I. INTRODUCTION

β-decay rates are an important ingredient in simulations of the astrophysical r process. Because parts of the r-process path are still not accessible to experiment, it is up to theoretical models to produce approximate rates for many relevant neutron-rich isotopes. Models could also help resolve the issues raised by Ref. [1], which argued that the flux of antineutrinos from nuclear reactors does not agree with the standard model. Reference [2] pointed out that the discrepancy could be attributable to an overly simple treatment of first-forbidden β decay in fission products.

Several schemes/methods for calculating β-decay rates across almost the entire nuclear chart have been devised. Besides the phenomenological semigross theory [3], they include the microscopic proton-neutron quasiparticle random-phase approximation (pnQRPA) with separable Gamow-Teller (GT) interaction [4] and an analogous scheme with no dynamic T = 0 pairing interaction [5] and its extension for the first-forbidden (FF) transitions added using the gross theory [6]. The self-consistent pnQRPA models were developed based on the extended Thomas-Fermi plus Strutinsky integral method [7], on the Fayans density functional [8], and, very recently, on the spherical pnQRPA [9]. For the interacting shell-model calculations, see Ref. [10] (in Refs. [8–10] both the GT and FF transitions were consistently included). An artificial neural network analysis can be found in Ref. [11]. Full β-decay tables for neutron-rich isotopes have been only published by Möller et al. [4,5] and Marketin et al. [9].

Many other authors have applied more sophisticated and/or computationally intensive methods to smaller sets of nuclei, focusing on some of those important for the r process.
they are independent of the time-even functional. In much of what follows, therefore, we do not assume that the functional results from mean-field theory with an interaction and so will have to fit a significant number of parameters. After presenting our methods and assessment, we display the (summarized) results of a full table of β− rates, computed with the pNpFAM, for even-even nuclei in all medium-mass and heavy isotopic chains. We use a simple but apparently accurate model to quantify and extrapolate theoretical uncertainty.

This article is organized as follows. Section II is a brief overview of the theoretical background, and Sec. III details our computational approach and parameter-fitting procedures. In Sec. IV, we assess the quality of our results, comparing them to earlier work and to experimental data where available. Section V contains conclusions.

II. THEORETICAL BACKGROUND

A. Finite-amplitude method

The finite-amplitude method (FAM), a formulation of the random-phase approximation that speeds the computation of nuclear response functions, was introduced in nuclear physics in Ref. [21]. It was later generalized to the QRPA in Ref. [22] and to the relativistic QRPA in Ref. [23]. In Ref. [18] we applied the method to charge-exchange transitions, in particular allowed and first-forbidden β decay.

The FAM solves equations for the amplitude of the linear response to a small but finite perturbation. As a result, the method does not directly yield the poles and residues of the response, which are the central objects in the matrix version of the QRPA. However, if the goal of the computation is to get transition strength functions in a large model space the FAM can yield results in orders of magnitude less CPU time than matrix QRPA.

The FAM offers another advantage for β decay: The weighted sums or integrals of transition strength can be expressed as contour integrals. This fact was first exploited by Hinohara et al. [24], who evaluated the response at a relatively small number of complex frequencies to compute sum rules. In Ref. [18] we used the idea to evaluate the more complicated β-decay phase-space-weighted integrals, which are not analytic and contain interference terms between first-forbidden operators. With the FAM we can thus use typical supercomputer resources to calculate many observables in a large number of nuclei. We are able to extend systematic Skyrme parameter fitting from mean-field calculations to deformed QRPA calculations, at least in a preliminary way.

B. Model parameters and fitting targets

Our starting point in the particle-hole channel is a generic Skyrme EDF:

\[
E = \sum_{i=0,1} \sum_{r_{ij}=-\infty}^{+t} dr [\mathcal{H}_{r_{ij}}^{\text{even}}(r) + \mathcal{H}_{r_{ij}}^{\text{odd}}(r)].
\]

Here \(\mathcal{H}_{r_{ij}}^{\text{even}}\) contains products of time-even local densities only, with coefficients fixed by fitting to masses and perhaps a few other quantities, and \(\mathcal{H}_{r_{ij}}^{\text{odd}}\) is given by

\[
\mathcal{H}_{r_{ij}}^{\text{odd}}(r) \equiv C_i^r [\rho_{00}(r) s_{r_{ij}} + C_i^s s_{r_{ij}} \cdot \nabla s_{r_{ij}} + C_i^{j_3^2} j_{r_{ij}}^3 + C_i^f s_{r_{ij}} \cdot T_{r_{ij}} + C_i^{\alpha^2} s_{r_{ij}} \cdot \nabla \times J_{r_{ij}} + C_i^{C} s_{r_{ij}} \cdot F_{r_{ij}} + C_i^{\gamma^2} (\nabla \cdot s_{r_{ij}})^2],
\]

with the spin density \(s_{r_{ij}}\), the current density \(j_{r_{ij}}\), the spin-kinetic density \(T_{r_{ij}}\), and the tensor-kinetic density \(F_{r_{ij}}\), defined, e.g., in Ref. [25]. If one requires the functional to be the mean-field expectation value of a Skyrme interaction, EDF coupling constants are completely determined by the (fewer) parameters that specify the interaction, as discussed in Refs. [25,26] and mentioned above. In this work, we adopt the view that the effective “interaction” comes from the EDF rather than the other way around. Consequently, we are free to fit all the time-odd coupling constants without spoiling the mass fits generated by the time-even couplings. We do, however, adopt the values obtained from the Skyrme parametrization as our starting point for the fits, unless we note otherwise.

