Chiral Effective Field Theory Calculations of Weak Transitions in Light Nuclei

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We report Quantum Monte Carlo calculations of weak transitions in $A \leq 10$ nuclei, based on the Norfolk two- and three-nucleon chiral interactions, and associated one- and two-body axial currents. We find that the contribution from two-body currents is at the 2–3% level, with the exception of matrix elements entering the rates of 8 Li, 8 B, and 8 He beta decays. These matrix elements are suppressed in impulse approximation based on the (leading order) Gamow Teller transition operator alone; two-body currents provide a 20–30% correction, which is, however, insufficient to bring theory in agreement with experimental data. For the other transitions, the agreement with the data is satisfactory, and the results exhibit a negligible to mild model dependence when different combinations of Norfolk interactions are utilized to construct the nuclear wave functions. We report a complete study of two-body weak transition densities which reveals the expected universal behavior of two-body currents at short distances throughout the range of A=3 to A=10 systems considered here.

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I. INTRODUCTION

In this work, we present Quantum Monte Carlo (QMC) calculations, including both variational Monte Carlo (VMC) and Green's function Monte Carlo (GFMC) calculations, of Gamow-Teller (GT) matrix elements entering beta-decay and electron-capture rates in A = 3-10nuclei. These observables are experimentally known (in most cases) at the sub-percent level, and are used here primarily to validate our microscopic theoretical modeling of the nucleus as a system of nucleons interacting with each other via effective interactions, and with electroweak probes via effective currents. Specifically, this modeling is based on local two- and three-nucleon interactions formulated in configuration space, and derived from a chiral effective field theory (χEFT) that retains, in addition to nucleons and pions, Δ -isobars as explicit degrees of freedom [1–4]. They are referred to below as the Norfolk interactions and are denoted as NV2+3. Accompanying these interactions are one- and two-body axial currents—local in configuration space—derived within the same χ EFT formulation [3–5]. In the calculations to follow, we include up to tree-level contributions at nextto-next-to-next-to-leading order (N3LO) in the chiral expansion, and disregard subleading corrections involving loops and higher order contact terms.

An analogous study was recently reported by some of the present authors in Ref. [6]. There, the QMC calculations were based on the Argonne- v_{18} (AV18) two-nucleon [7] and Illinois-7 (IL-7) three-nucleon [8] interactions, in combination with the axial currents of Ref. [5]. We found agreement with the experimental Gamow-Teller matrix elements at the $\approx 2\%$ level for the A=6

and 7 systems, and at the $\approx 10\%$ level in 10 C. The large uncertainty in the A=10 transition is primarily systematic and results from the narrow energy separation between the first two $J^{\pi}=1^+$ states in 10 B, which makes it hard to precisely disentangle them [6]. The study of Ref. [6] found that two-body currents generate an additive contribution of the order of $\approx 3\%$ and concluded that the agreement with the data is mainly attributable to the use of fully correlated nuclear wave functions, rather than two-body effects in the currents.

In the meantime, no-core shell-model calculations of weak matrix elements based on chiral interactions and currents [9] found the sign of the overall correction generated by two-body currents to be opposite to that obtained in Ref. [6] for the same systems (but in agreement with a hybrid calculation of the A=6 decay reported in Ref. [10]). This discrepancy was attributed to the hybrid nature of the calculation of Ref. [6], *i.e.*, to the mismatch between the two- and three-body correlations implemented to construct the nuclear wave functions—and induced by the AV18 and IL7—and those entering the axial currents which were instead derived from χ EFT.

In this work, by reexamining the evaluation of these weak matrix elements with the NV2+3 chiral interactions [1–4], in combination with consistent chiral axial currents at tree-level [5], we aim to address and explore the aforementioned claim. We investigate the sensitivity of the calculated matrix elements with respect to different choices of regulators and to different strategies adopted to constrain the three-body Norfolk interactions (NV3). This latter aspect is important in order to understand the interplay between these interactions and the axial currents, since the strength of the contact current and

that of the three-body interaction of one-pion-range are rigorously related to each other by the symmetries imposed in the χ EFT formulation [11–13].

This study has several merits. First, we report new GFMC results of the energy spectra of $A \leq 10$ nuclei based on two classes of NV2+3 interactions. In addition to the systems studied in Ref. [6], we study weak transitions in A=8 nuclei where we find that two-body axial currents provide a large correction to the one-body results. Finally, we provide the first calculations of two-body weak transition densities which shed light on the role of short-range physics in these observables.

