High precision analytical description of the allowed β spectrum shape

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A fully analytical description of the allowed β spectrum shape is given in view of ongoing and planned measurements. Its study forms an invaluable tool in the search for physics beyond the standard electroweak model and the weak magnetism recoil term. Contributions stemming from finite size corrections, mass effects, and radiative corrections are reviewed. Particular focus is placed on atomic and chemical effects, where the existing description is extended and analytically provided. The effects of QCD-induced recoil terms are discussed, and cross-checks were performed for different theoretical formalisms. Special attention was given to a comparison of the treatment of nuclear structure effects in different formalisms. Corrections were derived for both Fermi and Gamow-Teller transitions, and methods of analytical evaluation thoroughly discussed. In its integrated form, calculated *f* values were in agreement with the most precise numerical results within the aimed for precision. The need for an accurate evaluation of weak magnetism contributions was stressed, and the possible significance of the oft-neglected induced pseudoscalar interaction was noted. Together with improved atomic corrections, an analytical description was presented of the allowed β spectrum shape accurate to a few parts in 10⁻⁴ down to 1 keV for low to medium *Z* nuclei, thereby extending the work by previous authors by nearly an order of magnitude.

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I. INTRODUCTION

The study of β decay played a pivotal role in uncovering the nature of the left-handed V-A weak interaction, and by extension the electroweak sector of the standard model (SM) (Weinberg, 2009), half a century ago. Over the following decades, the continuous development of new experimental techniques has kept research on β decay at the forefront of revealing the structure of the SM and its inner workings (Severijns, Beck, and Naviliat-Cuncic, 2006; Vos, Wilschut, and Timmermans, 2015). Probing an array of different observables, such as correlation coefficients or the $\mathcal{F}t$ value, allows for a sensitive investigation of several beyond standard model (BSM) influences. Deviations from pure V-A theory can occur as manifestations of exotic interactions, such as right-handed currents involving new heavy particles. Through investigation of sidereal variations of experimental observables it additionally allows for a study of Lorentz invariance violation (Noordmans, Wilschut, and Timmermans, 2013).

In the LHC era, competitive results can be extracted from $pp \rightarrow e + MET + X$ reaction channels (Khachatryan *et al.*, 2015). In the past decade, the development of so-called effective field theories has seen a tremendous amount of

work and interest where, through inclusion of a new physics scale $\Lambda_{BSM} \gg \Lambda_{LHC}$, both high- and low-energy experimental results can be interpreted in the same theoretical framework (Bhattacharya *et al.*, 2012; Cirigliano, González-Alonso, and Graesser, 2013). The Lee-Yang Hamiltonian (Lee and Yang, 1956) is generalized to a form where couplings with higher dimensional combinations of SM field operators are categorized according to their transformation properties; see, e.g., the work by Bhattacharya *et al.* (2012). Several comparative reviews have been presented in the past few years (Cirigliano, Gardner, and Holstein, 2013; Naviliat-Cuncic and González-Alonso, 2013; Holstein, 2014a, 2014b; Severijns, 2014).

Recently, renewed interest has arisen in the β spectrum (Kuzminov and Osetrova, 2000; Towner and Hardy, 2005; Dawson *et al.*, 2009; Loidl *et al.*, 2010; George *et al.*, 2014; Mougeot and Bisch, 2014; Severijns, 2014; Severin *et al.*, 2014; Mougeot, 2015; Huyan *et al.*, 2016) as a tool for precision measurements searching for exotic currents beyond the standard electroweak model and investigating QCD-induced form factors. The latter are related to the fact that the quark involved in β decay is not a free particle but is embedded in a nucleon. Beyond standard model scalar and tensor coupling constants appear in the mathematical description of the β spectrum shape via the so-called Fierz interference term (Jackson, Treiman, and Wyld, 1957)

$$b_{\text{Fierz}} \simeq \pm \frac{1}{1+\rho^2} \left[\text{Re}\left(\frac{C_s + C'_s}{C_V}\right) + \rho^2 \text{Re}\left(\frac{C_T + C'_T}{C_A}\right) \right], \quad (1)$$

with the upper (lower) sign for electron (positron) decay, respectively, and $C_{S,T}^{(t)}$ are the coupling constants for possible scalar- or tensor-type weak interactions. Further, $C_{V,A}^{(t)}$ are the coupling constants for the SM vector and axial-vector interactions and $\rho = C_A M_{GT} / C_V M_F$ is the ratio of the Gamow-Teller and Fermi fractions in the β decay, with M_{GT} and M_F being the respective nuclear matrix elements. For pure Fermi and Gamow-Teller transitions this reduces to, respectively,

$$b_{\text{Fierz},F} \simeq \pm \operatorname{Re}\left(\frac{C_S + C'_S}{C_V}\right),$$
 (2a)

$$b_{\text{Fierz},GT} \simeq \pm \text{Re}\left(\frac{C_T + C'_T}{C_A}\right),$$
 (2b)

i.e., searches for beyond standard model currents become independent (to first order) of the nuclear matrix elements. In order to improve existing limits on these scalar or tensor coupling constants (Severijns and Naviliat-Cuncic, 2011; Holstein, 2014a; Vos, Wilschut, and Timmermans, 2015) a precision of typically 0.5% is required when determining b_{Fierz} (Severijns and Naviliat-Cuncic, 2013).

As the required experimental precision increases, QCDinduced form factors have to be taken into account in order not to limit the sensitivity to BSM physics (Pitcairn *et al.*, 2009; Wauters *et al.*, 2009, 2010; Severijns and Naviliat-Cuncic, 2013; Soti *et al.*, 2014; Sternberg *et al.*, 2015; Fenker, 2016). These form factors are part of the SM and are dominated by the so-called weak magnetism term $b_{\rm wm}$. Its inclusion is typically sufficient to be accurate below the 10^{-3} level, while higher-order terms can be incorporated should the need arise. Conversely, similar experimental precision has to be reached in order to test these corrections.

The Fierz and weak magnetism terms modify the shape of the β spectrum as follows:

$$N(W)dW \propto pW(W_0 - W)^2 \times \left(1 + \frac{\gamma m_e}{W} b_{\text{Fierz}} \pm \frac{4}{3} \frac{W}{M} b_{\text{wm}}\right) dW,$$
(3)

with *p*, *W*, and *W*₀ the *β* particle momentum, its total energy, and total energy at the spectrum end point, respectively. Additionally, $\gamma = \sqrt{1 - (\alpha Z)^2}$ with α the fine-structure constant, *Z* the atomic number of the daughter nucleus, and *m*_e and *M* the rest mass of the electron and the average mass of the mother and daughter nuclei, respectively. It is clear that, in order to search for new physics or when trying to establish the effect of weak magnetism, the *β* spectrum has to be sufficiently well understood theoretically in order to avoid other SM effects mimicking a nonzero Fierz term or behaving like a weak magnetism contribution. A description of the *β* spectrum reliable at the 10⁻⁴ level is then required.

In the past a detailed description of the β spectrum was provided by Behrens and collaborators (Bühring, 1963, 1965; Bühring and Schülke, 1965; Behrens and Jänecke, 1969; Behrens and Bühring, 1971, 1982; Behrens *et al.*, 1978), based on the formalism by Stech and Schülke (Schülke, 1964; Stech and Schülke, 1964), by Holstein (Holstein, 1971, 1972a, 1972b, 1974a, 1974c; Holstein, Shanahan, and Treiman, 1972; Holstein and Treiman, 1971;) and more recently by Wilkinson (1982, 1989, 1990, 1993a, 1993b, 1993c, 1995a, 1995b, 1997). The work by Behrens *et al.*, while rigorous and complete, relies on numerical solutions of the Dirac equation for the outgoing leptons, whereas that by Wilkinson provided analytical parametrizations of the dominant effects.

In this work special attention has been given to improve the analytical description of atomic effects. The theoretical work performed by Bühring concerning screening effects was combined with high precision atomic potentials to guarantee good behavior for all Z values. An analytical fit is proposed for the atomic exchange effect, based on state-of-the-art numerical calculations. Its contribution can easily exceed

20% in the lowest energy regions for higher Z, and so is not to be neglected. The effects of shakeup and shakeoff have been reviewed, as well as their influence on the aforementioned atomic effects. Because of the aim for precision in this work, molecular effects have been explored and discussed. Limited analytical work is presented which can act as a guideline for estimating the error associated with a neglect of these influences.

The effects of nuclear structure and spatial variations of the final state wave functions inside the nuclear volume have been reviewed and we propose a new correction combining the transparency of the Holstein formalism with the rigor of the Behrens-Bühring formalism. Bundling all this information, the analytical atomic β spectrum shape presented here is expected to be accurate at the few parts in 10⁴ level down to 1 keV for low to medium *Z* nuclei.

In the following, the β spectrum shape is discussed by first providing an overview of the full description in Sec. II. Several sections treat the different electromagnetic and kinematic corrections, discussing the Fermi function (Sec. III), followed by the effects of the finite size and mass of the nucleus (Sec. IV), radiative corrections (Sec. V), and finally different atomic and chemical effects (Sec. VII). Another important part deals with corrections related to nuclear structure (Sec. VI), where considerable attention has been given to its correct evaluation. To this end a comparison was made between different approaches. It is here the weak magnetism contribution is reviewed and complete expressions are given for both Fermi and Gamow-Teller decays. Significant attention was given to the correct evaluation of the matrix elements in the nuclear structure dependent terms. Finally, the precision required to search for scalar and tensor weak interactions and to study the effect of weak magnetism will be discussed. In the appendixes a comparison is given between the electromagnetic corrections in the Behrens-Bühring and Holstein formalisms and further elaborated on the correct evaluation of nuclear structure dependent effects.

II. COMPLETE EXPRESSION

Apart from the electromagnetic corrections to the β spectrum shape, several other smaller corrections are to be included when a precision at the 10^{-4} level is required. The detailed description of the allowed β spectrum shape, including these smaller corrections, is given by

$$N(W)dW = \frac{G_V^2 V_{ud}^2}{2\pi^3} F_0(Z, W) L_0(Z, W) U(Z, W) D_{\text{FS}}(Z, W, \beta_2) R(W, W_0) R_N(W, W_0, M) \times Q(Z, W) S(Z, W) X(Z, W) r(Z, W) C(Z, W) D_C(Z, W, \beta_2) p W(W_0 - W)^2 dW \equiv \frac{G_V^2 V_{ud}^2}{2\pi^3} K(Z, W, W_0, M) A(Z, W) C'(Z, W) p W(W_0 - W)^2 dW.$$
(4)

Here Z is the proton number of the daughter nucleus, $W = E/m_ec^2 + 1$ is the total β particle energy in units of the electron rest mass, W_0 is the total energy at the spectrum end point, $p = \sqrt{W^2 - 1}$ is the β particle momentum in units of $m_e c$, G_V is the vector coupling strength in nuclei, and $V_{ud}^2 = \cos^2 \theta_C$, with θ_C the Cabibbo angle, is the square of the *up-down* matrix element of the Cabibbo-Kobayashi-Maskawa quark-mixing matrix.

The factor $F_0(Z, W)$ is the point charge Fermi function that takes into account the Coulomb interaction between the β particle and the daughter nucleus. The product $L_0(Z, W)U(Z, W)D_{\rm FS}(Z, W, \beta_2)$ describes the required corrections to this Fermi function after evaluation at the origin, which depend on the size and shape of the daughter nucleus (Sec. IV). Whereas previous effects are electrostatic in origin, $R(W, W_0)$ takes into account radiative corrections calculated using quantum electrodynamics (QED) (Sec. V). Moving from an infinitely massive nucleus to one of finite mass introduces further kinematical corrections described by R_N and Q. All these factors are combined into the factor $K(Z, W, W_0, M)$. The nuclear decay occurs in an atomic environment, meaning additional atomic corrections have to be taken into account. Here S(Z, W) is the screening correction (Sec. VII.A), X(Z, W) takes into account the so-called atomic exchange effect (Sec. VII.B) while r(Z, W) accounts for the atomic mismatch (Sec. VII.D). These effects are combined into A(Z, W). Finally, the nuclear structuresensitive effects are written as C(Z, W), with D_C its corresponding nuclear deformation correction. These are extensively discussed in Sec. VI.

We comment here on the different effects encompassed by the name "finite size effects" used by different authors. For this, we must first realize the Fermi function comes about by extracting the electron amplitude at either the origin or the nuclear radius from the transition amplitude. We will perform the former in this work. As the nucleus is an object of finite size and the electron wave function is not a constant within this surface, this extraction requires corrections from convoluting its wave function with that of initial and final states. As the extracted Fermi function is typically written down in analytical form for a point charge through F_0 , this too requires corrections stemming from the finite size and shape of the daughter nucleus. We call these effects "electrostatic finite size" corrections in order to clearly distinguish their origin and describe them mathematically through L_0 , U, and D_{FS} . This simply amounts to the extraction of a more correct electron wave function evaluated at the origin. We still require a convolution of the correct wave function through the nuclear volume via initial and final nuclear states contributing to the decay. This involves a convolution with all relevant operators contributing to the decay, which we do not artificially separate but write completely as C. As this depends on the electron wave function behavior inside the nucleus, Coulomb effects are present in the calculation thereof. In the approach by Holstein (1974b), Calaprice and Holstein (1976), and others these are artificially separated into nuclear structure and Coulomb sensitive factors when describing the spectral functions. Together with the electrostatic finite size effects previously defined, these are collectively called finite size corrections. In the works inspired by Behrens and Bühring (1982), on the other hand, only the part involving the leptonic convolution is typically referred to as the finite size correction. Others still refer to only electrostatic finite size effects. By specifying the electrostatic origin of these corrections, we hope to put these confusions to rest. As the nuclear structuresensitive correction C is nonzero even for point nuclei, we refrain from calling these finite size effects altogether even though we recognize the finite nuclear wave function influences these results. Appendix E, in particular, aims to further discuss the overlap and differences in the different formalisms commonly found in the literature.

In the entire treatment natural units suited for β decay are used, i.e., $c = \hbar = m_e = 1$. In all formulas presented we define Z as a positive quantity and distinguish between β^- and β^+ explicitly unless mentioned otherwise. When estimates of the magnitude of different corrections are given, it represents the relative change in the phase space integral after inclusion of the effect unless mentioned otherwise. As the detailed description of β decay was developed over a period of several decades with several different formalisms being used, we also attempt here to relate different theoretical results and trace their origin in order to avoid double counting issues. The appendixes further elaborate on this idea.

Finally, all formulas presented have been implemented in a custom C++ program (Hayen and Severijns, 2018). Based on simple configuration files describing the decay parameters and basic nuclear properties, the allowed β spectrum shape and corresponding (anti)neutrino spectrum can be automatically calculated. This includes the nuclear structure-sensitive terms, allowing for its evaluation in the extreme single-particle approximation as well as through connections with a shell model of mean-field software packages. The former approach is shown here to work very well once one moves to a properly deformed potential rather than a spherical harmonic oscillator potential. Results specific to nuclear structure related parameters utilizing this code are discussed elsewhere (Severijns *et al.*, 2018).

III. FERMI FUNCTION

The Hamiltonian that governs all types of β decay must include not only the weak interaction responsible for the actual decay, but also the electromagnetic interaction of the β particle with its surroundings. As the latter is much stronger than the former, it cannot be treated in perturbation theory (Halpern, 1970; Holstein, 1979). Writing down the transition matrix element to first order in the weak Hamiltonian we find

$$M_{fi} = -2\pi i \delta(E_f - E_i) \langle f | T \left[\exp\left(-i \int_0^\infty dt \mathcal{H}^Z(t)\right) \right] \\ \times \mathcal{H}_\beta(0) T \left[\exp\left(-i \int_{-\infty}^0 dt \mathcal{H}^{Z'}(t)\right) \right] | i \rangle$$
(5)

with *T* ensuring a time-ordered product and Z'(Z) the charge of the mother (daughter). The typical approximation made in dealing with β decay equates initial and final Coulomb interactions, replacing it instead with only the final. This is then corrected for by including radiative corrections and atomic final state influences discussed later. We choose to follow this approach, such that Eq. (5) corresponds to using the solution of the Dirac equation with an electrostatic potential rather than a plane wave for the electron (Roman, 1965). It is this change that requires the introduction of the socalled Fermi function. Denoted by F(Z, W), it takes into account the distortion of the electron radial wave function by the nuclear charge, i.e., the Coulomb interaction between the β particle and the daughter nucleus. Many (Konopinski and Uhlenbeck, 1941; Behrens and Jänecke, 1969) start from the point charge Fermi function introduced by Fermi (1934).¹

$$F_0(Z, W) = 4(2pR)^{2(\gamma-1)} e^{\pi y} \frac{|\Gamma(\gamma + iy)|^2}{[\Gamma(1+2\gamma)]^2},$$
 (6)

with

$$\gamma = \sqrt{1 - (\alpha Z)^2}, \qquad y = \pm \alpha Z W/p.$$
 (7)

Here R is the cutoff radius in the evaluation of the Dirac equation for the electron or positron necessitated by its slight divergence at the origin for a fictitious point charge. This cutoff radius is taken to represent the radius of the daughter nucleus, although it has in fact no real physical significance.

Throughout the vast literature of β decay many different definitions of the Fermi function have appeared. We attempt to provide a short overview of the most frequently used and stress that no formulation is in itself superior. Each requires additional corrections stemming from the finite size of the nucleus and its precise shape. These have been estimated to various orders of precision and are discussed in the next sections. This is not to say the Fermi function as written, for example, in Eq. (6) carries no merit. It absorbs most of the Z and W dependence of the tree-order Coulomb interaction, whereas additional corrections are typically limited to at most a few percent.

Before we summarize the different Fermi functions, we briefly discuss the relativistic solution of an electron inside a spherical Coulomb potential. The seminal work by Rose (1961) elaborates on a fully analytical solution of the Dirac equation for an electron in a hydrogenic approximation. In general the solution can be written as

$$\Psi_{\kappa} = \begin{pmatrix} \operatorname{sign}(\kappa) f_{\kappa}(r) \sum_{\mu} \chi^{\mu}_{-\kappa} \\ g_{\kappa}(r) \sum_{\mu} \chi^{\mu}_{\kappa} \end{pmatrix}, \qquad (8)$$

with χ_{κ}^{μ} the traditional spin-angular functions, and κ the eigenvalue of the operator² $K = \beta(\boldsymbol{\sigma} \cdot \boldsymbol{L} + 1)$. For allowed decay all states with j = 1/2 contribute, meaning $\kappa = -1$ for $s_{1/2}$ and $\kappa = 1$ for $p_{1/2}$ orbitals. All Coulombic information is then encoded in the radial wave functions $f_{\kappa}(r)$ and $g_{\kappa}(r)$. When calculating the transition matrix element, one typically extracts the dominant terms of the square of the electron wave function throughout the nucleus, evaluated at some radius *r*, to define³

$$F(Z, W, r) = \frac{f_1^2(r) + g_{-1}^2(r)}{2p^2}.$$
(9)

Historically, this has been evaluated either at the origin or at the nuclear radius. Using the analytical solutions of the Dirac equation in a central Coulomb potential, one finds the factor 4 in Eq. (6) replaced by $(1 + \gamma)/2$. This was noted by Konopinski and Uhlenbeck (1941) and has since been adopted by Blatt and Weisskopf (1952), Fano (1952), Konopinski (1966), and Wilkinson (1989). Others (Greuling, 1942; Dismuke *et al.*, 1952), including the seminal textbooks of Schopper (1966) and Behrens and Bühring (1982), adhere to the formulation of Eq. (6). The factor $(1 + \gamma)/2$ is then absorbed into an L_0 correction, which is also responsible for modifications due to the finite nuclear size. To remain consistent with the latter works, we opted to do the same and use Eq. (6) as our Fermi function and discuss among others the L_0 correction factor in the following sections.

Additional definitions of the Fermi function have attempted to extend Eq. (6) by including finite size corrections, through evaluation at the origin or both.⁴ A frequent formulation that performs both of these is that by Behrens and Jänecke (1969)

$$F_{BJ}(Z,W) = F_0(Z,W) \left(1 \mp \frac{13}{15} \alpha ZWR + \cdots \right), \quad (10)$$

discussed by Calaprice and Holstein (1976). The additional term can easily be traced from the power expansions of f_1 and g_{-1} inside the nucleus by (Huffaker and Laird, 1967)

$$\begin{cases} g_{-1}(r) \\ f_{1}(r) \end{cases} \propto [p^{2}F_{0}(Z,W)]^{1/2} \\ \times \left\{ 1 - \left[\frac{13}{30} + \frac{1}{2}\left(\frac{r}{R}\right)^{2}\right] \times \alpha ZWR - \frac{1}{6}(pr)^{2} \right\}$$
(11)

and understood as a rough electrostatic finite size correction. The additional correction stemming from the convolution of the leptonic wave functions with the weak charge distribution then brings about nuclear structure-sensitive $\langle r^2 \rangle$ terms. Calaprice and Holstein (1976) treated this with slightly higher accuracy. Since the expansion performed by Huffaker and Laird (1967) neglects terms of the order of $(\alpha Z)^2$, the aforementioned factor $(1 + \gamma)/2$ is absent. The error introduced by this is of the order of 0.5% for Z = 20.

As mentioned, we split up all corrections in the current work. The finite size corrections are split up into an *electrostatic* finite size correction, stemming from the electric potential difference, and a *convolution* finite size correction from an integration over the nuclear volume and sensitive to nuclear structure. A discussion of the latter is postponed until Sec. VI.

¹An English summary of the Fermi theory was presented by Konopinski and Uhlenbeck (1935) and Konopinski (1943).

²Here 1 is a 4 × 4 unit matrix, $\vec{\sigma}$ stands for the Pauli matrices in four dimensions, \vec{L} is the orbital angular momentum operator, and β is the equal to the γ_0 Dirac matrix. For *s* orbitals this evaluates to $\kappa = -1$.

³Additional corrections stemming from the "small" terms f_{-1} and g_1 can be written as a multiplication of Eq. (9) and some power expansion in pr and αZ and are included in the convolution finite size correction discussed in Sec. VI.

⁴Tabulations by Bhalla and Rose (1961) and Bhalla (1964) have been shown to contain errors for β^+ decay and have issues with double counting (Huffaker and Laird, 1967), thereby requiring a large correction (Calaprice and Holstein, 1976). We therefore do not consider the Bhalla-Rose Fermi function.

In order to facilitate the discussion of the correction terms on Eq. (6), we take Eq. (9) one step further. The results obtained by Fermi (1934) and Konopinski and Uhlenbeck (1941)) used analytical solutions of the Dirac equation in a simple Coulomb potential for a point charge. Once we move away from this simple picture to include the finite nuclear size or atomic screening, no such analytical solutions are available. Even when this is impossible, however, the behavior of the electron radial wave functions can be developed in a power expansion close to the origin (Behrens and Bühring, 1971)

$$\begin{cases} f_{\kappa}(r) \\ g_{\kappa}(r) \end{cases} = \alpha_{\kappa} \{ (2|\kappa|-1)!! \}^{-1} (pr)^{|\kappa|-1} \times \sum_{n=0}^{\infty} \begin{cases} a_{\kappa n} \\ b_{\kappa n} \end{cases} r^{n}, \quad (12)$$

where the electrostatic information is now contained in the socalled "Coulomb amplitudes" α_{κ} , and $a_{\kappa n}$, $b_{\kappa n}$ are iteratively defined parameters. In this picture, we are not any more hampered in evaluating the electron wave function at the origin and write instead (Behrens and Bühring, 1982)

$$F_0 L_0 = \frac{\alpha_{-1}^2 + \alpha_1^2}{2p^2}.$$
 (13)

This result is valid, regardless of the charge distribution of the nucleus even though only a point charge solution is analytically solvable.

IV. FINITE MASS AND ELECTROSTATIC FINITE SIZE EFFECTS

Attributing a finite mass to the nucleas opens up the decay kinematics from a two-body to a three-body process. This has consequences for both the outgoing energy of the lepton fields and the electromagnetic corrections where one considered the nuclear Coulomb potential as static. Giving the nucleus now a finite size introduces several more corrections on the S matrix as its nuclear volume is not any more a simple delta function. As discussed in the previous section the traditional Fermi function is calculated based on an infinitely heavy pointcharge model of the nucleus. As a consequence the electron radial wave functions diverge slightly at the origin and are instead evaluated at the nuclear radius R. This requires several corrections, stemming from the finite size of the nucleus. We stress again the ambiguity in the available literature when discussing the so-called finite size corrections. For this reason, we explicitly separated this into electrostatic and convolution parts. The former originates from a difference in the electrostatic potential when moving to a more realistic nuclear shape, whereas the latter takes into account the integration of the leptonic wave functions throughout the nuclear volume and includes the small Coulomb terms f_{-1} and g_1 . We focus here on the electrostatic finite size corrections and leave the convolution part for Sec. VI.

A. $L_0(Z,W)$, U(Z,W), and $D_{FS}(Z,W,\beta_2)$: Electrostatic finite nuclear size corrections

The deviation from a point-charge distribution inherently changes the electron density inside the nucleus and as such introduces additional electromagnetic corrections to the Fermi function. We discuss the consequence on Eq. (13) from moving to a uniformly charged sphere through L_0 , and go even further to a Fermi distribution in U. Wilkinson (1990, 1993b) discussed the former effects in detail and provided analytical expressions for each of them. Finally, we discuss the electrostatic effect of nuclear deformation $D_{\rm FS}$.

1. $L_0(Z, W)$

When considering instead of a point nucleus one of finite size, the electron and positron wave functions become finite at the center of the nucleus. Any other description but a point charge is, however, not analytically solvable, and so an approximate correction factor $L_0(Z, W)$ is introduced to be used in combination with the analytical Fermi function. In order to remain consistent with the formulation of the original Fermi function in Eq. (6), the nucleus is presented by a uniformly charged sphere of radius *R* that is adjusted to give the experimental $\langle r^2 \rangle^{1/2}$ of the daughter nucleus, i.e.,

$$R = \sqrt{5/3} \langle r^2 \rangle^{1/2}.$$
 (14)

According to Eq. (13), L_0 is defined as embracing all information that is not contained in the Fermi function of Eq. (6) when evaluating the dominant electron components at the origin. As discussed in the previous section, this entails the fact that even when considering a point charge, L_0 differs from unity since $L_0 \sim (1 + \gamma)/2$. Analogous to the expansion of f_1 and g_{-1} in Eq. (11), Behrens and Bühring (1982) performed an expansion to higher order [see Eqs. (12) and (13)] and found after a straightforward calculation

$$L_0 \simeq \frac{1+\gamma}{2} \left[1 \mp \alpha ZWR + \frac{7}{15} (\alpha Z^2) - \frac{1}{2} \gamma \frac{\alpha ZR}{W} \right]$$
$$\simeq 1 \mp \alpha ZWR + \frac{13}{60} (\alpha Z^2) - \frac{1}{2} \frac{\alpha ZR}{W}$$
(15)

for *Z* sufficiently low. Here $1 - \gamma$ was approximated as $\frac{1}{2}(\alpha Z^2)$ and γ as unity. More precise numerical calculations were tabulated by Behrens and Jänecke (1969) using R = $1.20A^{1/2}$ fm close to the valley of stability. This was extended by Wilkinson (1990) to cover all isotopes between the proton and neutron drip lines for $Z \leq 60$. Together with a slight generalization of the prefactors in Eq. (15) (keeping terms proportional to γ alive rather than approximate them to unity), numerical results were fitted after inclusion of simple power expansions. The result was presented in analytical form as

$$L_0(Z, W) = 1 + \frac{13}{60} (\alpha Z)^2 \mp \frac{\alpha ZWR(41 - 26\gamma)}{[15(2\gamma - 1)]}$$

$$\mp \frac{\alpha ZR\gamma(17 - 2\gamma)}{[30W(2\gamma - 1)]} + a_{-1}\frac{R}{W} + \sum_{n=0}^5 a_n (WR)^n$$

$$+ 0.41(R - 0.0164)(\alpha Z)^{4.5}, \qquad (16)$$

for electrons, while for positrons the 0.41 in the last term of this equation is to be replaced by 0.22. The a_n values (n = -1, 0, 1, ..., 5) are given by the parametrization

TABLE I. Coefficients for the parametrization of $L_0(Z, W)$ for electrons. Reproduction of Table 1 in Wilkinson (1990).

	b_1	b_2	b_3	b_4	b_5	b_6
a_{-1}	0.115	-1.8123	8.2498	-11.223	-14.854	32.086
a_0	-0.00062	0.007 165	0.018 41	-0.53736	1.2691	-1.5467
a_1	0.024 82	-0.5975	4.841 99	-15.3374	23.9774	-12.6534
a_2	-0.14038	3.649 53	-38.8143	172.1368	-346.708	288.7873
a_3	0.008 152	-1.15664	49.9663	-273.711	657.6292	-603.7033
a_4	1.214 5	-23.9931	149.9718	-471.2985	662.1909	-305.6804
<i>a</i> ₅	-1.563 2	33.419 2	-255.1333	938.5297	-1641.2845	1095.358

TABLE II. Coefficients for the parametrization of $L_0(Z, W)$ for positrons. Reproduction of Table I in Wilkinson (1990), with small modifications as discussed in the text. The signs used in odd powers of Z have been flipped to agree with the convention used in this work.

	b_1	b_2	b_3	b_4	b_5	b_6
a_{-1}	0.070 1	-2.572	27.597 1	-128.658	272.264	-214.925
a_0	-0.002308	0.066 463	-0.6407	2.63606	-5.6317	4.0011
a_1	0.079 36	-2.09284	18.454 62	-80.9375	160.8384	-124.8927
a_2	-0.938 32	22.025 13	-197.00221	807.1878	-1566.6077	1156.3287
a_3	4.276 181	-96.82411	835.265 05	-3355.8441	6411.3255	-4681.573
a_4	-8.2135	179.0862	-1492.1295	5872.5362	-11038.7299	7963.4701
<i>a</i> ₅	5.458 3	-115.8922	940.830 5	-3633.9181	6727.6296	-4795.0481

$$a_n = \sum_{x=1}^{6} b_{x,n} (\alpha Z)^x,$$
(17)

with $b_{x,n}$ (x = 1, ..., 6) being listed in Tables 1 and 2 of Wilkinson (1990) for electrons and positrons, respectively, and reproduced here in Tables I and II.⁵ [Note that in Table II 0.066 483 under b_2 was replaced by 0.066 463 and 2.836 06 under b_4 was replaced by 2.636 06 (Wilkinson, 1993b).] These parametrizations yield $L_0(Z, W)$ accurately to 1 part in 10⁴ for $p \le 45$ and for $|Z| \le 60$. The effect of $L_0(Z, W)$ ranges from a few 0.1% up to several % and so is everywhere highly significant.

2. U(Z,W)

In the previous section we reported on the finite size correction when the nuclear shape was assumed to be uniformly spherical with a fixed radius R. In reality the nuclear charge distribution is smeared out over a certain distance. This in turn introduces an additional, smaller, correction term from replacing the uniform spherical charge distribution that defines $L_0(Z, W)$ with a more realistic one having the same $\langle r^2 \rangle^{1/2}$ (Behrens and Bühring, 1970, 1972). Throughout time several different distributions have been proposed, ranging from a simple Yukawa form to Fermi functions and even Gudermannian combinations (Asai and Ogata, 1974; Ogata and Asai, 1974). An extensive overview of oft-encountered charge distributions in nuclear physics is given by Andrae (2000), together with some basic properties. We explicitly consider here the modified Gaussian and Fermi potentials. The first is given by

$$\rho_{MG}(r) = N_0 \left\{ 1 + A \left(\frac{r}{2}\right)^2 \right\} e^{-(r/a)^2}, \tag{18}$$

where

$$N_0 = \frac{8}{2+3A} a^{-3} \pi^{-1/2} \tag{19}$$

is a normalization constant and *a* is constrained by $\langle r^2 \rangle = 3/5R^2$ to find

$$a = R[\frac{5}{2}(2+5A)/(2+3A)]^{-1/2}.$$
 (20)

Here A is a free parameter. The second is a basic Fermi distribution

$$\rho_F(r) \sim \{1 + \exp\left[(r-c)/a\right]\}^{-1}.$$
(21)

The former agrees well with charge distributions of low *p*-shell and *sd*-shell nuclei, while the latter more accurately resembles the shape of heavier nuclei.