The subset \(\{C_i^s, C_i^j, C_i^f\}\) of time-odd coupling constants maps directly to the parameters of the Landau-Migdal interaction for infinite homogeneous nuclear matter with tensor terms [27]. In that sense, these couplings are intimately related to the bulk properties of the nuclear matter. The constant \(C_i^s\) is purely tensor in character, and it determines the tensor term in the Landau-Migdal interaction. The spin-density coupling constant \(C_i^j\) strongly affects the GT strength distribution and can be fit to the locations of GT resonances [26]. The last term \(C_i^f\) maps to a linear combination of the Landau parameters of both the tensor term and a term that depends on the scattering angle of the Landau-Migdal quasiparticles.

Two other parameters have a large effect in the QRPA: the strengths of the residual particle-particle (or pairing) interaction between protons and neutrons,

\[
V_{\text{pp}} = (V_0 \hat{\Pi}_{T=0} + V_1 \hat{\Pi}_{T=1}) \left[ 1 - \alpha \frac{\rho_0(r)}{\rho_s} \right] \delta(r),
\]

where \(\rho_s = 0.16 \text{ fm}^{-3}\) is the saturation density of nuclear matter and \(\alpha \in [0,1]\) controls the pairing density-dependence (throughout this work we use mixed pairing, i.e., \(\alpha = 0.5\)). The isovector \((T = 1)\) proton-neutron pairing mainly affects Fermi β decay, which plays almost no role in heavy nuclei; we simply set its strength \(V_1\) to be the average of the neutron-neutron and proton-proton pairing strengths [both fixed in the Hartree-Fock-Bogoliubov (HFB) part of the calculation]. The isoscalar \((T = 0)\) pairing is a different story; it has a strong effect on GT decay, and determining a reasonable value for its strength is a common issue in single and double β-decay computations. The HFB mean field is independent of the \(T = 0\) pairing term as long as explicit proton-neutron mixing is neglected. We are thus free to fit \(V_0\) in the QRPA.

We include three types of observables in our fitting procedure: Gamow-Teller resonance energies, spin-dipole resonance energies, and total β-decay rates, all in both spherical and deformed isotopes. Although we pick several sets of target nuclei for fitting the decay half-lives, each set includes a large range of mass values so that our fits can be global. To assess the success of the fits and to compare them to earlier work in
that are too short lived to allow measurement. The uncertainty in our predicted half-lives, particularly in isotopes Here we attempt to provide reasonable estimates for the analysis of theoretical uncertainty is growing in importance. For very different models, we also compute the following metrics of each computed \( \log_{10} t \) value (where \( t \) is the half-life),

\[
r = \log_{10} \left( \frac{t_{th}}{t_{exp}} \right),
\]

(4)

the average of the residuals,

\[
M_r = \frac{1}{n} \sum_{i=1}^{n} r_i
\]

(5)

(where we have used an index \( i \) to indicate that there is an \( r \) for every nucleus), and the standard deviation of the residuals around the average,

\[
\sigma_r = \sqrt{\frac{1}{n} \sum_{i=1}^{n} [r_i - M_r]^2}.
\]

(6)

Reference [11] contains an excellent compilation of these quantities.

C. Uncertainty analysis

As theoretical approaches grow more sophisticated, the analysis of theoretical uncertainty is growing in importance. Here we attempt to provide reasonable estimates for the uncertainty in our predicted half-lives, particularly in isotopes that are too short lived to allow measurement.

The standard prescription for assigning a theoretical (statistical) uncertainty \( \Delta O \) to a computed observable \( O \) is [28]

\[
\Delta O = \sqrt{\sum_{ab} \frac{\partial O}{\partial x_a} C_{ab} \frac{\partial O}{\partial x_b} \bigg|_{x=x_0}},
\]

(7)

where \( C \) is the covariance matrix

\[
C = (J^T J)^{-1}
\]

(8)

and \( x = (x_1, \ldots, x_N) \) are the \( N_x \) parameters of the model. The partial derivatives of all the observables \( \{O_a\} \) with respect to all the parameters evaluated at the result of the fit \( x_0 \) form the Jacobian \( J \):

\[
J_{ab} = \left. \frac{\partial O_a}{\partial x_b} \right|_{x=x_0}.
\]

(9)

When not analytically accessible, the needed partial derivatives can be estimated through finite central differences. In general, the theoretical uncertainty related to the Jacobian must be supplemented by numerical and experimental uncertainties. We have assumed that the theoretical uncertainty is much larger than the other two, which we therefore neglect.

To use Eq. (7) to assign an uncertainty to every \( \beta^- \)-decay rate in our table, we would need to evaluate the Jacobian in Eq. (9) for (the logarithms of) all the half-lives \( t \) in our table. Unfortunately, the required \( 2N_t \) full decay-table computations are still not possible in a reasonable amount of computer time, even with the efficiency of the FAM. We therefore attempt to gauge the uncertainties and their \( Q \) dependence in a more naive way. We construct a simple few-parameter model for the uncertainties that we can then fit to the observed differences between our numerical results and known experimental values. The resulting approach is agnostic about how the decay is actually calculated; it treats the nuclear model as a black box that produces predictions for \( Q \) values and \( \beta^- \)-decay rates. It does, however, make several assumptions about both the calculated and the experimental strength distributions that are only approximately correct.