Searches for physics beyond the Standard Model (BSM) via beta decay are the focus of current and planned experimental programs carried out at the Facility for Rare Isotope Beams (FRIB), the the University of Washington, and Argonne National Laboratory (see, e.g., Ref. [14] and references therein). Among the targets under consideration are ⁶He, ⁸Li, ⁸B and ¹⁰C. A systematic study of axial-current matrix elements in these systems is a prerequisite for all further investigations and BSM searches in beta decay. Beta decays are ideal processes to assess the validity of the dynamical inputs of ab inito calculations, namely many-body correlations and weak currents. The latter also impact calculations of neutrinoless double beta decay matrix elements, whose knowledge is critical to the neutrinoless double beta decay experimental program [15].

This paper is structured as follows: a brief review of the QMC computational method and Norfolk interactions is given in Secs. II and III. The many-body axial currents used in this work are reported in Sec. IV. Results and conclusions are provided in Secs. V and VI.

II. QUANTUM MONTE CARLO METHOD

The Quantum Monte Carlo methods used in this study have been described in detail in several review articles, the most recent of which being Refs. [16, 17]. Here, we sketch the computational procedure and refer the interested reader to Refs. [16, 17] and references therein.

We seek accurate solutions of the many-nucleon Schrödinger equation

$$H\Psi(J^{\pi}; T, T_z) = E\Psi(J^{\pi}; T, T_z) , \qquad (1)$$

where J^{π} are the total angular momentum and parity of the state, and T and T_z are the total isospin and its projection, respectively. We use the Hamiltonian

$$H = \sum_{i} K_{i} + \sum_{i < j} v_{ij} + \sum_{i < j < k} V_{ijk} , \qquad (2)$$

where K_i is the non-relativistic kinetic energy, and v_{ij} and V_{ijk} are the NV2 and NV3 local chiral interactions [1–4], collectively denoted as NV2+3.

The VMC trial function $\Psi_V(J^{\pi}; T, T_z)$ for a given nucleus is constructed from products of two- and three-body

correlation operators acting on an antisymmetric singleparticle state of the appropriate quantum numbers. The correlation operators are designed to reflect the influence of the interactions at short distances, while appropriate boundary conditions are imposed at long range [18–21]. The $\Psi_V(J^\pi;T,T_z)$ contains variational parameters that are adjusted to minimize the expectation value

$$E_V = \frac{\langle \Psi_V | H | \Psi_V \rangle}{\langle \Psi_V | \Psi_V \rangle} \ge E_0 , \qquad (3)$$

which is evaluated by Metropolis Monte Carlo integration [22]. The lowest value for E_V is then taken as the approximate ground-state energy of the exact lowest eigenvalue of H, E_0 , for the specified quantum numbers.

A good trial wave function is given by

$$|\Psi_V\rangle = \mathcal{S} \prod_{i< j}^A \left[1 + U_{ij} + \sum_{k \neq i, j}^A \widetilde{U}_{ijk}^{TNI} \right] |\Psi_J\rangle.$$
 (4)

The Jastrow wave function Ψ_J is fully antisymmetric and has the $(J^{\pi}; T, T_z)$ quantum numbers of the state of interest, while U_{ij} and $\widetilde{U}_{ijk}^{TNI}$ are two- and three-body correlation operators.

The GFMC method [16] improves on the VMC wave functions by acting on Ψ_V with the operator $\exp[-(H-E_0)\tau]$. The operator is applied in a sequence of small imaginary-time steps $\Delta\tau$ to produce a propagated wave function

$$\Psi(\tau) = e^{-(H - E_0)\tau} \Psi_V = \left[e^{-(H - E_0)\Delta\tau} \right]^n \Psi_V . \tag{5}$$

Obviously $\Psi(\tau=0) = \Psi_V$ and $\Psi(\tau \to \infty) = \Psi_0$. Quantities of interest are evaluated in terms of a "mixed" expectation value between Ψ_V and $\Psi(\tau)$:

$$\langle O(\tau) \rangle_M = \frac{\langle \Psi(\tau) | O | \Psi_V \rangle}{\langle \Psi(\tau) | \Psi_V \rangle},$$
 (6)

where the operator O acts on the trial function Ψ_V . The desired expectation values would, of course, have $\Psi(\tau)$ on both sides; by writing $\Psi(\tau) = \Psi_V + \delta \Psi(\tau)$ and neglecting terms of order $[\delta \Psi(\tau)]^2$, we obtain the approximate expression

$$\langle O(\tau) \rangle = \frac{\langle \Psi(\tau)|O|\Psi(\tau) \rangle}{\langle \Psi(\tau)|\Psi(\tau) \rangle}$$

$$\approx \langle O(\tau) \rangle_M + [\langle O(\tau) \rangle_M - \langle O \rangle_V] , \qquad (7)$$

where $\langle O \rangle_{\rm V}$ is the variational expectation value.

