of emulators beyond their calibration domains. To this end, we train the emulators on the *internal* region of the entire FOM dataset, and then extrapolate into the *external* region. Table I summarizes the ranges of the parameters α used and the total number of FOM evaluations for training and testing.

We first analyze the performance of EM1 in reproducing the strength functions in $^{180}\mathrm{Yb}$ and $^{80}\mathrm{Ni}$. Figure 5(a,c) shows the relative accuracy $\chi^2_{S(i)}$, defined by rewriting Eq. (27) as

$$\chi_S^2 = \sum_{i=1}^{N_{\text{FOM}}} \chi_{S(i)}^2.$$
 (29)

We employ $n_1 = 13$ in both cases. For isovector dipole response calculations in $^{180}{\rm Yb}$, $n_0 = 8$ eigenvectors and eigenvalues are retained, while for the Gamow-Teller response in $^{80}{\rm Ni}$ we take $n_0 = 12$. The error is very low in the internal training region (indicated by a black rectangle), between 0.01%-0.2%, and remains small in the testing region with most values of χ_S^2 less than 1% and a few points where χ_S^2 remains below 2%. In panels (b) and (d) we compare the emulated and FOM strength functions for three selected parameter values, one inside the training region and two points in the testing region, including also the worst performing value for both nuclei. Although the shape of the strength function changes significantly throughout the dataset, with a noticeable fragmentation on the edges of the testing region, the emulator captures its behavior effectively.

Having established the ability of EM1 to reproduce the strength function, we now compare the performance of both emulators in reconstructing the two observables of interest: α_D and $T_{1/2}$. In Fig. 6, we inspect the overall predictions across the parameter range. Both observables present high sensitivity to changes in the parameters, with challenging features for the emulators to learn, such as α_D showing highly non-linear behavior near $d_{TV} \sim 0.2 - 0.7$ and $b_{TV} \sim 3.5 - 4.5$ fm², and $T_{1/2}$ varying by orders of magnitude. Despite only being trained in the region shown by a black boundary, both emulators are able to capture well the response of the observables in the test-extrapolation region. EM1 can reproduce the behavior of α_D with a discrepancy of only a few percent, while EM2 struggles more in the strong non-linear region, having a larger discrepancy of around 10%. For $T_{1/2}$, both emulators reproduce the response, but EM2 shows some deviations of more than 50% on a

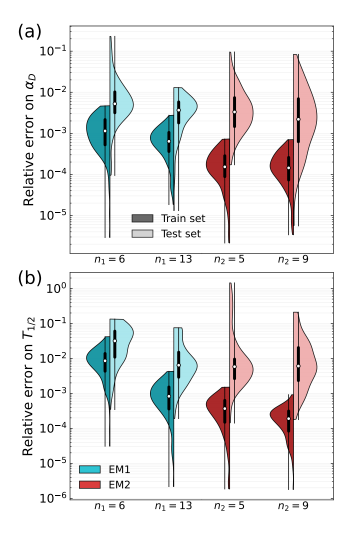


FIG. 7. Violin plots [93] benchmarking the performance of EM1 and EM2 on the training and test sets for two choices of their dimensions n. Panels (a) and (b) show the relative error on α_D in ¹⁸⁰Yb and $T_{1/2}$ of ⁸⁰Ni, respectively. For each emulator, we display two model dimensions for EM1 ($n_1=6$ and 13) and for EM2 ($n_2=5$ and 9). In each split violin, the left (darker) half corresponds to the training set and the right (lighter) half to the test set. The white dot marks the median, the thick black bar indicates the middle 50% of the data, and thin segments show the full range of the relative error.

small region around $g_0 \sim 0$ and $V_0^{is} > 1.5$. Considering that EM2 only used observables as training data, its predictive power in the extrapolation region is remarkable and showcases its ability to reproduce system's dynamics through the eigensystems of parametric matrices.