The first assumption is that the final states that contribute significantly to a decay rate lie not too far from the ground state in a relatively small window of excitation energy, so that we can reasonably approximate them, again either in our calculation or in the real world, by one effective state with an effective \( Q \) value \( q_{\text{eff}} \) (that is not too different from the ground-state \( Q \) value):

\[
q_{\text{eff}} = \frac{\sum_k C_k f_k(q_k + 1, Z_f) q_k}{\sum_k C_k f_k(q_k + 1, Z_f)}
\]

(10)

Here \( C_k \) is the standard integrated shape function for the transition to the final state \( k \)—for an allowed state decay this is simply the transition strength, and for a nonunique forbidden decay it encapsulates several terms—and \( f_k \) is the usual allowed phase-space integral. The quantity \( q_k \) is the \( Q \) value of the decay to the state \( k \) divided by \( m_e c^2 \), and \( Z_f \) is the charge of the daughter nucleus.

With these definitions, we can proceed to define an effective shape factor \( C_{\text{eff}} \) as

\[
t = \frac{\sum_k C_k f_k(q_k + 1, Z_f)}{C_{\text{eff}} f_{\text{eff}}(q_{\text{eff}} + 1, Z_f)} = \frac{\kappa}{C_{\text{eff}} f_{\text{eff}}(q_{\text{eff}} + 1, Z_f)}.
\]

(11)

The \( C_k \) depend on the \( q_k \) for forbidden transitions, but the dependence is weak compared to that of the corresponding phase-space integral, and so we neglect it in our effective shape factor.

All these definitions can be made independently for the experimental strength distribution and the theoretical one. The quantity we wish to understand is the ratio \( r \) of the theoretical and experimental lifetimes,

\[
r = \log_{10} \frac{t_{th}}{t_{exp}} = \log_{10} \frac{C_{\text{exp}}^{\text{eff}} t_{th}}{C_{\text{eff}}^{\text{exp}} t_{exp}} + \log_{10} f(q_{\text{exp}} + 1, Z_f) - \log_{10} f(q_{\text{eff}}^{\text{th}} + 1, Z_f).
\]

(12)

where the meanings of \( d_{\text{exp}}^{\text{exp}} \) and \( q_{\text{eff}}^{\text{th}} \) and the corresponding quantities \( C_{\text{exp}}^{\text{exp}} \) and \( C_{\text{eff}}^{\text{exp}} \) should be clear. We omit the nucleus index \( i \) in them for brevity. Because the phase space grows quickly with decay energy, the effective \( Q \) values will usually be close to ground-state-to-ground-state \( Q \) value. We therefore expand both logarithms in Eq. (12) of the phase-space factors about the theoretical ground-state-to-ground-state \( Q \) value \( q_{\text{g.s.}}^{\text{th}} \) to first order in \( q \):

\[
\log_{10} f(q + 1, Z_f) \approx \log_{10} f(q_{\text{g.s.}}^{\text{th}} + 1, Z_f) + \frac{d \log_{10} f(q + 1, Z_f)}{dq} \bigg|_{q = q_{\text{g.s.}}^{\text{th}}} (q - q_{\text{g.s.}}^{\text{th}})
\]

\[
= \log_{10} f(q_{\text{g.s.}}^{\text{th}} + 1, Z_f) + f'(q_{\text{g.s.}}^{\text{th}} + 1, Z_f) q - q_{\text{g.s.}}^{\text{th}} \ln 10.
\]

(13)
This approximation is best when the $Q$ value is high, because the curvature of $\log_{10} f(q+1,Z_f)$ is small at high $q$. For $Q$ values lower than about 2–3 MeV the first-order approximation is poor, so we exclude such data points from our analysis.

Replacing the two logarithms in Eq. (12) with the first-order expressions, we find that several terms cancel in the difference, so that

$$ r \approx \log_{10} \frac{C_{\text{th}}^{\text{exp}}}{C_{\text{th}}^{\text{th}}} + \frac{f'(q_{\text{th}}^{\text{g.s.}} + 1,Z_f)}{f(q_{\text{th}}^{\text{g.s.}} + 1,Z_f)} q_{\text{th}}^{\text{exp}} - q_{\text{th}}^{\text{th}}. \quad (14) $$

We now make another set of assumptions, this time about the distribution of the errors in the theoretical values: First, we assume that the relative error in the effective shape factor is normally distributed with a slight systematic component. That is, we assume that for each nucleus, we have

$$ \log_{10} \frac{C_{\text{th}}^{\text{exp}}}{C_{\text{th}}^{\text{th}}} = c + \delta c, \quad (15) $$

where $c$ is a constant, independent of the nucleus in which the decay occurs, that captures the systematic error and the set of all nucleus-dependent $\delta c$'s is normally distributed with standard deviation $\Delta c$, which is a measure of statistical error that is independent of decay energy. Similarly, we assume that the errors in the theoretical effective $Q$ value follow a normal distribution, so that

$$ q_{\text{th}}^{\text{exp}} - q_{\text{th}}^{\text{th}} = q + \delta q, \quad (16) $$

where $q$ is now a nucleus-independent systematic error in the effective $Q$ value and the set of $\delta q$'s (again, one for each nucleus) is normally distributed around zero with the standard deviation $\Delta q$ that represents statistical error, again independent of $Q$. (Both $q$ and $\delta q$ are expressed in units of $m_e c^2$, and the factor $1/\ln 10$ is absorbed into $q$ and $\delta q$ for convenience.) Finally, we assume that the errors $\delta q$ and $\delta c$ are independent.