For off-diagonal matrix elements required by the transitions we are interested here, the generalized mixed estimate is given by the expression

$$\frac{\langle \Psi^{f}(\tau)|O|\Psi^{i}(\tau)\rangle}{\sqrt{\langle \Psi^{f}(\tau)|\Psi^{f}(\tau)\rangle}\sqrt{\langle \Psi^{i}(\tau)|\Psi^{i}(\tau)\rangle}}$$

$$\approx \langle O(\tau)\rangle_{M_{i}} + \langle O(\tau)\rangle_{M_{f}} - \langle O\rangle_{V} , \qquad (8)$$

\tilde{g}_A	h_A	F_{π}	c_1	c_2	c_3	c_4	$b_3 + b_8$
1.29	2.74	184.80	-0.57	-0.25	-0.79	1.33	1.40

TABLE I: Values of (fixed) low energy constants (LECs) used in this work: \tilde{g}_A and h_A are adimensional, $F_{\pi} = 2f_{\pi}$ is given in MeV, and the remaining LECs are given in GeV⁻¹. See text for explanation.

where

$$\langle O(\tau) \rangle_{M_f} = \frac{\langle \Psi^f(\tau) | O | \Psi_V^i \rangle}{\langle \Psi^f(\tau) | \Psi_V^f \rangle} \sqrt{\frac{\langle \Psi_V^f | \Psi_V^f \rangle}{\langle \Psi_V^i | \Psi_V^i \rangle}} , \qquad (9)$$

and $\langle O(\tau)\rangle_{M_i}$ is defined similarly. For more details see Eqs. (19)–(24) and the accompanying discussions in Ref. [23].

III. NORFOLK INTERACTION MODELS

We base our calculations of weak transitions in A=6-10 on the local NV2 and NV3 interactions developed in Refs. [1–4]. The NV2 model has been derived from a χ EFT that uses pions, nucleons and Δ 's as fundamental degrees of freedom. It consists of a long-range part, $v_{ij}^{\rm L}$, mediated by one- and two-pion exchanges, and a short-range part, v_{ij}^{S} , described in terms of contact interactions with strengths specified by unknown low-energy constants (LECs). The strength of the long-range part is fully determined by the nucleon and nucleon-to- Δ axial coupling constants g_A and h_A , the pion decay amplitude F_{π} , and the LECs c_1 , c_2 , c_3 , c_4 , and $b_3 + b_8$, constrained by reproducing πN scattering data [24]. The LECs entering the contact interactions are fixed by fitting nucleon-nucleon scattering data from the most recent and up-to-date database collected by the Granada group [25–27]. The value for h_A is taken from the large N_c expansion or strong-coupling model [28]. The value of the nucleon axial coupling constant used to construct the nucleon-nucleon interaction accounts for the Goldberger-Treiman discrepancy [29, 30] and, to distinguish it from the experimental value of $g_A = 1.2723(23)$ [31] entering the axial currents, we denote it with \tilde{g}_A . For completeness, Tables I and II report the values of these constants, along with the pion and nucleon masses, the Δ -nucleon mass difference, the electron mass, and the fine structure constant α used in the NV2 interactions (these last two characterize the electromagnetic part of the NV2s [1]).

The contact terms are implemented using a Gaussian representation of the three dimensional delta function, with R_S denoting the Gaussian parameter [1–4]. The pion-range operators are strongly singular at short range in configuration space, and are regularized by a radial function characterized by a cutoff R_L [1–4]. There are two classes (I and II) of NV2s, differing only in the range of energy over which they are fitted to the database—class I up to 125 MeV, and class II up to 200 MeV. For

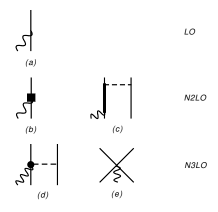


FIG. 1: Diagrams illustrating the contributions to the axial current up to N3LO used in this work. Nucleons, Δ -isobars, pions, and external fields are denoted by solid, thick-solid, dashed, and wavy lines, respectively. The square in panel (b) represents relativistic corrections, while the dot in panels (d) denotes a vertex induced by subleading terms in the π -nucleon chiral Lagrangian [3].

each class, two combinations of short- and long-range regulators have been used, namely (R_S, R_L) =(0.8, 1.2) fm (models NV2-Ia and NV2-IIa) and (R_S, R_L) =(0.7, 1.0) fm (models NV2-Ib and NV2-IIb). Class I (II) fits about 2700 (3700) data points with a $\chi^2/\text{datum} \lesssim 1.1$ ($\lesssim 1.4$) [1, 2].