We can now provide an analytical approximation of the induced effect by calculating the L_0 correction for different charge distributions for low Z. The calculation is analogous to that required to obtain Eq. (15). Canceling common normalization factors and constants, we find

$$\frac{L'_0}{L_0} = \frac{\alpha_1'^2 + \alpha_{-1}'^2}{\alpha_1^2 + \alpha_{-1}^2} \tag{22}$$

$$=\frac{B_{1}^{\prime-2}+B_{-1}^{\prime-2}}{B_{1}^{-2}+B_{-1}^{-2}},$$
(23)

where we neglected terms of the order of $\mathcal{O}(WR^2)$, and the primes stand for the more advanced charge distributions. Now $B_{+1}^{(\prime)}$ is given by

⁵Note that the signs of odd powers of *Z* have been flipped for positrons to remain consistent with our initial notation. Specifically, *Z* is always defined as a positive quantity, with the upper (lower) sign for β^- (β^+) decay.

$$B_{\pm 1}^{(\prime)} = \pm f_{\pm 1}^{I}(R)g_{02}(R) - g_{\pm 1}^{I}(R)f_{02}(R).$$
(24)

Here the superscript *I* denotes the wave functions inside the nucleus, and f_{02} and g_{02} are radial Coulomb functions that depend on the sign of κ . The functions inside the nucleus can be found through a series expansion of the solution according to Eq. (12) and matching powers in the Dirac equation. This gives a recursive relationship for a_n and b_n . The last quantity needed depends on the order of the potential. In the case of a uniformly charged sphere, i.e., a second order dependence, the large wave functions must be expanded up to n = 6. In general this is n = 2(m + 1), where *m* is the largest, relevant radial order of the potential. The radial expansion of the Coulomb functions f_{02} and g_{02} is known and can be plugged in. Expanding results up to order $\mathcal{O}(\alpha ZR)$, $\mathcal{O}((WR)^2)$, and $\mathcal{O}((\alpha Z)^2)$, we find after a tedious but straightforward calculation

$$U(Z, W) \equiv \frac{L'_0}{L_0}$$

$$\approx 1 + \alpha Z W R \Delta_1 + \frac{\gamma}{W} \alpha Z R \Delta_2$$

$$+ (\alpha Z)^2 \Delta_3 - (W R)^2 \Delta_4, \qquad (25)$$

where

$$\Delta_1 = \frac{4}{3}\Delta v_0 + \frac{17}{30}\Delta v_2 + \frac{25}{63}\Delta v_4, \tag{26a}$$

$$\Delta_2 = \frac{2}{3}\Delta v_0 + \frac{7}{12}\Delta v_2 + \frac{11}{63}\Delta v_4, \tag{26b}$$

$$\Delta_3 = \frac{1}{3}\Delta v_0^2 + \frac{1}{15}\Delta v_2^2 + \frac{1}{35}\Delta v_4^2 + \frac{1}{6}\Delta v_2 v_0 + \frac{1}{9}\Delta v_4 v_0 + \frac{1}{20}\Delta v_4 v_2 + \frac{1}{5}\Delta v_2 + \frac{1}{7}\Delta v_4, \quad (26c)$$

$$\Delta_4 = \frac{4}{3}\Delta v_0 + \frac{4}{5}\Delta v_2 + \frac{4}{7}\Delta v_4,$$
 (26d)

taking into account even-*r* potentials up to fourth order and defining $\Delta v_n = v'_n - v_n$. Here we define v_n through

$$V(r) = \sum_{n=0}^{\infty} V_n r^n = \sum_{n=0}^{\infty} \left(-\frac{\alpha Z}{R^{n+1}} \right) v_n r^n.$$
(27)

As an example, the electrostatic potential of the modified Gaussian distribution of Eq. (18) is

$$V(r) = -\alpha Z \left[\frac{\operatorname{erf}(r/a)}{r} - \frac{2A}{(2+3A)a\sqrt{\pi}} \exp[-(r/a)^2] \right],$$

where erf is the error function. Expanding all terms we find

$$v_0 = \sqrt{\frac{5}{2}} \frac{4(1+A)(2+5A)^{1/2}}{\sqrt{\pi}(2+3A)^{3/2}},$$
 (28a)

$$v_2 = -\frac{4}{3(3A+2)\sqrt{\pi}} \left(\frac{5(2+5A)}{2(2+3A)}\right)^{3/2},$$
 (28b)

$$v_4 = \frac{2 - 7A}{5(3A + 2)\sqrt{\pi}} \left(\frac{5(2 + 5A)}{2(2 + 3A)}\right)^{5/2},$$
 (28c)

compared to $v_0 = 3/2$, $v_2 = -1/2$, and $v_4 = 0$ for the uniformly charged sphere. For sufficiently low Z, Eq. (25) can be employed for any charge distribution.

For higher Z the contributions from higher-order terms neglected in Eq. (25) cannot be any more ignored, and computational methods must be employed. Wilkinson (1993b) has done this for the Fermi distribution of Eq. (21) with $a \approx 0.55$ fm as it agrees quite well with Hartree-Fock calculations and experimental data. The correction is then written as

$$U(Z, W) = 1 + \sum_{n=0}^{2} a_n p^n,$$
 (29)

where

$$a_0 = -5.6 \times 10^{-5} \mp 4.94 \times 10^{-5}Z + 6.23 \times 10^{-8}Z^2,$$

$$a_1 = 5.17 \times 10^{-6} \pm 2.517 \times 10^{-6}Z + 2.00 \times 10^{-8}Z^2,$$

$$a_2 = -9.17 \times 10^{-8} \pm 5.53 \times 10^{-9}Z + 1.25 \times 10^{-10}Z^2 \quad (30)$$

in natural units. The effect of U(Z, W) amounts to some 0.1% for medium-high masses and so cannot be neglected. As the precise shape of the nucleus remains model dependent, some uncertainty remains. Experimental electron scattering data have, however, been fitted and tabulated by De Vries, De Jager, and De Vries (1987), allowing one to make a very good estimate. Much work in mean-field theories has also been presented over the years (Anni and Co', 1995; Anni, Co', and Pellegrino, 1995; Möller *et al.*, 2016).

Interesting to note is the applicability of Eq. (25) when used in combination with Eq. (29), for example, in describing the correction due to the oft-used three-parameter Fermi function, also known as the "wine-bottle" distribution (Andrae, 2000; Towner and Hardy, 2015)

$$\rho(r) = \rho_0 (1 + w(r/c)^2) \{ 1 + \exp[(r - c)/a] \}^{-1}, \quad (31)$$

where *w* describes the central depression. As this distribution closely resembles that of the normal Fermi distribution for small *w*, Eq. (29) can be used as a first approximation, after which Eq. (25) describes the difference in two Fermi distributions. As the difference in v_n will be small, Eq. (25) remains relevant even for higher *Z*.

3. $D_{FS}(Z,W,\beta_2)$

As in the previous section wherein we considered the change in the Fermi function due to a more realistic, although still spherically symmetric, charge distribution, we must account for a possible nuclear deformation. We limit ourselves here to axially symmetric deformations, writing the surface of our ellipsoid as⁶

⁶This is merely an approximation, as our factor unity should be replaced by $a_0 = 1 - \beta_2^2/4\pi$ to conserve the total volume of the undistorted sphere to second order. This effect is negligible, however, and we continue with Eq. (32).

$$R(\theta, \phi) = R_0 [1 + \beta_2 Y_2^0(\theta, \phi)],$$
(32)

with β_2 the traditional measure of quadrupole deformation (Davidson, 1968; Eisenberg and Greiner, 1975). In the case $\beta_2 > 0$, the nucleus has a prolate shape while it is oblate for $\beta_2 < 0$. This deformation gives rise to a nonzero intrinsic electric quadrupole moment typically parametrized as

$$Q_0 = 3\sqrt{\frac{\alpha}{5\pi}} R_0^2 Z \beta_2 (1 + 0.16\beta_2)$$
(33)

in our natural units. This implies experimental β_2 values can be obtained from measured electric quadrupole measurements or from theoretical models such as the work by Möller *et al.* (2016). Following the approach of Wilkinson (1994), we define *a* and *b* to be the axes perpendicular and along the symmetry axis, respectively, such that b/a > 1 for prolate deformations. Combining Eq. (33) with Eq. (25) from Wilkinson (1994) we find

$$\frac{b}{a} = \sqrt{\frac{1+C}{1-C/2}} \tag{34}$$

with

$$C = \frac{5}{3} \sqrt{\frac{5}{\pi}} \beta_2 (1 + 0.16\beta_2).$$
(35)

Together with the requirement that

$$\langle r^2 \rangle = \frac{1}{5} (b^2 + 2a^2),$$
 (36)

a and *b* are uniquely determined. Defining an angle-averaged charge distribution as

$$\rho(r) = \begin{cases} \rho_0 & \text{for } 0 \le r \le a, \\ \rho_0 \left(1 - \frac{b}{r} \left[\frac{r^2 - a^2}{b^2 - a^2} \right]^{1/2} \right) & \text{for } a < r \le b, \end{cases}$$
(37)

where

$$\rho_0 = \frac{3}{4\pi} \frac{Z}{a^2 b},\tag{38}$$

we transitioned from a deformed to a spherically symmetric charge distribution with which we can continue.⁷ We define ρ_0 such that the total charge is equal to Z. Because of the expected smallness of the correction, we continue in an analytical fashion and consider the charge distribution as a continuous superposition of uniform spheres of radius *r* and density $d\rho/dr$. We then write a modified finite size correction



FIG. 1. Several examples of $D_{\rm FS}$ for different momenta, using $\beta_2 = 0.2$. Interesting to note is the reversal of the energy dependence when switching between β^- and β^+ decay. Important to note is that even though $D_{\rm FS}$ can grow to several parts per 10³, it has to be combined with D_C from Sec. VI.F, which decreases the overall effect.

$$L_0(Z, W, \beta_2)^* = \frac{4}{3} \pi R^{2(1-\gamma)} \\ \times \int_0^\infty r^3 \frac{d\rho}{dr} L_0(Z, W, r) r^{2(\gamma-1)} dr, \quad (39)$$

where $L_0(Z, W, r)$ denotes the use of Eq. (16) with *R* replaced by the continuous variable. In the case of a uniformly charged sphere we have $d\rho/dr = (3/4\pi R^3)\delta(R)$ with $\delta(x)$ the Dirac delta function, trivially satisfying Eq. (39). The ratio

$$D_{\rm FS}(Z, W, \beta) = \frac{L_0(Z, W, \beta_2)^*}{L_0(Z, W)}$$
(40)

then constitutes the deformed nuclear shape correction to the Fermi function. Figure 1 shows the magnitude of the effect for different Z at several momenta for a reasonably large deformation $\beta_2 = 0.2$. Important to note is that this has to be combined with the other deformation-dependent effects, discussed in the nuclear structure of Sec. VI.F. This decreases the overall effect and cannot be neglected.

B. $R_N(W, W_0, M)$ and Q(Z, W, M): Finite nuclear mass corrections

Up to now we approximated β decay as a two-body phase space, whereas it should in fact be a three-body phase space opened up by the recoil of the nucleus. It was thus previously assumed that the nucleus is infinitely massive. The deviation from an infinitely massive nucleus introduces a kinematic

⁷Equation (37) is valid for prolate deformations. For an oblate nucleus, one simply interchanges a and b in the r ranges and reverses the signs in the square brackets. We assume prolate deformations unless otherwise specified.

recoil correction R_N and a subsequent small electromagnetic correction Q. We briefly discuss both in part.

1. $R_N(W, W_0, M)$

The effect of the recoil after β decay of a nucleus of finite mass M ($M = AM_{nucleon} - B$, with $M_{nucleon} = 1837.4$ the mass of the nucleon in units of m_ec^2 , and B the binding energy) is to multiply the phase space by a factor⁸ $R_N(W, W_0, M)$ (Horowitz, Kofoed-Hansen, and Lindhard, 1948; Kofoed-Hansen, 1948; Shekhter, 1959; Wilkinson, 1982, 1990)

$$R_N(W, W_0, M) = 1 + r_0 + r_1/W + r_2W + r_3/W^2, \quad (41)$$

where for vector decay (Fermi transitions)

$$r_0^V = W_0^2 / (2M^2) - 11/(6M^2), \qquad (42a)$$

$$r_1^V = W_0 / (3M^2), \tag{42b}$$

$$r_2^V = 2/M - 4W_0/(3M^2), \tag{42c}$$

$$r_3^V = 16/(3M^2), \tag{42d}$$

and for axial decay (Gamow-Teller transitions)

$$r_0^A = -2W_0/(3M) - W_0^2/(6M^2) - 77/(18M^2), \qquad (43a)$$

$$r_1^A = -2/(3M) + 7W_0/(9M^2), \tag{43b}$$

$$r_2^A = 10/(3M) - 28W_0/(9M^2), (43c)$$

$$r_3^A = 88/(9M^2). \tag{43d}$$

This is a small effect of the order of 10^{-5} to 10^{-3} at most [see Fig. 1 by Wilkinson (1990)]. For mixed transitions these corrections can simply be added together with the appropriate weighting factors

$$\frac{1}{1+\rho^2}$$
 and $\frac{1}{1+\rho^{-2}}$ (44)

for Fermi and Gamow-Teller transitions, respectively, and with $\rho = C_A M_{GT} / C_V M_F$ as defined before.

2. Q(Z, W, M)

A final consequence of the finite nuclear mass and consequent recoil is a change in the Coulomb field in which the departing electron or positron moves. It is not fixed in space but is itself recoiling against the combined lepton momenta so that the field experienced by the β particle differs with time from what it would have been if the nucleus would not be recoiling, as is assumed for the Fermi function in Eq. (6). Wilkinson (1982, 1993b) calculated the effect of this retreat of the source of the Coulomb field from the combined lepton momenta for a pure vector (Fermi) transition. This can be generalized for mixed Fermi and Gamow-Teller transitions to (Wilkinson, 1982)

$$Q(Z, W, M) \simeq 1 \mp \frac{\pi \alpha Z}{M} \frac{1}{p} \left(1 + a \frac{W_0 - W}{3W} \right), \quad (45)$$

with *a* the $\beta - \nu$ correlation coefficient, $a = (1 - \rho^2/3)/(1 + \rho^2)$ (Jackson, Treiman, and Wyld, 1957). For pure vector transitions one has a = 1, while for pure Gamow-Teller transitions we find a = -1/3 (Jackson, Treiman, and Wyld, 1957). The size of this correction amounts at most to a few percent of the typical error in the phase space factor *f* due to the uncertainty in the Q_{EC} value and so is typically negligible, although Wilkinson retained it in the calculation of *f* values for the superallowed Fermi transitions (Wilkinson, 1993b).

V. RADIATIVE CORRECTIONS

We started this work with a discussion concerning the Fermi function, a consequence of the electrostatic interaction of the daughter nucleus with the departing β particle. Within the context of QED, the continuous exchange of photons is not the only radiative process occurring after the decay. These radiative corrections to the β spectrum shape have a long history (Berman, 1958; Källén, 1967) and took a leap forward with the work by Sirlin (1967). A distinction was made between "inner" and "outer" radiative corrections, where the former is sensitive to the actual underlying weak interaction whereas the latter carries some nuclear dependence. The inner corrections can be calculated to high precision with standard electroweak methods and are typically incorporated into effective coupling constants. The outer corrections, on the other hand, include energy-dependent terms and are our main concern here. Writing the uncorrected β spectrum as $d\Gamma_0/dW$ we have then

$$\frac{d\Gamma}{dW} = \frac{d\Gamma_0}{dW} (1 + \Delta_R^{V/A}) [1 + \delta_R(W, W_0)], \qquad (46)$$

where $\Delta_R^{V/A}$ and $\delta_R(W, W_0)$ stand for inner and outer radiative corrections, respectively. The former has played an important role in establishing the universality of the electroweak interaction through comparison of the decay strength of muon decay (Sirlin and Ferroglia, 2013). Its calculation was addressed twice by Marciano and Sirlin (1986, 2006), the second time with improved precision, leading to the value $\Delta_R^V = (2.361 \pm 0.038)\%$. For the axial-vector case the inner radiative correction is simply incorporated into the experimental value of g_A from neutron decay.

As mentioned, we are concerned here only with the outer radiative corrections, seeing as to how they are both energy and nucleus dependent. These corrections, contributing on top of those already included in the Fermi function, typically amount to a few percent and clearly cannot be neglected. Contributions are treated in a Feynman-diagram fashion and concern both virtual photon exchange as well as one or more real photons in the final state. Seeing as to how these are

⁸Note that we added the subscript *N* to this factor originally defined as $R(W, W_0, M)$ in Wilkinson (1989) to differentiate it from the radiative corrections.



FIG. 2. Feynman diagrams for the order α corrections. The final two diagrams display the inner bremsstrahlung corrections, discussed separately (Sec. V.C). This entails the fact that when the final state photon goes undetected or can be distinguished from the β particle, the spectrum of the latter is appropriately corrected for.

experimentally relevant due to their possible detection, we briefly elaborate on the so-called inner bremsstrahlung. As this section contains no new results, we limit ourselves to listing the results obtained through decades of intensive study by a series of different authors. Excellent reviews have been provided by Wilkinson (1995b, 1997), Towner and Hardy (2008), and Sirlin and Ferroglia (2013). For completeness, we briefly comment on the radiative corrections for the neutrino due to its relevance in the reactor neutrino oscillation studies.

A. Total spectral influence

We discuss here the energy-dependent part of the outer radiative correction to various orders in $Z^n \alpha^m$. Outside of the terms for which m = n which are already contained in the Fermi function, it was found that *m* must always be larger than *n* (Bég, Bernstein, and Sirlin, 1972). This is a crucial result, as $Z^2\alpha$ terms can easily exceed unity and overthrow the conserved vector current (CVC) hypothesis. Great attention has been given to the three lowest m = n + 1 terms, each written as δ_m . The outer radiative correction is typically expressed as

$$R(W, W_0) = 1 + \delta_R(W, W_0).$$
(47)

Previously (Wilkinson, 1982, 1997; Towner and Hardy, 2002; Hardy and Towner, 2005a, 2005b), δ'_R was simply the sum of δ_1 , δ_2 , and δ_3 . Recently, however, significant improvements have been made by Marciano and Sirlin (1986) and Czarnecki, Marciano, and Sirlin (2004), where the leading order contributions $\mathcal{O}(\alpha^n \ln^n(M_N/2W_0))$ have been summed via renormalization group analysis. This allows one to write (Towner and Hardy, 2008; Sirlin and Ferroglia, 2013)

$$1 + \delta_R = \left\{ 1 + \frac{\alpha}{2\pi} \left[g(W_0, W) - 3 \ln \frac{m_p}{2W_0} \right] \right\} \\ \times \{ L(2W_0, m_p) + \delta_2 + \delta_3 \},$$
(48)

where m_p is the proton mass,

$$L(2W_0, m_p) = 1.026725 \left[1 - \frac{2\alpha}{3\pi} \ln 2W_0 \right]^{9/4}, \quad (49)$$

and $g(W_0, W)$ is described later. We discuss each of the δ_m separately.

1. Order α correction

The lowest order radiative correction corresponds to one (virtual-)photon Feynman diagrams. This includes a renormalization of the weak vertex, electron, and proton propagators but also internal brehmsstrahlung, discussed next. The relevant Feynman diagrams are shown in Fig. 2. Because of the zero photon mass, both the renormalization and inner brems-strahlung processes individually create an infrared divergence which is canceled when both contributions are added. Usually δ_1 is noted with the well-known $g(W_0, W)$ function (Sirlin, 1967; Sirlin and Ferroglia, 2013)

$$g(W_0, W) = 3\ln(m_p) - \frac{3}{4} + \frac{4}{\beta}L_s\left(\frac{2\beta}{1+\beta}\right) + 4\left(\frac{\tanh^{-1}\beta}{\beta} - 1\right) \left[\frac{W_0 - W}{3W} - \frac{3}{2} + \ln[2(W_0 - W)]\right] + \frac{\tanh^{-1}\beta}{\beta} \left[2(1+\beta^2) + \frac{(W_0 - W)^2}{6W^2} - 4\tanh^{-1}\beta\right]$$
(50)

with \tanh^{-1} the inverse hyperbolic tangent function, M_N the nucleon mass, $\beta = p/W$, and

$$L_{s} = \int_{0}^{x} \frac{\ln(1-t)}{t} dt = -\sum_{k=1}^{k=\infty} \frac{x^{k}}{k^{2}} \equiv -\text{Li}_{2}(x), \quad (51)$$

the Spence function, also known as the dilogarithm. Its large W_0 limit is (Wilkinson, 1995b)

$$g(W_0 \to \infty, W) = \left[3\ln\left(\frac{M_N}{2W_0}\right) + \frac{81}{10} - \frac{4\pi^2}{3}\right],$$
 (52)

which is dominated by the first term. Equation (50) is universal in the sense that is the same for both electrons and positrons, and Fermi and Gamow-Teller decays independent of the nucleus. It is exact except for small terms of the order of $\mathcal{O}(\alpha(W/M) \ln(M/W))$ and $\mathcal{O}(\alpha q/M)$, where q is the momentum transferred to the (anti)neutrino.

Finally, we comment on a peculiar logarithmic divergence for $W = W_0$ in Eq. (50). In the integration over the phase space this is clearly not a problem, but it points to a possible shortcoming in the analysis. It was shown that it is related to the emission of soft real photons (Repko and Wu, 1983). Two possibilities have been proposed to remove this divergence due to its relevance in the end-point-sensitive tritium β decay. The first (Repko and Wu, 1983) proposes to sum these soft real photon contributions to all orders of perturbation theory, leading to the replacement

$$t(\beta)\ln(W_0 - W) \to (W_0 - W)^{t(\beta)} - 1,$$
 (53)

where

$$t(\beta) = \frac{2\alpha}{\pi} \left[\frac{\tanh^{-1}\beta}{\beta} - 1 \right].$$
 (54)

The change in the tritium β spectrum shape is negligible due to its low end point; however, for higher energies the change in the *ft* value can become several parts in 10⁴ and so cannot always be neglected. The second possibility of resolving the divergence takes into account the finite detector resolution (Gardner, Bernard, and Meißner, 2004). In the treatment by Sirlin, the outgoing β particle and γ are always distinguishable, which is equivalent to an energy resolution of zero. If we instead consider a finite energy resolution, the distinguishability of the real photon depends on its energy, implying the same conclusion to the β spectrum correction. This assumes an absorption of the γ particle, however, an effect which is completely negligible in most experimental setups. We thus use Eq. (53) in our current description.

2. Order $Z\alpha^2$ correction

Spurred on by the high precision measurements of superallowed Fermi decays, two pioneering papers were published by Jaus and Rasche (1970) and Jaus (1972) discussing the higher-order radiative corrections. In subsequent years some tension arose between numerical and analytical methods (Sirlin and Zucchini, 1986), until finally both approaches agreed after a correction in the former (Jaus and Rasche, 1987). Using the notation of the latter one writes

$$\delta_2(Z, W) = Z\alpha^2 \sum_{i=1}^4 \Delta_i(W).$$
(55)

The first three terms in the sum concern the graphs shown in Fig. 3, with Δ_3 coming specifically from the axial-vector component. All of these depend on the nucleus considered as was to be expected from any correction involving interaction with the nucleus. The explicit weak interaction dynamics is, however, contained in the inner radiative correction, and the terms in Eq. (55) depend only on the shape and radius of the charge distribution. The Feynman diagrams for the fourth term, on the other hand, are not explicitly shown, as they are nonleading and present the interaction of the photon with only the electron through vacuum polarization and renormalization of the β particle propagator.

The terms of leading order in the nucleon mass M_N called $Z\alpha^2\Delta_1$ in Eq. (55) are difficult to evaluate and depend on the nuclear form factor $F(q^2)$. Using the trivial identity F = 1 + (F - 1), Δ_1 was split up into $\Delta_1 = \Delta_1^0 + \Delta_1^F$, where



FIG. 3. Dominant Feynman diagrams for the order $Z\alpha^2$ corrections. The first of these already contains a correction present in the product $F_0\delta_1$ and has to be explicitly subtracted. Three more diagrams contribute to the order $\mathcal{O}(Z\alpha^2)$ correction, but are nondominant. These can be found in Sirlin and Zucchini (1986) and discuss the vacuum polarization of the brehmsstrahlung photon and the two possibilities for the renormalization of the electron propagator.

the first term is now energy dependent but nucleus independent and vice versa for the second. The first is then typically combined with that other "purely QED" term Δ_4 . Expressions have been derived for Δ_1^0 in both the nonrelativistic (assuming zero β particle momentum) and extreme-relativistic approximation [neglecting terms of $\mathcal{O}(W/M)$] so that (Sirlin and Zucchini, 1986)

$$\Delta_{1}^{0} + \Delta_{4} \stackrel{\text{NRA}}{=} \ln M_{N} - \frac{2}{3}\ln(2W) + \frac{35}{9} + \frac{\pi^{2}}{6} - 6\ln 2$$
$$\stackrel{\text{ERA}}{=} \ln M_{N} - \frac{5}{3}\ln(2W) + \frac{43}{18}.$$
(56)

Results based on the extreme-relativistic approximation were compared to the numerical results of Jaus and Rasche (1987) by Sirlin (1987) and a general good agreement was found. Differences in the decay rate for Z = 26 were found to be on the few 10⁻⁴ level. For higher Z then, we expect a better agreement using a proper interpolation between the nonrelativistic and extreme-relativistic results should this be needed. The remaining terms have been evaluated using different models for the charge distribution $\rho(r)$. We list here the integral forms together with their evaluation for a uniformly charged sphere. Normalizing ρ as $\int \rho(r)r^2 dr = 1$, we find (Sirlin, 1987)

$$\Delta_1^F = 1 - \gamma_E - 4\pi \int_0^\infty \rho(r) r^2 \ln(M_N r) dr - (8/M_N)$$
$$\times \int_0^\infty \rho(r) r [1 + \gamma_E + \ln(M_N r)] dr, \tag{57}$$

$$\Delta_2 = (4/M_N) \int_0^\infty \rho(r) r \left[1 - \frac{\pi}{4M_N r} \right],$$
(58)

$$\Delta_3 = \frac{8g_A g_M}{M_N} \int_0^\infty \rho(r) r \left[\gamma_E + \ln(M_N r) - \frac{1}{2} + \frac{\pi}{8M_N r} \right].$$
(59)

The other Δ_i have been evaluated using different models for the charge distribution $\rho(r)$. For the uniformly charged sphere of radius $R = (5/3\langle r^2 \rangle)^{1/2} = \sqrt{10}/\Lambda$, one has (Sirlin, 1987)

$$\Delta_{1}^{F} = \ln\left[\frac{\Lambda}{M_{N}}\right] - \kappa_{2} - \frac{3}{\sqrt{10}\pi}\frac{\Lambda}{M_{N}} \times \left[\frac{1}{2} + \gamma_{E} + \ln\sqrt{10} + \ln\left[\frac{M_{N}}{\Lambda}\right]\right],$$
(60)

$$\Delta_2 = \frac{3}{2\sqrt{10}\pi} \frac{\Lambda}{M_N} \left[1 - \frac{\pi}{2\sqrt{10}} \frac{\Lambda}{M_N} \right],\tag{61}$$

$$\Delta_3 = \frac{3}{\sqrt{10}\pi} g_A g_M \frac{\Lambda}{M_N} \left[\gamma_E - 1 + \ln\sqrt{10} + \ln\frac{M_N}{\Lambda} + \frac{\pi}{4\sqrt{10}} \frac{\Lambda}{M_N} \right],\tag{62}$$

with $\kappa_2 \equiv \gamma_E - 4/3 + \ln \sqrt{10} = 0.395$, $g_A = 1.270$, $g_M = 4.706$, and $\gamma_E \approx 0.5772$. Further, $\Lambda \equiv \sqrt{6}/\sqrt{\langle r^2 \rangle}$ with the rms nuclear charge radius $\langle r^2 \rangle$ in natural units, with Λ ranging from about 400 at A = 10 to about 160 at A = 250.

For the modified Gaussian model for the charge distribution $\rho(r)$ (Sirlin, 1987) more extended expressions are obtained. Both models give, however, similar results for the superallowed Fermi transitions ranging from ¹⁴O to ⁵⁴Co corresponding to differences in the averages $\langle \delta_2(E) \rangle$ over the energy spectrum of less than 10⁻⁴ (Sirlin, 1987). Using a Fermi distribution for heavier nuclei, the difference can then be expected to be even smaller.

3. Order $Z^2 \alpha^3$ correction

The reasoning for the order $Z^2 \alpha^3$ corrections is analogous to that discussed, with the relevant Feynman diagrams shown in Fig. 4. Technically its evaluation is extremely challenging, however, and Sirlin (1987) proposed only a so-called "heuristic" correction. This is written as

$$\delta_3^{he} = Z^2 \alpha^3 \left[a \ln\left(\frac{\Lambda}{W}\right) + bf(W) + \frac{4\pi}{3}g(W) - 0.649\ln(2W_0) \right],\tag{63}$$

where

$$a = \frac{\pi}{3} - \frac{3}{2\pi}$$
 [footnote 10 in Sirlin (1987)], (64)

$$b = \frac{4}{3\pi} \left(\frac{11}{4} - \gamma_E - \frac{\pi^2}{6} \right), \tag{65}$$

$$f(W) = \ln(2W) - 5/6,$$
 (66)

$$g(W) = \frac{1}{2} [\ln^2(R) - \ln^2(2W)] + \frac{5}{3} \ln(2RW).$$
 (67)



FIG. 4. Dominant Feynman diagrams for the order $Z^2 \alpha^3$ corrections.

4. Higher-order corrections

Because of the computational complexity, little is known about the higher-order corrections of the type $Z^m \alpha^{m+1}$. Based on the leading order behavior of the previously discussed terms, Wilkinson (1997) has put forward a very approximate estimate

$$\bar{\delta}_{Z^n \alpha^m} \approx Z^n \alpha^m K_{nm} \overline{\ln^{m-n} \frac{\Lambda}{W}},\tag{68}$$

and suggested for K_{nm} an average value of 0.50 from these results. If K_{nm} is given a fixed value the summed effect of all the higher-order terms is (Wilkinson, 1997)

$$\bar{\delta}_{\text{higher}} \approx \sum_{n=3}^{n=\infty} \bar{\delta}_{Z^n \alpha^{n+1}} = \bar{\delta}_{Z^3 \alpha^4} / (1 - Z\alpha), \tag{69}$$

with $\delta_{Z^3\alpha^4}$ evaluated using Eq. (68). It is clear these corrections become relevant for higher Z.

B. Neutrino radiative corrections

Even though the outgoing (anti)neutrino has no direct interaction with the surrounding electric field, it is indirectly influenced through virtual photon exchange and energy conservation from inner brehmsstrahlung. Because of the recent interest in the proper conversion of the cumulative β spectrum emerging from a nuclear reactor, these have been treated explicitly by Sirlin (2011). The former case is essentially unchanged relative to the e^{\mp} spectrum, after substitution of $W_e \rightarrow W_0 - W_{\nu}$ in all relevant quantities. The treatment of the internal bremsstrahlung differs, however, and the subsequent total $\mathcal{O}(\alpha)$ radiative correction is much smaller. One finds

$$R_{\nu}(Z, W, W_0) = 1 + \frac{\alpha}{2\pi} \left\{ 3\ln\left(\frac{m_p}{m_e}\right) + \frac{23}{4} - \frac{8}{\hat{\beta}} \text{Li}_2\left(\frac{2\hat{\beta}}{1+\hat{\beta}}\right) + 8\left(\frac{\tanh^{-1}\hat{\beta}}{\hat{\beta}} - 1\right)\ln\left(\frac{2\hat{W}\hat{\beta}}{m_e}\right) + 4\frac{\tanh^{-1}\hat{\beta}}{\hat{\beta}} \left[\frac{7+3\hat{\beta}^2}{8} - 2\tanh^{-1}\hat{\beta}\right] \right\},$$
(70)

where analogous to the electron we have

$$\hat{W} = W_0 - W_{\nu}, \qquad \hat{p} = \sqrt{\hat{W}^2 - m_e^2}, \qquad \hat{\beta} = \frac{\hat{p}}{\hat{W}}, \quad (71)$$

with W_{ν} the energy of the outgoing (anti)neutrino.

C. Radiative β decay: Internal bremsstrahlung

The radiative corrections previously discussed include the effects of inner bremsstrahlung, otherwise known as radiative β decay where an additional photon is created in the final state

$$n \to p + e^- + \bar{\nu}_e + \gamma,$$
 (72)

where the energy of the γ ray follows a continuous spectrum thereby reducing the β particle energy. This is typically understood as a two-step process, where an electron is first ejected with energy W', and subsequently emits a photon with energy ω . Classically, this can simply be understood from the sudden acceleration of the β particle during which it emits radiation. This effect has been extensively described by Bloch (1936) and Knipp and Uhlenbeck (1936) and is commonly referred to as the KUB theory. Using results from classical electrodynamics one can write the photon ejection probability as (Schopper, 1966)

$$\Phi(W,\omega) = \frac{\alpha p}{\pi \omega p'} \left\{ \frac{W^2 + W'^2}{W' p} \ln(W+p) - 2 \right\}, \quad (73)$$

with $W = W' - \omega$ and $p' = \sqrt{W'^2 - 1}$ its correspondingly redefined momentum. In order to find the photon spectrum we must average over all intermediate virtual states to find

$$S(\omega) = \int_{1+\omega}^{W_0} N(W') \Phi(W', \omega) dW', \qquad (74)$$

where ω is the final photon energy and N(W') is the β spectrum. Equations (73) and (74) immediately reveal the dependence on the end-point energy of the β transition and its rapid falloff toward higher photon energies.

In this initial approach Coulomb effects with the daughter nucleus were neglected, however, and all results were evaluated using plane waves. Nilsson (1956), Lewis and Ford (1957), and Spruch and Gold (1959) introduced this correction and obtained expressions for its energy spectrum and angular correlation. This roughly translates to introducing the Fermi function into Eq. (74). Felsner (1963) has produced more rigorous work on this and shows a much improved agreement with experimental data. Recent theoretical work was performed by Ivanov et al. (2014) describing the QED tree-level contributions to the internal bremsstrahlung spectrum of ³⁵S. This amounts to changing Eq. (73) by Eq. (16) in the aforementioned work, where additional terms of order $\mathcal{O}(\omega/W)$, $\mathcal{O}(\alpha^2 Z^2/W^2)$, and $\mathcal{O}(W/M)$ are introduced. Further research was performed specifically for the neutron to investigate the influence of recoil order terms and weak magnetism (Ivanov, Pitschmann, and Troitskaya, 2013; Ivanov et al., 2017). Even though its inclusion shows up stronger in the internal brehmsstrahlung compared to the regular branching ratio, its magnitude is still far below the currently available experimental precision.

Detection of this radiation reveals underlying weak interaction physics and is both correlated with the outgoing lepton momenta and carries a degree of circular polarization (Cutkosky, 1954; Berman, 1958; Kinoshita and Sirlin, 1959; Batkin and Churakova, 1992; Mao *et al.*, 2011). Consequently, its experimental detection has been an interesting observable for several decades (Boehm and Wu, 1954; Goldhaber, Grodzins, and Sunyar, 1957; Basavaraju, 1983; Budick, Chen, and Lin, 1992; Khalil, 2011a, 2011b; Singh and Dhaliwal, 2014, 2015). Recent experimental branching ratios for the neutron find BR_{$\beta\gamma$} ~ 3 × 10⁻³ (Bales *et al.*, 2016), and also for higher Z significant branching ratios were found such as for ³²P at 2×10^{-3} (Berenyi and Varga, 1969). It was studied intensively following electron capture, reviewed extensively by Bambynek *et al.* (1977). Its relevance in the analysis of the tritium β spectrum measurements has been theoretically discussed by Gardner, Bernard, and Meißner (2004) and continues to garner interest through the study of correlation parameters (Gardner and He, 2012, 2013) looking for *T* and *CP* violation.