With these assumptions, we then have

$$ r = c + \frac{f'(q_{\text{th}}^{\text{g.s.}} + 1,Z_f)}{f(q_{\text{th}}^{\text{g.s.}} + 1,Z_f)} q + \delta r, \quad (17) $$

where $\delta r$ is a random error, the set of which must be normally distributed at each $q_{\text{th}}^{\text{g.s.}}$ with width

$$ \Delta r(q_{\text{th}}^{\text{g.s.}})^2 = \Delta c^2 + \left[ \frac{f'(q_{\text{th}}^{\text{g.s.}} + 1,Z_f)}{f(q_{\text{th}}^{\text{g.s.}} + 1,Z_f)} \right]^2 \Delta q^2. \quad (18) $$

We can now use Eqs. (17) and (18) to determine the values of $c$, $q$, $\Delta c$, and $\Delta q$. We obtain the first two through a fit to Eq. (17), which expresses $r_i$ as a function of $f'/f$ (and thus of $q_{\text{th}}^{\text{g.s.}}$), with the set of ratios $r_i$ given by our calculations (and experiment) and $\delta r_i$ set to zero. Finally, we insert the fit values of $c$ and $q$ into the square of Eq. (17), which then expresses $\delta r_i^2$ as a function of $(f'/f)^2$, and determine the values of $\Delta c$ and $\Delta q$ by requiring that the line for $\Delta r(q_{\text{th}}^{\text{g.s.}})^2$ as a function of $(f'/f)^2$ that expresses Eq. (18) goes through the middle of the data, with as many points below $\Delta r^2$ as above it. A second least-squares fit of $\Delta c$ and $\Delta q$ in the right-hand side of Eq. (18) to the set of $\delta r_i^2$'s accomplishes the task nicely. The explicitly $Q$-dependent $\Delta r$ can then be extrapolated to large $Q$ values where there are no data.

For both fits, in practice, we only include data points with $Q \geq 3$ MeV and exclude any data points with $|r_i| > 4$ as obvious outliers. We also adopt the Primakoff-Rosen approximation [29] for the phase-space integral in the uncertainty model, making the expression $f'/f$ a simple ratio of two polynomials, independent of $Z_f$:

$$ f'(x,Z_f) \approx \frac{5x^4 - 20x + 15}{x^3 - 10x^2 + 15x - 6}. \quad (19) $$

Once the four parameters of the uncertainty model have been determined, the uncertainties of the theoretical predictions are obtained the inverse relation of (4)

$$ t_{\exp} = t_{\text{th}} \times 10^r, \quad (20) $$

with

$$ r = r(q_{\text{th}}^{\text{g.s.}}) \pm \Delta r(q_{\text{th}}^{\text{g.s.}}). \quad (21) $$

### III. Skyrme Functional and Computational Method

The first step in our computational procedure is the construction of ground states in the doubly even mother nucleus with hfbtho, a well-established HFB solver working in (transformed) harmonic-oscillator basis [30]. We cut off the single-particle space at 60 MeV to avoid divergences from our zero-range pairing. For each nucleus, we search for a prolate, an oblate, and a spherical solution and take the most bound of these to be the ground state.

Because the $\beta$-decay rates are very sensitive to the $Q$ value of the decay, we look for a modern Skyrme functional that reproduces $Q$ values well. Because we do not explicitly treat odd nuclei, we use the prescription of Ref. [31] to approximate the $Q$ value; we have checked the prescription against odd-$A$ calculations in the equal filling approximation, and the two procedures generally agree to within about 0.5 MeV. Of the several functionals we examine, SkO' [32] (with the strengths of proton-proton and neutron-neutron pairing fit to the experimental pairing gaps of ten isotopes picked in a wide mass range $50 \leq A \leq 230$) does the best job with $Q$ values, producing errors for ground-state-to-ground-state $Q$ values that are normally distributed, with an average systematic error of 0.154 MeV and statistical error of 0.576 MeV. Figure 1 compares the $Q$ values produced by SkO' with those of the next best functional, SV-min.

To compute $\beta$-decay rates and resonance energies, we use the code pnfam, an implementation of the charge-changing FAM presented in Ref. [18]. Built to work together with hfbtho, pnfam allows us to compute properties of axially-deformed nuclei, including both allowed and first-forbidden $\beta$ decay.

We obtain our most robust fits by fixing all but two of the time-odd coupling constants of the functional at values implied by its interpretation as an interaction. (The mapping of the Skyrme coupling constants to the EDF coupling constants is explicitly discussed in Refs. [25,26].) The exceptions are $C_1^\gamma$ and $C_1^{\Delta A}$. We set the latter to zero to avoid known finite-size...
instabilities [33] which, in the case of HFBTHO and PNFAM, manifest themselves as divergences in the iterative solution. That leaves Cs1, which, along with the isoscalar pairing strength V0, we fit to a set of GT resonance energies, spin-dipole resonance energies, and β-decay rates selected from a wide mass range with no particular region favored. We use the code POUNDERS, based on a derivative-free algorithm [34] designed for optimizing computer-time-consuming penalty functions, to efficiently minimize the weighted sum of the least squares, simultaneously fitting both parameters. We take Cs1 to be independent of the density and V0 to have the same density dependence as the proton and neutron pairing. The axial-vector coupling constant gA is known to be quenched in nuclei, but the source and magnitude of the quenching is an open problem; a variety of very different values for an effective gA have been used. For lack of a better prescription, we use the commonly adopted quenched value gA = 1.0 in the GT channel, while applying no quenching in the first-forbidden channels (we also might have tried a similarly quenched value for the first-forbidden channels). We weight the three types of observables—two kinds of energies and a rate—in the least-squares fit so as to approximately normalize the total penalty function \( \chi^2 \) to the number of degrees of freedom, following the recommendation in Ref. [28]. We assume that the theoretical error dominates the experimental error and thus assign equal weight to an observable of the same type. We select these weights based on how well the different observables are reproduced in an initial test fit, so that each type of observable is approximately equally weighted in the actual fit. Typical fits then take 10 000–20 000 CPU hours, and we use XSEDE supercomputers [35] to carry them out.