The emulator dimension did not appreciably affect computational times, with EM1 taking around twice as much time per evaluation as EM2, yet both showing appreciable speedups of 10^6-10^7 with respect to the FOM's computational time. For α_D , the median relative error for all emulators ranges between 0.06%-0.1% for the training set, and between 0.2%-0.5% for the testing set, while for $T_{1/2}$ it shows a bigger spread between

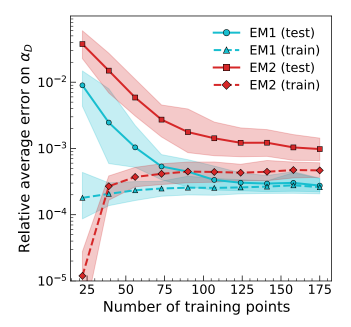


FIG. 8. The relative average error on α_D on the training set (triangles: EM1, diamonds: EM2) and test set (circles: EM1, squares: EM2) as a function of the number of training points. The comparison is made for both EM1 (cyan) with $n_1 = 8 \, (n_0 = 4)$ and EM2 (red) $n_2 = 5$. Both emulators are trained with the same data and randomly initialized 100 times, with mean values denoted by solid and dashed lines, and 95% coverage intervals marked by the shaded areas. Similar results have been obtained for $T_{1/2}$.

0.02% - 1% for training and 0.6% - 3% for testing. EM1 shows more consistency between its training and testing errors, with the testing error decreasing for both observables when the emulator dimension is increased from 6 to 13. In contrast, EM2 shows a bigger discrepancy between its training and testing errors, a difference that becomes larger when its intrinsic dimension is increased from $n_2 = 5$ to 9, with training errors being reduced while testing errors remain almost unchanged. However, the relative error distribution significantly broadens. These two behaviors suggest that, for these values of dimensions we explored, EM1 better captures the intrinsic dynamics of the systems, with good generalization to extrapolating to unobserved parameters, while EM2 shows signs of overfitting, with almost two orders of magnitude spread between its training and testing errors for $T_{1/2}$ with $n_2 = 9$.

Finally, we study how the performance of both emulators in reproducing α_D is affected by the amount of FOM training data. The main results are shown in Fig. 8 for EM1 with $n_1 = 8 \, (n_0 = 4)$ and EM2 with $n_2 = 5$. While Figs. 5, 6, and 7, illustrate extrapolation capabilities, in Fig. 8 we follow a setup more likely to be used in practical calculations, where the emulator is purposely trained to cover the expected evaluation region in parameter space. For testing purposes we select 50 FOM cal-

culations drawn randomly from the points of Fig. 5(a,c). From the remaining 175 points, we retain a varying fraction of training points ensuring that the i-th partition contains all the points of the previous one. This is done to minimize the impact on the results of the specific parameter locations of FOM evaluations, which is known to affect the quality of trained emulators [51, 94]. In all training partitions all four corner points spanning the edges of the parameter ranges in Table I are included, together with the central point α_C . To take into account the variability on results associated with the initialization of their hyperparameters we perform the training 100 times, each time initializing matrix elements of matrices and vectors in EM1 and 2 with different random seeds. The solid and dashed lines in the figure shows the mean error while the bands encompass 95% of the obtained curves.

The test accuracy of EM1 improves steeply at the beginning and then plateaus around 125 training points, while the test accuracy of EM2 continues to slightly improve as we use more data. The opposite trend can be observed for the relative error on the training set, as expected since more points are being included that the emulators must reproduce, yet their complexity (dimension) remains fixed. Based on the performance of the test set, we see that EM1 can achieve an accuracy of almost an order of magnitude better than EM2 for a fixed number of training points, and its performance in the train and test sets converges to the same value. We attribute these observations to the fact that EM1 is reproducing the strength function $S(\omega)$ as an intermediate step in the calculation, highlighting the impact that physics-informed decisions in the construction of data-informed emulators can have on their final performance. The fact that EM2 is able to achieve sub-percent accuracy with only ~ 50 training points is quite impressive, given that each FOM evaluation only gives one observable, which is used to train 51 free parameters. This observation further showcases PMMs [67] as a flexible and robust machine learning approach capable of learning complicated mappings directly from the data.

V. CONCLUSIONS

We have developed and benchmarked two emulators, EM1 and EM2, that are able to reproduce full QRPA strength-function calculations with sub-percent accuracy while offering a speedup of 6-7 orders of magnitude compared to the the original full-order solver. As representative benchmarks, we examined two key nuclear observables: (i) the electric dipole polarizability of $^{180}\mathrm{Yb}$ as a function of the two isovector couplings $(b_{TV},d_{TV}),$ and (ii) the Gamow-Teller β -decay half-life of $^{80}\mathrm{Ni}$ as a function of the isoscalar pairing strength and the Landau-Migdal coupling $(V_0^{is},g_0).$