VI. NUCLEAR STRUCTURE EFFECTS: THE SHAPE FACTOR

In the foregoing, nuclear structure effects have been ignored and we focused instead on electromagnetic corrections and kinematics. When considering the nucleus as a nontrivial system with a finite size, nuclear structure and spatial variations of leptonic wave functions become deeply intertwined. The exact treatment of this fact has undergone careful study by several authors and different formalisms over several decades (Konopinski and Uhlenbeck, 1941; Rose and Osborn, 1954a; Weidenmüller, 1961; Bühring, 1963; 1965; Huffaker and Greuling, 1963; Schülke, 1964; Bühring and Schülke, 1965; Konopinski, 1966; Schopper, 1966; Huffaker and Laird, 1967; Behrens and Jänecke, 1969; Behrens and Bühring, 1970, 1982; Blin-Stoyle, 1973; Morita, 1973; Holstein, 1974a, 1974b, 1974c; Calaprice, Chung, and Wildenthal, 1977; Kleppinger, Calaprice, and Holstein, 1977; Behrens et al., 1978). In essence, we have finally arrived at the heart of *nuclear* β decay. Our starting point lies in the elementary particle approach and from this we branch out into the Behrens-Bühring and Holstein formalisms. We utilize the accurate calculational machinery of the former and transform our results to the clean notation of the latter.

A. Introduction

The β decay Hamiltonian is traditionally constructed as a current-current interaction

$$H_{\beta}(x) = \frac{G\cos\theta_{C}}{\sqrt{2}} [J^{\dagger}_{\mu}(x)L^{\mu}(x) + \text{H.c.}], \qquad (75)$$

where θ_C is the Cabibbo angle and $J_{\mu}(x)$ and $L^{\mu}(x)$ are the nuclear and lepton currents, respectively. This is plugged into the *S* matrix of Eq. (5) as a first-order perturbation. As we discussed in Sec. III, the introduction of electromagnetism requires a change in the lepton current L^{μ} exchanging the β particle wave function. So the strong interaction requires a change in the nuclear current from its pure *V*-*A* shape. We have then the generalizations

$$J^{\dagger}_{\mu}(x) = \langle f | V_{\mu}(x) + A_{\mu}(x) | i \rangle, \qquad (76)$$

$$L^{\mu}(x) = i\bar{\phi}_{e}(x)\gamma^{\mu}(1+\gamma_{5})v_{\nu}(x), \qquad (77)$$

where ϕ_e is the solution of the β particle wave function in a Coulomb potential. The transition matrix element (TME) constructed from this Hamiltonian is constrained by angular momentum coupling rules for the spin transition $J_i^{\pi_i} \rightarrow J_f^{\pi_f}$,

allowing decays of different multipole orders K within the vector triangle (J_i, J_f, K) . It is natural then to write the TME as a sum over all possible K with complementary projection operators acting on both lepton and nuclear spaces. This approach allows for the definition of form factors F_K that absorb all the nuclear structure information. These form factors are a function of $q^2 \equiv (p_f - p_i)^2$, the only Lorentz invariant scalar available. With form factors the treatment can continue without any model dependence, which is also known as the elementary particle approach practiced by many (Kim and Primakoff, 1965; Armstrong and Kim, 1972b; Holstein, 1974a; Behrens and Bühring, 1982). Specifically this translates into an expression of the following type (Schülke, 1964; Schopper, 1966):

$$\langle f | V_{\mu} - A_{\mu} | i \rangle \propto \sum_{KM} \sum_{s,L=K-1} (-1)^{J_{f} - M_{f} + M} (-i)^{L} \\ \times \sqrt{4\pi} \sqrt{2J_{i} + 1} \begin{pmatrix} J_{f} & K & J_{i} \\ -M_{f} & M & M_{i} \end{pmatrix} \\ \times \frac{(qR)^{L}}{(2L+1)!!} F_{KLs}(q^{2}),$$
 (78)

where *K* corresponds to the multipole order of the transition and *K*, *L*, and *s* have to form a vector triangle. The form factors $F_{KLs}(q^2)$ contain the actual nuclear information and can be expanded as a function of $(qR)^2$. Since for β decay this quantity is very small, the expansion can typically be stopped after the first order. A similar expression is obtained in the Holstein formalism [see Holstein (1974a) for a discussion and comparison]. In the case of a nontrivial lepton current, a similar expansion⁹ is made such that the TME can be written down to different orders of precision. In the aforementioned formalisms, all information other than the phase space factor and Fermi function are typically grouped in a so-called shape factor C(Z, W):

$$N(W)dW \propto pW(W_0 - W)^2 F(Z, W)C(Z, W).$$
(79)

In some formalisms the Fermi function and/or the finite size effect is also included in C(Z, W). Here we, however, adhere to the notation in Eq. (79). Now C(Z, W) effectively includes a combination of two effects: The first is the spatial variation of the leptonic and nuclear wave functions inside the finite nuclear volume, the second is the effect of raw nuclear structure. It should be understood that all additional corrections discussed in the previous sections are not included in this original formulation and can be considered higher-order corrections. Before we continue with an analytical description, we discuss the nature and evaluation of the form factors $F_{KLs}(q^2)$.

B. Form factors

1. Introduction

The form factors describe rather generally the nuclear structure and allow for a model-independent analysis of β decay observables. These can then be deduced from experimental measurement, and in theory we could stop the discussion. Using the impulse approximation we can, however, continue and express these form factors in terms of nuclear matrix elements. This entails that we treat the nucleus as an assembly of noninteracting nucleons that all couple to the β decay Hamiltonian as if they were free particles. Generally speaking, we have then a collection of one-body operators O_{KLs} that can be written as

$$O_{KLs} = \sum_{\alpha,\beta} \langle \alpha | O_{KLs} | \beta \rangle a_{\alpha}^{\dagger} a_{\beta}, \qquad (80)$$

where a_{α}^{\dagger} creates a nucleon in state α , and a_{β} annihilates a nucleon in state β . We have then

$$\langle f|O_{KLs}|i\rangle = \sum_{\alpha,\beta} \langle \alpha|O_{KLs}|\beta\rangle \langle f|a^{\dagger}_{\alpha}a_{\beta}|i\rangle.$$
(81)

The last factors in this equation are called the one-body density matrix elements which can be calculated using the shell model. In special cases this formulation can be further simplified and allow for immediate analytical evaluation. An example will be given. After the brief discussion of the impulse approximation, we are able to explicitly write the form factors in terms of nuclear matrix elements [see Sec. 6.2 in Behrens and Bühring (1982) for a more thorough discussion]. As an example, consider Table III where the correspondence between form factor coefficients¹⁰ and nuclear matrix elements is shown, as well as the possible spin changes and "forbidden" transitions for which the form factor coefficient is identically zero. For simplicity, induced currents (see Sec. VI.B.2) have been neglected here. A more extensive overview of $F_{KLs}^{(n)}$ with selection rules can be found in Behrens and Jänecke (1969).

2. Induced currents

Before we continue with the explicit calculation of the shape factor C(Z, W), we address the issue of induced currents. Because the decaying nucleon sits inside a nuclear potential, influences from QCD seep into the weak vertex. Assuming the weak interaction to be purely *V*-*A*, several Lorentz invariant terms can be added that transform in the same way. For the simple neutron decay this is written in the Behrens-Bühring (BB) (Behrens and Bühring, 1982) and Holstein (HS) (Holstein, 1974b) formalisms as¹¹

⁹Nontrivial here means Coulomb corrected wave functions rather than simple free Dirac spinors. In this case an expansion can be made in terms of r, αZ , $m_e R$, and $W_e R$. This is discussed in the next section.

¹⁰These are related to the more general form factors via the expansion $F_{KLs}(q^2) = \sum_n A(n, L)(qR)^{2n} F_{KLs}^{(n)}$, where A(n, L) is a trivial prefactor. As $F_{KLs}^{(0)}$ and $F_{KLs}^{(1)}$ are typically of similar magnitude, it is clear that $F_{KLs}(q^2)$ is dominated by the former.

¹¹Here $\mathbf{q} = -i(\nabla_f - \nabla_i)$ when written in operator form. In the presence of an electromagnetic field gauge invariance requires $q_{\mu} \rightarrow \partial/\partial \chi_{\mu} \rightarrow \partial/\partial \chi_{\mu} - ieA_{\mu}$ where $A_{\mu} = (\mathbf{A}, i\phi)$ is the potential of the electromagnetic field; see Behrens and Bühring (1971) and Holstein (1974a).

TABLE III. Definitions of the nuclear matrix elements (Behrens *et al.*, 1978) for allowed transitions. Here the form factors are written in their symbolic integral notation. The possible spin changes are shown explicitly, together with "forbidden" transitions for which the form factor coefficient is automatically zero. Here α is constructed with the Pauli matrices, and σ on its off-diagonal elements. We have neglected second-class contributions and terms of order $\mathcal{O}(g_M/(M_N R)^2)$.

Form factor (BB)	Cartesian form	ΔJ	Forbidden	Туре	
$VF_{000}^{(0)}$	$+g_V\int 1$	0			
${}^{A}F_{101}^{(0)}$	$\mp g_A \int {oldsymbol \sigma}$	0,1	0-0	Allowed	
$VF_{000}^{(1)}$	$+g_V\int (\frac{r}{R})^2$	0			
${}^{A}F_{101}^{(1)}$	$\mp g_A \int \boldsymbol{\sigma}(\frac{r}{R})^2 \pm \frac{g_P}{2(M_N R)^2} \int \boldsymbol{\sigma}$	0,1	0 - 0	Main correction terms	
${}^{A}F_{121}^{(0)}$	$\mp g_A \frac{3}{\sqrt{2}} \int \frac{(\boldsymbol{\sigma} \boldsymbol{r}) \boldsymbol{r} - (1/3) \boldsymbol{\sigma} \cdot \boldsymbol{r}^2}{R^2} \pm \frac{g_P 5 \sqrt{2}}{(2M_N R)^2} \int \boldsymbol{\sigma}$	0,1	0 - 0	Main correction terms	
$VF_{011}^{(0)}$	$+g_V \int i \frac{\alpha r}{R}$	0			
${}^{V}F_{111}^{(0)}$	$-g_V\sqrt{\frac{3}{2}}\int \frac{\alpha \times \mathbf{r}}{R} - \frac{g_M - g_V}{2M_N R}\sqrt{3}\int \boldsymbol{\sigma}$	0,1	0 - 0	Relativistic correction terms	
${}^{A}F_{110}^{(0)}$	$\pm g_A \sqrt{3} \int \gamma_5 \frac{ir}{R} \pm \frac{g_P \sqrt{3}}{(2M_N R)^2} \left[W_0 R \pm \frac{6}{5} \alpha Z \right] \int \boldsymbol{\sigma}$	0,1	0 - 0		

$$J_{\mu}^{BB} = i \langle \bar{u}_{p} | C_{V} \gamma_{\mu} - f_{M} \sigma_{\mu\nu} q^{\nu} + i f_{S} q_{\mu} - \frac{C_{A}}{C_{V}} \gamma_{\mu} \gamma_{5}$$
$$- f_{T} \sigma_{\mu\nu} \gamma_{5} q^{\nu} + i f_{P} \gamma_{5} q_{\mu} | u_{n} \rangle, \qquad (82)$$

$$J_{\mu}^{\rm HS} = i \langle \bar{u}_{p} | g_{V} \gamma_{\mu} - \frac{g_{M} - g_{V}}{2M} \sigma_{\mu\nu} q^{\nu} + i \frac{g_{S}}{2M} q_{\mu} + g_{A} \gamma_{5} \gamma_{\mu} - \frac{g_{II}}{2M} \sigma_{\mu\nu} \gamma_{5} q^{\nu} + i \frac{g_{P}}{2M} \gamma_{5} q_{\mu} | u_{n} \rangle,$$
(83)

where ${}^{12}\sigma_{\mu\nu} = -(i/2)[\gamma_{\mu},\gamma_{\nu}]$ and all *C* and *g* are functions of q^2 . It is here the elementary particle treatment shines, as the entire formulation in terms of form factors can be retained simply by redefining them and including these induced currents (Bühring, 1965). The two approaches, while inherently different, are completely equivalent,¹³ and the redefinitions of the different $F_{KLs}^{(n)}$ can be found in Bühring and Schülke (1965) and Behrens and Bühring (1971, 1982). As an example ${}^{V}F_{000}^{(0)}$ now becomes

$${}^{V}F_{000}^{(0)} \to {}^{V}\mathcal{M}_{000}^{(0)} \pm \frac{f_{S}}{R} \left(W_{0}R \pm \frac{6}{5}\alpha Z \right) {}^{V}\mathcal{M}_{000}^{(0)}, \quad (84)$$

where \mathcal{M}_{KLs} are the actual nuclear matrix elements as written in the second column of Table III. When neglecting induced currents it is clear that both notations coincide. Just as the work of Behrens and collaborators continues with the more general formulation using ${}^{V/A}F_{KLs}(\mathbf{q}^2)$, it introduces Holstein general form factors. Here the total TME is expanded into a combination of form factors labeled *a*, *b*, *c*, *d*, *e*, *f*, *h*, *g*, and *j*_{2,3}, all of which have some nontrivial q^2 dependence¹⁴ (Holstein, 1971, 1974c; Holstein and Treiman, 1971). These are now the Fermi (*a*), weak magnetism (*b*), Gamow-Teller (*c*), induced tensor (*d*), induced scalar (*e*), and further higher-order corrections, respectively. When expanding in terms of q^2 the influence of higher-order terms can be neglected except for the leading Fermi and Gamow-Teller form factors written as

$$a(q^2) = a_1 + a_2 q^2 + \cdots,$$
(85)

$$c(q^2) = c_1 + c_2 q^2 + \cdots$$
 (86)

Each of these form factors is a combination of the BB form factors discussed previously, with the additional advantage that unlike the BB form factors these are manifestly covariant and more clearly reveal underlying symmetries.

Table IV lists these form factors in terms of the nuclear matrix elements which are listed in Table V, as predicted by the impulse approximation. These again have to be provided by the shell model unless we can treat them approximately as is explained in Sec. VI.D.

3. Validity of the impulse approximation

In the previous section we discussed the possibility of transforming the model-independent form factors to one-body matrix elements using the impulse approximation. The latter assumes that all nucleons in the nucleus interact with the weak vertex as if they were free, thereby neglecting many-body effects such as meson exchange and core polarization. Armstrong and Kim (1972a) showed that meson exchange need not even be invoked to show the breakdown using only the partially conserved axial current (PCAC). It comes that as no surprise that the impulse approximation breaks down once sufficient accuracy is required. For the vector matrix element

¹²The original results by Holstein are written using the conventions by Bjorken and Drell (1964), which differs in both the metric used and the sign of q. We have written Eq. (83) in the metric and notation by Behrens and Bühring (1982) to show the correspondence in the definition of the constants.

¹³This can be seen from Eqs. (7) and (8) in Holstein (1974c) and is thoroughly discussed in Sec. 9.2 in Behrens and Bühring (1982).

¹⁴A careful distinction must be made as the form factors are expanded as a function of 3-momentum \mathbf{q}^2 in the BB formalism, while it is performed with the 4-momentum $q^2 = W_0^2 - \mathbf{q}^2$ in that of Holstein.

TABLE IV. Summary of the a, e, b, c_1, c_2, d , and h form factors and their relation to the nuclear matrix elements defined in Table V. Here CVC and SCC refer to conserved vector current hypothesis and second-class currents (see Sec. VI.C), respectively. In this table the impulse approximation is given only to first order, and the relativistic matrix elements are neglected as is done by Holstein (1974c).

Form factor	Formula in impulse approximation	Remark	Туре
a e b	$a \cong g_V \mathcal{M}_F$ $e \cong g_V (\mathcal{M}_F \pm Ag_S)$ $b \cong A(g_M \mathcal{M}_{GT} + g_V \mathcal{M}_L)$	$g_V = 1$ (CVC) (Ademollo and Gatto, 1964) e = 0 (CVC, SCC) $g_M \cong 4.706$	Vector
$\begin{array}{c} c_1 \\ c_2 \end{array}$	$c_1 \cong g_A \mathcal{M}_{GT}$ $c_2 \cong \frac{1}{6} g_A [\mathcal{M}_{\sigma r^2} + \frac{1}{\sqrt{10}} \mathcal{M}_{1v}]$	$g_A \rightarrow g_{A,\text{eff}} = 1$ (Towner, 1987) $c_2 \sim R^2$	
d	$d \cong A(g_A \mathcal{M}_{\sigma L} \pm g_{II} \mathcal{M}_{GT})$ $= d^I + d^{II}$	$g_{II} \sim g_T \cong 0 \text{ (SCC)}$ $d^I = 0 \text{ (analog states)}$	Axial vector
<i>h</i>	$h \cong \frac{-2}{\sqrt{10}} M^2 g_A \mathcal{M}_{1y} - A^2 g_P \mathcal{M}_{GT}$	$g_{P,\text{free}} \approx -229 \rightarrow g_{P,\text{eff}} = \text{uncertain}$	

TABLE V. Definitions of the nuclear matrix elements by Calaprice *et al.* (1977). Here τ denotes the typical isospin ladder operator, and an explicit sum over all nucleons is included. Note that the BB matrix elements from Table III are dimensionless due to appropriate powers of *R*, unlike those of Holstein presented here.

Matrix element	Operator form
$\overline{\mathcal{M}_F}$	$\langle \beta \ \Sigma \tau_i^{\pm} \ lpha angle$
\mathcal{M}_{GT}	$\langle \beta \ \Sigma \tau_i^{\pm} \vec{\sigma}_i \ \alpha \rangle$
\mathcal{M}_L	$\langle \beta \ \Sigma \tau_i^{\pm} \vec{l}_i \ lpha angle$
$\mathcal{M}_{\sigma r^2}$	$\langle eta \ \Sigma au_i^{\pm} ec{\sigma}_i r_i^2 \ lpha angle$
$\mathcal{M}_{\sigma L}$	$\langle eta \ \Sigma au_i^{\pm} i ec{\sigma}_i imes ec{l}_i \ lpha angle$
\mathcal{M}_{Ky}	$\frac{(\frac{16\pi}{5})^{1/2}\langle\beta\ \Sigma\tau_i^{\pm}r_i^2C_{12k}^{nn'k}\sigma_{\rm in}Y_2^{n'}(\hat{r}_i)\ \alpha\rangle}{(1-1)^{1/2}\langle\beta\ \Sigma\tau_i^{\pm}r_i^2C_{12k}^{nn'k}\sigma_{\rm in}Y_2^{n'}(\hat{r}_i)\ \alpha\rangle}$

 ${}^{V}F_{000}$ the impulse approximation is a valid approach, as the current is conserved. Next we discuss the divergence of the axial current and its relation to the pion field, for which we have to calculate QCD matrix elements. Corrections to the simple shell model methods are historically split up into two categories: meson exchange effects and "nuclear" effects, the latter of which comprises core polarization, relativistic effects, and configuration mixing stemming from an insufficient knowledge of the nuclear wave functions. The distinction is artificial, however, as the nuclear potential relies on an incorporation of mesonic degrees of freedom. This fact was emphasized by Wilkinson (1974). The study of mesonic contributions has a long history and includes pioneering works by Blin-Stoyle, Gupta, and Primakoff (1959), Chemtob and Rho (1971), Delorme and Rho (1971), and Rho (1974), but falls outside of the scope of this work. Reviews can be found by Blin-Stoyle (1978), Ejiri and Fujita (1978), and Towner (1987). Effects of core polarization, on the other hand, result from a truncation of the set of basis states by considering the nucleus as an inert core with some valence nucleons (Shimizu, Ichimura, and Arima, 1974). In many cases this is not an appropriate approximation, and significant corrections occur from allowing particle-hole excitations across shell gaps (Barroso and Blin-Stoyle, 1975; Koshigiri, Ohtsubo, and Morita, 1981). This is a complicated matter, however, as severe cancellations occur between core polarization and meson exchange effects, while remaining careful for double counting (Towner and Khanna, 1979). Individual effects can easily range to 40%, but due to cancellations in allowed decays these largely correspond to the impulse approximation results (Morita, 1985). For forbidden decays, however, this cancellation is much less pronounced and significant deviations are encountered when using the impulse approximation (Baumann *et al.*, 1998).

For allowed decays then one typically remedies the situation in an *ad hoc* way by renormalizing the coupling constants to produce effective couplings (Wilkinson, 1973a, 1973b, 1974; Towner, 1992; Warburton and Towner, 1992). Based on this reasoning, we expect the effective coupling constant to decrease for increasingly heavy nuclei. This is the case for the sd shell, where g_A is set to 1.1 and to unity in fp-shell nuclei (Martínez-Pinedo et al., 1996). New experimental measurements of highly forbidden decays push this value even farther downward to $g_A \approx 0.9$ (Haaranen, Kotila, and Suhonen, 2017). Statistical methods have been applied by Deppisch and Suhonen (2016), where an overview of the quenching of g_A is provided for different methods in different mass regions. Because of the g_A^4 dependence of the $0\nu\beta\beta$ cross section, its study has received renewed appreciation (Kostensalo, Haaranen, and Suhonen, 2017). Interesting to note here is the use of the β spectrum as the prime means of deducing g_A as its shape depends only on the relative values of the involved nuclear matrix elements.

Less known is the subsequent renormalization of the induced pseudoscalar coupling, for which the original interaction is typically seen as an exchange of a virtual pion (Ericson, Figureau, and Thévenet, 1973). The subsequent interaction then depends on the pion propagator, which is however modified by its interaction with the surrounding nuclear matter. This is a delicate matter and contains severe model dependence (Delorme *et al.*, 1976). In an extreme case the pseudoscalar coupling constant can be quenched by as much as 80%, which is significantly larger than the connected quenching of g_A . Great care is then required when evaluating matrix elements containing a pseudoscalar component.

Finally, we stress the model dependence of the quenching of the coupling constants, as it originates from a failure to directly take into account many-body effects and correlations. This can, for example, clearly be seen from a difference in effective g_A when using the shell model or the interacting boson model (Haaranen, Kotila, and Suhonen, 2017). *Ab initio* methods such as Green's function Monte Carlo provide a more systematic framework and circumvent the need for quenching by taking into account many-body correlations (Brida, Pieper, and Wiringa, 2011). This falls outside of the scope of this review and the interested reader is referred to Pieper and Wiringa (2001) and Carlson *et al.* (2015).

C. Symmetries

In dealing with allowed β decay, one finds that results are significantly constrained and simplified through the application of symmetries. The first we discuss here is the CVC hypothesis (Feynman and Gell-Mann, 1958), after which we briefly comment on second-class currents. Discussions on *T* invariance and partially conserved axial current can be found in Holstein (1974b) and Behrens and Bühring (1982).

1. Conserved vector current

As with the electromagnetic interaction, the conservation of the vector current allows us to derive simple relations for different form factors. In the most trivial example one finds

$$\begin{aligned} \langle J_f M_f | V_0(0) | J_i M_i \rangle &= {}^{V} F_{000}(q^2) \delta_{J_i J_f} \delta_{M_i M_f} \\ &= \langle T_f T_{3f} | T_{\mp} | T_i T_{3i} \rangle \delta_{J_i J_f} \delta_{M_i M_f} \\ &= \sqrt{(T_i \pm T_{3i})(T_i \mp T_{3i} + 1)} \times \delta_{J_i J_f} \delta_{M_i M_f} \end{aligned}$$

$$(87)$$

with V_0 the temporal part of the vector current and T_3 the third component of the isospin vector, meaning

$$VF_{000} = \sqrt{(T_i \pm T_{3i})(T_i \mp T_{3i} + 1)} = V\mathcal{M}_{000}^{(0)}.$$
 (88)

Comparing this to Eq. (84), we find that CVC excludes any induced scalar currents.

Further, this can be used to find relationships between different vector form factors, as discussed by Behrens and Bühring (1971). A relationship relevant to our further discussion is the following¹⁵:

$$-2N^{V}F_{011}^{N-1} = [W_{0} \mp (m_{n} - m_{p})]R^{V}F_{000}^{N} + \pm \alpha Z \left\{ \int \left(\frac{r}{R}\right)^{2N} U(r)Y_{0}^{0} \right\}.$$
(89)

Here m_n and m_p stand for the neutron and proton masses, respectively, and the integration is performed with the nuclear wave functions which were omitted for notational convenience. Further, U(r) is defined via $V(r) = -\alpha Z/RU(r)$ with V(r) the electronic potential, and Y_0^0 is the constant spherical harmonic function coming from the expansion of the lepton current.

Through its likeness with the electromagnetism, a deeper connection can be constructed between weak and electromagnetic observables. As both divergenceless currents behave quasi-identically, matrix elements can be interrelated. Behrens and Bühring (1982) discussed this in our adopted form factor formalism and showed that ${}^{V}F_{KK0}(q^2)$ and ${}^{V}F_{KK1}(q^2)$ are related to elastic electron scattering reduced transition strengths for electric and magnetic types, respectively. This entails that we can evaluate these matrix elements using the *charge* distribution rather than the weak charge distribution.

The weak magnetism form factor, mentioned in the previous section, provides the largest contribution of the form factors induced by the strong interaction. Its main contribution comes from the ${}^{V}F_{111}(q^2)$ form factor which is related to a magnetic matrix element, hence the name. In the general nuclear current decomposition as written by Holstein (1974b), this is now written as $b(q^2)$ rather than g_M . For β transitions between analog states the CVC hypothesis relates this form factor to electromagnetic properties of the β transition, allowing one to calculate *b* on the basis of experimental data. Thus, for the so-called mirror β transitions between isospin T = 1/2 mirror nuclei we write (Calaprice and Holstein, 1976)

$$b^{\mp} = A \sqrt{\frac{J+1}{J}} \mu^{\mp}, \qquad \mu^{\mp} = \mp (\mu_1 - \mu_2), \qquad (90)$$

where A is the mass number, J is the angular momentum, and μ_1 and μ_2 are the magnetic moments of the mother and daughter nuclei, respectively.

Further, for the pure Gamow-Teller transitions within isospin triplet states the CVC hypothesis allows calculating the weak magnetism form factor from the width Γ_{M1}^{iso} of the analog *isovector* part of the γ transition with *M*1 multipolarity via (Calaprice and Holstein, 1976)

$$b_{\gamma}^2 = \eta 6 \frac{\Gamma_{M1}^{\rm iso} M^2}{E_{\gamma}^3 \alpha},\tag{91}$$

with E_{γ} the energy of the γ transition, M the average nuclear mass of mother and daughter, α the fine-structure constant, and η a constant. The latter is unity if the final states of β^{\pm} and γ processes are equal, while it becomes $(2J_i^{\gamma} + 1)/(2J_f^{\gamma} + 1)$ when reversed to correct for the proper degeneracy in the phase space of the M1 transition.

A new compilation of experimental data and subsequent discussion on weak magnetism behavior throughout the mass range was performed by Severijns *et al.*, 2018.

2. Partially conserved axial current

The application of partially conserved axial current typically results in the Goldberger-Treiman relation (Goldberger and Treiman, 1958), relating the pion-nucleon coupling to g_A . This is generally not of much use unless one knows the pion matrix element. This relation can, however, be translated into a condition for g_P :

$$g_P(q^2) = -g_A(0)\frac{(2M_n)^2}{m_\pi^2 - q^2}.$$
(92)

Simply using experimental results for nucleon and pion masses at $q^2 = 0$ one obtains $g_P(0) \approx -229$. This expression is only valid assuming isospin invariance, and much work has been done [see Gorringe and Fearing (2003) and Bhattacharya

¹⁵This formula is a simplification of the general form factor coefficient ${}^{V}F_{011}^{N-1}(k_e, m, n, \rho)$ which can be found in Behrens and Bühring (1971).

et al. (2012) and references therein] to provide higher-order corrections to this result. These corrections are small and change g_P on the 5% level. Experimentally measurements have been performed for muon capture, for which $q^2 = 0.88m_{\mu}^2$, and so can be roughly translated using Eq. (92). Currently, the most accurate results from chiral perturbation theory are in sufficient agreement with experimental results to warrant using the PCAC results (Gorringe and Fearing, 2003). We again point to the importance of the quenching of g_P due the meson exchange effects in the nuclear medium. Thus, Eq. (92) is valid only for the free nucleon, and significant care must be taken when evaluating pseudoscalar currents for nuclear decays.

3. Second-class currents

We can classify terms in the expansion of the nuclear current following their transformation properties under the Gparity operation $G = Ce^{i\pi T_y}$, i.e., the product of a charge conjugation operation C and a rotation by π around the y axis in isospin space (Weinberg, 1958). Only two terms in Eq. (83) transform differently compared to their main vector or axialvector analogs. These are called second-class currents, as opposed to their first-class counterparts. The first is the induced scalar interaction, which was already eliminated because of CVC. The second is the induced tensor interaction, for which no similar constraint is available. As can be seen from Table IV the axial tensor form factor d contains both a so-called first-class term, i.e., d^{I} , and a second-class term d^{II} . If both the vector and axial-vector weak nucleon currents V_{μ} and A_{μ} have a definite G parity the induced terms (and thus also the one related to the d form factor) are expected to hold the G symmetry, that is, the decay of a proton and a neutron in a nucleus should be symmetric. Experimentally no indication for the existence of second-class currents has been found as yet (Grenacs, 1985; Shiomi, 1996; Wilkinson, 2000; Minamisono et al., 2001, 2011; Sumikama et al., 2011), so that second-class currents can be ignored, i.e., $d^{II} \equiv 0$. The d form factor then reduces to its first-class part, i.e., $d = d^{I}$, which is only nonzero for transitions between nonanalog states (i.e., belonging to a different isotopic multiplet), as the matrix element $M_{\sigma L}$ in d^{I} can be shown to vanish for transitions between analog states (Holstein, 1974a).

D. Analytical matrix elements

In the introduction of Sec. VI.B.1 we mentioned the decomposition of a general operator O_{KLs} into its singleparticle constituents. Using the form factor formalism, this now has a more concrete meaning, and using the impulse approximation we can build on this foundation. Following the work of De-Shalit and Talmi (1963) and its discussion by Behrens and Bühring (1982), we can write many-particle matrix elements in terms of single-particle matrix elements by introducing a factor C(K) that absorbs all many-particle effects and which depends only on the tensor rank K. For a simple configuration with two nucleons in initial and final states in an orbital with total angular momentum jone finds

$$\langle \phi(j^2, J_f T_f T_{3f}) | \left| \sum_{n=1,2} \{ O_{KLs} t_- \}_n \right| | \phi(j^2, J_i T_i T_{3i}) \rangle$$

$$= C(K) \langle j || O_{KLs} || j \rangle,$$

$$(93)$$

where C(K) is a simple combination of spin and isospin variables. For a nontrivial system of an actual nucleus, we typically employ the shell model and consider the interacting nucleons to be in different orbitals outside a fixed core. Writing the total wave function for a nucleus with spin *J*, isospin *T*, and *N* active particles as

$$\Psi(NJT) = \sum_{k} a_k \phi_k(NJT) \tag{94}$$

with $\phi_k(NJT)$ antisymmetric single-particle wave functions, we find Eq. (93) generalizes directly to

$$\langle \Psi_f(NJ_fT_f) || O_{KLs} || \Psi_i(NJ_iT_i) \rangle = \sum_{k,l} a_k a_l C_{kl}(K) \langle j_l || O_{KLs} || j_k \rangle,$$
(95)

where C_{kl} describes the coefficients of fractional parentage relevant for kl configuration (De-Shalit and Talmi, 1963). Comparing this to Eq. (81) we identify the prefactors as the one-body density matrix elements. In the following discussion we are not concerned with the absolute calculation of matrix elements, but rather in the ratio between two different ones. When taking a ratio of two manyparticle matrix elements of rank K, we see the complex many-body dynamics described by C(K) in Eq. (93) drops out if both operators are of the same rank K. If the shell model state of Eq. (94) is dominated by a single configuration this result holds approximately. We will extensively use this fact in the analytical calculation of the shape factor C(Z, W).

It remains then to write down the shape factor in an analytical form. Here we stand before a crossing, as we can attempt to split up C(Z, W) into a purely leptonic convolution part and a nuclear structure part, or make no attempt at decoupling and stick with the full formulation. The former is done by Holstein, whereas in the seminal work by Behrens and Bühring (1982) the two parts remain coupled. We continue here with this approach as it provides the greatest precision results, but present the final results in the notation of Holstein. This allows for easy inspection and a clear connection to the aforementioned symmetries. In Appendix E we compare both approaches.