Following the fit, we proceed to compute the β-decay rates of all even-even neutron-rich nuclei with 28 \( \leq Z \leq 110, A \geq 50, \) all the way to the neutron drip line, omitting just a few very stable isotopes for which the \( Q \) value is negative in our HFB calculations.

### IV. RESULTS AND DISCUSSION

#### A. Fit results

To assess how sensitive our fit is to the set of target β-decay rates and resonance energies, we repeat the process with four sets of rates, summarized in Table 1. Each of these sets spans a large range of masses. Set A contains β-decay isotopes with relatively short half-lives only, set B relatively long half-lives only, and set C a wide range of half-lives. (Short half-lives should be less sensitive to details in nuclear structure, whereas long half-lives, despite being less reasonable, allow us to see

<table>
<thead>
<tr>
<th>Set</th>
<th>GT resonances</th>
<th>SD resonances</th>
<th>β-decay half-lives</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>(^{208}\text{Pb}, \ ^{112}\text{Sn}, \ ^{76}\text{Ge}, \ ^{150}\text{Te}, \ ^{90}\text{Zr}, \ ^{48}\text{Ca})</td>
<td>None</td>
<td>(^{48}\text{Ar}, \ ^{60}\text{Cr}, \ ^{72}\text{Ni}, \ ^{82}\text{Zn}, \ ^{92}\text{Kr}, \ ^{102}\text{Sr}, \ ^{112}\text{Ru}, \ ^{126}\text{Cd}, \ ^{134}\text{Sn}, \ ^{149}\text{Ba})</td>
</tr>
<tr>
<td>B</td>
<td>Same as A</td>
<td>None</td>
<td>(^{52}\text{Ti}, \ ^{74}\text{Zn}, \ ^{92}\text{Sr}, \ ^{114}\text{Pd}, \ ^{134}\text{Te}, \ ^{150}\text{Sm}, \ ^{180}\text{Yb}, \ ^{200}\text{Pt}, \ ^{228}\text{Rn}, \ ^{242}\text{U})</td>
</tr>
<tr>
<td>C</td>
<td>Same as A</td>
<td>None</td>
<td>(^{52}\text{Ti}, \ ^{74}\text{Zn}, \ ^{92}\text{Sr}, \ ^{114}\text{Ru}, \ ^{134}\text{Te}, \ ^{150}\text{Sm}, \ ^{180}\text{Yb}, \ ^{204}\text{Pt}, \ ^{226}\text{Rn}, \ ^{242}\text{U})</td>
</tr>
<tr>
<td>D</td>
<td>Those of A and (^{150}\text{Nd})</td>
<td>None</td>
<td>(^{58}\text{Ti}, \ ^{78}\text{Zn}, \ ^{98}\text{Kr}, \ ^{126}\text{Cd}, \ ^{152}\text{Ca}, \ ^{166}\text{Gd}, \ ^{204}\text{Pt})</td>
</tr>
<tr>
<td>E</td>
<td>Same as D</td>
<td>(^{90}\text{Zr}, \ ^{208}\text{Pb})</td>
<td>(^{58}\text{Ti}, \ ^{78}\text{Zn}, \ ^{98}\text{Kr}, \ ^{126}\text{Cd}, \ ^{152}\text{Ca}, \ ^{166}\text{Gd}, \ ^{226}\text{Rn})</td>
</tr>
</tbody>
</table>

TABLE I. The sets of fitting targets used in this work. The β-decay half-lives in set A range from 0.069 s (\(^{102}\text{Sr}\)) to 1.84 s (\(^{92}\text{Kr}\)), in set B from 95.6 s (\(^{74}\text{Zn}\)) to 45360 s (\(^{200}\text{Pt}\)), in set C from 0.54 s (\(^{114}\text{Ru}\)) to 9399.6 s (\(^{92}\text{Sr}\)), and in set E from 0.046 s (\(^{98}\text{Kr}\)) to 444 s (\(^{226}\text{Rn}\)). The nuclei selected for fitting the β-decay half-lives in sets D and E all exhibit an excitation spectrum clearly associated with either a spherical or a well-deformed shape; set E only consists of open-shell nuclei. The experimental GT resonance energies are from Refs. [36–40], the spin-dipole resonance energies are from Refs. [41,42], and the half-lives are from Ref. [43].
FIG. 2. Reproduction of target data in those of our fits that use the target set E. The four-parameter fits 3A and 3B yield almost the same results as the two-parameter fits 1D and 1E, with only a small decrease in the penalty function.

how sensitive our fits are to the selection of fitting targets). Set D contains only nuclei that are known to be rather rigid, with an excitation spectrum characteristic of a spherical or a well-deformed nucleus. (The QRPA, which is based on a single mean field, should work best in rigid nuclei). In set E, we include only open-shell rigid nuclei (for which isoscalar pairing should be most effective), swapping out $^{204}$Pt for $^{226}$Rn, and including two spin-dipole resonances. Figure 2 shows the quality of the fits for set E both with computed (1D) and experimental (1E) $Q$ values (see Table II for definitions of the number-letter combinations). The two procedures yield very similar results. The comparison in Fig. 3, which shows the results of the two-parameter fits (along with those of more-parameter fits discussed shortly) when the resulting functionals are applied to the set of all measured even-$\beta$-decay half-lives, shows that all these two-parameter fits (1A through 1E) yield the same level of predictive accuracy.