For EM1, we followed an approach inspired by datadriven operator learning methods that approximate lowdimensional representations of operators using FOM evaluations [61, 62, 89]. We created a low-dimensional representation of $S(\omega)$ through the sum of a few effective Lorentzians, and learned the necessary governing equations as a function of the underlying parameters α from FOM. The forms of these equations were inspired by the structure of the full QRPA pipeline, with the governing matrix $\mathcal{M}(\alpha)$ following the structure of the PMM framework [67]. We built EM2 in a fully data-driven approach by using the PMM framework to directly connect the underlying parameters α with the observables. EM1 showed better accuracy on the observables for testset parameters, reproducing strong non-linear features in α_D and the variation over two orders of magnitude in β -decay half-lives. EM2 only required data from the final QRPA observables to be trained, and usually showed better accuracy in the training region, while still being able to reproduce part of the response of the observables in the test set. Besides the half-life and polarizability, EM1 was able to also provide a good reproduction of the strength function $S(\omega)$, which can be used to calculate other quantities of interest. The web-based tools deployed alongside this work [74] can run EM1 in real time to explore how $S(\omega)$ changes as the interaction parameters are tuned, yielding precise and rapid insight on parameter influence while avoiding repeated FOM runs.

By restructuring the training region to represent a more realistic case, we have demonstrated that EM1 is able to achieve sub-percent accuracy on the test set being trained with only 22 data points, over which the strength function varies considerably. On the other hand, due to its simplicity, EM2 can be straightforwardly applied to learn observables directly, with the tradeoff of requiring more training data than EM1.

In the current work we focused on varying two of the controlling parameters α of the EDF as a first step to test the viability of our outlined framework to create QRPA emulators. The high computational cost associated with ORPA has limited the calibration of nuclear EDFs informed by excited state data. By extending these methods to incorporate all the relevant EDF parameters we will be in position to offer a possible future path for the implementation of QRPA-based methods in Bayesian optimization of EDFs and subsequent uncertainty quantification. The reduction in variability, paired with methods to estimate the errors of the emulators during the on-line (deployment) phase, will provide the robustness needed to incorporate the emulators into calibration and large studies pipelines. Since our approach is agnostic to the specific physical system, it should be straightforward to extend to problems across physics and chemistry areas that apply linear-response theory, thus opening the door to scalable Bayesian uncertainty quantification in broad applications for scientific discovery.

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Appendix A: Acronyms and Symbols

Tables II and III summarize the acronyms and symbols used.

TABLE II. Glossary of acronyms used in the paper

	0 1 1		
DFT	Density Functional Theory		
EDF	Energy Density Functional		
EM1	Emulator 1 (physics-informed surrogate)		
EM2	Emulator 2 (data-driven PMM surrogate)		
FAM	Finite-Amplitude Method (linear-response		
	solver)		
FOM	Full Order Model (high-fidelity calculation)		
GT	Gamow–Teller (charge-exchange) transition		
HF(B)	Hartree–Fock(-Bogoliubov) framework		
PMM	Parametric Matrix Model		
(Q)RPA	(Quasiparticle) Random Phase		
,	Approximation		
RHB	Relativistic Hartree–Bogoliubov framework		
TDDFT	Time-Dependent Density Functional Theory		

TABLE III. Symbols and parameters used

Titbee iii. Symbolo and parameters about				
α	Vector of controlling parameters			
$lpha_C$	Central parameter point (training cen-			
	ter) (19)			
α_D	Electric dipole polarizability (11)			
η	Lorentzian folding width (10)			
$\mathcal{B}_{lpha;i}$	Reduced transition probability (9)			
b_{TV}, d_{TV}	Isovector EDF couplings			
$E_{\alpha;i}$	QRPA excitations (8)			
$\mathcal{F}(oldsymbol{lpha})$	External field			
$f(E_0, Z, A)$	Fermi function			
$\mathcal{M}(oldsymbol{lpha})$	QRPA matrix (6)			
\mathcal{N}	QRPA metric (5)			
N_d	Rank of the QRPA matrix			
n_0, n_1, n_2	Emulator latent dimensions			
$n_{2\mathrm{qp}}$	Number of two–quasiparticle states			
O	Observables			
$\hat{\mathcal{O}}$	Emulated observables			
$\delta \mathcal{R}(\omega)$	Linear response amplitude (2)			
$S(\alpha; \omega)$	Strength function (7)			
$T_{1/2}$	β -decay half-life (13)			
V_0^{is}, g_0	Isoscalar pairing strength and Lan-			
₩ · = '	dau-Migdal coupling			

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