E. Isospin invariant shape factor

In the beginning of this section we introduced the concept of a shape factor denoted by C(Z, W). This is not to be confused with the many-particle coupling coefficients in Eq. (93). It encompasses the information introduced by the expansion of the nuclear and lepton current in the transition matrix element, and as such for a large part determines the shape of the spectrum. In the formalism by Behrens and Bühring (1982) C(Z, W) is written as

$$C(Z, W) = \sum_{k_e, k_\nu, K} \lambda_{k_e} \bigg\{ M_K^2(k_e, k_\nu) + m_K^2(k_e, k_\nu) - \frac{2\mu_{k_e}\gamma_{k_e}}{k_e W} M_K(k_e, k_\nu) m_K(k_e, k_\nu) \bigg\},$$
(96)

where

$$\lambda_{k_e} = \frac{\alpha_{-k_e}^2 + \alpha_{+k_e}^2}{\alpha_{-1}^2 + \alpha_{+1}^2},$$
(97)

$$\mu_{k_e} = \frac{\alpha_{-k_e}^2 - \alpha_{+k_e}^2}{\alpha_{-k_e}^2 + \alpha_{+k_e}^2} \frac{k_e W}{\gamma_{k_e}},$$
(98)

are Coulomb functions depending on α_k [see Eq. (13)], while $M_K(k_e, k_\nu)$ and $m_K(k_e, k_\nu)$ contain the convolution of leptonic wave functions and nuclear structure information encoded as form factors, discussed in the previous section. The integers k_e and k_ν are defined as $|\kappa_{e,\nu}|$, where $\kappa_{e,\nu}$ is related to the angular momenta in the usual way.¹⁶ It is a well-known fact that the solution to the Dirac equation does not contain a definite parity, such that we consider the outgoing leptonic wave functions as spherical waves. The integer *K* corresponds to the multipolarity of the transition and must form a vector triangle with j_e and j_ν as well as with the nuclear spins J_i and J_f . For allowed transitions we have then $|J_i - J_f| \le K \le J_i + J_f$ from the nuclear vector triangle.

In this coupled approach all leptonic information is still contained within M_K and m_K . Even though the nuclear decompositions are completely equivalent in both the HS and BB formalisms, the treatment of the leptonic current is not. In the latter, a rigorous expansion of the radial wave functions is made in r^2 , $(m_e R)^a$, $(WR)^b$, and $(\alpha Z)^c$ thereby introducing a function $I(k_e, m, n, \rho; r)$ that is sensitive to nuclear shape information.¹⁷ This function is tabulated both in the general case and for a uniformly charged sphere in Tables 4.2 and 4.3 in Behrens and Bühring (1982), respectively. As it also includes nuclear structure information, this is typically combined with the original form factors $F_{KLs}^{(n)}$ and is written as¹⁸ $F_{KLs}^{(n)}(k_e, m, n, \rho)$.

This now allows to better understand the structure of C(Z, W) and continue with its analytical formulation. In the notation by Behrens and Bühring, capital letters are used for large components, while lower case terms represent small components. Developing M_K and m_K in terms of WR we can after a tedious but straightforward calculation write a general shape factor (Behrens *et al.*, 1978; Behrens and Bühring,

1982). In the approach by BB, all constant factors are divided out and one arrives at

$$C(Z, W) = 1 + aW + b\frac{\mu_1\gamma}{W} + cW^2,$$
 (99)

where *a*, *b*, and *c* are given by Eqs. (14.117)–(14.119) in Behrens and Bühring (1982). We will, however, divide out only the main matrix elements ${}^{V}F_{000}^{(0)} \equiv g_{V}\mathcal{M}_{F}$ and ${}^{A}F_{101}^{(0)} \equiv \mp g_{A}\mathcal{M}_{GT}$ and adjust our shape factor accordingly.

This leaves us with a series of other, often more complicated, nuclear matrix elements which require evaluation somehow. Assuming isospin invariance and CVC, however, we can link these matrix elements to *electromagnetic* matrix elements. This entails that instead of using initial and final state nuclear wave functions, one can use the full charge distribution as discussed in Sec. VI.C. This corresponds to using F1111 = 27/35, F1221 = 57/70, F1222 = 233/210, and F1211 = -3/70. This approach can be improved when using a more realistic charge distribution. Several of these possible replacements have been discussed in the electrostatic finite size corrections in Sec. IV, specifically when discussing the U correction factor. This is elaborated upon in Appendix A.

1. Superallowed $0^+ \rightarrow 0^+$ Fermi decay

In the case of superallowed Fermi decay, only terms with K = 0 contribute. We deal then only with different form factor coefficients of the form ${}^{V}F_{000}^{1}(1, m, n, \rho)$. We use the expansion of $M_0(1, 1)$ and $m_0(1, 1)$ valid to orders $(\alpha Z)^2$, R^2 , αZR to calculate C(Z, W) from Eq. (96). Here the Coulomb function μ_1 can be safely assumed to correspond to unity to our current order of precision (Behrens and Jänecke, 1969). After extraction of ${}^{V}F_{000}^{0}$ we can write the shape factor C(Z, W) in the following expansion:

$${}^{V}C(Z,W)_{0} \simeq 1 + {}^{V}C_{0} + {}^{V}C_{1}W + {}^{V}C_{-1}/W + {}^{V}C_{2}W^{2},$$
 (100)

where

$$^{V}C_{0} = -\frac{233}{630}(\alpha Z)^{2} - \frac{1}{5}(W_{0}R)^{2} \mp \frac{6}{35}\alpha ZW_{0}R,$$
 (101a)

$${}^{V}C_{1} = \mp \frac{13}{35} \alpha ZR + \frac{4}{15} W_{0}R^{2}, \qquad (101b)$$

$${}^{V}C_{-1} = \frac{2}{15} W_0 R^2 \pm \frac{1}{70} \alpha ZR, \qquad (101c)$$

$${}^{V}C_2 = -\frac{4}{15}R^2. \tag{101d}$$

2. Pure Gamow-Teller decay

In pure Gamow-Teller decay the situation becomes more complicated, and we now have contributions from $K \ge 1$ terms. For notational convenience and clarity we first introduce the Holstein variables and the translation used between Holstein's and the Behrens-Bühring formalism in which the calculations were performed. We have used

¹⁶Here κ is the eigenvalue of the operator $K = \beta(\boldsymbol{\sigma} \cdot \boldsymbol{L} + 1)$, such that $k = |\kappa| = j + 1/2, \kappa = -l - 1$ if l = j + 1/2, and $\kappa = l$ if l = j - 1/2.

¹⁷Here m = a + b + c represents the total power of mR, WR, and αZ , n = b + c is the total power of WR and αZ , and $\rho = c$ is the power of αZ .

¹⁸This is presented in Eqs. (6.159)–(6.166) in Behrens and Bühring (1982). As $I(k_e, m, n, 0) = 1$, we have $F_{KLs}^{(n)}(k_e, m, n, 0) = F_{KLs}^{(n)}$.

$${}^{A}F_{101}^{(0)} = \mp c_1, \tag{102a}$$

$${}^{V}F_{111}^{(0)} = -\sqrt{\frac{3}{2}}\frac{1}{MR}b,$$
 (102b)

$${}^{A}F_{110}^{(0)} = -\frac{\sqrt{3}}{2MR}d,$$
 (102c)

where the relevant Holstein form factors to the appropriate order in \mathbf{q}^2 are defined in Table IV. Extracting now the main Gamow-Teller form factor ${}^{A}F_{101}^{(0)}$ we can derive a similar result¹⁹

$${}^{A}C(Z,W)_{0} \simeq 1 + {}^{A}C_{0} + {}^{A}C_{1}W + {}^{A}C_{-1}/W$$
$$+ {}^{A}C_{2}W^{2} + \Phi \mathcal{P}(Z,W), \qquad (105)$$

where

$${}^{A}C_{0} = -\frac{1}{5}(W_{0}R)^{2} + \frac{4}{9}R^{2}\left(1 - \frac{1}{20}\Lambda\right) + \frac{1}{3}\frac{W_{0}}{Mc_{1}}(\mp 2b + d) \pm \frac{2}{5}\alpha Z\frac{1}{MRc_{1}}(\pm 2b + d) + \frac{2}{35}\alpha ZW_{0}R(1 - \Lambda) - \frac{233}{630}(\alpha Z)^{2}, \qquad (106a)$$

$${}^{A}C_{1} = \pm \frac{4}{3} \frac{b}{Mc_{1}} + \frac{4}{9} W_{0} R^{2} \left(1 - \frac{1}{10} \Lambda \right) \mp \frac{4}{7} \alpha Z R \left(1 - \frac{1}{10} \Lambda \right),$$
(106b)

$${}^{A}C_{-1} = -\frac{1}{3Mc_{1}}(\pm 2b + d) - \frac{2}{45}W_{0}R^{2}(1 - \Lambda) \mp \frac{\alpha ZR}{70}, \quad (106c)$$

$${}^{A}C_{2} = -\frac{4}{9}R^{2}(1 - \frac{1}{10}\Lambda), \tag{106d}$$

and

$$\Lambda = \frac{\sqrt{2}}{3} 10 \frac{\mathcal{M}_{121}^{(0)}}{\mathcal{M}_{101}^{(0)}}.$$
 (107)

¹⁹Care must be taken when precisely comparing the evaluation of h and ${}^{A}F_{121}$. The latter can be written as

$${}^{A}F^{0}_{121} = \mp g_{A}\mathcal{M}^{0}_{121} \mp \frac{g_{P}}{2M_{n}R} 5\sqrt{\frac{2}{3}}\mathcal{N}^{0}_{110}, \qquad (103)$$

where upon careful evaluation we find $\mathcal{M}_{121}^0 \propto -\mathcal{M}_{1y}$, and \mathcal{N}_{110}^0 is the relativistic matrix element

$$\mathcal{N}_{110}^0 = \sqrt{3} \int \beta \gamma_5 \frac{i\mathbf{r}}{R}.$$
 (104)

We use either a Foldy-Wouthuysen transformation (Foldy and Wouthuysen, 1950; Bjorken and Drell, 1964; Roman, 1965) or reduce f(r) using the nonrelativistic limit of the Dirac equation to reduce $\beta\gamma_5$. Converting the angular momenta coupling, we finally obtain $\mathcal{N}_{110}^0 = -(\sqrt{3}/2M_NR)\mathcal{M}_{101}^0$. In order to precisely show the influence of the oft-ignored induced pseudoscalar current, we separate them in Eqs. (106a)–(106d).

In order to stress its importance, we separated out one part of Eq. (105), written as $\Phi \mathcal{P}(Z, W)$. Here Φ is

$$\Phi = \frac{g_P}{g_A} \frac{1}{(2M_N R)^2}$$
(108)

and represents the typically ignored induced pseudoscalar contribution. Using the free nucleon values obtained from PCAC we find $\Phi \approx -0.13$ for a typical medium-*Z* nucleus with $R \sim 0.01$. This raises serious questions about the validity of neglecting induced pseudoscalar effects when high precision is required, even when assuming a strongly quenched g_P value.²⁰ Now $\mathcal{P}(Z, W)$ is a polynomial arising from the contributions of ${}^{A}F_{101}^{1}(k_e, m, n, \rho)$, ${}^{A}F_{110}^{0}(k_e, m, n, \rho)$, and ${}^{A}F_{121}^{0}(k_e, m, n, \rho)$ as can be seen from Table IV. The generalization from the bare form factors ${}^{A}F_{1Ls}$ to the form factor coefficients is straightforward and is tabulated in Table 7 by Behrens and Bühring (1971). The nonrelativistic approximation is derived using standard methods, and \mathcal{P} is then given by

$$\mathcal{P}(Z,W) = P_0 + P_1 W + P_{-1}/W, \tag{109}$$

where

$$P_0 = \pm \frac{2}{25} \alpha Z W_0 R + \frac{51}{250} (\alpha Z)^2, \qquad (110a)$$

$$P_1 = \pm \frac{2}{25} \alpha ZR, \qquad (110b)$$

$$P_{-1} = -\frac{2}{3}\gamma W_0 R^2 \pm \frac{26}{25}\alpha Z R\gamma.$$
(110c)

Many of the factors in these equations can be large enough to be significant on the few parts in 10^4 or even more. Clearly, care has to be taken when performing high precision spectrum shape measurements, and, in particular, when extracting ftvalues from mirror transitions. The quenching of g_P in nuclear matter is expected to be nucleus dependent, just like g_A , such that its inclusion can introduce shifts in the overall $\mathcal{F}t$ values.

Contributions coming from $K \ge 2$ contribute only at the few 10^{-6} level, such that for our current precision we do not take it into account.

F. Isospin breakdown and nuclear structure

The results presented in the previous section depended on isospin invariance to replace the nuclear wave functions in the matrix elements by the complete charge distribution. Isospin is, however, not an exact symmetry and its breakdown is significant on the level of precision we aim for. This breakdown manifests itself in the form of an isovector correction and further requires the explicit calculation of several singleparticle matrix elements in the Gamow-Teller shape factor.

1. Isovector correction

The use of CVC and isospin invariance links weak matrix elements to electromagnetic variables, specifically to the

²⁰See also González-Alonso and Martin Camalich (2014). Important to note is the distinction between g_P as it is defined here and that of the fundamental pseudoscalar interaction described there.

so-called isovector component. Because of the breakdown of isospin, we introduce a further modification called the isovector correction based on the approach by Wilkinson (1993b). In order to put the correction on a clearer footing, we take a look at the problem before using a perturbative expansion of the wave functions. This can more easily be seen in the notation of Hardy and Towner (2005b) where M_K is written as

$$M_{K}(k_{e},k_{\nu}) = \frac{\sqrt{4\pi}}{\hat{K}\hat{J}_{i}} \sum_{Ls} (-)^{K-L} \langle j_{\alpha} || F(r)\hat{T}_{KLs} || j_{\beta} \rangle, \quad (111)$$

with F(r) a combination of lepton wave functions. The dominant contributions for the Fermi and Gamow-Teller matrix elements come from its temporal and spatial components, respectively. In this notation, both are proportional to

$$L_L = C \int_0^\infty R_\alpha(r) F(r) R_\beta(r) r^2 dr, \qquad (112)$$

where all constants related to angular momentum coupling have been absorbed by *C* and $R_{\alpha(\beta)}$ stands for the initial (final) nuclear wave function. These states tend to have a higher $\langle r^2 \rangle$ value compared to the entire charge distribution. Combined with the decrease of the leptonic wave functions inside the nuclear volume, this results in an overestimation of the ratio of form factors in Eqs. (100) and (105). We generalize the correction proposed by Wilkinson (1993b) and write

$$C(Z,W)_I = \frac{\left[\int_0^\infty dr \Psi_e \Psi_\nu \rho_w\right]^2}{\left[\int_0^\infty dr \Psi_e \Psi_\nu \rho_{\rm ch}\right]^2},\tag{113}$$

where ρ_w is the nuclear density distribution participating in the decay process, the so-called *weak* charge distribution, and ρ_{ch} the charge distribution. An expansion of the lepton wave functions gives rise to terms of order r^2 , such that estimates of the weak rms radii enter in the equation, denoted by $\langle r^2 \rangle_w$. We write the radial part of the weak charge distribution as a product of initial and final wave functions, each consisting of a sum of weighted harmonic oscillator functions²¹

$$\rho_{w}(r) = C_{N} \sum_{n l n' l'} \alpha_{n l} \beta_{n' l'} R_{n l}(r) R_{n' l'}(r), \qquad (114)$$

where C_N is a normalization constant, and $R_{nl}(r)$ is the harmonic oscillator radial wave function for quantum numbers *n* and *l*, such that $\int R_{nl}^2 dr = 1$. In closed form these are written as

$$R_{nl}(r) = Nr^{l+1} \exp\left(-\nu r^2\right) L_k^{l+1/2}(2\nu r^2), \qquad (115a)$$

$$N = \left[\left(\frac{2\nu^3}{\pi} \right)^{1/2} \frac{2^{k+2l+3}k!\nu^l}{(2k+2l+1)!!} \right]^{1/2}, \qquad (115b)$$

$$k = n - 1, \tag{115c}$$

where $L_n^{(\alpha)}(x)$ is the generalized Laguerre polynomial, and ν is a free parameter. An additional advantage of these functions is the availability of a closed formula for the radial integrals which will be necessary in the following treatment (Nilsson, 1955):

$$\langle n_f l_f | r^L | n_i l_i \rangle = (-1)^{n_i + n_f} (2\nu)^{L/2}$$

$$\times \sqrt{\frac{\Gamma(n_i)\Gamma(n_f)}{\Gamma(n_i + t - \tau_i)\Gamma(n_f + t - \tau_f)}} \tau_i! \tau_f!$$

$$\times \sum_{\sigma} \left[\frac{\Gamma(t + \sigma + 1)}{\sigma!(n_i - \sigma - 1)!(n_f - \sigma - 1)!} \right]$$

$$\times \frac{1}{(\sigma + \tau_i - n_i + 1)!(\sigma + \tau_f - n_f + 1)!} \right],$$

$$(116)$$

where

$$\tau_i = \frac{1}{2}(l_f - l_i + L), \tag{117a}$$

$$\tau_f = \frac{1}{2}(l_i - l_f + L), \tag{117b}$$

$$t = \frac{1}{2}(l_i + l_f + L + 1), \tag{117c}$$

with the upper and lower limits of σ :

$$\max(n_i - \tau_i - 1, n_f - \tau_f - 1) \le \sigma \le \min(n_i - 1, n_f - 1).$$
(118)

For the specific case when n = n' and l = l', we find the well-known relation

$$\langle r^2 \rangle_{nl} = \frac{1}{4\nu} (4n + 2l - 1).$$
 (119)

The free parameter ν is related to the traditional oscillator parameter through $b = \sqrt{2\nu}$. It can now be constrained through its relation with the nuclear radius. Summing the contribution of Eq. (119) for all nucleons and equating it to the nuclear rms radius, one finds $b = 2^{7/6}3^{-1/6}5^{-1/2}RA^{-1/6} \approx$ $0.836RA^{-1/3}$. This corresponds to the traditional approximation $\hbar\omega = 41A^{-1/3}$ MeV when using $R = r_0A^{1/3}$.

The C_I correction as defined in Eq. (113) can be calculated using a more explicit formulation of the lepton wave functions. In the nonrelativistic approximation we write the latter as the large part of the radial behavior. For the $j = 1/2 \beta$ particle and (anti)neutrino this is $g_{-1}(r)$ and $j_0(qr)$, respectively, where j_0 is a spherical Bessel function and q is the (anti)neutrino momentum. In the standard way of expanding in powers of r and equating terms, one finds to order r^2 :

²¹The harmonic oscillator wave functions can be trivially extended to include a spin-1/2 contribution and leave the radial part of the Hamiltonian unchanged. The possible l(l') will thus be restricted to $j \pm 1/2$ of initial and final particle states. In case *j* is not a good quantum number any more, such as an axially deformed potential, more *l* values enter. This is discussed in Sec. VI.H.

$$\Psi_e(r) \approx 1 - \frac{1}{6} [(W \pm V_0)^2 - 1]r^2,$$
 (120a)

$$\Psi_{\nu}(r) \approx 1 - \frac{1}{6} (W_0 - W)^2 r^2, \qquad (120b)$$

for a uniformly charged sphere, with $V_0 = \pm 3\alpha Z/(2R)$ the electrostatic potential at the origin. Equation (113) then reduces to

$$C(Z,W)_{I} = C_{N} \sum_{nln'l'} \alpha_{nl}^{2} \beta_{n'l'}^{2} I_{nl}^{n'l'} (I_{nl}^{n'l'} - 2\xi \langle r^{2} \rangle_{nl}^{n'l'}) \times \left(1 + \frac{6}{5} \xi R^{2}\right)$$
(121)

to first order in ξ , where $I_{nl}^{n'l'} = \int_0^\infty R_{nl} R_{n'l'} dr$ and

$$\xi = \frac{1}{6}[(W_0 - W)^2 + (W + V_0)^2 - 1].$$
(122)

Here we used our definition that $\langle r^2 \rangle = (3/5)R^2$ for any charge distribution. In the extreme single-particle approximation in *j*-*j* coupling (1*s*, 1*p*, 1*d*_{5/2}, 2*s*, ...) this expression drastically simplifies and the sum in Eq. (121) disappears. Further assuming the initial and final particles to be in the same *nl* state, we find

$$C_I = 1 - \frac{1}{2\nu}\xi(4n + 2l - 1) + \frac{6}{5}\xi R^2.$$
 (123)

When considering the modified Gaussian function of Eq. (18) as a charge distribution, we can relate ν to the *A* fit parameter using the same idea as that of Wilkinson (1993b). We can interpret the constant term in Eq. (18) as the l = 0 contribution while the $A(r/a)^2$ then necessarily stands for $l \neq 0$. Assuming n = 1 and l = 0 as done there, we must have $3/(4\nu) = 3a^2/2$ after proper normalization of the l = 0 part of ρ_{MG} . Reordering terms we find then from Eq. (123) his result

$$C(Z, W)_I = 1 - \frac{8}{5} w \xi R^2 \frac{1}{5A' + 2},$$
 (124)

where

$$w = (4n + 2l - 1)/5.$$
(125)

Our more general expression [Eq. (121)] does not rely on a specific charge distribution and should employ the same charge distribution as was used to evaluate the $V/AF1mn\rho$ terms in the previous section. This is discussed further in Appendix A.

In case of vector transitions we can simply apply C_I to the entire shape factor ${}^{V}C$ of the previous section. It is slightly more complicated for Gamow-Teller transitions as we have assumed the nuclear wave functions to correspond to the charge distribution only for form factors of the type ${}^{A}F_{101}^{(0)}(1, m, n, \rho)$, analogous to the vector result. In order to avoid double counting then we can split up ${}^{A}C(Z, W)$ into a nuclear structure dependent part ${}^{A}_{ns}C(Z, W)$ containing all terms proportional to *b*, *d*, and Λ , and all other shape dependent terms ${}^{A}_{sh}C(Z, W)$. We properly apply the isovector correction using

$${}^{A}C(Z,W) \to C_{I}(Z,W){}^{A}_{\mathrm{sh}}C(Z,W) + {}^{A}_{\mathrm{ns}}C(Z,W).$$
(126)

The terms corresponding to ${}^{A}_{sh}C$ are written by Wilkinson (1990) and discussed in Appendix A.

2. Single-particle matrix elements

In the nuclear structure shape function of pure Gamow-Teller decay ${}^{A}_{ns}C$ we are sensitive to several ratios of nontrivial form factors. The weak magnetism contribution, denoted by *b*, can be constrained by CVC for specific transitions, while *d* can be shown to vanish for analog transitions. We have no such constrictions for Λ , defined in Eq. (107), and so we must rely on the explicit calculation of the matrix elements. Specifically, we need to calculate the ratio between $\mathcal{M}^{(0)}_{121}$ and $\mathcal{M}^{(0)}_{101}$. Explicit expressions have been provided in Appendix B. As an example, we write the ratio $\mathcal{M}^{(0)}_{121}/\mathcal{M}^{(0)}_{101}$ here for several allowed scenarios

$$\frac{\mathcal{M}_{121}^{(0)}}{\mathcal{M}_{101}^{(0)}} = \begin{cases} -l\frac{\sqrt{2}}{2l+3}\frac{\langle r^2 \rangle}{R^2} & j_f = j_i = l + \frac{1}{2}, \\ -(l+1)\frac{\sqrt{2}}{2l-1}\frac{\langle r^2 \rangle}{R^2} & j_f = j_i = l - \frac{1}{2}, \\ \frac{1}{2^{3/2}}\frac{\langle r^2 \rangle}{R^2} & j_f = l \pm \frac{1}{2}j_i = l \mp \frac{1}{2}. \end{cases}$$
(127)

We can test the validity of Eq. (127) in the simpler cases where we expect our single-particle approximation through *j*-*j* coupling to work. This can be done by comparing results for the weak magnetism contributions for which experimental data are available. As an example, we consider the mirror decays of six isotopes throughout the lower mass range [see Severijns *et al.*, 2018 and references therein] In the extreme single-particle approximation for mirror nuclei the weak magnetism current is easily calculated, and one finds $b/Ac_1 =$ $1/g_A(l+1+g_M)$ and $-1/g_A(l-g_M)$ for $j_f = j_i = l \pm 1/2$, respectively. Results are shown in Table VI.

We conclude that, for this investigated mass range where the single-particle behavior can be reasonably well assumed, the nonrelativistic impulse approximation evaluation yields results consistent within 15% except for the $d_{3/2}$ orbital in the

TABLE VI. Examples of T = 1/2 mirror transitions for which we compare experimental b/Ac_1 values to their pure single-particle analogs throughout the mass range 3 < A < 59. We have some freedom in choosing g_A for the higher lying states. For ³H we used $g_A = 1.2723$ while for all *sd* nuclei we used $g_A = 1.1$ based on standard works (Wildenthal, 1984; Brown and Wildenthal, 1988) and $g_A = 1$ for the *fp* shell (Martínez-Pinedo *et al.*, 1996; Siiskonen, Hjorth-Jensen, and Suhonen, 2001).

Nucleus	J^{π}	$(b/Ac_1)_{exp}$	$(b/Ac_1)_{\rm sp}$
³ H	1/2+	4.2212(24)	4.48
¹³ N	$1/2^{-}$	3.1816(75)	3.37
³¹ S	$1/2^+$	5.351(14)	5.18
³⁹ Ca ^a	$3/2^+$	1.2349(28)	2.46
⁴⁵ V	7/2-	7.51(23)	8.70
⁵⁹ Zn	3/2-	6.68(33)	6.71

^aThe electromagnetic moments of ³⁹Ca do not agree very well with the Schmidt values (Minamisono *et al.*, 1976; Matsuta *et al.*, 1999), implying larger deviations were expected to occur.

case of ³⁹Ca. From its magnetic moment and that of its related neighbor ³⁷Ar (Pitt *et al.*, 1988), we could however already conclude that a single-particle approach is not justified. Indeed, despite having only one hole outside a double magic nucleus, core polarization and meson exchange effects are significantly enlarged (Barroso and Blin-Stoyle, 1975).

We can conclude this to be a very good agreement for a simple approach. This assumes a careful study of the magnetic moments and ground state properties, however, taking into account possible orbital reversals through strong deformations. We assume the matrix element evaluation in Eq. (127) to perform similarly, justifying our single-particle approach in this mass range given the previously discussed requirements.

G. Relativistic terms in superallowed decays

In the previous section, when dealing with superallowed $0^+ \rightarrow 0^+$ decay, we ignored contributions from relativistic matrix elements, as is typically done. For the Gamow-Teller decay we made no such approximation and immediately implemented their properties. To recapitulate, by relativistic we mean matrix operators directly or indirectly containing γ_5 . This is because these operators connect the small *f* and large *g* radial wave functions when treating the transforming nucleon relativistically as in Eq. (8). Specifically, we calculate matrix elements of the form

$$\int_0^\infty g_f(r)\phi(r)f_i(r)r^2dr,\qquad(128)$$

where $\phi(r)$ is equal to $j_L(qr)$ or $(r/R)^{L+2N}I(k_e, m, n, \rho; r)$, and f and g can be interchanged. One then typically introduces a further nonrelativistic approximation (Brysk, 1952; Talmi, 1953; Rose and Osborn, 1954a, 1954b; Eichler, 1963; Lipnik and Sunier, 1966; Hardy and Towner, 2005a). In this nonrelativistic approach the small f component can be transformed such that $g_{\kappa}(r)$ is a solution of the Schrödinger equation, as done in footnote 19 when discussing Eq. (105). Typically, a factor $1/M_N R \approx 1/20$ appears, suppressing the relativistic influence significantly. In the simple case of superallowed Fermi decay we can write

$$C(Z, W) = C(Z, W)_{\rm NR} + \frac{{}^{V}F_{011}^{(0)}}{{}^{V}F_{000}^{(0)}}f_2(W) + \frac{{}^{V}F_{011}^{(1)}}{{}^{V}F_{000}^{(0)}}f_3(W) + \cdots,$$
(129)

in the notation of Behrens and Buhring (1967), where $f_i(W)$ are simple functions of the lepton momenta and slowly varying Coulomb functions, proportional to *R*, and where the subscript NR stands for nonrelativistic. These Coulomb functions are described for Fermi transitions in Appendix D. Using the results in our discussion of CVC, we can approximate Eq. (89) to write²² (Fujita, 1962)

$$2N^{V}F_{011}^{(N-1)} = -\{[W_{0} \mp (m_{n} - m_{p})]R \pm \frac{6}{5}\alpha Z\}^{V}F_{000}^{(N)}.$$
 (130)

It is then immediately clear why we have initially chosen to neglect these matrix elements, as the prefactor is exactly zero in the case of isospin invariance. Here $6\alpha Z/5R$ is the difference in Coulomb energy assuming a uniformly charged sphere. Isospin is, however, not an exact symmetry nor is the nucleus a perfectly homogeneous sphere, and Wilkinson (1993b) has treated this with much care, providing upper limits on the magnitude of the effect averaged over the full spectrum. This was again done using the assumption of an extreme single-particle interaction in the nucleus. In this case this results in a conservative estimate, as the effect increases for higher lying orbitals. For the heaviest nucleus considered ⁵⁴Co, the upper limit was set at 0.01%, while for the lightest ¹⁴O, it was put at 0.001%. The end-point energy for the former is 8243.12 keV, so that an average slope would be in the 10^{-5} MeV⁻¹ range as the dominant terms in $f_2(W)$ are linear in W. This gives a general idea of the involved magnitudes. As is done for Gamow-Teller decays discussed, we can attempt to evaluate these matrix elements explicitly using single-particle values. This is elaborated upon in Appendix D.

H. Nuclear deformation

As previously discussed in Sec. IV.A.3, a deviation in the spherical shape of the nucleus introduces profound effects on the β decay rate on the few 10⁻⁴ level. This influence extends toward the terms sensitive to nuclear structure previously discussed.

1. Leptonic convolution

The most obvious change occurs in the convolution of the lepton and nuclear wave functions, where we initially assumed a uniform charge density. Here we used the rough expansion of the lepton wave functions near the origin of Eqs. (120a) and (120b). This uniform density is then replaced by that of Eq. (37), from which we can calculate the electric potential at the center of the nucleus, yielding (Wilkinson, 1994)

$$V_{0d} = \frac{3\alpha Z}{2a} (\Phi^2 - 1)^{-1/2} \ln\left[(\Phi^2 - 1)^{1/2} + \Phi\right]$$
(131)

for the prolate case and

$$V_{0d} = \frac{3\alpha Z}{2a} (1 - \Phi^2)^{-1/2} \arcsin(1 - \Phi^2)^{1/2}$$
(132)

for the oblate case, where

$$\Phi = \frac{b}{a} \tag{133}$$

as discussed before. We follow the approach of Wilkinson (1994) and write

$$\zeta_i = \frac{1}{10} [(W_0 - W)^2 + (W \pm V_{0i})^2 - 1], \qquad (134)$$

such that we define

²²This relation is not exact and can in fact vary significantly, as discussed by Damgaard and Winther (1966). An exact treatment can be found in Behrens and Buhring (1967). We are here, however, only interested in an order of magnitude estimation.

$$D_C(Z, W, \beta_2) = \frac{1 - \zeta_d R^2}{1 - \zeta_s R^2},$$
 (135)

where $V_{0s} = \pm 3\alpha Z/(2R)$. The effect of D_C largely compensates that of D_{FS} defined in Sec. IV.A.3, so that the effect is typically on the order of a few 10^{-4} for extreme deformations.

Finally, we discuss the influence of deformation on our socalled isovector correction C_I . Here we account for the fact that the decaying nucleon sits in an orbital with limited overlap with the full charge distribution. We considered this nucleon to sit in the highest occupied orbital according to the typical *j*-*j* coupling. In the deformed case, however, *j* is not any more a good quantum number and the sum in Eq. (121) extends over several *l* rather than just $j \pm 1/2$. This can create significant deviations in the expectation values for $\langle r^2 \rangle_w$. We briefly discuss the deformed harmonic oscillator in the following section and comment on the modification of C_I in the final overview and cross-check in Sec. VIII.

2. Deformed single-particle matrix elements

In the case of Gamow-Teller transitions we are sensitive to matrix elements which cannot be reduced to their electromagnetic analogs through the use of CVC and are calculated using single-particle matrix elements. In the case of nuclear deformation, our initial harmonic oscillator wave functions are not any more good approximations of the nuclear wave function, as l is not a good quantum number. The single-particle wave function is then a normalized linear combination (Davidson, 1968)

$$\chi_{\Omega}(\mathbf{r}') = \sum_{j} C_{j\Omega} \phi(j\Omega)$$
(136)

with Ω the projection of the single-particle angular momentum on the symmetry axis of the deformed nucleus, and $\mathbf{r'}$ the radial coordinate in the body-fixed frame. The coefficients $C_{j\Omega}$ are tabulated by Nilsson (1955) and Davidson (1968) for different values of β_2 . The $\phi(j\Omega)$ are solutions of the spherical single-particle orbital with the proper spin-angular functions

$$\phi(j\Omega) = g(r',\kappa)\chi^{\Omega}_{\kappa} \tag{137}$$

such that we can keep using the harmonic oscillator radial wave functions in the calculation of matrix elements. We use the result of Behrens and Bühring (1982) to find

$$\langle \phi(J_{f}K_{f};\Omega_{f})||O_{KLs}||\phi(J_{i}K_{i};\Omega_{i})\rangle$$

$$= \sqrt{\frac{(2J_{f}+1)(2J_{i}+1)}{(1+\delta_{K_{f}0})(1+\delta_{K_{i}0})}} \sum_{j_{f}j_{i}} C_{j_{f}\Omega_{f}}C_{j_{i}\Omega_{i}}$$

$$\times \left\{ (-1)^{J_{f}-K_{f}+j_{f}-\Omega_{f}} \begin{pmatrix} J_{f} & K & J_{i} \\ -K_{f} & \Omega_{f}-\Omega_{i} & K_{i} \end{pmatrix} \right.$$

$$\times \begin{pmatrix} j_{f} & K & j_{i} \\ -\Omega_{f} & \Omega_{f}-\Omega_{i} & \Omega_{i} \end{pmatrix} + \begin{pmatrix} J_{f} & K & J_{i} \\ K_{f} & -\Omega_{f}-\Omega_{i} & K_{i} \end{pmatrix}$$

$$\times \begin{pmatrix} j_{f} & K & j_{i} \\ \Omega_{f} & -\Omega_{f}-\Omega_{i} & \Omega_{i} \end{pmatrix} \right\} \langle j_{f}||O_{KLs}||j_{i}\rangle, \qquad (138)$$

where the last factor is again the single-particle matrix element, and K is the projection of the combined angular momentum I = L + j on the symmetry axis, with L the angular momentum of the deformed core, and j that of the single particle. In the rotational ground state for an odd-A nucleus $K = \Omega$. In both even-even and odd-odd nuclei the active particles couple to K = 0 in the ground state. Equation (138) imposes an additional selection rule as $K \ge |K_f - K_i|$. As an example, consider the weak magnetism form factor b/Ac_1 for the heavily deformed ¹⁹Ne mirror isotope. Its ground state is $1/2^+$, consistent with a deformation of $\beta_2 = 0.269$ of Möller *et al.* (2016), meaning the unpaired nucleon sits in the $1/2^+[220]$ orbital. From Davidson (1968), we find the wave function is dominated by the $1d_{5/2}$ harmonic oscillator function, combined with parts from $2s_{1/2}$ and $1d_{3/2}$. From our deliberations of Eq. (127) we know that only cross-terms with $l_i = l_f$ will survive. As ¹⁹F has a nearly identical deformation, we assume $C_{j_f\Omega_f} = C_{j_i\Omega_i}$. We find then $b/Ac_1 = 5.55$ rather than 7.01 when assuming a pure $d_{5/2}$ occupation. This is much closer to the value we get from CVC $(b/Ac_1)_{CVC} = 4.9129(25)$ and entails that even for very strong deformations, we are able to predict these quantities in a single-particle estimate within 15%. This is more extensively discussed elsewhere (Severijns et al., 2018).