Can we do better by including some of the other time-odd coupling constants in our fit? To find out, we refit the four-parameter set $\{V_0, C_1^T, C_1^F, C_1^V\}$, which determines the Landau parameters $\{g_0^{\prime}, g_1^{\prime}, h_0\}$ and thus allows us to incorporate infinite-nuclear-matter stability conditions [27] as constraints. The parameter $C_1^F$ introduces a time-odd tensor term that is not present in the two-parameter fits. The result, however, improves the description of the fitting targets only marginally, as the points labeled 3A and 3B in Fig. 2 show, and actually worsens the agreement with half-life measurements overall (as Fig. 3 shows). The situation gets even worse when we use the results of this fit as a starting point to fit three more time-odd coupling constants, $\{C_1^T, C_1^V, C_1^{IV}\}$ (fit 4). Then the $\beta$-decay rates to which we fit are reproduced better, but the agreement with all measured rates deteriorates.

Improvement in the fitting targets accompanying a deterioration in overall agreement with data is a symptom of overfitting. To better understand why this happens, we evaluate the Jacobian matrix (9) at the parameter values produced by the two-parameter fit 1E. The Jacobian appears in Table III, with the values of the coupling constants in natural units following the prescription of Ref. [44] and the natural scale of isoscalar pairing taken to be the strength of isovector pairing. A clear column structure appears in both the resonance energies and the half-lives, signaling that the members of each individual set move largely in unison when the parameters are varied. Thus, there are essentially just two meaningful degrees of freedom that we can expect to fix with this experimental data: $V_0$ and $C_1^T$.

To see this in more detail, we carry out a singular value decomposition of the Jacobian. The largest singular value, 122.53, corresponds to a vector pointing nearly in the direction of $C_1^T$ in parameter space, and the second largest, 10.85, to a vector pointing nearly in the direction of $V_0$. The third-largest value, 1.648, is almost two orders of magnitude smaller than the largest, and corresponds mostly to $C_1^F$, with many other directions mixed in. The charge-changing data we have available—GT and spin-dipole resonances and half-lives—are not enough to reasonably constrain more than the parameters $V_0$ and $C_1^T$ in our initial fit.

Figure 3, besides containing the results of our fits, contains results from other work: Refs. [3,6,11] and [9]. Of all the these computations, the one by Homma et al. [3] seems to best reproduce the known $\beta^-$ half-lives, even though it neglects nonunique first-forbidden decay and uses simple separable interactions. As Fig. 4 shows, in our computation the nonunique $1^-$ contribution is quite important (even dominant) in many experimentally inaccessible nuclei, so it is far from clear how the various calculations will fare with data in the future. In any event, the most striking fact is that all the computations manage to reproduce existing data at roughly the same level of precision. It may not be possible to do much better without moving beyond Skyrme QRPA, at least while using a global parameter set, as we have done here.
TABLE II. Summary of the various fits in this work. The functional listed for each fit dictates the values of the coupling constants that are not fit, except for that of $C_i^{-1}$, which is set to zero everywhere to avoid finite-size instabilities. The units of $V_0$ and $C_i^{-1}$ are MeV fm$^3$, and the units of the other coupling constants are MeV fm$^3$.

<table>
<thead>
<tr>
<th>Fit</th>
<th>Starting point</th>
<th>Target set</th>
<th>$Q$ values</th>
<th>fitted parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>SKO'</td>
<td>A</td>
<td>Comp.</td>
<td>$V_0 = -173.176, C_1^{-1} = 128.279$</td>
</tr>
<tr>
<td>1B</td>
<td>SKO'</td>
<td>B</td>
<td>Comp.</td>
<td>$V_0 = -176.614, C_1^{-1} = 133.038$</td>
</tr>
<tr>
<td>1C</td>
<td>SKO'</td>
<td>C</td>
<td>Comp.</td>
<td>$V_0 = -176.097, C_1^{-1} = 126.966$</td>
</tr>
<tr>
<td>1D</td>
<td>SKO'</td>
<td>E</td>
<td>Comp.</td>
<td>$V_0 = -209.384, C_1^{-1} = 129.297$</td>
</tr>
<tr>
<td>1E</td>
<td>SKO'</td>
<td>E</td>
<td>Exp.</td>
<td>$V_0 = -159.397, C_1^{-1} = 99.8479$</td>
</tr>
<tr>
<td>2</td>
<td>SV-min</td>
<td>D</td>
<td>Comp.</td>
<td>$V_0 = -165.567, C_1^{-1} = 132.271$</td>
</tr>
<tr>
<td>3A</td>
<td>SKO'</td>
<td>E</td>
<td>Comp.</td>
<td>$V_0 = -195.174, C_1^{-1} = 144.833, C_2^{-1} = -20.1618, C_3^{-1} = -10.3125$</td>
</tr>
<tr>
<td>3B</td>
<td>SKO'</td>
<td>E</td>
<td>Exp.</td>
<td>$V_0 = -165.158, C_1^{-1} = 120.27, C_2^{-1} = -17.4345, C_3^{-1} = -17.9902$</td>
</tr>
<tr>
<td>4</td>
<td>Fit 3A</td>
<td>E</td>
<td>Comp.</td>
<td>$C_1^{-1} = 54.5, C_i^{-1} = -78.7965, C_i^{-1} = -87.5$</td>
</tr>
<tr>
<td>5</td>
<td>SkO'</td>
<td>E</td>
<td>Comp.</td>
<td>$V_0 = -191.875, C_1^{-1} = 146.182, C_i^{-1} = -86.4276$</td>
</tr>
</tbody>
</table>

B. Extrapolation to neutron-rich isotopes

Figure 4 displays the relative contribution to the decay rate from each multipole. Except in the immediate vicinity of the valley of stability, the changes appear quite gradual as a function of $Z$ and $N$. In nuclei with large $Q$ values, the details of single-particle structure are less important than in isotopes for which transitions to only a few low-energy states are possible.

Figure 4 also demonstrates the importance of going beyond the allowed approximation. In many heavy nuclei, the computed rates are dominated by the first-forbidden channel. Towards the drip line, both allowed and forbidden channels are important for all masses. The figure also shows that the nonunique $1^-$ channel is usually the most important of the forbidden multipole. Thus, any quenching of the (unique) $2^-$ channel and antiquenching of the (nonunique) $0^-$ channel from meson-exchange currents [45] would not have a significant impact on our overall results. In the $1^-$ channel the contributions of several different operators makes the effects of quenching hard to estimate.

In Fig. 5 we compare our half-lives to those of Möller et al. [5] in all medium and heavy even-even isotopes. Our half-lives tend to be longer than those of Ref. [5] close to the valley of stability in light nuclei and somewhat shorter in heavy nuclei with significant forbidden contributions. Approaching the neutron drip line, the two computations yield similar results up to a constant offset in those of Ref. [5] in even-even nuclei. All models can expect to do better near the drip line, where a significant fraction of the total $\beta$-decay strength can be below threshold.

Because our naive model for uncertainties is based on several assumptions that are only approximately correct or cannot easily be verified, we check its predictions where there are enough data to do so. Figure 6 shows the ratios of our half-lives to those of experiment together with the uncertainty model’s mean value and one- and two-standard-deviation

TABLE III. The Jacobian matrix, evaluated at the result of the two-parameter fit 1E. All parameters except for the strength of isoscalar pairing are expressed in natural units. The strength of isoscalar pairing has been scaled by the strength of isovector pairing. The derivatives of the $\log_{10}$ values are hence dimensionless and those of the resonance energies are in the units of MeV.

<table>
<thead>
<tr>
<th>$O$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dV_0$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dC_i^{-1}$</th>
<th>$dO/dC_i^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{208}$Pb $E_{GTR}$</td>
<td>57.261</td>
<td>-0.000</td>
<td>2.434</td>
<td>5.869</td>
<td>0.429</td>
<td>-1.002</td>
<td>0.000</td>
<td>0.143</td>
<td></td>
</tr>
<tr>
<td>$^{112}$Sn $E_{GTR}$</td>
<td>29.498</td>
<td>-1.032</td>
<td>1.432</td>
<td>2.863</td>
<td>0.286</td>
<td>-0.573</td>
<td>0.000</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>$^{76}$Ge $E_{GTR}$</td>
<td>45.115</td>
<td>-7.225</td>
<td>2.004</td>
<td>4.295</td>
<td>0.429</td>
<td>-1.145</td>
<td>0.000</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>$^{130}$Te $E_{GTR}$</td>
<td>53.790</td>
<td>-3.096</td>
<td>2.434</td>
<td>5.297</td>
<td>0.429</td>
<td>-1.002</td>
<td>0.143</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>$^{90}$Zr $E_{GTR}$</td>
<td>29.498</td>
<td>-1.032</td>
<td>1.288</td>
<td>2.720</td>
<td>0.429</td>
<td>-1.002</td>
<td>-0.143</td>
<td>0.143</td>
<td></td>
</tr>
<tr>
<td>$^{48}$Ca $E_{GTR}$</td>
<td>32.968</td>
<td>-0.000</td>
<td>1.432</td>
<td>3.149</td>
<td>0.573</td>
<td>-1.288</td>
<td>0.000</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>$^{208}$Pb $E_{PDR}$</td>
<td>52.055</td>
<td>-0.000</td>
<td>2.291</td>
<td>4.008</td>
<td>0.286</td>
<td>-1.575</td>
<td>-0.143</td>
<td>-0.143</td>
<td></td>
</tr>
<tr>
<td>$^{90}$Zr $E_{SBR}$</td>
<td>29.498</td>
<td>-0.000</td>
<td>1.575</td>
<td>2.004</td>
<td>0.286</td>
<td>-1.432</td>
<td>-0.286</td>
<td>-0.143</td>
<td></td>
</tr>
<tr>
<td>$^{80}$Tl $log_{10}$</td>
<td>0.14</td>
<td>-4.318</td>
<td>0.203</td>
<td>0.445</td>
<td>0.045</td>
<td>-0.109</td>
<td>-0.011</td>
<td>-0.002</td>
<td></td>
</tr>
<tr>
<td>$^{76}$Zn $log_{10}$</td>
<td>0.14</td>
<td>-2.922</td>
<td>0.256</td>
<td>0.589</td>
<td>0.164</td>
<td>-0.382</td>
<td>0.253</td>
<td>-0.025</td>
<td></td>
</tr>
<tr>
<td>$^{90}$Kr $log_{10}$</td>
<td>0.14</td>
<td>-3.252</td>
<td>0.265</td>
<td>0.599</td>
<td>0.050</td>
<td>-0.116</td>
<td>-0.012</td>
<td>-0.003</td>
<td></td>
</tr>
<tr>
<td>$^{126}$Cd $log_{10}$</td>
<td>0.14</td>
<td>-4.641</td>
<td>0.252</td>
<td>0.496</td>
<td>0.017</td>
<td>-0.050</td>
<td>0.001</td>
<td>0.007</td>
<td></td>
</tr>
<tr>
<td>$^{152}$Ce $log_{10}$</td>
<td>0.14</td>
<td>-2.474</td>
<td>0.293</td>
<td>0.540</td>
<td>0.051</td>
<td>-0.120</td>
<td>0.003</td>
<td>-0.009</td>
<td></td>
</tr>
<tr>
<td>$^{166}$Gd $log_{10}$</td>
<td>0.14</td>
<td>-2.924</td>
<td>0.250</td>
<td>0.497</td>
<td>0.035</td>
<td>-0.132</td>
<td>-0.007</td>
<td>-0.010</td>
<td></td>
</tr>
<tr>
<td>$^{208}$Pt $log_{10}$</td>
<td>0.14</td>
<td>-3.340</td>
<td>-0.015</td>
<td>0.160</td>
<td>-0.018</td>
<td>-0.316</td>
<td>-0.076</td>
<td>0.026</td>
<td></td>
</tr>
</tbody>
</table>
FIG. 3. Comparison of the mean and standard deviation of the log$_{10}$ values in our fits with those of previous work. The labels for our fits correspond to those of Table I. The results from prior work are contained in [5] (Mö), [3] (Ho), [6] (Na), [11] (Co), and [9] (Ma). Only even-even isotopes are considered.