I. Summary

We presented in a fully analytical form the so-called shape factor correction to the β spectrum, both for Fermi and for the first time for Gamow-Teller decays. Calculations were performed in the Behrens-Bühring formalism and coupled to the work of Holstein. This allows one to obtain the highest precision results due to an improved treatment of the lepton wave functions while maintaining notational clarity. In doing so, we expounded on the symmetry properties that underlie the relevant form factors and discussed their calculation, focusing on both the reduction to single-particle matrix elements and their careful evaluation. As the move toward sub-per mille precision requires, we further discussed the evaluation of deformation influences, the breakdown of isospin invariance, and pointed to the relevance of induced interactions. We attempted to do this while preserving transparency in our calculations in the complex and at times convoluted methods required for the shape factor evaluation. The oft-neglected induced pseudoscalar interaction was kept alive in our formulation, and we showed the need for its careful evaluation. Despite nontrivial quenching in the nuclear medium, its significance cannot be underestimated when high precision results are required.

The move toward the formulation of Holstein allows for an easier interpretation of and use with experimental results. In the appendixes we showed the correspondence between our improved results and the earlier conclusions of Holstein and others to put aside any possibility for double counting. We believe we presented a more complete, transparent, and clearcut integration of the final state Coulombic effects without the artificial separation of nuclear structure and leptonic convolution effects.

For many experimentally interesting cases we have no use for CVC to derive precise values for the induced currents, raising the question of the required precision on these observables. As many spectral shape measurements are intent on measuring a possible Fierz contribution, the main induced spectral distortions come from the term linear in W. Given values of $b/Ac_1 \sim \mathcal{O}(5)$, the slope is influenced mainly by the weak magnetism contribution, averaging around 0.4% MeV⁻¹. The next largest contribution comes from the $2\alpha ZRx/35$ term, where we have to carefully evaluate both the $\mathcal{M}_{121}^{(0)}$ matrix element and the induced pseudoscalar contribution [Eq. (107)]. For both weak magnetism and Λ of Eq. (107), an extreme single-particle calculation can be sufficient to bring the required precision into the 10^{-4} domain, given that it is a valid approximation. We have seen in Sec. VI.F.2 that this is not always the case. Its legitimacy can be deduced from a study of the magnetic moments. When we are not able to confidently trust the extreme single-particle limit, however, one requires shell model input. Care then has to be taken to avoid double counting with deformation effects. Spectral measurements, in particular, have an advantage in that all terms linear in W in the shape factor have a strongly suppressed influence from Λ , putting the focus on b/Ac_1 for which explicit experimental data are available.

VII. ATOMIC AND CHEMICAL EFFECTS

The nucleus cannot be completely separated from its orbiting electrons as the decay is governed by the total Hamiltonian. Even though the interaction point lies within the nucleus, the emitted β particle undergoes continuous interaction with the atomic electrons that surround it. The screened Coulomb field both changes the interaction density within the nuclear volume and opens possibilities for more discrete interactions such as shake-off and exchange effects. These discrete processes all originate from a nonorthogonality between initial and final atomic states. We discuss in turn the screening correction, the exchange effect, shake-up and shake-off processes, the atomic mismatch correction, possible bound state β decay, and explore the molecular effects on the β spectrum and decay rate. The bulk of the effects is typically located in the low-energy region, as expected from the Compton wavelength of the outgoing electron. However, experiments sensitive to explicit energy dependences in the β spectrum shape cannot ignore atomic effects even at higher energies.

The description of screening has been known for several decades and is now combined with numerical results from atomic physics. In addition, a new analytical fit of the exchange effect is proposed here, based on precise numerical calculations.

Finally, we discuss and review molecular effects on the outgoing β spectrum. As the decaying atom can reside in many different chemical environments, the consequent molecular effects have to be evaluated in a case by case manner. We discuss general features and point to possible problematic areas in an analytic fashion.

A. Screening corrections

When the β particle is created inside the nucleus, it sees not only the nuclear charge but also the atomic electrons surrounding it. The electromagnetic potential is altered at the site, enhanced by the nonzero probability of finding an s electron inside the nucleus. The effective charge as seen by the β particle is screened because of atomic electrons. Whereas the inclusion of the Fermi function is a first-order Coulomb correction, effects coming from a screened nuclear charge are higher-order effects. This is because the largest deviation in the screened electronic potential lies at the atomic rather than nuclear radius. Considering the Fermi function equivalent to switching the electron radial wave function in the S matrix from plane wave to Coulomb function, an analytical treatment of screening requires an analytical solution of the Dirac equation in a screened Coulomb potential. Unfortunately, this is inherently impossible (Rose, 1961). The Schrödinger and Klein-Gordon equations allow for an analytical solution in a simple screened Coulomb potential (Durand, 1964), but the validity of this can be questioned when considering a high precision description.

To overcome this hurdle, initial results were obtained by Rose (1936) using the Wentzel-Kramers-Brillouin (WKB) argument, arguing a simple rescaling of the total β particle energy

$$\tilde{W} = W - V_0, \qquad \tilde{p} = \sqrt{\tilde{W}^2 - 1},$$
 (139)

where V_0 is the potential shift due to screening at the origin. The screening correction is then $\tilde{p} \, \tilde{W} \, \tilde{F} / p W F$. Even though this approach is strictly valid only for energies going toward infinity, good behavior was nevertheless found when comparing to numerical results (Matese and Johnson, 1966). However, due to the rescaling of the energy the correction is not defined for energies lower than V_0 , resulting in discontinuities for energies typically lower than 10 keV. Further study of electronic screening has been a subject of intensive research over several decades (Longmire and Brown, 1949; Good, 1954; Brown, 1964; Durand, 1964; Bühring, 1965, 1984b; Wilkinson, 1970; Lopez and Durand, 1988), including using the Feynman diagram technique (Durand and Lopez, 1987). There appears to be some confusion, however, in the evaluation of V_0 , specifically whether it corresponds to the potential shift relative to the daughter or mother atom as it was not clearly stated in the initial article by Rose (1936). As described by different authors, V_0 is defined as (Lopez and Durand, 1988; Saenz and Froelich, 1997a)

$$V_0 = \pm \alpha \langle \chi | \sum_{i=1}^{Z_p} \frac{1}{r_i} | \chi \rangle, \qquad (140)$$

where Z_p and $|\chi\rangle$ correspond to the charge and electronic wave function of the *parent* atom, rather than the daughter atom. This was noted as well by Bühring (1984a). The result of Eq. (140) is obtained after applying the sudden approximation, i.e., the charge change of the nucleus is instantaneous compared to the static electronic distribution and closure for the final atomic states. To first order in αZ they then found

$$S(Z,W) = 1 + \frac{WV_0}{p^2} + \frac{V_0}{W},$$
(141)

which corresponds to the result initially obtained by Rose (1936). Instead of performing a Lippmann-Schwinger expansion of the Coulomb interaction, one can move beyond perturbation theory, as discussed by Halpern (1970). The latter shows that Coulomb interaction before and after the β decay has the effect of changing the β wave functions from a plane wave to an exact solution of the Dirac equation in a screened Coulomb field. This fact will be used when deriving a more elaborate screening correction.

Of paramount importance for a consistent description of screening is an accurate atomic potential in the region of the nucleus. Several potentials were proposed over the past few decades, many of which are based on the Thomas-Fermi or Thomas-Fermi-Dirac potential describing the atom with a nonrelativistic statistical model (Fermi, 1927; Thomas, 1927; Dirac, 1930; Bonham and Strand, 1963; Gross and Dreizler, 1979; Bethe and Jackiw, 1997). These potentials typically fail, however, for regions of large potential gradients and low electron densities. This makes them unreliable for distances close to but also far away from the nucleus (Salvat *et al.*, 1987), precisely the region where screening will be most important.

Of particular theoretical importance due to its simplicity is the Hulthén screening potential

$$V_H(r) = \mp \alpha Z \lambda [\exp(\lambda r) - 1]^{-1}, \qquad (142)$$

with a single parameter λ that depends on Z_p , the charge of the parent nucleus. This dependence is typically written as (Bühring, 1983a, 1983b, 1984a)

$$\lambda(Z_p) = 2C(|Z_p|)\alpha |Z_p|^{1/3}m_e,$$
(143)

with $C(|Z_p|)$ a slowly increasing function of Z_p of order unity.²³ This potential has been used in several theoretical treatments of screening, with the analysis by Bühring (1984b) being of significant interest to us. In contrast to the work by Durand (1964), here the Dirac equation was used to investigate the effect.

This is made possible because of a distinct advantage in the analysis of electronic screening. The wave functions have to be well known only in the nuclear region rather than the entire space. The birth of the β particle happens inside the nucleus, so that the decay rate is affected by the depth of the potential well at this site. As electromagnetism is conservative, this means any potential energy lost while traveling through the atom will be regained at infinity. It is exactly this reason why the electronic screening effect is a second-order effect compared to the Fermi function. This locality of the interaction allows for a power expansion of the wave function near the origin. In this way an analytical description for the electron radial wave function over the full space can be avoided and one does not have to rely on approximate WKB methods. Such a power expansion was performed by Bühring (1984b)

using the Hulthén potential defined in Eq. (142). The screening correction was found to be

$$S(Z, W) = X(W/W) |\Gamma(\gamma + i\tilde{y})/\Gamma(\gamma + iy)|^{2}$$

$$\times |\Gamma(\gamma + 2i\tilde{p}/\lambda)/\Gamma(1 + 2ip/\lambda)|^{2}$$

$$\times \exp(-\pi y)(2p/\lambda)^{2(1-\gamma)}, \qquad (144)$$

where²⁴

$$X = [1 + \frac{1}{4}(\lambda/p)^2]^{-1} \{1 + \frac{1}{8}[(\tilde{W} + \gamma)/\tilde{W}](\lambda/p)^2 + \frac{1}{2}\gamma^2[1 + (1 \mp \alpha Z\lambda/(W + 1))^{1/2}]^2 \times [(W - 1)/\tilde{W}](\lambda/p)^2 \times [1 - \frac{1}{8}(1 - \gamma)(1/\gamma)(\lambda/p)^2]\},$$
(145)

and

$$\tilde{W} = W \mp \frac{1}{2} \alpha Z \lambda, \qquad y = \pm \frac{\alpha Z W}{p},$$
$$\tilde{p} = \frac{1}{2} p + \frac{1}{2} [p^2 \mp 2\alpha Z \tilde{W} \lambda]^{1/2}, \qquad \tilde{y} = \pm \frac{\alpha Z \tilde{W}}{\tilde{p}}.$$
(146)

Here $(1/2)\alpha Z\lambda$ is the potential shift due to screening at the origin. This choice of potential has several inherent weaknesses however. First, using only a single parameter severely limits its utility over the entire space, but the main obstacle lies in the evaluation of λ from Eq. (143). The correct calculation of $C(|Z_p|)$ is problematic, since estimates vary wildly, as discussed by Bühring (1984b) and references therein.

The potential used in this work is that by Salvat *et al.* (1987). The screening potential is numerically calculated using Dirac-Hartree-Fock-Slater techniques for Z = 1-92 and is then fitted with a sum of three Yukawa potentials

$$V_{\mathcal{S}}(r) = \mp \frac{\alpha Z}{r} \sum_{i=1}^{3} \alpha_i \exp(-\beta_i r).$$
(147)

It includes relativistic effects and obtains high precision results. Even though it provides a better agreement with the spatial electronic distribution than aforementioned potentials, it cannot fully account for the oscillatory features of individual orbitals. The influence hereof can, however, safely be assumed to be negligible as its dominant contribution occurs within the nucleus.

The potential shift due to screening for $r \to 0$ is then

$$\pm \alpha Z \sum_{i=1}^{3} \alpha_i \beta_i, \tag{148}$$

where α_i and β_i belong to the parent atom, such that via the substitution $\lambda = 2\sum_i \alpha_i \beta_i$ the entire machinery developed by Bühring (1984b) and Eqs. (144) and (146) transfers identically. Figure 5 shows the comparison of precise numerical calculations using Eq. (147) and analytical descriptions for

²³Here we implicitly used the sudden approximation in using Z_p rather than $Z_p \pm 1$ of the daughter atom since the atomic electrons are considered to be in parental orbitals.

²⁴Note that $X = 1 + \mathcal{O}(\lambda^2)$ as $\lambda \to 0$.



FIG. 5. Comparison of the screening correction for four different methods: Rose correction, screening evaluated with Eq. (143) (Hulthén) showing the area of uncertainty due to the uncertainty in C(|Z|), screening evaluated with Eq. (148) (Salvat), and numerical results using the potential from Eq. (147).

both screening strengths. It was originally noted by Bühring (1984b) that setting X [Eq. (145)] equal to unity in Eq. (144) gives better results at low energies for Z = 80. We checked this for a series of isotopes after comparison to numerical calculations, and this appears to indeed give a better description. We therefore assume X = 1 to be the better approximation. This could be understood by considering the limitations of the analytical description relative to the numerical procedure. In the former, the influence of the atomic potential is limited to the value at the origin through Eq. (148). While this is certainly the dominant effect, a further radial dependence of the screening potential induces additional change of the wave function at the origin. This effect can easily be seen to be small as $\alpha_i \beta_i^2 \sim \mathcal{O}(\alpha^2)$. The combination of Eqs. (25) and (147) can be utilized to further show the smallness of the effect. Analogous to this, the effect of screening on the spatial integration over the lepton wave functions contained in the shape factor C(Z, W) previously discussed can be seen to be of similar magnitude. Further comments can be found in Appendix A.

Recently a different screening correction by Mougeot and Bisch (2014) has come forward, proposing instead

$$S(Z,W) = \frac{f_{\rm sc}}{f_{\rm unsc}},\tag{149}$$

where f is defined as²⁵

$$f = \frac{\overline{g_{\kappa}^2}}{\overline{g_{\kappa}^2} + \overline{f_{-\kappa}^2}},$$
(150)

where the bar stands for the average over the entire space. Starting rigorously from the β decay Hamiltonian and the

corresponding S matrix, it is seemingly impossible to arrive at an expression resembling Eq. (149). Further, the proposed energy dependence is completely opposite in sign to that of Eqs. (141) and (144) and of larger magnitude. As applying this correction for superallowed Fermi decays is in direct violation of the CVC hypothesis and Cabibbo-Kobayashi-Maskawa (CKM) unitarity, a critical comment is necessary. This screening correction was originally proposed as a possible explanation in order to reproduce the measured spectra of ⁶³Ni and ²⁴¹Pu, the first measurements ever of such high precision at low energy, referring only to the interpretation of f as the probability of an electron emerging in an s state after decay when $\kappa = -1$. While this is approximately true for f evaluated at the nuclear radius, the same cannot be said for an average over the entire space as the electron's creation lies within the nucleus. As it is ejected from the nucleus as a j = 1/2 particle, both $\kappa = \pm 1$ states should contribute since the solution of the Dirac equation (8) does not have a defined orbital angular momentum. A screening correction must then also have contributions from both, analogous to the Fermi function, Eq. (9). It is noteworthy that the exchange correction in Mougeot and Bisch (2014) considered only the contribution from the atomic $\kappa = -1$ orbitals. Good agreement with the measured spectra of ⁶³Ni and ²⁴¹Pu appears then as a compensation of the neglect of atomic exchange effects with $\kappa = 1$ orbitals, as well as further contributions from nuclear structure and atomic corrections discussed later. Therefore, the use of this screening correction is not recommended for high precision studies of β decays.

B. Atomic exchange effects

The nonorthogonality of initial and final state atomic wave functions in β decay allows for additional indirect processes through which electrons can be emitted into a continuum state. In case of the exchange effect, this nonorthogonality leaves a possibility for a β particle to be emitted directly into a bound state of the daughter atom, thereby expelling an initially bound electron into the continuum. Experimentally it is impossible to distinguish this indirect process from regular β^- decay, so that an additional correction to the experimentally measured spectrum is required (Bahcall, 1962, 1963c, 1963a, 1963b; Haxton, 1985).

1. Formalism and procedure

We present here an analytical parametrization of this exchange correction fitted to precise numerical data. The formalism laid out here follows that of Harston and Pyper (1992). Whereas in principle all occupied orbitals in the mother atom can contribute, we consider here only allowed β decay, so that only contributions from *s* and $\bar{p} \equiv p_{1/2}$ orbitals play a part. In this case, the exchange effect is written as

$$X(E) = 1 + \eta_{ex}^{T}(E), \tag{151}$$

with

$$\eta_{ex}^{T}(E) = f_{s}(2T_{s} + T_{s}^{2}) + (1 - f_{s})(2T_{\bar{p}} + T_{\bar{p}}^{2}).$$
(152)

Here f_s is defined as

²⁵A typing error is present in the original work (Mougeot and Bisch, 2014) in the sign of κ for f_{κ} , as it does otherwise not agree with the formalism by Harston and Pyper (1992) on which it is based and loses its interpretation as the probability of emerging in a *s* state when evaluated at the nuclear radius for $\kappa = -1$.

$$f_s = \frac{g_{-1}^c(R)^2}{g_{-1}^c(R)^2 + f_1^c(R)^2}$$
(153)

and

$$T_s = -\sum_{ns' \in \gamma} \langle Es' | ns \rangle \frac{g_{n,-1}^b(R)}{g_{-1}^c(R)},$$
(154)

$$T_{\bar{p}} = -\sum_{n\bar{p}'\in\gamma} \langle E\bar{p}' | n\bar{p} \rangle \frac{f^{b}_{n,1}(R)}{f^{c}_{1}(R)}$$
(155)

sum over the different occupied *s* and \bar{p} shells in the parent state, denoted by γ . We implicitly used that κ is -1 and 1 for *s* and \bar{p} orbitals, respectively.²⁶ The quantities in Eqs. (153)–(155) are evaluated at the nuclear radius *R* using bound (superscript *b*) and continuous (superscript *c*) solutions of the Dirac equation, as written in Eq. (8). Continuum wave functions are normalized to unity. The factor $\langle Es' | ns \rangle$ stands for the integration of the overlap of bound and continuum state wave functions over the full space. In the nonrelativistic hydrogenic approximation the general overlap integral can be written down analytically as (Harston and Pyper, 1992)

$$\begin{split} \langle Z_f, El | Z_i n l \rangle &= \frac{(Z_i - Z_f) e^{2i(l+1-n) \arctan(p/\bar{Z}_i)}}{(-1)^{n-l-1} (2l+1)! p(p^2 + \bar{Z}_i^2)^{l+2}} \\ &\times 2^{2l+3} (p\bar{Z}_i)^{l+3/2} |\Gamma(l+1-i(Z_f/p))| \\ &\times e^{\pi Z_f/2p} e^{-(2Z_f/p) \arctan(p/\bar{Z}_i)} \\ &\times {}_2F_1(l+1+i(Z_f/p), l+1-n, \\ &2l+2, -4ip\bar{Z}_i/(\bar{Z}_i-ip)^2), \end{split}$$
(156)

where $Z_i (Z_f)$ is the initial (final) effective charge seen by the participating orbitals, $\overline{Z}_i = Z_i/n$, and $_2F_1$ is the confluent hypergeometric function. Some examples for l = 0 are given by Harston and Pyper (1992) and will not be repeated here. A quick glance at Eq. (156) reveals the downward trend for increasing *n*, meaning the exchange process is mainly dominated by the 1*s* orbital as is intuitively expected. While a useful first estimate, both approximations made to achieve Eq. (156) are much too coarse to provide the precision we are after. We move from a hydrogenic approximation to a proper atomic potential and use the solution of the Dirac equation in this potential to further include relativistic effects.

As will be shown explicitly, the choice of potential is of crucial importance for a consistent description of the exchange effect. Here it was chosen to use the potential described by Behrens and Bühring, in the explicit formulation by Mougeot and Bisch (2014). It is a combination of a spherical intranuclear part, a numerical screening potential by Salvat *et al.* (1987), and an exchange potential as was introduced historically by Slater (1951). The latter is evaluated using the same electron density functions described in Salvat *et al.* (1987) and

combined with a more elaborate approach to the approximation by Latter (1955) to ensure good behavior for $r \rightarrow \infty$. The exchange potential, introduced to remove electrostatic selfinteraction and ensure antisymmetry of the fermion wave function, is not to be confused with the exchange effect described here. One free parameter in this potential was varied to give agreement with numerically calculated binding energies²⁷ within 0.1% or 0.1 eV. These binding energies can easily be interchanged with experimental data when available. Results for screened and unscreened Coulomb functions tabulated by Behrens and Jänecke (1969) were cross-checked and showed an overall excellent agreement.

The electron radial wave functions $rq_{\kappa}(r)$ and $rf_{\kappa}(r)$ are calculated numerically using a power expansion for both singular points of the Dirac equation at $r = 0, \infty$ and connected via rescaling of the inner solution with α_{κ} [see Eqs. (12) and (13)]. The general approach is outlined by Behrens and Bühring (1982) and more specifically by Salvat, Fernfindez-varea, and Williamson (1995). The Fortran 77 package RADIAL (Salvat, Fernfindez-varea, and Williamson, 1995) was slightly modified and interfaced with a custom code to calculate the exchange effect. As the free and bound state wave functions are typically solved on different grids, a common grid is constructed and evaluated using exact results or via Lagrangian three point interpolation. The integral $\langle Es' | ns \rangle$ in Eqs. (154) and (155) can then be evaluated directly. Care has to be taken to make the grid sufficiently dense such that interpolation on the common grid does not introduce systematic errors.

As an example, the full exchange correction to the β^- decay of ⁴⁵Ca is presented in Fig. 6, explicitly showing the contributions from different orbitals. At very low energies of typically about 1 keV or lower, contributions from higherlying orbitals can become negative, lowering the total correction by several percentage points, as seen at the extreme left in Fig. 6. It is clear from the magnitude of the effect at the lowest energies (few keV region) that an accurate description is a necessity for any work performed in the low-energy region. For low end-point energy transitions the phase space integrals can be significantly altered, e.g., by up to 30% for transitions such as ²⁴¹Pu. The effect drops off quickly, however, and is typically negligible at energies of several hundred of keV.

2. Contribution from exchange with $p_{1/2}$ orbitals

As the electron is ejected with j = 1/2 after allowed β^- decay, the exchange effect occurs with all bound j = 1/2 states. Typically the $p_{1/2}$ exchange contribution is ignored for small to medium masses (Harston and Pyper, 1992; Mougeot and Bisch, 2014), however, this approximation is not any more viable for the precision we aim for over the full nuclear chart. Examining the behavior of Eq. (156) for different *l* reveals the overlap integrals to be of similar magnitude, using

²⁶Here κ is the eigenvalue of the $K = \beta(\boldsymbol{\sigma} \cdot \boldsymbol{L} + 1)$ operator. Further $\boldsymbol{\sigma}$ is the typical combination of Pauli matrices, \boldsymbol{L} is the orbital angular momentum operator, and 1 is a 4 × 4 unit matrix.

²⁷Here we prefer the precise relativistic local density approximation calculations by Kotochigova *et al.* (1997a, 1997b) tabulated up to Z = 92. From this point onward we switch to those calculated by Desclaux (1973), which appear to be in agreement with the former within 1% for the innermost orbitals.



FIG. 6. Exchange correction to β^- decay of ⁴⁵Ca showing the explicit contributions from different *s* orbitals. The total effect rises to nearly 10% in the few keV region and drops significantly in the 100 eV region due to negative contributions from higher-lying orbitals, in this case 2*s*.

 $_{2}F_{1}(\alpha, \beta, \gamma, z) \approx 1 + (\alpha\beta/\gamma)z$ for $|z| \ll 1$. The \bar{p} contribution is fundamentally suppressed by the factor $1 - f_{s}$ with f_{s} close to unity, since T_{s} and $T_{\bar{p}}$ are of similar magnitude. To our knowledge, this contribution has not been explicitly discussed in the literature. As an example we consider the exchange correction to the β^{-} decay of ²⁴¹Pu by all standards a high mass decay. The result is shown Fig. 7 together with the specific contributions from the occupied $p_{1/2}$ orbitals.

The contribution of the latter is highly relevant in the lowest energy regions, where its contribution can grow up to 2%, or nearly 10% of the full exchange effect. It continues to be nonnegligible for our purposes throughout the entire decay



FIG. 7. Total exchange correction to the β^{-} decay of ²⁴¹Pu showing the explicit contributions from the different orbitals (top), together with the specific contribution of the $p_{1/2}$ orbitals (bottom). The former reaches ~29% at 100 eV, while the latter rises to nearly 2% at 100 eV, indicating that it is definitely not to be neglected. The $p_{1/2}$ influence continues to be felt over a range of tens of keV at the precision we are aiming for.

spectrum. As an example of a light mass nucleus, the maximum contribution of exchange with $p_{1/2}$ for the decay of ⁴⁵Ca never exceeds a few parts in 10⁵, leaving it completely negligible. We include the contribution, small though it may be, in all following results.

3. Influence of the atomic potential

As mentioned, the choice of potential is a critical component in the description of the exchange correction. Three different potentials were compared, with increasing steps of complexity. Initial theoretical results used simple Coulomb solutions of Schrödinger (Harston and Pyper, 1992) and Dirac (Mougeot et al., 2012) equations and introduced a rough effect of screening by assuming an effective charge for each orbital. The simplest potential utilized here considers a simple exponentially screened field. The screening strength can be adjusted to give the best agreement with bound state energies, with decreasing importance for increasing main quantum numbers.²⁸ A second potential was constructed as a slight extension of the first, consisting of three Yukawa potentials, the coefficients of which were fitted to numerical data from Dirac-Hartree-Fock-Slater calculations (Salvat et al., 1987) as in the previous section. Finally, the complete potential previously described was used. For the latter the optimization of the exchange term was turned both on (optimized) and off (unoptimized). Results are shown in Fig. 8 for the exchange corrections thus obtained for the β decay of ⁶³Ni. The general trend is replicated by all potentials, apart from the very lowest energies where some orbitals can give large negative contributions. The total net magnitude is a delicate quantity in the lowest energy regions, as even for the most complex potential an optimization of the exchange potential can have significant effects. Without this optimization the binding energies for higher-lying orbitals can be seriously in error and consequently give incorrect contributions to the exchange correction.

The choice of the atomic potential reveals another key ingredient when looking at Eqs. (154) and (155) in the nuclear radius. It is imperative to know how strongly the exchange corrections depend on its precise value, as many charge radii are not experimentally known (Angeli and Marinova, 2013). For this purpose, we take up the case of ⁶³Ni as in the previous section and vary the radius by 10% in both directions. This is far larger than what can be predicted using current methods (Bao et al., 2016), where the uncertainty is estimated to be around 0.03 fm, or about 0.8% in the case of ⁶³Ni. The result is shown in the bottom half of Fig. 8. The discrepancy grows toward lower energies and only crosses 10^{-4} at roughly 1 keV. In the extreme case of ²⁴¹Pu, spectral differences cross only the 10^{-4} level at roughly 5 keV, but introduce a constant offset that can reach several parts in 10^{-4} for the same procedure. The difference shows a reasonably linear behavior on ΔR ,

 $^{^{28}}$ It can frequently occur in this approach that tuning the screening parameter to match the 1*s* binding energies results in unbound higher *ns* states. As the lowest *s* states give the highest contribution throughout most of the spectrum, we choose to neglect this detrimental effect in this rough method.



FIG. 8. Comparison of the exchange effect for ⁶³Ni shown for different atomic potentials (top), and absolute differences in the total exchange correction when varying the nuclear radius by 10% with the optimized potential (bottom). Results vary wildly, stressing the need for an accurate atomic potential. Particularly interesting is the large discrepancy stemming from the optimization of the binding energy. The different potentials used are defined in the text.

such that extrapolation to reasonable uncertainties brings it to below the few 10^{-5} effect.

4. Analytical parametrization

The calculations required to arrive at the results shown in Fig. 6 are involved and in the spirit of this work we would like to have a completely analytical description of the β spectrum shape. The immediate issue with the case at hand resides in the integration in Eqs. (154) and (155). Whereas for most other effects, the radial wave functions have to be known only near the origin, thereby allowing power expansions even for nontrivial potentials. The exchange correction requires knowledge of the radial wave functions of both continuum and bound states for the entire space and for arbitrary potentials. The results in Fig. 8 show that a hydrogenic approach with simple screening potentials is not sufficient when high precision is required, and the analytical results of Eq. (156) for pure Coulomb fields do not allow for easy insertion of fit parameters. This is further hindered by the evaluation of the confluent hypergeometric function, the evaluation of which is in itself a significant hurdle. More importantly, the magnitude of the exchange effect is a complex interplay between atomic shell effects (through the binding energy sensitivity; see Fig. 8), spatial increase of the potential, and spatial extension of the bound states for different *n*. An example of this can be seen in Fig. 9. Here the argon, krypton, and xenon shell closures have been specified. This complex Z dependence poses a significant challenge to a sufficiently precise analytical fit valid over the full Z and W range. Other external parameters based on tabulated values would have to be provided as input to the fit function. We therefore choose to tabulate the required fit parameters for each Z individually. The analytical fit for the exchange correction as a function of W contains nine fit parameters and is written as



FIG. 9. Demonstration of atomic shell effects on the exchange contributions from the 1s and 2s orbitals evaluated at 3 keV. The increased binding energy at higher Z reduces the spatial extension of the bound states and thus decreases the spatial overlap between bound and continuum electron radial wave functions at low energies.

$$X(W) \approx 1 + a/W' + b/W'^{2} + c \exp(-dW') + e \sin[(W - f)^{g} + h]/W^{i}, \qquad (157)$$

with $W' \equiv W - 1$. The fit parameters can be found in the Supplemental Material and show excellent agreement over the full tested energy range [338]. The latter was chosen at 1 MeV, where all contributions dip below the 10^{-4} level. The maximal differences are on the order of a few 10^{-4} , with average residuals located at the $< 10^{-5}$ level. The influence on the phase space integrals agrees typically on the order of a few 10^{-5} in absolute terms. Special care was taken to avoid steep slopes over small energy ranges in the relative differences to allow for precise measurements of the energy dependence in the β spectrum shape.

C. Shakeoff and shakeup processes: The end-point shift

The change of atomic orbitals due to the aforementioned effects also has a direct consequence on the internal house-keeping of the atom. Initial and final states belonging to different quantum numbers are not any more orthogonal, allowing discrete excitations into higher allowed states (shakeup) and even the continuum (shakeoff). This then reduces the available energy for the lepton pair and changes the final state interaction with the β particle. In deriving the results in the previous sections, these effects have typically been ignored on the basis of their relative importance compared to the magnitude of the higher-order effect. For the precision aimed at in this work, this is not always valid. The probabilities for shakeup and shakeoff depend strongly on Z, with the former (latter) becoming subsequently less (more) important for increasing Z.

1. Shakeup

Instead of double ionization of the shakeoff process, atomic electrons in the final states can simply be excited into higher states. The probability of this process can typically be reduced to integrals of the radial wave functions of initial and final bound states, where the shakeoff probability can simply be calculated as the deviation from unity when summing all aforementioned probabilities. Decays of few-electron systems are particularly prone to this effect, as for instance in the tritium system. Here the single electron in the final state has a significant probability to end up in the 2*s* orbital or even higher ones (Williams and Koonin, 1983; Arafune and Watanabe, 1986; Hargrove, Paterson, and Batkin, 1999). Additionally, the recoil momentum of the daughter nucleus can induce excitations due to the sudden acceleration. One can show (Feagin, Merzbacher, and Thompson, 1979) that the average excitation energy due to the aforementioned effect is equal to

$$\Delta E_R = \frac{1}{2} Z m_e v_R^2 = Z E_R \frac{m_e}{M}, \qquad (158)$$

with *M* the nuclear mass and v_R and E_R the recoil velocity and energy, respectively. For Z = 50 and a maximum recoil energy of 1 keV, there is an average excitation energy of roughly 0.25 eV. Considering atomic excitations are on the order of tens of eV, it is clear then that incomplete overlap between initial and final atomic states is the dominant effect in the full shakeup picture.

The Q value of the decay is subsequently reduced by the mean excitation energy

$$\overline{\Delta E}_{\rm ex} = \sum_{f} P_f E_f, \tag{159}$$

where E_f is the excitation energy of the final state f and P_f the probability to populate that level.

a. Effect on screening and exchange corrections

Possible excitations in the final state also change the screening correction, where instead of Eq. (141), we should write now in the notation of Saenz and Froelich (1997a)

$$S(Z, W) = 1 - \left(\frac{W}{p^2} + \frac{1}{W}\right) \alpha \sum_n \langle \varphi_0 | \phi_n \rangle^2 \langle \phi_n | \sum_{i=1}^{Z_p} \frac{1}{r_i} | \varphi_0 \rangle$$
$$= 1 - \left(\frac{W}{p^2} + \frac{1}{W}\right) V'_0, \tag{160}$$

where $|\phi_{n'}\rangle$ represents the *n*'th final electronic state, and $|\varphi_0\rangle$ is the initial electronic state. The value for the screening potential can change by as much as 20% in the special case of tritium β^- decay, simply because the atomic Coulomb interaction is limited to a single electron (Hargrove, Paterson, and Batkin, 1999). For systems with a higher number of electrons, the relative change in the electronic distribution decreases. The magnitude of the effect is, however, also proportional to *Z*, such that errors introduced from neglecting shakeup are of the order of 1×10^{-4} .

In the case of the exchange correction the situation is slightly more complicated. For medium to high Z nuclei, the probability for shakeup is negligible at the 10^{-3} level, as

explicitly calculated by Harston and Pyper (1992). For lower Z nuclei shakeup probabilities can be significant (e.g., around 25% in tritium β^- decay), and exchange with higher-lying *ns* orbitals should be taken into account even if they were not occupied in the initial state (Harston and Pyper, 1993). In the extreme case of tritium, this inclusion changes the exchange correction at the few 10⁻⁴ level down to 0.5 keV. Except for this special case then, we can safely neglect shakeup influences on the exchange correction.

2. Shakeoff

Just as in the previous section, nonorthogonality of initial and final atomic states allows for excitations in the final state, including the continuum. Whereas in the tritium system the probability for additional ionization is an order of magnitude smaller than that of shakeup, this balance quickly changes when going to higher Z. One of the underlying reasons for this is the following: The possibility for shakeup depends on atomic overlap integrals between shells with differing main quantum numbers. Because of the Pauli principle, excitations are possible only to unfilled shells, for which the overlap integrals quickly disappear for inner electrons. Outer shells are close to the continuum, and electronic excitations are typically unbound for higher Z.

The description of shakeoff has seen significant theoretical (Levinger, 1953; Schwartz, 1953; Green, 1957; Stephas and Crasemann, 1967, 1971; Carlson et al., 1968; Freedman, 1974; Law and Suzuki, 1982; Suzuki and Jaw, 1982; Frolov and Ruiz, 2010; Ruiz, 2013; Ruiz, Margraf, and Frolov, 2013) and experimental (Snell and Pleasonton, 1957; Carlson, Pleasonton, and Johnson, 1963; Carlson, 1963a, 1963b; Schupp and Freedman, 1980; Scielzo et al., 2003; Couratin et al., 2012, 2013) effort. For practical reasons, this has mainly focused on the calculation of electron ejection from the Kshell, which is a small effect in all nuclei²⁹ and not in our current interests. The seminal work by Carlson et al. (1968), even though theoretically significant improvements have been made, illustrates the general principles we are concerned with as follows: (i) For a given main quantum number the shakeoff probability decreases with increasing Z, due to the reasoning mentioned earlier. (ii) For a given atom, the shakeoff probability increases with increasing main quantum number. (iii) The total shakeoff probability is reasonably independent of Z, occurring for approximately 20%–30% on all decays.³⁰

Neglecting shakeup and collisional ionization (Carlson, 1963a; Freedman, 1974), this can be described by the oneelectron ionization probability (Couratin *et al.*, 2013)

$$p_{i} \simeq 1 - \sum_{n' \le n_{\max}} \{ |\langle \phi_{n'l_{i}}^{D} | \phi_{n_{i}l_{i}}^{M} \rangle|^{2} + K^{2} |\langle \phi_{n'l_{i}\pm 1}^{D} | \mathbf{r} | \phi_{n_{i}l_{i}}^{M} \rangle|^{2} - K^{2} \operatorname{Re} \langle \phi_{n'l_{i}}^{D} | \phi_{n_{i}l_{i}}^{M} \rangle^{*} \langle \phi_{n'l_{i}}^{D} | r^{2} | \phi_{n_{i}l_{i}}^{M} \rangle \},$$
(161)

²⁹The probability for K ejection for ³H decay is on the order of 2×10^{-3} , while for higher Z it is of the order of a few 10^{-4} .

³⁰With exception to the lightest systems such as ³H and ⁶He, where shakeup and nondissociative resonances are much more important; see the previous section.

with $\phi_{nl}^{D(M)}$ the daughter (mother) wave function with primary quantum numbers (n, l) and $K = \sqrt{2E_R/M}$ for a nucleus with recoil energy E_R and mass M. Rather than calculate the probabilities of excitation into the continuum, it is easier to simply consider the atomic overlap and approximate all deviations from unity as coming from shakeoff. Holes created by shakeoff are filled through emission of x rays or Auger electrons. The rate of these two competing processes is regulated by so-called fluorescence yields and is reviewed by Krause (1979), Hubbell *et al.* (1994), and Schönfeld and Janßen (1996).

For simplicity, we first briefly discuss the single ionization process with a β particle and an electron in the final state. In the case of β^- decay, the two outgoing electrons are indistinguishable, which has to be reflected in the decay amplitude through a coherence term (Law and Campbell, 1972a, 1972b). The decay is properly treated as a single step such that the full energy available has to be shared between all final states. The probability for shakeoff is thus dependent on the β energy and decreases for increasing energy. This was first emphasized by Stephas and Crasemann (1967), even though one typically forgets about the phase space dependent effects and instead determines the integrated ejection probability. The mean energy of the ejected electron is similar to its binding energy (Feinberg, 1941), such that the largest spectral change occurs in the low-energy region [see, e.g., the Kurie plots in Law and Campbell (1972b)]. In case the ejection occurs from one of the outer shells, the typical binding energy is on the order of a few tens of eV, such that to first order the change in spectral shape can be approximated as an additional decrease in W_0 by the mean excitation energy weighted with the probability for ejection. From the results of Couratin et al. (2013) combined with the binding energies of Desclaux (1973), we found the mean energy loss due to shakeoff per decay around 5 eV for chlorine.³¹ Because of the arguments listed, we expect this effect to be of similar magnitude for all Z.

a. Effect on screening and exchange corrections

The screening and exchange corrections presented [Eqs. (144) and (157)] depend strongly on the final atomic potential. In the case of shakeoff electron ejection, at least one additional hole is created through a single-step process, thereby altering the electronic density at the nucleus as well as the wave functions of all remaining occupied orbitals. For both corrections, the magnitude of the contribution decreases with increasing n, while the probability for shakeoff is proportional with n. We can then reasonably approximate the shakeoff influence as coming from events originating from the outer shell as we have done before. Using the screening potential as defined in Eq. (160) [see also, e.g., Pyper and Harston (1988)], the change in screening due to a hole in the outer shells is

$$\frac{V_0 - V_{\rm SO}}{V_0} = \frac{\langle f | r^{-1} | i \rangle - \sum_{f'} \langle f' | i \rangle^2 \langle f' | r^{-1} | i \rangle}{\langle f | r^{-1} | i \rangle} \\\approx \frac{\sum_{f'} \langle f' | i \rangle^2 (1 - \Delta Z_{\rm eff}) \langle nl | r^{-1} | nl \rangle}{\langle f | r^{-1} | i \rangle}, \quad (162)$$

where $|f'\rangle$ is the final state with a hole in an orbital denoted by n and l. We assumed the inner orbitals to remain unchanged in the event of shakeoff in the outer shell and allowed for a difference in effective charge seen by the remaining electron in the outer shell through ΔZ_{eff} . Using results by Law *et al.*, this change is approximately 0.3–0.4. We use the general result for hydrogenic orbitals $\langle r^{-1} \rangle = Z_{\text{eff}}/n^2$, such that Eq. (162) transforms to

$$\frac{V_0 - V_{\rm SO}}{V_0} \approx \frac{1}{\langle f | r^{-1} | i \rangle} \sum_n \frac{\langle f' | i \rangle^2 Z_{\rm eff} (1 - \Delta Z_{\rm eff})}{n^2}.$$
 (163)

Effective charges seen by orbitals can be calculated from the mean radii as noted in Harston and Pyper (1992). For a specific example of Ru^{1+} , Eq. (163) yields approximately 0.1%, which subsequently pushes the relative change in spectral shape to the few 10^{-5} level in the lowest energy range. These approximations are crude, but have to yield results precise only within a factor of 2 for us to neglect it.

For exchange the situation is again more complicated, because of both the magnitude of the effect and the sensitivity to the wave function over the entire space. Rigorously, we have to combine the approach taken by Law and Campbell (1972a, 1972b) with that of Harston and Pyper (1992), by introducing the indistinguishability of the outgoing electrons from the former approach in the exchange terms described by the latter. Harston and Pyper (1993) described this situation for the tritium system, but this was ignored in their former work (Harston and Pyper, 1992). Exchange can occur only because of the combination of the indistinguishability of electrons and the Pauli principle, such that in allowed β^- decay only exchange with bound states having angular momentum j = 1/2 can occur. In the medium to high Z nuclei, the probability for shakeoff to occur in an $s_{1/2}$ or $p_{1/2}$ state is smaller than 0.1% (Harston and Pyper, 1992). The effect of shakeoff on exchange then decreases for increasing Z. For lower Z the effect of a single shakeoff electron of energy W'can be approximated by including the following term in Eq. (152), using the notation of Harston and Pyper (1992):

$$\chi_{\text{ex}}^{\text{cont}}(E) = \int_{E'=0}^{E_{\text{max}}} \phi(E, E') dE', \qquad (164)$$

where

$$\phi(E, E') = \sum_{A} \left(-\langle \gamma_u | \gamma'_v \rangle \langle \gamma'_v(A \to E's) | \gamma_u \rangle \frac{g_{E',s}^c}{g_{E,s}^c} + \langle \gamma'_v(A \to E's) | \gamma_u \rangle^2 \left[\frac{g_{E',s}^c}{g_{E,s}^c} \right]^2 \right) \frac{W'}{p'},$$
(165)

where the summation over A runs over all s states, neglecting the $p_{1/2}$ state. When making the approximation that shakeoff

³¹Charge states created due to emission of Auger electrons do not contribute to our correction as they emerge with a fixed energy independent of the β decay. The mean energy is then obtained by summing over the holes created for different charge states, using $\tilde{E}_{SO} \approx 1.8B_{nl}$ with B_{nl} the binding energy (Feinberg, 1941).

mainly occurs for the outer shell, we can limit the sum over A to the final *ns* shell. The evaluation of the integral in Eq. (164) is explained in Harston and Pyper (1993). For the tritium decay, χ_{ex}^{cont} has a maximum value of -0.01% at 1 keV. Using again the decay of ³⁵Ar as an example (Couratin *et al.*, 2013) for hole creation in *ns* states in the final state, we can safely assume this correction can be neglected at the current order of precision.

D. Atomic overlap: Alternative atomic excitation correction

The β decay of a nucleus results in a sudden change of the nuclear potential, due to both a charge difference and a recoil effect. Both effects on the spectrum of the emitted β particle are described elsewhere. As the eigenstates for initial and final states belong to slightly different Hamiltonians, the initial and final atomic orbital wave functions only partially overlap. This allows for discrete effects such as shakeoff and shakeup discussed in the previous section, which decrease the decay rate as the phase space becomes smaller.

The description of the effect can in a first approach be reduced to a difference in atomic binding energies. This analysis was first performed by Bahcall (1963c, 1963b, 1965) and is also recently included in detailed $\mathcal{F}t$ analyses (Hardy and Towner, 2009). A correction is constructed by looking at the relative change in the spectrum shape when changing W_0 with $W_0 - \overline{\Delta E}_{ex}$. In this formalism, the correction can then be written as

$$r(Z,W) = 1 - \frac{1}{W_0 - W} \frac{\partial^2}{\partial Z^2} B(G), \qquad (166)$$

where B(G) is the total atomic binding energy for a neutral atom with $Z \pm 1$ protons in the case of β^{\mp} decay. The second derivative of this quantity is related to the average excitation energy due to orbital mismatch via

$$\overline{\Delta E}_{\text{ex}} = -\frac{1}{2} \frac{\partial^2}{\partial Z^2} B(G).$$
(167)

It can easily be parametrized as a function of Z using numerical values by Carlson *et al.* (1970) and Desclaux (1973) or Kotochigova *et al.* (1997b),

$$\frac{\partial^2}{\partial Z^2} B(G) = 44.200 Z^{0.41} + 2.3196 \times 10^{-7} Z^{4.45} \text{ eV}.$$
(168)

Bambynek *et al.* (1977) reviewed the approach by Bahcall and its effect on electron capture ratios and found generally good agreement. Discussion and improvements have been presented by Faessler *et al.* (1970) and Vatai (1970) who are mainly concerned with proper evaluation of the overlap integrals in the case of holes in the final state.

For completeness we also discuss a small correction to r(Z, W). In the analysis performed by Bahcall (1963c), one implicitly uses the *sudden approximation*, i.e., the change in the nuclear charge is instantaneous compared to atomic orbital velocities. For higher Z this is no longer true for low-lying electrons, as K electrons can achieve relativistic speeds. This was treated by Feagin, Merzbacher, and Thompson (1979),

who introduced an additional quantity to be combined with $\overline{\Delta E}_{\rm ex}$ previously discussed. This is written as

$$\overline{\Delta E}_K = 2(C_0 + C_1), \tag{169}$$

where C_0 and C_1 are described in detail by Wilkinson (1993a). Its influence on the Q value is on the order of a few eV (Feagin, Merzbacher, and Thompson, 1979). In the notation of Wilkinson we extend the parametrization of K(Z) to include the full Z range

$$K(Z) = -0.872 + 1.270Z^{0.097} + 9.062 \times 10^{-11}Z^{4.5}.$$
 (170)

The correction r(Z, W) then becomes

$$r(Z,W) = 1 - \frac{2}{W_0 - W} \left(\frac{1}{2} \frac{\partial^2}{\partial Z^2} B(G) + 2(C_0 + C_1) \right).$$
(171)

Its influence is felt mainly near the end point of the transition, where it cannot be neglected, as $(\partial^2/\partial Z^2)B(G)$ can become as large as a few hundred of eV. It is thus particularly important for low-energy transitions such as ⁶³Ni (end-point energy 67.2 keV) and ²⁴¹Pu (end-point energy 20.8 keV). For the former the correction reaches 1% at 15 keV before the end point and increases rapidly from that point onward to the end point.

E. Bound state β decay

Finally, we comment on the possibility of two-body bound state β decay. As the electron is created inside an electronic potential well, there exists a possibility for the β decay to be captured inside the potential well and produce an electron in a bound atomic state, effectively reducing the decay to a twobody problem. This was first studied by Daudel, Jean, and Lecoin (1947) and was later expanded upon by Bahcall (1961) and Kabir (1967) and finally by Budick (1983) and Pyper and Harston (1988). As this is a relevant issue in the analysis of $\mathcal{F}t$ values, we briefly comment on it here. In the notation by Bahcall the ratio of probabilities can be written as

$$\frac{\Gamma_b}{\Gamma_c} = \frac{\pi (\alpha Z)^3}{f(Z, W_0)} (W_0 - 1)^2 \Sigma,$$
(172)

where Σ can be found in Bahcall (1961) and depends on the binding energy of the bound orbital, atomic overlap integrals and the orbital wave functions evaluated at the nuclear surface. For free neutron decay, this ratio is approximately 4.2×10^{-6} and is completely negligible. For tritium decay, however, this ratio becomes roughly 1% for T^+ and 0.5% for T in the initial state. Higher W_0 values lead to smaller ratios, as the phase space integral f is roughly proportional to W_0 to the fifth power. The kinematic dependence for Γ_b/Γ_c then approximately follows a W_0^{-3} behavior. Nuclei studied in $0^+ \rightarrow 0^+$ superallowed decays have Q values of several MeV, rendering the bound state decay probability completely negligible. For low-energy decays and (partially) ionized initial states, this correction can grow significantly, however. This does not affect the β spectrum shape, as it is a separate final state in the calculation of the S matrix. It enters the equation when considering the $\mathcal{F}t$ analysis, and so cannot always be neglected.

F. Chemical influences

In many experiments, the decaying atom is bound within a molecule. Therefore, the electronic structure is modified as electrons rearrange themselves in molecular orbitals. The presence of additional electrons and spectator nuclei influences the Coulombic final state interactions, while rotational and vibrational states open up more possibilities for energy transfer to the molecular final state (Cantwell, 1956). These effects can be considered even as higher-order corrections, but prove essential in the determination of the antineutrino mass in the tritium system. After comments by Bergkvist (1971), an extensive amount of literature was produced on the atomic and molecular effects on the endpoint energy of tritium (Law, 1981; Kaplan, Smutny, and Smelov, 1982; Budick, 1983; Williams and Koonin, 1983; Strobel, Pearson, and Chao, 1984; Fackler et al., 1985; Jeziorski et al., 1985; Kolos et al., 1985, 1988; Arafune and Watanabe, 1986; Lindhard and Hansen, 1986; Durand and Lopez, 1987; Szalewicz et al., 1987; Lopez and Durand, 1988; Claxton, Schafroth, and Meier, 1992; Froelich et al., 1993; Harston and Pyper, 1993; Froelich and Saenz, 1996; Glushkov, Khetselius, and Lovett, 2009). Driven by experimental discrepancies (Backe et al., 1993; Belesev et al., 1995; Lobashev et al., 1999; Weinheimer et al., 1999), this culminated in the seminal works by Saenz and Froelich (Saenz and Froelich, 1997a, 1997b; Jonsell, Saenz, and Froelich, 1999; Saenz, Jonsell, and Froelich, 2000; Doss et al., 2006; Doss, 2007), describing in an *ab initio* and analytical way the influence of additional molecular electrons and spectator nuclei. There the effects of all electrons and nuclei within the molecule were treated on equal footing. The main results are summarized here.

1. Recoil corrections

In the case of β decay inside a molecule, the recoiling daughter nucleus moves inside the molecular potential rather than a vacuum. This potential is typically described with a Born-Oppenheimer energy curve, which can be modeled using a Lennard-Jones-type potential. When placed inside, the recoiling daughter nucleus kicks the molecule in a (predissociative) rovibrational state. Dissociation can occur for rovibrational states with a total energy larger than the dissociation energy, or through electronic excitation into a resonant continuum state. The typical energy scale of the former is of the order of a few eV (Jeziorski et al., 1985; Jonsell, Saenz, and Froelich, 1999) while for the latter is on the order of tens to a few hundred of eV (Kaplan, Smutny, and Smelov, 1982; Claxton, Schafroth, and Meier, 1992; Saenz and Froelich, 1997b; Jonsell, Saenz, and Froelich, 1999; Saenz, Jonsell, and Froelich, 2000; Doss et al., 2006). The integrated probability of the molecular continuum depends on the end-point energy of the transition, as a higher end-point energy implies a higher recoil energy, thereby typically ending up in the dissociative regime (Cantwell, 1956). In the case of a

including atomic tritium (Kaplan, Smutny, and Smelov, 1982). The spectral shape depends on W_0^2 , such that the relative error goes like $2\sigma_Q/Q$. For the study of $\mathcal{F}t$ values, this dependence is even heightened as the statistical rate function depends on Qthrough a fifth power. The required precision on Q should then be studied case by case to determine the required accuracy. In the lowest energy transitions, such as those of tritium, ⁶³Ni, or 241 Pu, a substantial error on the Q value on the few 10^{-4} to 10^{-3} level can be crudely expected for σ_0 on the order of a few tens of eV. The change in the decay rate was treated approximately by Pyper and Harston (1988), who found

final bound state with some angular momentum J, the energy of the β particle is reduced due to the rovibrational excitation and the center-of-mass movement of the entire molecule rather than merely that of the recoiling atom. For isotropic emission of the β particle relative to the molecular orientation, the majority of the energy deposited will be placed into rovibrational motion rather than center-of-mass of movement. We assume then that the mass used in the recoil corrections does not significantly change. At the precision aimed for in this work we neglect this effect, as it forms a small correction on already small corrections (see Sec. IV.B.1 for R_N and Sec. IV.B.2 for Q).

In case the recoiling daughter atom is to be detected (e.g., in the measurement of the β -neutrino correlation $a_{\beta\nu}$), the situation is not so simple. As the recoil energy is inversely proportional to its mass, the probability for dissociation decreases with increasing Z. There is also a clear dependence on the $\beta - \nu$ angular correlation, as higher recoil momenta tend to populate higher rotational and vibrational bands and vice versa. This intuitive picture is also confirmed for diatomic molecules as shown by Cantwell (1956). Its evolution in the molecular potential has the effect of partially randomizing the outgoing angle, as well as an energy transfer to its molecular partners. This effect can at least partially be included by simulating the response of a recoiling nucleus in a Lennard-Jones potential using Monte Carlo techniques, as was done by Vorobel et al. (2003) and Vetter et al. (2008). In the most precise analysis, this effect introduces a systematic error of 0.05%, pointing to the need for a more detailed understanding of molecular and dynamic effects in the future.

Whereas previously the Q value of the decay was decreased

because of shakeup and shakeoff within the atom, inclusion of

molecular effects adds a further decrease because of rotational

and vibrational excitations. Numerical results have mainly

been published for the ditritium molecule T_2 , due to the high precision required in the determination of the antineutrino

mass. The molecular influence then has the effect of changing the excitation possibilities to a broad continuum with reso-

nances, and provide further broadening through population of

rovibrational states. As mentioned, the width of the populated

rovibrational energy spectrum lies in the few eV region and

can typically be neglected. Excitation into the continuum now

becomes nontrivial, however, as was explicitly demonstrated

in the case of T_2 by Doss et al. (2006). Averaged over

the entire continuum spectrum, differences between several

tritium-substituted molecules is on the order of a few eV.

2. Influence on the *Q* value

$$\frac{\Delta\lambda}{\lambda} \approx \frac{3\Delta W_0}{W_0} \left(1 + \frac{1}{6} \gamma W_0 \right), \tag{173}$$

where ΔW_0 is the difference in mean end-point energy after averaging over all final states between two chemical states.

3. Molecular screening

In the approach by Saenz and Froelich (1997a), all Coulombic effects are treated equally to first order. This effect is typically split into an electronic and nuclear part, where the former can be written down as in Eq. (140) with the sum extending over all molecular electrons, and the latter as

$$P_{\rm nuc}^{(0,1)}(p) = \sum_{n} \alpha |\langle \varphi_0 | \phi_n \rangle|^2 \left[Z \left(\frac{8W}{\pi p} + \frac{8p}{3\pi W} \right) + \sum_{S} Z_S \left(\frac{W}{p^2} + \frac{1}{W} \right) \langle \xi_{000}^0 | \frac{1}{R_S} | \xi_{000}^0 \rangle \right], \quad (174)$$

where the sum *n* extends over all final states, the sum *S* takes into account all spectator nuclei in the molecule, $|\xi_{000}^0\rangle$ is the rovibronic ground state of the molecule, and R_s is the distance operator between the decaying atom and the spectator nucleus *S*. The first part in this equation represents the Fermi function to first order and is larger by a factor *p* than the effects of the spectator nuclei simply from the prefactor. We could stop here and let the influences of molecular electrons and spectator nuclei be calculated numerically using standard quantum chemical calculations (Yamanouchi, 2001).

In the spirit of this work, however, we intend to arrive at an analytical approximation to the required order of precision. In order to obtain an estimate for the change in screening due to molecular effects, we ignore the energy difference in final atomic states and use closure to perform the sum over n (which is to first order corrected for by using the atomic mismatch correction, see Sec. VII.D). For each atom in the molecule, we can consider it to have an inert atomic inner structure for its electronic configuration, coupled with a shared wave function describing the valence electrons participating in chemical bonds.³² The full electronic part of the molecular wave function with *i* atoms can be written as

$$\psi_{\text{Mol}}^{\text{el}} = |\text{valence}\rangle \times \prod_{i} |\text{inert}\rangle_{i},$$
(175)

where we implicitly used the Born-Oppenheimer approximation, and i = 0 corresponds to the decaying nucleus. For notation, we write $|\text{inert}\rangle \equiv |i\rangle$ and $|\text{valence}\rangle \equiv |v\rangle$. The latter can be described in a typical quantum chemical treatment using molecular orbitals (Roothaan, 1951; Lichten, 1967; Bransden and Joachain, 1983; Levine, 2000). Using the clamped-nuclei approximation³³ for the rovibrational ground state we write

$$\langle \xi_{000}^0 | \frac{1}{R_S} | \xi_{000}^0 \rangle \approx \frac{1}{R_{S,e}}$$
 (176)

with $R_{S,e}$ the equilibrium distance between decaying and spectator nucleus *S*, such that we can write the total Coulombic influence as

$$P_{\text{tot}}^{(0,1)}(p) \approx \alpha \left\{ Z \left(\frac{8W}{\pi p} + \frac{8p}{3\pi W} \right) + \left(\frac{W}{p^2} + \frac{1}{W} \right) \left[\sum_{S} (Z_S - Z_{S,\text{in}}) \frac{1}{R_{S,e}} - {}_0 \langle i | \sum_{Z_{\text{in}}} \frac{1}{r} | i \rangle_0 - \langle v | \sum_{Z_{\text{val}}} \frac{1}{r} | v \rangle \right] \right\},$$
(177)

where we used that

$$_{i>0}\langle i|\frac{1}{r}|i\rangle_{i>0}\approx \frac{1}{R_{S,e}}.$$

We now explicitly introduced a screened spectator nucleus with charge $Z_S - Z_{S,in}$ which can be easily evaluated when $R_{S,e}$ can be estimated to sufficient accuracy. The influence of molecular charge distributions can then be written as

$$\Delta S_{\text{Mol}} = \alpha \left(\frac{W}{p^2} + \frac{1}{W}\right) \left[\sum_{S} (Z_S - Z_{S,\text{in}}) \frac{1}{R_{S,e}} - Z_{\text{eff}} \langle r^{-1} \rangle_{\text{Val}}\right],$$
(178)

where $Z_{\rm eff} = Z_{\rm Val} - (Z - Z_{\rm in})$ and $\langle r^{-1} \rangle_{\rm Val}$ corresponds to the average inverse distance of all valence electrons relative to the decaying nucleus. Then $\Delta S_{\rm Mol}$ should be added with the regular electronic screening correction of Eq. (144).

As an example, consider the decay of ⁴⁵Ca bound inside CaCl₂, with calcium doubly oxidized Ca(II). The bond length for CaCl was measured to be 2.437 Å, which we can use as a rough estimate, while $Z_s = 17$ and $Z_{s,in} = 16$ as only one electron participates in the bonding. In natural units we have then $R_{Cl,e} \sim 600$, such that the first term in Eq. (178) is roughly proportional to 3×10^{-5} , i.e., 2 orders of magnitude smaller than V_0 . In the evaluation of the second term keep in mind that it constitutes both the valence electrons and the absence of the valence electrons of the decaying atom. We have then $Z_{eff} = 2$, while $\langle r^{-1} \rangle_{Val}$ will be of the same order of magnitude as the bond length since the valence electrons will be located mainly near the Cl atoms. The effect from the second term will thus also be on the order of 2×10^{-5} with opposite sign to that of the spectator nuclei. The deviation

³²We are able to make this approximation because of three restrictions on the participating states, the first two of which depend on the properties of the overlap integral (Atkins, 1984): (i) they must have the same rotational symmetry around the internucleus axis, (ii) they cannot be too diffuse nor too compact, and (iii) the energies must be similar.

³³This entails treating the nuclei in a molecular system as infinitely massive compared to the surrounding electrons. Effectively this also means a decoupling of nuclear and electronic motion as in the Born-Oppenheimer approximation; see, e.g., Bransden and Joachain (1983).

from molecular structure on the electronic screening potential is then of the order of 2×10^{-5} . As the screening correction is typically on the (sub)percent level at its maximum, molecular deviations are expected to have an upper limit at the few 10^{-4} level.

4. Molecular exchange effect

The atomic exchange process represents the possibility for a direct β decay into a bound orbital of the final electronic state. In the case of a molecule, the electronic phase space is greatly enlarged and perturbed relative to the single atomic electronic state. In particular, electrons can now reside in molecular, rather than atomic orbitals, while binding energies for other orbitals can change. As shown explicitly in Eqs. (154) and (155), the probability for the exchange process into a certain orbital depends on the spatial overlap between continuum and bound state. The continuum state tends to oscillate rapidly for distances large compared to its Compton wavelength. Because of this oscillatory nature, however, we are much more sensitive to the shape of the wave function as compared to the previous section, where we were interested only in the $\langle r^{-1} \rangle$ matrix element. This in turn allowed us to approximate the electronic distribution as in Eq. (175). At the level of precision we are aiming at, combined with the magnitude of the exchange effect at low energies, this is no longer sufficient. Molecular wave functions for all but the trivial H_2^+ system have to be calculated numerically, however, which leaves us with limited possibilities in an analytical description. We will nonetheless introduce the approximation of Eq. (175), and qualitatively describe the behavior of the valence electrons. An important feature to aid us in this matter is the symmetry group of the molecule.

Molecular orbitals can in a qualitatively enlightening way be constructed from a linear combination of atomic orbitals (LCAO)³⁴ (Bransden and Joachain, 1983). In general we can write all electron wave functions as a LCAO and minimize the coefficients to provide the lowest energy in a Hartree-Fock scheme. For argument sake, however, we introduce the additional approximation that the internal orbitals for all atoms are the same in the molecular as in the atomic case. The molecular valence orbitals are then a combination of occupied atomic valence orbitals and energetically close atomic excited states.³⁵ We assume then the contributions of the inner orbitals to the exchange effect are unchanged, while that of the final occupied orbital will be perturbed. As the electron density near the decaying atom can both increase and decrease in a molecular bond, the overlap integral in Eqs. (154) and (155) will do the same. Here we reach the limit of the analytical description, and the molecular bonding possibilities are too extensive to provide parametrizations for the effect. When the valence electron is nearly fully removed in an ionic bond, the exchange effect will be approximately zero, as the internuclear distances are much larger than the β Compton wavelength where the wave function rapidly oscillates. In the reverse extreme situation, the valence electron density is doubled. This then naively doubles the exchange correction. Conservatively, then, we treat the contribution of the last orbital with a 100% error bar. This way, in all but the most extreme ionic bonds, we additionally absorb small errors coming from small binding energy changes in the inner atomic orbitals. Taking Fig. 6 as an example, this introduces a $<1 \times 10^{-4}$ error from 15 keV onward and grows to a 0.5% error at 1 keV. For ²⁴¹Pu, the valence orbital participating in exchange is 7s, whose contribution drops to below 10^{-4} after 3 keV while its maximum quickly grows to 1% in the first 0.5 keV. An estimate for the effect can be obtained from the analytical results first obtained by Harston and Pyper (1992) using screened hydrogenic orbitals. The effects of the change of the wave function in a lattice versus that of a gas have been studied by Kolos et al. (1988) in the case of tritium, and results showed no significant change. Because of our conservative error bar, this effect is completely absorbed and can be neglected.

VIII. OVERVIEW AND CROSS-CHECK

Table VII shows all effects included in our description of the β spectrum shape with corresponding references to the equations used. It remains then to be considered how well our description stacks up against others results from the literature. There exists limited information on precise shape factor evaluation, other than what has been performed by Behrens *et al.* (1978), which is, however limited to the shape factor evaluation in an approximate manner.

This is an interesting moment to compare what we have done so far to precise $\mathcal{F}t$ values for superallowed Fermi transitions calculated by Hardy and Towner (2015) and Towner and Hardy (2015). A ratio of f values calculated within our framework relative to the results of Hardy and Towner (2015) is shown in Fig. 10.

In general we have excellent results, and all residuals are in the few 10^{-4} region. This is particularly interesting, as for the heaviest nuclei in question we have moved far away from stability, with ⁷⁴Rb being 11 neutrons away from its closest stable isotope and subject to strong deformations and shape coexistence. The daughter nuclei of the last four nuclei investigated, ⁶²Ga, ⁶⁶As, ⁷⁰Br, and ⁷⁴Rb, have deformations with β_2 equal to 0.195, 0.208, -0.307, and 0.401, respectively³⁶ (Möller *et al.*, 2016). We conclude that as we are in the aimed-for range for even these extremely exotic nuclei, and

 $^{^{34}}$ For a quantitative description one has to rely on Hartree-Fock and Kohn-Sham calculations with large basis sets; see, e.g., Jansík *et al.* (2009) and Ide, Kato, and Yamanouchi (2014). This is beyond the scope of this review, and we must content ourselves with a qualitative outset.

³⁵This is a reasonable approximation, except for when there is degeneracy in the different atomic orbitals. When the same atom is present several times in the molecule and is related through symmetry, different LCAO change the spatial behavior of the wave functions while the energy stays the same.

³⁶Clearly this is an interesting and challenging region, as experimental evidence is often not in agreement with theoretical predictions. Specifically in the case of ⁷⁰Se, the daughter nucleus of our largest deviation, the sign of β_2 is contested (Hurst *et al.*, 2007). The total influence of deformation on the *ft* value is, however, symmetric relative to β_2 [see Fig. 2 by Wilkinson (1994)], such that we are not as sensitive to the sign. Either way, the last proton will end up in the $g_{9/2}$ orbital in the Nilsson model.

TABLE VII. Overview of the features present in the β spectrum shape [Eq. (4)], and the effects incorporated into the beta spectrum generator code (Hayen and Severijns, 2018). Here the magnitudes are listed as the maximal typical deviation for medium *Z* nuclei with a few MeV endpoint energy. Some of these corrections fall off very quickly (e.g., the exchange correction *X*) but can be sizable in a small energy region. Varying *Z* or W_0 can obviously allow for some migration within categories for several correction terms.