FIG. 4. The contributions of different allowed and first-forbidden multipoles to the total computed $\beta$-decay rates. Only even-even nuclei are considered.
bands, all as a function of ground-state $Q$ value. We can discount the model at very low $Q$ but it appears to work well above $Q \approx 4$ MeV. Of the 72 nuclei, 48 (66.7%) fall within one standard deviation of the mean and 71 (98.6%) within two. These numbers are consistent with what one would expect from a normal distribution. The model quantifies our statement above that calculations are more accurate close to the drip line, where $Q$ is generally large.

A recent radioactive isotope beam factory measurement [46] of 110 neutron-rich isotopes, 40 of them previously unknown, allows us to test the reliability of our predictions and especially our model for theoretical uncertainties. Because the data are so recent, we did not include them in any of our fits, and hence we are effectively using older data to predict the results of these new measurements. We have 28 even-even nuclei with which to compare rates; for half of these there are earlier data in the ENSDF set (Fig. 7). Our predictions agree with experiment to within our theoretical uncertainty (though our error bars may be a bit too large here). Our uncertainty model thus appears to be reasonable.

V. CONCLUSIONS

We have explored the ability of the axially-deformed Skyrme QRPA to provide a global description of $\beta^-$-decay rates in even-even neutron-rich nuclei. With experimental rates and charge-exchange resonance energies as fitting targets we have found that among time-odd couplings, only those multiplying the isoscalar-pairing and the spin-density parts of the functional are well constrained; attempts to fit more than these two constants lead to overfitting. The tensor contributions to the EDF are, in particular, not well constrained by this data. To get more accurate Skyrme-QRPA predictions, one can resort to local fits, i.e., $A$-dependent couplings. The recent work of Ref. [9], for example, attaches a sensible $A$ dependence to the strength of isoscalar pairing. In addition, including neutron separation energies as additional fitting targets could help constrain the couplings somewhat better. Furthermore, including quenching of the FF operators might make a difference for heavier nuclei.

The level of agreement between our calculations and data throughout the isotopic chart is similar to that produced by other recent computations, in spite of our consistent inclusion of deformation, tensor terms in the functional, etc. It could be difficult to do much better without an account of multiphonon effects, which have been found to noticeably affect the GT strength distribution (e.g., in Ref. [47]) and consequently the $\beta^-$-decay rates in Ref. [48].

The most glaring shortcoming of our work here is the restriction to even-even nuclei. An extension of the FAM to odd-mass nuclei will be the subject of a future publication [49]. For the moment, we make our results for the 1387 even-even neutron-rich nuclei, with crudely estimated theoretical uncertainties, available as Supplemental Material to this article [50].

ACKNOWLEDGMENTS

We thank W. Nazarewicz, J. Dobaczewski, and S. Wild for useful discussions. Support for this work was provided through the Scientific Discovery through Advanced Computing (SciDAC) program funded by US Department of Energy,
Office of Science, Advanced Scientific Computing Research and Nuclear Physics, under Contract No. DE-SC0008641, ER41896. This work used the Extreme Science and Engineering Discovery Environment (XSEDE), which is supported by National Science Foundation Grant No. ACI-1053575.