Item	Effect	Formula	Magnitude
1 2	Phase space factor Traditional Fermi function	$pW(W_0 - W)^2$ F_0 [Eq. (6)]	Unity or larger
3	Finite size of the nucleus	$L_0 [Eq. (16)]$	10 ⁻¹ -10 ⁻²
4	Radiative corrections	<i>R</i> [Eq. (47)]	
5	Shape factor	<i>C</i> [Eqs. (100) and (105)]	
6	Atomic exchange	<i>X</i> [Eq. (157)]	
7	Atomic mismatch	<i>r</i> [Eq. (171)]	
8	Atomic screening	S [Eq. (144)] ^a	10 ⁻³ -10 ⁻⁴
9	Shakeup	See item 7 and Eq. (160) ^b	
10	Shakeoff	See item 7, Eq. (163), and χ_{ex}^{cont} [Eq. (164)] ^c	
11	Isovector correction	C_I [Eq. (113)]	
12	Distorted Coulomb potential due to recoil	Q [Eq. (45)]	
13	Diffuse nuclear surface	U [Eqs. (25) and (29)]	
14	Nuclear deformation	D_{FS} [Eq. (40)] and D_C [Eq. (135)]	
15	Recoiling nucleus	R_N [Eq. (41)]	
16	Molecular screening	ΔS_{Mol} [Eq. (178)]	
17	Molecular exchange	Case by case	
18	Bound state β decay	Γ_b/Γ_c [Eq. (172)] ^d	Smaller than 1×10^{-4}
19	Neutrino mass	Negligible	
20	Forbidden decays	Not incorporated	

^aHere the Salvat potential of Eq. (147) is used with X [Eq. (145)] set to unity.

^bThe effect of shakeup on screening was discussed in Sec. VII.C.1 with Eq. (160).

^cShakeoff influences on screening and exchange corrections were discussed separately in Sec. VII.C.2. This has to be evaluated in a case by case scenario.

^dThis does not affect the spectral shape, as discussed in Sec. VII.E, but does enter the $\mathcal{F}t$ analysis.



FIG. 10. Ratio of f values for all superallowed $0^+ \rightarrow 0^+$ Fermi decays up to mass 74 included in the analysis by Hardy and Towner (2015). Uncertainties mainly result from the uncertainties on Q values to illustrate the importance of possible deviations on the V_{ud} analysis. For the heaviest nuclei we show results for both the spherical shell and a deformed shell filling. These last four nuclei (62 Ga, 66 As, 70 Br, and 74 Rb) all have their valence nucleons outside the N = Z = 28 shell closure. These nuclei are, however, extremely exotic and show strong deformations and shape coexistence. In the Nilsson model a complex interplay between the $2p_{1/2}$, $1f_{5/2}$, and $1g_{9/2}$ orbitals arises. A reversal between the first two would not influence our isovector correction as w is equal. The same is not true in case the $1g_{9/2}$ becomes filled, as is the case for 70 Br.

we can trust the validity of our approach for pure Fermi decays in a completely analytical manner without the need for additional computation.

An additional measure that also tests the Gamow-Teller parts can be found in the compilation of mirror decays. Its importance cannot be overstated (Severijns *et al.*, 2008), as experimental precision is ever improving. Here the final $\mathcal{F}t$ value depends on the mixing ratio ρ and the ratio of vector and axial-vector f values

$$\mathcal{F}t^{\text{mirror}} \equiv \frac{2\mathcal{F}t^{0^+ \to 0^+}}{1 + (f_A/f_V)\rho^2}.$$
 (179)

The separate f values have been calculated by Towner and Hardy (2015) using the same methods as those used in Fig. 10. We have now two possibilities to compare our results. As these are all mirror nuclei, the Holstein form factor d vanishes identically and CVC allows us to precisely calculate the weak magnetism term from existing experimental data (Severijns *et al.*, 2018). On the other hand, we are able to approximate the latter using single-particle matrix elements as discussed in Sec. VI.F.2. Both results are shown in Fig. 11.

As expected from the results for superallowed decays, the agreement in the vector sector is exquisite, with all differences smaller than 4×10^{-4} . For the axial-vector part the general agreement is good, but there are some distinct features. We see that for cases where the extreme single-particle approach is justified, the deviation is in the 10^{-4} range for both CVC and



FIG. 11. Comparison of pure f_V (top) and f_A/f_V (bottom) values for mirror decays as calculated by Towner and Hardy (2015) and by using our formalism described here. We see an excellent agreement for the f_V values, with differences being smaller than 4×10^{-4} , as expected from the results in Fig. 10. For the calculation of f_A/f_V , the weak magnetism contributions were calculated using CVC in one case and the single-particle estimates discussed in Sec. VI.F.2. Overall a good agreement is found except for special cases where the spherical harmonic oscillator evaluation breaks down, i.e., for Z = 17 and 18. The importance of an accurate representation of weak magnetism is underlined by these ³³Cl and ³⁵Ar isotopes, for which the shell model is also unable to correctly calculate b/Ac_1 .

single-particle results. In the cases where this is absolutely not the case, such as the outliers at ³³Cl and ³⁵Ar, the disagreement reaches 1%. A similar failure is then expected in the evaluation of the ratio of matrix elements in Λ , defined in Eq. (107). Important to note, however, is that the shell model has issues pinning down the right values in these cases as well (Severijns et al., 2018). The accuracy with which the shell model is able to explain these values is found to be on the order of 10% (Severijns et al., 2018), meaning deviations away from unity in our comparison do not necessarily stem from a failure in our evaluation of Λ and are at least in part due to the dominant weak magnetism correction. Judging from the comparison between the CVC and single-particle results, one would conclude there to be a complete failure of the extreme single-particle model to accurately predict the weak magnetism contribution. A surprising yet pleasant result is found, however, when one moves from a spherical harmonic oscillator to a deformed Woods-Saxon (DWS) potential but retains the extreme single-particle approximation (Severijns *et al.*, 2018). Excellent agreement is found throughout the entire region for which experimental results are available, and even the outliers can be nicely reproduced when introducing strong oblate deformations based on mean-field results (Möller et al., 2016). Using this approach to calculate all relevant matrix elements, we show the new results for f_A/f_V in Fig. 12. While not any more analytically available, it shows a remarkable



FIG. 12. Comparison of f_A/f_V values as calculated by Towner and Hardy (2015) and those by using our formalism described here, with the addition of a DWS result as described in the text. We retained the same vertical scale of Fig. 11 to show the vast improvement over the spherical harmonic oscillator results. The calculations of the nuclear matrix elements, while still evaluated in an extreme single-particle fashion, are now not any more analytically available. The single-particle wave function is expanded in a spherical harmonic oscillator basis for which the coefficients have to be calculated by numerical diagonalization of the deformed Hamiltonian. This is performed automatically by the C++ code accompanying this work (Hayen and Severijns, 2018). As shell model calculations reach only a slightly better precision in estimating b/Ac_1 (around 10%) (Severijns et al., 2018), the observed deviations from unity when using CVC results can be attributed to this fact.

agreement with much more advanced shell model calculations while using the *extreme single-particle* approximation.

Remaining uncertainties can safely be attributed to differences in the exact value of weak magnetism, for which the shell model is shown to perform only marginally better than our deformed single-particle approach (Severijns *et al.*, 2018). Combined with the excellent agreement in the vector sector through a comparison of f_V , these results show both the potential of the extreme single-particle method and the consistency of the developed formalism described here.

On the other hand, this raises important questions on the accuracy of the f_A/f_V calculations when experimental results enter this domain. Seeing as to how differences in weak magnetism predictions can shift these values by several parts in 10^3 , this undermines the claims of reaching 0.01% in theoretical calculations (Towner and Hardy, 2015). The currently most precisely measured mirror $\mathcal{F}t$ isotope ¹⁹Ne is seen to behave rather well under the single-particle approximation and is not so much affected by this concern. Experimental campaigns are under way, however, for precision measurements of the $\beta - \nu$ correlation of ³²Ar, ¹⁹Ne, ³⁵Ar (Couratin et al., 2013; Liénard et al., 2015; Severijns and Blank, 2017) and the β -asymmetry parameter of ³⁵Ar (Severijns and Blank, 2017) and 37 K (Fenker, 2016). For the extraction of V_{ud} to be valid, a significant amount of attention needs to be given to a precise evaluation of the Λ factor in Eq. (107). When this can be done reliably, the formalism developed here can be combined with the experimentally determined weak magnetism contribution. Following this, we comment once more on the significance of the induced pseudoscalar component in Eq. (107). Its contribution, assuming the free nucleon value $g_P = -229$, is comparable to that of the ratio of matrix elements, meaning strong deviations are expected to occur. It is possible that shifts can occur even on the per mille level. When ignoring this contribution, a reasonable error must be attributed to all f_A/f_V calculations until a way is found to accurately account for this effect.

The results shown when moving to a deformed Woods-Saxon potential were obtained by the custom code that calculates the full β and (anti)neutrino spectrum shape. It allows for a calculation of all nuclear relevant matrix ${}^{V/A}\mathcal{M}_{KLs}$ in an extreme single-particle fashion, with several options for customization in order to correctly specify the single-particle state. It allows for a coupling with more advanced shell model and mean-field codes by expressing the transition matrix elements in a single-particle basis. It is properly discussed in a separate publication and will be publicly available (Hayen and Severijns, 2018). Results concerning weak magnetism including contributions from this custom code are discussed elsewhere (Severijns *et al.*, 2018).

IX. β SPECTRUM SHAPE SENSITIVITY TO WEAK MAGNETISM AND FIERZ TERMS

In the previous section we showed our independent correspondence with the currently best numerically calculated f values in the work by Towner and Hardy (2015). We can then reliably use the expressions found in this document and use them as the basis in looking for new physics results at the per mille level. As mentioned in the Introduction, investigation of the Fierz term has the important advantage of looking for deviations linear in new, exotic coupling constants. The β spectrum shape is an ideal observable for this as the sensitivity to the Fierz term varies as 1/E while the effect of weak magnetism on the β spectrum shape varies proportional to E.

A. Spectral sensitivity

The weak magnetism and a possible nonzero Fierz interference term both modify the shape of the β spectrum in an energy-dependent fashion. Historically, this was investigated experimentally by defining a so-called shape factor after which one defines the slope of the resultant ratio [see, e.g., Calaprice and Holstein (1976)]. Experimentally an unnormalized shape factor is obtained by dividing the number of counts observed for each β -particle energy by the quantity $K(Z, W, W_0, M)pW(W - W_0)^2$ [see Eq. (4)] which is then normalized to unity at some energy W_{norm} by dividing these unnormalized values by the value at W_{norm} to yield the normalized shape factor³⁷ (Calaprice and Holstein, 1976)

$$S(W) = C(Z, W)/C(Z, W_{\text{norm}}).$$
(180)



FIG. 13. Example of the effects of a weak magnetism and Fierz term on the β spectrum shape. Both effects are normalized at 100 keV, i.e., a ratio is constructed as in Eq. (180). This shows the advantage of measuring low-energy transitions for Fierz, while high-energy end-point transitions are favorable for weak magnetism studies.

The slope dS/dW then provides the physics information. A similar ratio can be constructed for the Fierz term. A typical value for b/Ac_1 is about 5 (Wauters *et al.*, 2010). For b_{Fierz} a precision of 1% or better is typically required in view of current constraints on $C_S^{(l)}$ or $C_T^{(l)}$ coupling constants (Severijns, Beck, and Naviliat-Cuncic, 2006; Naviliat-Cuncic and González-Alonso, 2013; Vos, Wilschut, and Timmermans, 2015). Moreover, as was shown recently, at a precision of about 0.1% measurements of b_{Fierz} in nuclear β decay and neutron decay remain competitive with direct searches for new bosons related to scalar and tensor type weak interactions at the LHC collider in the channel $p + p \rightarrow$ e + MET + X (with MET standing for the missing transverse energy) which has an underlying dynamics similar to β decay at the parton level (Naviliat-Cuncic and González-Alonso, 2013; Khachatryan et al., 2015). To generate some perspective on the magnitude of this slope when introducing a Fierz or weak magnetism term, we show in Fig. 13 an example shape factor S(W) for $b/Ac_1 = 5$ and 3, and $b_{\text{Fierz}} = 0.005$ and 0.001. This clearly stresses the need for an accurate value of b/Ac_1 when extracting Fierz information. Further, this requires an excellent theoretical description of the β spectrum shape, as has been presented here. The slope for $b_{\rm Fierz}=0.005$ over the first 250 keV interval is equal to -0.33% MeV⁻¹. The energy-dependent information of the theoretical spectrum must thus be accurate enough to guarantee all remaining slope artifacts to be <0.1% MeV⁻¹. Special attention has been given when describing atomic corrections to avoid any residual energy-dependent slopes.

At the lowest energies, care has to be taken when interpreting experimental results, as the Fierz term spectral modification is approximately linear in this regime. Experiments are then sensitive to $\sim (b_{\rm wm} - b_{\rm Fierz})E$, and there is no way to decouple the separate contributions from their energy-dependent behavior. This again underlines the required precision and accuracy in evaluating the weak

³⁷To remain consistent with our notation, we wrote the shape factor as *C* instead of the spectral function h_1 as per Holstein and coauthors.

magnetism contribution when searching for new physics (Gonzalez-Alonso and Naviliat-Cuncic, 2016).

B. Effective field theory approaches

The traditional search for exotic interactions in β decay is described by introducing additional coupling constants and currents in the β decay Hamiltonian. This makes direct comparison with results from high-energy colliders such as the LHC difficult. It is here that effective field theories can bridge the gap. Assuming the mass scale of new physics Λ_{BSM} to be beyond what is probed right now at the LHC, both highand low-energy experiments can be directly compared in an effective field theory. Here additional coupling constants are introduced that are sensitive to exotic currents in higher-order operators. A significant amount of effort has been put into its development, showing that both low- and high-energy frontiers can be competitive in their current experimental stages (Erler and Ramsey-Musolf, 2005; Ramsey-Musolf and Su, 2008; Bhattacharya et al., 2012; Cirigliano, Gardner, and Holstein, 2013; Cirigliano, González-Alonso, and Graesser, 2013; Naviliat-Cuncic and González-Alonso, 2013; Holstein, 2014a; Vos, Wilschut, and Timmermans, 2015). The classical β correlation terms of Jackson, Treiman, and Wyld (1957) have been written in terms of these extra coupling terms, allowing both energy frontiers to provide an upper limit on the same parameters.

Care has to be taken to avoid overlap when combining the formalism presented here with that of Bhattacharya *et al.* (2012). As an example, the Fierz term b_{Fierz} for neutron decay presented therein carries a nonzero SM contribution [Eq. (B15) in Bhattacharya *et al.* (2012)]

$$b^{\rm SM} = -\frac{m_e}{M_N} \frac{1 + 2\mu_V \lambda + \lambda^2}{1 + 3\lambda^2},$$
 (181)

where μ_V is the difference between proton and neutron magnetic moment, and $\lambda = g_A/g_V$. The full correction term is written as $m_e b^{\text{SM}}/E_e$. The presence of μ_V points to a weak magnetism contribution, and consequently the term can indeed be retraced to the equivalent terms in Eq. (105).

X. CONCLUSIONS AND OUTLOOK

In this review, we presented a fully analytical description of the β spectrum shape, for the first time combining kinematical, electromagnetic, nuclear, and atomic corrections to a few parts in 10⁴. These corrections are expected to hold from 1 keV to the end point. The importance of the atomic corrections was underlined, showing that although the largest deviations are found at low energy, their influence can be seen at high energy. The theoretical framework for the description of atomic screening was combined with precise Dirac-Hartree-Fock-Slater calculations and showed excellent agreement with numerical results down to the lowest energies. This avoids the ambiguous evaluation of the screening exponent as described before. Atomic exchange calculations were performed for the entire atomic number chain, and a fit was provided to enable analytical evaluation with high precision. The oft-neglected contributions stemming from exchange with atomic $p_{1/2}$ orbitals was shown to be significant and included into our analytical description. Further atomic corrections from different sources were combined with the work done by Wilkinson dealing with electromagnetic finite size corrections.

An overview of the corrections sensitive to nuclear structure was given and a full correction was derived for both Fermi and Gamow-Teller decay. We presented these results in the transparent notation by Holstein while maintaining the precision of the lepton wave function and nuclear current decomposition by Behrens and Bühring. Besides the weak magnetism contribution, we remain sensitive to nuclear structure information through the explicit evaluation of several matrix elements and the proposed isovector correction. We showed that the former can be analytically calculated to a sufficient precision given that the state is properly described in a spherical extreme single-particle manner. When this is not the case, it was shown that moving to a deformed Woods-Saxon potential while retaining the extreme single-particle approach provides excellent agreement. If wanted, more extensive methods such as the shell model can provide the relevant matrix elements which can directly be incorporated into the formulae presented here. Further, we note the influence of the induced pseudoscalar contribution to the Gamow-Teller shape factor and point to the importance of its correct evaluation when experimental precision for mirror decays reaches that of the superallowed transitions.

The formalism presented here was compared against the most precise numerical calculations of the total integral of the β spectra performed by Towner and co-authors. For both the superallowed and mirror decays, good agreement was found throughout the entire investigated mass range. Even for the most exotic cases where deformation effects provide a significant contribution, excellent agreement was found. The calculation of the Gamow-Teller spectrum is more difficult as here we are directly influenced by nuclear structure influences we can circumvent in the vector sector. Using the extreme single-particle approach, good results nevertheless were obtained for the cases where this is reasonably justified based on electromagnetic moment data. When this is not the case, significant deviations occur, as expected. Moving to a proper deformed potential alleviates these problems for the investigated cases. Important to note is, however, that the shell model calculations which were used to compare against cannot also reliably predict the weak magnetism contribution to the argued precision. This raises serious questions on the validity and accuracy of the presently available calculations.

We therefore conclude that the work presented here can accurately describe the β spectrum shape to the required precision given that the nuclear structure input can be reliably calculated, be that through use of CVC, the extreme single-particle approximation, or more advanced methods. Spectral shape measurements have an additional advantage here in that they are not as sensitive to nuclear structure effects outside of the weak magnetism effect, and can thus be the basis for experimental beyond standard model searches exploring the per mille regime.

When looking for Fierz contributions one must remain wary, since standard model effects such as the weak magnetism contribution and their corresponding uncertainties can interfere destructively with our ability to extract meaningful results. Great control in its evaluation is required in case CVC cannot be invoked and one must rely on alternate methods of calculating the relevant matrix elements. Ideally this value is known to about 10% in order to look for per mille level Fierz contributions, as discussed in the previous section. This is clearly a difficult task and is currently the largest uncertainty by almost an order of magnitude in the precise determination of the β spectrum shape, unless it can be determined through CVC arguments. Even so, one must be careful in the evaluation of the CVC result, as only the isovector component is of relevance.

The full calculation of the work presented here was automated in a custom code, discussed in a separate publication (Hayen and Severijns, 2018). Based on simple configuration files, it aims for flexibility and user friendliness in the use of this vast size of work.

Nevertheless, the importance of an accurate β spectrum shape cannot be understated in the search for beyond standard model physics in the electroweak sector due to the linearity of exotic coupling constants in the observables. Assuming no exotic currents it is conversely also a perfect tool to study the weak magnetism contribution. Particularly for higher masses little is known experimentally about about its magnitude. This also forms an important ingredient in the analysis of the reactor antineutrino anomaly (Mueller et al., 2011). In the current analysis, a constant value is assumed for weak magnetism throughout the entire fission fragment region. Clearly, this is not an optimal method of treating this complex effect, as can already be deduced from the discussion on single-particle matrix elements given in this document. Additionally, a correct treatment of higher-order effects in the translation from electron to antineutrino spectra forms the basis of a correct analysis procedure. The formulas presented here serve as the basis for this analysis, where we note the added complexity of the shape factor compared to its usual treatment. Atomic effects, though mainly confined to the lower energy regions, become transported to the end of the antineutrino spectrum and remain highly relevant. This further underlines the importance of a precise description of atomic effects at low energy and will no doubt be the subject of further research.

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APPENDIX A: GENERAL SHAPE FACTOR

In Sec. VI.E we gave results assuming isospin invariance in that we replaced the nuclear wave functions with the full charge distribution in the evaluation of the nuclear matrix elements. Doing so leaves us with the question of how to accurately treat said charge distribution. The general equations for superallowed Fermi decay are the following:

$${}^{V}C_{0} = -\frac{(W_{0}R)^{2}}{3} {}^{V}F1110 \mp \frac{2}{9} \alpha Z W_{0} R^{V}F1111 - \frac{(\alpha Z)^{2}}{3} {}^{V}F1222, \qquad (A1a)$$

$${}^{V}C_{1} = \frac{4}{9}W_{0}R^{2V}F1110 \mp \frac{2}{3}\alpha ZR({}^{V}F1221 - \frac{1}{3}VF1111),$$
 (A1b)

$${}^{V}C_{-1} = \frac{2}{9} W_0 R^{2V} F_{1110} \mp \frac{\alpha Z R}{3} {}^{V}F_{1211},$$
 (A1c)

$${}^{V}C_2 = -\frac{4}{9}R^{2V}F1110,$$
 (A1d)

using the notation by Wilkinson (1993b) to write

$${}^{V}Fk_{e}mn\rho = \frac{{}^{V}F_{000}^{(1)}(k_{e},m,n,\rho)}{{}^{V}F_{000}^{(0)}}.$$
 (A2)

Because of the property that $I(k_e, m, n, 0) \equiv 1$, we have simply

$${}^{V}F1110 = \frac{{}^{V}F_{000}^{(1)}}{{}^{V}F_{000}^{(0)}} = \frac{\langle r^{2} \rangle}{R^{2}}.$$
 (A3)

The results of Eq. (100) assumed a uniformly charged sphere with radius *R* such that $\langle r^2 \rangle_{exp} = 3R^2/5$. We can, however, do better than this. One method uses the modified Gaussian distribution of Eq. (18), where we now have one fit parameter *A*. This can be calculated using the method outlined by Wilkinson (1993c) from which the different ratios can be calculated using

$$F1111 = 0.757 + 0.0069[1 - \exp(-A/1.008)], \quad (A4)$$

$$F1221 = 0.844 - 0.0182[1 - \exp(-A/1.974)], \quad (A5)$$

$$F1222 = 1.219 - 0.0640[1 - \exp(-A/1.550)], \quad (A6)$$

such that we are consistent with our isovector correction employing a modified Gaussian charge distribution.

The Gamow-Teller shape factor was evaluated in the same spirit. Here the situation is more complicated, however, as we have more than one type of form factor contributing. The most general expression can be written as

$${}^{A}C_{0} = -\frac{1}{3}(W_{0}^{2} - 1)R^{2A}F_{101}^{(1)} \pm \frac{2}{3}\alpha Z \left(\sqrt{\frac{1}{3}}F_{110}^{(0)} + \sqrt{\frac{2}{3}}VF_{111}^{(0)}\right)$$

$$\mp \frac{2}{27}\alpha Z W_{0}R \left[-{}^{A}F_{101}^{(1)}(1, 1, 1, 1) + 2\sqrt{2}{}^{A}F_{121}^{(0)}(1, 1, 1, 1)\right]$$

$$-\frac{1}{3}(\alpha Z)^{2A}F_{101}^{(1)}(1, 2, 2, 2) - \frac{2}{3}W_{0}R \left(-\sqrt{\frac{1}{3}}F_{110}^{(0)} + \sqrt{\frac{2}{3}}VF_{111}^{(0)}\right)$$

$$+\frac{1}{9}R^{2} \left(\frac{11}{3}F_{101}^{(1)} - \frac{4}{3}\sqrt{2}{}^{A}F_{121}^{(0)}\right), \qquad (A7a)$$

$${}^{A}C_{1} = \frac{4}{3}\sqrt{2}R^{V}F_{111}^{(0)} - \frac{4}{27}W_{0}R^{2}(-5^{A}F_{101}^{(1)} + \sqrt{2}^{A}F_{121}^{(0)})$$

$$\mp \frac{2}{3}\alpha ZR[\frac{1}{9}F_{101}^{(1)}(1, 1, 1, 1, 1) - \frac{2}{9}\sqrt{2}^{A}F_{121}^{(0)}(1, 1, 1, 1) + {}^{A}F_{101}^{(0)}(1, 2, 2, 1)],$$
(A7b)

$$\label{eq:action} \begin{split} {}^{A}C_{-1} &= -\frac{2}{3}R \bigg(\sqrt{\frac{1}{3}}{}^{A}F_{110}^{(0)} + \sqrt{\frac{2}{3}}{}^{V}F_{111}^{(0)} \bigg) \\ &\quad + \frac{2}{27}W_{0}R^{2} (-{}^{A}F_{101}^{(1)} + 2\sqrt{2}{}^{A}F_{121}^{(0)}) \\ &\quad \pm \frac{\alpha ZR}{3}{}^{A}F_{101}^{(1)}(1,2,1,1), \end{split} \tag{A7c}$$

$${}^{A}C_{2} = -R^{2} \left(\frac{20A}{27}F_{101}^{(1)} - \frac{4}{27}\sqrt{2}F_{121}^{(0)}\right), \tag{A7d}$$

where we omitted a common division by ${}^{A}F_{101}^{(0)}$ in all terms for notational clarity. As mentioned in Sec. VI.F, the treatment of the Gamow-Teller shape factor is slightly more complex, since different types of form factors are present. There we split up the nuclear structure dependent parts, i.e., terms containing ${}^{V}F_{111}^{(0)}$, ${}^{A}F_{110}^{(0)}$, and ${}^{A}F_{121}^{(0)}$, from the terms containing ${}^{A}F_{101}^{(0)}(1, m, n, \rho)$. For the ratios of the latter we can use the same parametrization as for vector decay in Eqs. (A4)–(A6). The evaluation of the nuclear structure dependent terms can occur in the standard way via single-particle estimates or shell model calculations.

In this treatment we assumed only a nuclear charge distribution and neglected the influence of atomic electrons on the electrostatic potential. The largest effect was, however, already taken care of when introducing the screening correction in Sec. VII.A as the modification of the Fermi function, Eq. (9), due to screening. Further, due to the nature of the form factors coefficients all results are written in an expansion of the root mean square radius, which is matched to the experimental value for all chosen charge distributions. In order to estimate the relevance of the spatially averaged screening effect, we consider as an example the two largest contributors

$$I(k, 1, 1, 1; r) = (2k+1)r^{-2k-1} \int_0^r x^{2k} U(x) dx, \qquad (A8)$$

$$I(k, 2, 2, 1; r) = 2(2k+1)r^{-2}\int_0^r U(y)y^{-2k}$$
(A9)

$$\times \int_0^y x^{2k} U(x) dx dy, \tag{A10}$$

where U(x) is defined by

$$V(x) = -\frac{\alpha Z}{R}U(x).$$
 (A11)

Assuming now that the difference between the screened potential of Eq. (147) and that of a point charge holds also within the nucleus, the first-order expansion results in a difference $\Delta U = \sum_i \alpha_i \beta_i \sim \mathcal{O}(\alpha)$. It can be checked numerically that the shape factor of Eq. (100) changes by a few parts in 10⁴ up to large Z. In these high Z cases the order of perturbation to which all expressions are derived is insufficient to provide the highest level of precision, such that these changes are insignificant. In order to guarantee the best performance in this situation, a further expansion is required or replaced instead with a fully numerical approach.

APPENDIX B: SINGLE-PARTICLE MATRIX ELEMENTS

The Gamow-Teller shape factor contains several weak form factors which cannot always be transformed to their electromagnetic analogs through the use of CVC. In the notation of Behrens and Bühring these are ${}^{V}F_{111}^{(0)}$, ${}^{A}F_{120}^{(0)}$, and ${}^{A}F_{121}^{(0)}$. The former two are known as the weak magnetism and induced tensor contributions and can be related only to CVC results for specific transitions as discussed in Sec. VI.C. The latter needs to be evaluated in our definition of Λ in Eq. (107). For all these form factors we require their ratio with the main Gamow-Teller form factor ${}^{A}F_{101}^{(0)}$. As seen in Sec. VI.D, these form factors can be written as a combination of single-particle matrix elements when employing the impulse approximation. These results were compiled by Behrens and Bühring (1971) such that one has

$${}^{A}\mathcal{M}_{1L1}^{(0)} \approx \frac{\sqrt{2}}{\sqrt{2J_i + 1}} G_{1L1}(\kappa_f, \kappa_i) \frac{\langle r^L \rangle}{R^L}, \tag{B1}$$

$$^{A/V}\mathcal{M}_{11s}^{(0)} \approx \frac{\sqrt{2}}{\sqrt{2J_i + 1}} \left\{ \operatorname{sign}(\kappa_i) G_{11s}(\kappa_f, -\kappa_i) \times \langle g_f | \left(\frac{r}{R}\right) | f_i \rangle + \operatorname{sign}(\kappa_f) G_{11s}(-\kappa_f, \kappa_i) \langle f_f | \left(\frac{r}{R}\right) | g_i \rangle \right\},$$
(B2)

with s = 1 for ${}^{V}\mathcal{M}_{111}^{(0)}$ and s = 0 for ${}^{A}\mathcal{M}_{110}$. Here $G_{KLs}(\kappa_f, \kappa_i)$ capture all spin-angular information of the reduced matrix elements. These coefficients were introduced by Weidenmüller (1961) and are listed in Behrens and Bühring (1982). For ${}^{A}F_{121}^{(0)}$ and ${}^{V}F_{111}^{(0)}$, *K* and *s* are equal to those of ${}^{A}F_{101}^{(0)}$ such that the ratio considerably simplifies. For the former, this means the ratio can be written as

$$\frac{G_{121}(\kappa_f, \kappa_i)}{G_{101}(\kappa_f, \kappa_i)} = -\frac{C(l(\kappa_f)l(\kappa_i)2;00)}{C(l(\kappa_f)l(\kappa_i)0;00)} \begin{cases} 1 & 1 & 2\\ j_f & \frac{1}{2} & l(\kappa_f)\\ j_i & \frac{1}{2} & l(\kappa_i) \end{cases} \\ \times \begin{cases} 1 & 1 & 0\\ j_f & \frac{1}{2} & l(\kappa_f)\\ j_i & \frac{1}{2} & l(\kappa_f) \end{cases} ^{-1} \frac{\langle r^L \rangle}{R^L}, \quad (B3)$$

where $C(\dots)$ is a regular Clebsch-Gordan coefficient, and the quantities in brackets are Wigner-9*j* symbols. Here $l(\kappa) = \kappa$ if $\kappa > 0$ and $l(\kappa) = |\kappa| - 1$ if $\kappa < 0$.

For convenience, we list the relevant results from Appendix F of Behrens *et al.* (1978). These can be directly deduced from Eqs. (B1) and (B2) using the nonrelativistic approximation to express the small radial function f(r) as a function of g(r). These are categorized by the initial and final single-particle spins participating in the correction

$$\begin{split} j_{i} &= j_{f} = l + \frac{1}{2}, \\ \mathcal{M}_{101}^{0} &= \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{(l+1)(2l+3)}{2l+1} \right]^{1/2} I, \\ \mathcal{M}_{121}^{0} &= -\frac{\sqrt{2}}{\sqrt{2J_{i}+1}} l \left[\frac{2(l+1)}{(2l+1)(2l+3)} \right]^{1/2} \frac{\langle r^{2} \rangle}{R^{2}}, \\ \mathcal{M}_{111}^{0} &= -\frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[l + 1 \right] \left[\frac{6(l+1)(2l+3)}{2l+1} \right]^{1/2} \times \frac{1}{2M_{N}R} I, \\ \mathcal{M}_{110}^{0} &= -\frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3(l+1)}{(2l+1)(2l+3)} \right]^{1/2} \xi \frac{\langle r^{2} \rangle}{2R}, \\ j_{i} &= j_{f} = l - \frac{1}{2}, \\ \mathcal{M}_{101}^{0} &= -\frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{l(2l-1)}{2l+1} \right]^{1/2} I, \\ \mathcal{M}_{121}^{0} &= \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[l + 1 \right] \left[\frac{2l}{(2l-1)(2l+1)} \right]^{1/2} \frac{\langle r^{2} \rangle}{R^{2}}, \\ \mathcal{M}_{111}^{0} &= -\frac{\sqrt{2}}{\sqrt{2J_{i}+1}} l \left[\frac{6l(2l-1)}{2l+1} \right]^{1/2} \times \frac{1}{2M_{N}R} I, \\ \mathcal{M}_{110}^{0} &= -\frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3l}{(2l+1)(2l-1)} \right]^{1/2} \xi \frac{\langle r^{2} \rangle}{2R}, \\ j_{f} &= l \pm \frac{1}{2} \quad j_{i} = l \pm \frac{1}{2}, \\ \mathcal{M}_{101}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} 2 \left[\frac{l(l+1)}{2l+1} \right]^{1/2} I, \\ \mathcal{M}_{101}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} 2 \left[\frac{l(l+1)}{2(2l+1)} \right]^{1/2} \frac{\langle r^{2} \rangle}{R^{2}}, \\ \mathcal{M}_{111}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3l(l+1)}{2(2l+1)} \right]^{1/2} \times \frac{1}{M_{N}R} I, \\ \mathcal{M}_{110}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3l(l+1)}{2l+1} \right]^{1/2} \times \frac{1}{M_{N}R} I, \\ \mathcal{M}_{110}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3l(l+1)}{2l+1} \right]^{1/2} \times \frac{1}{M_{N}R} I, \\ \mathcal{M}_{110}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3l(l+1)}{2l+1} \right]^{1/2} \times \frac{1}{M_{N}R} I, \\ \mathcal{M}_{110}^{0} &= \pm \frac{\sqrt{2}}{\sqrt{2J_{i}+1}} \left[\frac{3l(l+1)}{2l+1} \right]^{1/2} \times \frac{1}{M_{N}R} I. \end{aligned}$$

with $I = \int g_f(r)g_i(r)r^2dr \approx 1$ in the nonrelativistic approximation. Here $\xi \langle r^2 \rangle / R^2$ is the evaluation of

$$\int_0^\infty g_f \{E_i - E_f - (V_i - V_f)\} \left(\frac{r}{R}\right)^2 g_i r^2 dr \qquad (B4)$$

for a spherical harmonic oscillator wave function to give

$$\xi = \frac{2\nu}{M_N} [2(n_i - n_f) + l_i - l_f]$$
(B5)

with ν the harmonic oscillator parameter defined as in Eq. (115a).

Assuming a spherical extreme single-particle approach, we then recover the results of Eq. (127). In the deformed case as discussed in Sec. VI.H.2, however, the situation is not as straightforward and we require full expressions for the different matrix elements.

APPENDIX C: MANY-PARTICLE MATRIX ELEMENTS IN *jj* COUPLING

The formulas presented in Secs. VI.F.2, VI.H.2, and Appendix B dealt with odd-A β decays, which in the extreme single-particle approach considers initial and final states consisting of one single nucleon. In Sec. VI.D we discussed that the many-particle angular momentum couplings reside in a factor C(K), which depends only on the tensor rank K of the operator. This fact is of great assistance, as it drops out completely when taking the ratio of two form factors with identical rank. In the spirit of the extreme single-particle approach, it is worthwhile to discuss odd-Z, odd-N (o-o) to even-Z, even-N (e-e) β decays and vice versa. For this we use the results by Rose and Osborn (1954a), written using the isospin formalism (Wilkinson, 1969). In line with our previous extreme single-particle methods, we consider now two particles in initial and final states, coupled to a core isospin. We then write for o-o to e-e transitions

$$\langle j_{2}j_{2}J_{f}M_{f}T_{f}T_{3f}| \sum_{n=1,2} \{O_{KLs}^{M}\tau_{\pm}\}_{n} |j_{1}j_{2}J_{i}M_{i}T_{i}T_{3i}\rangle$$

$$= \sqrt{\frac{\hat{J}_{i}\hat{J}_{f}\hat{T}_{i}\hat{T}_{f}}{1+\delta_{j_{1}j_{2}}}} (-)^{J_{f}-M_{f}} \begin{pmatrix} J_{f} & K & J_{i} \\ -M_{f} & M & M_{i} \end{pmatrix} (-)^{T_{f}-T_{3f}}$$

$$\times \begin{pmatrix} T_{f} & 1 & T_{i} \\ -T_{3f} & \pm 1 & T_{3i} \end{pmatrix} (-)^{K} \begin{cases} \frac{1}{2} & T_{f} & \frac{1}{2}(T_{i}+T_{f}) \\ T_{i} & \frac{1}{2} & 1 \end{cases}$$

$$\times \left[2 \begin{cases} j_{2} & J_{f} & j_{2} \\ J_{i} & j_{1} & K \end{cases} \langle j_{2} ||O_{KLs}||j_{1}\rangle + [1-(-)^{j_{1}+j_{2}}] \\ \\ \times \begin{cases} j_{2} & J_{f} & j_{1} \\ J_{i} & j_{2} & K \end{cases} \langle j_{2} ||O_{KLs}||j_{2}\rangle \delta_{j_{1}j_{2}} \end{bmatrix} \langle \frac{1}{2} ||t|| \frac{1}{2} \rangle, \quad (C1)$$

where the quantities in braces are Wigner-6*j* symbols. We explicitly extracted the results from the Wigner-Eckart theorem in both spin and isospin spaces. For a final matrix element one has to average over initial spin projections and sum over the final M_f . The isospin components can be directly evaluated, using $\langle 1/2 | |t| | 1/2 \rangle = \sqrt{3/2}$. An equivalent formula can be written down for *e-e* to *o-o* β decays.

In the case of deformation the angular momentum J is no longer a good quantum number, and Eq. (C1) has to be rewritten. Using the results of Berthier and Lipnik (1966), the spin-reduced matrix element for even-even to odd-odd decays is

$$\begin{split} \langle \phi(J_{f}K_{f};\Omega_{f}) | & \left| \sum_{n=1,2} \{ O_{KLs} \tau_{n}^{\pm} \} \right| | \phi(J_{i}K_{i}=0;\Omega_{i}=0) \rangle \\ &= \sqrt{\frac{\hat{J}_{i} \hat{J}_{f}}{2(1+\delta_{K_{f}0})}} \begin{pmatrix} J_{f} & K & J_{i} \\ -K_{f} & K_{f} & 0 \end{pmatrix} \times [1+(-1)^{J_{i}}] \\ &\times \sum_{j_{2}j_{1}} C_{j_{2}-\Omega_{2}} C_{j_{1}\Omega_{1}}(-1)^{j_{2}-\Omega_{2}} \begin{pmatrix} j_{2} & K & j_{1} \\ -\Omega_{2} & K_{f} & -\Omega_{1} \end{pmatrix} \\ &\times \langle j_{2} || O_{KLs} || j_{1} \rangle, \end{split}$$
(C2)

while the reverse case can be found in several publications (Berthier and Lipnik, 1966; Behrens and Bühring, 1982). Here $C_{j\Omega} = (-)^{1/2-j} \pi_{\Omega} C_{j-\Omega}$ with π_{Ω} the parity of the Ω orbital.

APPENDIX D: RELATIVISTIC COULOMB AMPLITUDES

In Sec. VI.G we discussed the influence of the relativistic matrix elements and concluded that for spectral shape measurements these are insignificant on the aimed-for level of precision. We discuss them briefly for completeness. These contributions consist of two parts, the nuclear structure embedded into the ratio of form factors and the electromagnetic influence through the slowly varying Coulomb functions. We specify here for Fermi transitions the energydependent functions using the older leptonic wave expansion by Behrens and Jänecke (1969) as shown in Table VIII.

The leading order Coulomb functions for pure Fermi transitions can be written as

$${}^{V}C(Z,W)_{\rm rel} = \frac{{}^{V}F_{011}^{(0)}}{{}^{V}F_{000}^{(0)}}{}^{V}f_{2}(W) + \frac{{}^{V}F_{011}^{(1)}}{{}^{V}F_{000}^{(0)}}{}^{V}f_{3}(W).$$
(D1)

The Coulomb functions f_2 and f_3 are written in terms of the elements in Table VIII as

$${}^{V}f_{2}(W) = -2(D_{1}+N_{1}) + 2\frac{\mu_{1}\gamma}{W}d_{1},$$
 (D2a)

$${}^{V}f_{3}(W) = -2(D_{3} + N_{1}H_{2} - N_{2}D_{1} - N_{3})$$

+ $2\frac{mu_{1}\gamma}{W}(d_{3} - N_{2}d_{1}).$ (D2b)

It is worth noting here that $D_1(1) + N_1(1) = \alpha Z/2 + W_0 R/3$ is independent of W. Typical values for $\overline{f_i(W)}$ for several $0^+ \rightarrow 0^+$ transitions can be found in Behrens and Buhring (1967) and are usually on the percent level of smaller. The relativistic matrix elements ${}^{V}F_{011}^{(N)}$ obey recursion relations after invoking CVC, specified in Eq. (89).

APPENDIX E: COMPARISON OF FINITE SIZE EFFECTS AND ELECTROMAGNETIC CORRECTIONS IN THE BEHRENS-BÜHRING AND HOLSTEIN FORMALISMS

As discussed in Sec. II, there is some confusion in the term "finite size effects" as it entails different things for different authors. Here we mainly based our approach on the rigorous work of Behrens and Bühring, culminating in the standard work by the same authors (Behrens and Bühring, 1982). We wrote some of these results in the more transparent formalism of Holstein. In light of transparency and their increasing importance in other fields not directly related to nuclear physics, we attempted to elucidate this more quantitatively.

1. Generalization including electrostatics

The simplest β decay Hamiltonian is written down as a simple current-current interaction

$ \begin{split} \bar{W}_e &= W_e \pm 3\alpha Z/(2R) \\ H_{2\sigma} &= D_{2\sigma+1} = 0 \text{ for } \sigma < 0 \end{split} $	
$H_0(k_e) = 1 H_2(k_e) = -\frac{(\bar{p}R)^2}{2(2k_e+1)}$	$ \begin{split} h_0(k_e) &= 0 \\ h_2(k_e) &= 0 \\ h_4(k_e) &= \frac{\pm a Z R}{2(2k_e+1)(2k_e+3)} \end{split} $
$D_1(k_e) = \frac{\bar{W}_e R}{2k_e + 1}$ $D_3(k_e) = -\frac{\bar{W}_e R(\bar{p}_e R)^2}{2(2k_e + 1)(2k_e + 3)} \mp \frac{\alpha Z}{2(2k_e + 3)}$	$d_1(k_e) = \frac{R}{2k_e+1}$ $d_3(k_e) = -\frac{R(\bar{p}_e R)^2}{2(2k_e+1)(2k_e+3)}$
$N_0(k_ u) = 1$ $N_2(k_ u) = -rac{(p_ u R)^2}{2(2k_ u+1)}$	$N_1(k_\nu) = \frac{p_\nu R}{2k_\nu + 1}$

$$H_{\beta} = \frac{G\cos\theta_C}{\sqrt{2}}\bar{u}(p)\gamma_{\mu}(1+\gamma_5)v(l)\langle f_{\mathbf{p}_2}|V^{\mu} + A^{\mu}|i_0\rangle + \text{H.c.},$$
(E1)

where plane waves have been used for the leptons, and \mathbf{p}_2 and $\mathbf{0}$ denote final and initial nuclear momenta. The *S* matrix is developed to first order and the corresponding spectrum is calculated using standard techniques. Given that the nucleus interacts electromagnetically with the outgoing leptons, one must incorporate the Coulomb interaction. As it is several times stronger than the weak interaction it cannot be treated perturbatively (Halpern, 1970). Neglecting the difference between initial and final Coulomb fields, the lepton wave functions are considered solutions of the Dirac equation in the final Coulomb field.³⁸

As discussed, the Behrens-Bühring formalism starts from the results of Halpern (1970). The approach taken by Holstein (1974a) and Calaprice and Holstein (1976) starts from the generalized matrix element as described by Armstrong and Kim (1972b) (Holstein, 1979)

$$\mathcal{M} = \frac{G\cos\theta_C}{\sqrt{2}} \int d^3 r \bar{\Psi}_e(\mathbf{r}, \mathbf{p}) \gamma_\mu (1 + \gamma_5) v(l) \\ \times \int \frac{d^3 k}{(2\pi^3)} e^{i\mathbf{r}\cdot\mathbf{k}} \langle f_{\mathbf{p}_2 + \mathbf{p} - \mathbf{k}} | V^\mu + A^\mu | i_0 \rangle, \qquad (E2)$$

where Ψ_e is the electron wave function in the presence of a nuclear Coulomb potential. Both results are identical up to this point after neglecting the difference in initial and final Coulomb potentials. In both the Holstein and Behrens-Bühring formalisms the nuclear current is now expanded using a series of form factors, replacing the impulse approximation results of Eqs. (83). Those of the former are useful for allowed transitions whereas those of the latter make no inherent distinction between the order of the transitions. Translation tables between both descriptions can be found in several publications (Behrens *et al.*, 1978; Behrens and Bühring, 1982) and will not be repeated here. Terms relevant

³⁸As discussed in Sec. III, neglecting this difference is corrected for in the radiative corrections of Sec. V.

for the discussion can be easily deduced from Tables III and IV.

Both approaches use as a starting point the so-called Behrens-Jänecke Fermi function, which corresponds to Eq. (9). In works by Holstein and derivatives thereof it is typically expressed as the point charge Fermi function F_0 times some correction factor as discussed in Sec. III. The crucial difference between both approaches, however, now lies in two points, both in favor of the Behrens-Bühring formulation. The first is the expansion of the electron wave function and the corresponding electrostatic finite size corrections discussed in Sec. IV. The second concerns the combination of the expansion of the nuclear current with that of the lepton wave function.

2. Approximations by Holstein and notes for the wary

We concern ourselves first with the expansion of the electron wave function. As mentioned in Secs. III and VI.E, the Behrens-Bühring approach expands these as a function of $(\alpha Z)^{\rho}$, $(WR)^{\nu-\rho}$, and $(m_e R)^{\mu-\nu}$ with the coefficients of these expansions encoded in the functions $I(|\kappa|, \mu, \nu, \rho; r)$. The latter are sensitive to the nuclear potential and examples can be found in Behrens and Bühring (1970). In the Holstein (1974a) approach one expands $\bar{\Psi}_e$ keeping only leading j = 1/2 and 3/2 terms, such that it becomes a function of f_{κ} , g_{κ} with $\kappa \in \{-2, -1, 1, 2\}$. Specific results of the expansion can be found in Armstrong and Kim (1972b) and Holstein (1974a, 1974c). Here the first difference occurs as the expansion of the latter is performed only for a uniformly charged sphere. Our discussion in Sec. IV shows that this is not sufficient, and effects of moving to a diffuse charge distribution cannot be ignored.

The calculation proceeds analogously to that of Armstrong and Kim (1972b) where a weak charge density is defined through the Fourier transform of the leading order expansion of the nuclear current. We have then

$$M = \frac{G\cos\theta_C}{\sqrt{2}} \int d^3 r \bar{\Psi}_e(\mathbf{r}, \mathbf{p}) \gamma_\mu (1 + \gamma_5) v(l) \rho^\mu(\mathbf{r}), \quad (E3)$$

where $\rho^{\mu}(\mathbf{r})$ is simply the Fourier transform of the involved form factors

$$\rho^{\mu}(\mathbf{r}) = \int \frac{d^3s}{(2\pi)^2} e^{i\mathbf{r}\cdot\mathbf{s}} \langle f_{\mathbf{p}_2+\mathbf{p}-\mathbf{k}} | V^{\mu} + A^{\mu} | i_0 \rangle.$$
(E4)

The second difference now appears in the expansion of the nuclear current and the consequent artificial split of nuclear structure terms and the Coulomb corrections in the Holstein approach. This is in part because Coulomb corrections to induced corrections were neglected (Holstein, 1974a), such that for Gamow-Teller decay it is approximated as

$$\langle f_{\mathbf{p}_2} | V^{\mu} + A^{\mu} | i_0 \rangle \approx -g^{\mu\kappa} c (W_0^2 - p_2^2) C_{J'1;J}^{M'\kappa;M},$$
 (E5)

with C a regular Clebsch-Gordan coefficient. The weak charge density is then typically normalized by extracting a factor c(0).

Here once again great care must be taken when comparing to the Holstein formalism, as the corrections are written differently in different papers (Holstein, 1974a, 1974c, 1979; Calaprice and Holstein, 1976) by the same authors. In particular, the Fermi function used differs by a factor $(1 + \gamma)/2$ depending on the publication, resulting in changes of order $(\alpha Z)^2$ when naively comparing formulas. Further, the redefinition of the so-called spectral functions taking into account Coulomb interactions has been written in two different ways. The first (Holstein, 1974c) replaces

$$h_i^{(0)}(W) \to F(Z, W)[h_i^{(0)}(W) + \Delta h_i(W)],$$
 (E6)

where $F(Z, W) = (1 + \gamma)/2F_0$, with $\Delta h_i(W)$ defined in those works. The later works (Calaprice and Holstein, 1976; Holstein, 1979) artificially split the transition matrix element of Eq. (E3) into a nuclear structure part and their Coulomb corrections

$$N(W)dW \propto \tilde{h}_{1}(W) \left[|A|^{2} + |B|^{2} + |C|^{2} + |D|^{2} + \frac{2}{3} \operatorname{Re}(A^{*}D + B^{*}C) + 2\frac{m_{e}}{W} \operatorname{Re}\left(A^{*}B + C^{*}D + \frac{1}{3}A^{*}C + \frac{1}{3}B^{*}D\right) - \frac{2}{9}\frac{p}{W} \operatorname{Re}(A^{*}F - B^{*}G + 3F^{*}D - 3G^{*}C) \right], \quad (E7)$$

with

$$\tilde{h}_{1}(W) = c_{1}^{2} - \frac{2}{3} \frac{W_{0}}{M} c_{1}(c_{1} + d \pm b) + \frac{2}{3} \frac{W}{M} [c_{1}(5c_{1} \pm 2b)] - \frac{m_{e}^{2}}{3MW} \left[2c_{1}^{2} + c_{1}(d \pm 2b) - c_{1}h \frac{W_{0} - W}{2M} \right], \quad (E8)$$

and A-G denoting integrals of the different parts of the leptonic expansion with the weak charge distribution of Eq. (E4). These are given by Holstein (1974c) and will not be repeated here for brevity. Here the Fermi function consistently refers to the so-called Behrens-Jänecke Fermi function, discussed in more detail in the following section. It is Eqs. (E7) and (E8) that should be compared against the combination F_0L_0UC in the current work. For clarity, we continue with the later results of Eqs. (E7) and (E8) and refer only to the older works when necessary in Appendix F.

Expanding Eq. (E5) to zeroth order will introduce the Fermi function, while the first order introduces the leptonic convolution. These will be discussed separately. Finally, the inclusion of the Coulomb corrections to induced currents is briefly touched upon.

3. Fermi function and finite size correction

The Coulomb corrections of Eq. (E7) depend on the precise evaluation of the integrals A-G, which are in turn dependent on the weak charge distribution of Eq. (E4) (Calaprice and Holstein, 1976). In a first approximation we define the Gamow-Teller form factor $c(q^2)$ as a constant, in which case

 $\rho(r)$ becomes a simple Dirac delta function. Now only A and B survive such that

$$|A|^{2} + |B|^{2} = \frac{1}{2}N^{2}\frac{2m_{e}}{W}\int d^{3}r\delta^{3}(\mathbf{r}) \times \left[g_{-1}^{2}(r) + \left(\frac{W}{p}\right)^{2}f_{1}^{2}(r)\right]$$

= $F_{0}L_{0}$ (E9)

when taking into account the differing normalization definitions of the wave functions in the Behrens-Bühring and Holstein formalisms. As f_1 and g_{-1} have been approximated only for a uniformly charged sphere for low Z in the Holstein approach, the expression obtained by Huffaker and Laird (1967) and Calaprice and Holstein (1976)

$$|A|^2 + |B|^2 \approx F_0[1 \mp \frac{13}{15}\alpha ZWR]$$
 (E10)

is clearly less precise than our Eq. (16) as was discussed already in Sec. III. The term can easily be recognized as a simplified version of that found in Eq. (16). Further, it contains no corrections stemming from a diffuse nuclear charge, for which we derived explicit expressions and analytical parametrizations in Sec. IV.A.2.

4. Nuclear and leptonic convolution

The finite nuclear size does not only affect the Coulomb potential as felt by the charged lepton but also the behavior of its wave function near the origin. The interaction volume becomes a sphere with radius R, and thus the matrix element requires an average for all the nucleon positions. This is clearly visible in Eq. (E3) and is inherently present in our definition of the *C* factor of Sec. VI.E. In the initial results by Huffaker and Laird (1967) the shape factor contains an additional term

$$C_0(W) = \frac{\langle r^2 \rangle}{R^2} \alpha Z W R, \qquad (E11)$$

as a remnant of explicit averaging of f_1 and g_{-1} inside the nuclear volume. For a constant nuclear wave function, i.e., a rectangle of width *R*, one finds $\langle r^2 \rangle = (3/5)R^2$ so that $C_0(W) = (3/5)\alpha ZWR$. The approach of Holstein and coauthors builds on this result and provides a more consistent discussion by introducing the elementary particle treatment. We move beyond the approximation of the previous section and now expand Eq. (E5) to first order in q^2 , $c(q^2) \approx$ $c_1 + c_2q^2$ to find

$$\rho \approx \int \frac{d^3k}{(2\pi)^3} e^{i\mathbf{r}\cdot\mathbf{k}} \left(1 + \frac{c_2}{c_1}q^2\right)$$
$$= \left(1 + \frac{c_2}{c_1}(W_0^2 + \nabla^2)\right) \delta^3(\mathbf{r}), \quad (E12)$$

where ∇ is the gradient operator. In this manner, all further expansions in q^2 will always result in a constant factor times $\delta^3(\mathbf{r})$ as $\mathcal{F}[q^n] \propto \nabla^n \delta^3(\mathbf{r})$. This entails that all leptonic radial wave functions will still be evaluated at the nuclear center, be it multiplied with some constant factor depending on nuclear structure. As such, all results will always be proportional to the Fermi function defined at the origin, such that it can always be meaningfully extracted as we have always done so far. Again the results presented by Holstein and colleagues have been listed only for uniformly charged spheres. Using Eq. (E12) we find only *A*, *D*, and *F* integrals survive, all of which are proportional to the square root of the Fermi function as expected (Calaprice and Holstein, 1976). Equations (13) and (A.16) in Calaprice and Holstein (1976) are then to be compared against Eqs. (E11) and (106d). A critical ingredient in the quantitative connection between the different formalisms lies in the evaluation of c_2/c_1 . In the impulse approximation it can be written as (Calaprice and Holstein, 1976)

$$\frac{c_2}{c_1} \approx \frac{1}{6} \frac{\langle \beta || \tau^{\pm} \sigma r^2 || \alpha \rangle}{\langle \beta || \tau^{\pm} \sigma || \alpha \rangle} + \frac{1}{6\sqrt{10}} \frac{\langle \beta || \tau^{\pm} [\sigma \times \sqrt{\frac{16}{5} \pi} Y_2(r)] || \alpha \rangle}{\langle \beta || \tau^{\pm} \sigma || \alpha \rangle}$$
(E13)

and is related to the shape of nuclear wave functions. In the initial isospin invariant formalism, these were always treated as rectangles with only nonzero values for r < R, i.e., a uniformly charged sphere with radius R. Using this, one arrives at $c_2/c_1 = (1/10)R^2$. The correction factor to the Behrens-Jänecke Fermi function (i.e., F_0L_0 in our formalism) in the Holstein formalism is then written as

$${}_{HS}C(Z,W)_0 \approx 1 + R^2/5 - (W_0R)^2/5 - \frac{9}{20}(\alpha Z)^2$$

$$\pm \frac{2}{15}\alpha ZW_0R \mp \frac{11}{15}\alpha ZWR$$

$$+ \frac{2}{5}W_0R^2W - \frac{2}{5}R^2W^2. \tag{E14}$$

If we use the older expansion of $\overline{\Psi}_e$ (Behrens and Jänecke, 1969; Wilkinson, 1990), for Gamow-Teller decay

$$_{old}C(Z,W)_0 = 1 + {}^{A}C_0 + {}^{A}C_1W + {}^{A}C_2W^2,$$
 (E15)

where

$${}^{A}C_{0} = -\frac{9}{20}(\alpha Z)^{2} - (W_{0}R)^{2}/5 \pm \alpha ZW_{0}R/15 + R^{2}/5,$$

$${}^{A}C_{1} = \mp 2\alpha ZR/3 + 4W_{0}R^{2}/9,$$

$${}^{A}C_{2} = -4R^{2}/9.$$
 (E16)

It is clear every term can be identified with a similar one in Eq. (E14), except for a factor of 2 difference in the αZW_0R term. All of these results are again valid only for a uniformly charged sphere, both in the expansion of the lepton wave function and in the calculation of the weak charge density. In Sec. A we looked at the deviation of the former due to a more realistic charge distribution. Further, to correct for the breakdown of the latter we introduced the isovector correction C_I in Sec. VI.F. Because of its large contribution this can for certain not be neglected. In combination with a more precise expansion of the lepton wave functions, it is clear the approach presented in this work is superior to that of Holstein.

5. Induced terms

Finally, the Holstein formalism requires an adjustment for the approximation introduced in Eq. (E5). Initially, Coulomb corrections to the induced terms in the nuclear current were neglected as the latter are already suppressed by a factor q/M. As discussed by Bottino, Ciochetti, and Ciocchetti (1973) and Bottino, Ciochetti, and Kim (1974), however, this approximation is not strictly valid. The reasoning starts from the result by Armstrong and Kim (1972b) showing that $\langle f(\mathbf{p}_f) |$ should be replaced by $\langle f(\mathbf{p}_f + \mathbf{p}_e - \mathbf{p}) |$, where \mathbf{p} is an internal momentum of the Fourier transform. This has the consequence that induced terms are now a function of q' = $(\mathbf{p}_f + \mathbf{p}_e - \mathbf{p}) - \mathbf{p}_i$ rather than $q = \mathbf{p}_f - \mathbf{p}_i$, thereby introducing Coulomb corrections to the induced terms. This can be seen from the fact that the additional term

$$\frac{1}{(2\pi)^3} \int d\vec{p} \left[\int d\vec{r} \left(\frac{\bar{\Psi}_e(\mathbf{r}, \mathbf{p}_e)}{\bar{u}_e(\mathbf{p}_e)} \right) e^{i\mathbf{r}\cdot\mathbf{p}} \right] \\ \times \frac{F_M(q')}{2m_p} \times \sigma^{\alpha\beta} (\mathbf{p}_e - \mathbf{p})_\beta$$
(E17)

becomes identically zero when $\bar{\Psi}_e$ reduces to $e^{-i\mathbf{r}\cdot\mathbf{p}_e}\bar{u}_e(\mathbf{p}_e)$ in the absence of Coulomb interactions. The correction factor given by Holstein (1974b) and Calaprice and Holstein (1976)

$$\delta h_1(Z) \approx \frac{\sqrt{10}}{6} \frac{\alpha Z}{MR} c_1 (2b + d^{II} \pm d^I \pm c_1) \quad \text{(E18)}$$

can be obtained from the results of Eqs. (9) and (13) in Bottino, Ciochetti, and Kim (1974) after proper conversion from F_M (F_A) to b (c) using the definition of the nuclear currents [Eq. (5) in Bottino, Ciochetti, and Kim (1974) and Eq. (1) in Holstein (1974a)], with an exception of the d^I and d^{II} terms. The results from (Bottino, Ciochetti, and Kim (1974) can easily be extended to include this result, and a quick glance toward the behavior of b and d terms leads directly to Eq. (E18). A discussion on the signs for electron and positron decay can be found in Holstein (1974a). As there is no artificial separation of nuclear structure and Coulombic terms and all terms of the nuclear current are retained, these corrections occur naturally in the C factor of the Behrens-Bühring formalism followed in this work.

6. Summary

While the Holstein formalism initially profits from a more transparent description of the different factors and their origins participating in the β spectrum shape, several approximations introduced along the way limit the usefulness of the quoted final formulas when requiring a high precision description. Several terms have to be added, such as Coulomb corrections to induced currents, thereby introducing confusion as to the origin of these terms and how they play with the other results. This is further hindered by a difference in definitions throughout several papers about the correct manner of introducing Coulomb corrections. We have looked at individual parts of the calculation and for each of them discussed the superiority of the formalism presented in this work.

To facilitate the interpretation of experimental results and reduce notational clutter, the formulas in Sec. VI.E, while calculated in the Behrens-Bühring formalism, are presented using the well-known Holstein form factors.

APPENDIX F: VALIDITY OF HARMONIC OSCILLATOR FUNCTIONS

Recently the issue of finite size corrections was addressed by Wang, Friar, and Hayes (2016), incorporating density functional theory (DFT) results to calculate the weak charge density. The analytical formulation to first order in αZ and Rby Holstein (1974a) was used and rewritten in terms of nuclear Zemach moments (Zemach, 1956)

$$\begin{split} \delta_{\mathrm{FS}}^{\mathrm{Wang}} &= -\frac{\alpha Z}{3} \left(4W \langle r \rangle_{(2)} + W \langle r \rangle_{(2)}^r - \frac{W_{\nu} \langle r \rangle_{(2)}^r}{3} \right. \\ &\left. + \frac{1}{W} (2 \langle r \rangle_{(2)} - \langle r \rangle_{(2)}^r) \right), \end{split} \tag{F1}$$

where $\langle r \rangle_{(2)}$ and $\langle r \rangle_{(2)}^r$ are defined as

$$\langle r \rangle_{(2)} = \int d^3 s s \int d^3 r \rho_{\rm w}(r) \rho_{\rm ch}(|\vec{r} - \vec{s}|),$$
 (F2)

$$\langle r \rangle_{(2)}^{r} = \int d^{3}ss \int d^{3}r \rho_{\rm w}(r) r \frac{\partial}{\partial r} \rho_{\rm ch}(|\vec{r} - \vec{s}|).$$
(F3)

Here $\rho_{\rm w}$ and $\rho_{\rm ch}$ are the weak and regular charge densities, respectively. As discussed in the previous section, care must be taken when looking to compare Eq. (F1) to those presented here. Holstein (1974a) defined the Coulomb correction to the spectral function through Eq. (E6), where $\Delta h_1(W)$ is given by a simplified version of the Coulomb terms of Eq. (E7) without the last term and subtracting the Fermi function $(1 + \gamma)/2F_0$. This is then developed to only first order in αZ to yield Eq. (25) in Holstein (1974a). It is thus a combination of electrostatic finite size effects contained in L_0 and the leptonic convolution. The previous section discussed the improvements made by both formalisms, and we will not spend any more time on explicitly showing the correspondence to the formulas presented there. We will instead treat the evaluation of the Zemach moments in a purely spherical harmonic oscillator fashion and compare the results with those of Wang, Friar, and Hayes (2016) obtained through DFT. This will serve as a benchmark for the validity of using harmonic oscillator wave functions in the evaluation of our expanded results.

As previously discussed, a uniform density is not appropriate for the weak charge density. As a result, we added the C_I correction discussed in Sec. VI.F. Here we introduced the rms radius of the weak charge density $\langle r^2 \rangle_w$ as the essential parameter in our correction. Wang, Friar, and Hayes (2016) calculated this quantity for a series of nuclei ranging from A = 14 to 120 using both DFT and uniformly charged radius results. We presented the way to analytically calculate these radii using a charge-insensitive harmonic oscillator distribution model. We used the charge radii listed by Wang, Friar, and Hayes (2016) to obtain an optimal comparison of our

TABLE IX. Comparison of DFT and analytical results using both uniformly charged sphere and harmonic oscillator (HO) functions as described in the text. We compare the weak charge radius $\langle r^2 \rangle_w$ used in our isovector correction [Eq. (113), column 3], together with the Zemach moments discussed by Wang, Friar, and Hayes (2016) for several β^- transitions. Here *BB* stands for the approximations listed in Eqs. (F4) and (F5). Units are in fm and fm². These were used together with a uniform and harmonic oscillator charge distribution.

	$\langle r^2 \rangle_{\rm w}^{\rm Wang}$	$\langle r^2 \rangle_{\rm w}^{\rm HO}$	$\langle r \rangle_{(2)}^{UniBB}$	$\langle r \rangle_{(2)}^{\text{HO}-BB}$	$\langle r \rangle_{(2)}^{\text{Wang}}$
$\frac{14}{14}C/14N$	8.78	7.77	3.52	3.67	3.66
²⁵ Na/ ²⁵ Mg	11.63	11.82	4.01	4.41	4.22
³⁵ S/ ³⁵ Cl	13.77	12.97	4.43	4.70	4.62
⁴⁵ Ca/ ⁴⁵ Sc	17.12	18.09	4.71	5.38	5.07
$^{61}Cr/^{61}Mn$	18.88	19.02	5.02	5.58	5.36
⁶⁴ Co/ ⁶⁴ Ni	18.87	18.97	5.12	5.60	5.41
100 Nb / 100 Mo	25.80	27.17	5.86	6.62	6.28
104 Nb/ 104 Mo	26.14	27.77	6.07	6.73	6.33
$^{121}Sn/^{121}Sb$	28.11	33.81	6.26	7.31	6.56

method. This was plugged into Eq. (115a) to evaluate $\langle r^2 \rangle_w$ for the last occupied transforming nucleon. We used the approximation termed as "Behrens-Bühring," which involves approximating the form factor $c(q^2) \approx c_1(1 + \langle r^2 \rangle_w/6)$. The Zemach moments are then reduced to

$$\langle r \rangle_{(2)} \approx \langle r \rangle_{\rm ch} + \frac{\langle r^2 \rangle_{\rm w} \langle r^{-1} \rangle_{\rm ch}}{3},$$
 (F4)

$$\langle r \rangle_{(2)}^r \approx \frac{2 \langle r^2 \rangle_{\rm w} \langle r^{-1} \rangle_{\rm ch}}{3}.$$
 (F5)

The last of these is a poor approximation, as can be seen from numerical results listed by Wang, Friar, and Hayes (2016). If we attempt to extend this approximation by expanding *c* to $\mathcal{O}(q^4)$, dimensional analysis tells us this factor will be proportional to $-\langle r^4 \rangle_w \langle r^{-3} \rangle_{ch}$, which is not convergent. We thus keep in mind that Eq. (F5) tends to overestimate the integral by about 30%. Table IX shows a comparison of our analytical methods with DFT results for the relevant parameters in our corrections.

While the agreement is fair for lower masses, differences appear for the highest masses that were investigated. The reason for this is twofold.

- (1) The approximation we used by considering only the final decaying neutron is not any more valid for the higher mass A > 100 regions. Here the difference between quantum numbers for proton and neutron orbitals becomes too large and Eq. (115a) is an unsustainable estimate. A more correct approach is then to calculate the actual overlap integral $\int dr r^2 R_{nl}^{\alpha} R_{n'l'}^{\beta} / \int dr R_{nl}^{\alpha} R_{n'l'}^{\beta}$ where α and β are the neutron and proton harmonic oscillator wave functions.
- (2) Our harmonic oscillator functions are charge insensitive, while the DFT results are not. We expect then an additional effect arising from the breakdown of isospin invariance, introducing the so-called nuclear mismatch δ_C (Towner and Hardy, 2010). This has a strong Z dependent behavior as is to be expected, and is a

worthy subject in its own right. This is, however, beyond the scope of the work here.

Application of point 1 yields results accurate to within 10% of the DFT results. The remaining discrepancy we attribute to inherent limitations of the harmonic oscillator treatment and to a lesser extent the nuclear mismatch problem of point 2. The latter can be assumed to drop out to first order when taking ratios of matrix elements suffering from the same issue. This precision is, however, sufficient for our current purposes, and we conclude the harmonic oscillator wave functions yield appropriate results. An extension of the latter to deformed nuclei is straightforward, as discussed in several places throughout this text. Results obtained in a deformed Woods-Saxon potential using harmonic oscillator basis functions show excellent agreement with experimental weak magnetism data, extensively discussed by Severijns et al., 2018. This allows us to put our trust into the approach presented here.

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