The NV2 models were found to provide insufficient attraction in GFMC calculations of the binding energies of light nuclei [2]. To remedy this shortcoming, a consistent three-body interaction was constructed up to N2LO in the chiral expansion. It consists of a long-range part mediated by two-pion exchange and a short-range part parametrized in terms of two contact interactions [32, 33] proportional to the LECs c_D and c_E . These LECs have been obtained by fitting either observables that involve exclusively strong interactions [34–37] or a combination of observables that involve both strong and weak interactions [4, 12, 38]. This last strategy is feasible because of the relation established in χ EFT [11] that links c_D with the LECs entering the contact axial current at N3LO [12, 13, 38] (see next section for details).

In Ref. [37], c_D and c_E were determined by simultaneously reproducing the experimental trinucleon ground-state energies and nd doublet scattering length. These first-generation NV2+3 interactions, denoted with NV2+3-Ia/b and NV2+3-IIa/b, have been implemented in both VMC and GFMC codes and used to study static properties of light nuclei [2, 17, 37, 39–41], and in auxiliary-field diffusion Monte Carlo (AFDMC) [42], Brueckner-Bethe-Goldstone (BBG) [43, 44] and Fermi hypernetted chain/single-operator chain (FHNC/SOC) [45, 46] approaches to investigate the equation of state of neutron matter [47, 48].

In more recent work [4], c_D and c_E were constrained by fitting, in addition to the trinucleon energies, the empir-

m_{π_0}	$m_{\pi_{\pm}}$	M_n	M_p	$m_{\Delta N}$	m_e	α^{-1}
134.9766	139.5702	939.56524	938.27192	293.1	0.510999	137.03599

TABLE II: Values of charged and neutral pion masses, proton and neutron masses, Δ -nucleon mass difference, and electron mass (all in MeV), and of the (adimensional) fine structure constant α . Note that $\hbar c$ is taken as 197.32697 MeV fm.

	Ia (Ia*)	Ib (Ib*)	IIa (IIa*)	IIb (IIb*)
c_D	3.666 (-0.635)	-2.061 (-4.71)	1.278 (-0.61)	-4.480 (-5.25)
c_E	$-1.638 \ (-0.090)$	-0.982 (0.55)	-1.029 (-0.35)	-0.412 (0.05)
z_0	0.090 (1.035)	2.013(2.881)	0.615(1.03)	2.806(3.059)

TABLE III: Adimensional c_D and c_E values of the contact terms in the NV3 interactions obtained from fits to i) the nd scattering length and trinucleon binding energies [37]; and ii) the central value of the 3 H GT matrix element and the trinucleon binding energies (starred values) [4]. The adimensional z_0 values are obtained using the relation given in Eq. (16).

ical value of the GT matrix element in tritium β decay. These second-generation NV2+3 interactions were designated as NV2+3-Ia*/b* and NV2+3-IIa*/b*. These two different procedures for fixing c_D and c_E produced rather different values for these LECs. They are reported in Table III.

IV. χ EFT AXIAL CURRENTS

Many-body axial currents have been first examined within χ EFT by Park and collaborators in Ref. [49]. In that work, the authors retained pions and nucleons in their effective theory and calculated the two-body axial currents up to one-loop terms. The derivation neglected, for example, pion-pole contributions. More recently, two-body axial currents with pions and nucleons have been derived by the Bonn group [50] using the unitary transformation method, and by the JLab-Pisa group using time-ordered perturbation theory [3, 5]. The two derivations differ in the treatment of reducible diagrams. When calculating box diagrams entering the electromagnetic charge and current operators [51–55], the two methods lead to results that are in agreement. However, as discussed at length in Refs. [3, 56], the two groups find different results for the box diagrams in the two-body axial current operator at N4LO. The numerical impact of this difference has been investigated in Refs. [3, 4], where both the JLab-Pisa and Bonn versions of the N4LO current operators have been implemented to calculate the GT matrix element in triton beta decay. In those studies, it was found that the corrections generated by the JLab-Pisa and Bonn N4LO operators are qualitatively in agreement (they both quench the GT matrix element at leading order), and provide, respectively, a $\approx 6\%$ and $\approx 4\%$ contribution to the total GT matrix element.

Here, we consider two-body axial currents derived within the same χEFT used to construct the NV2+3 in-

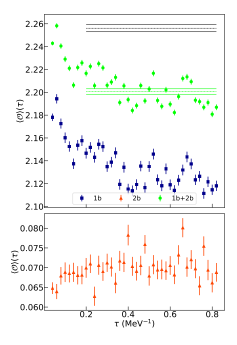


FIG. 2: (Color online) Propagation of the $^6\mathrm{He} \to ^6\mathrm{Li}$ transition matrix element as function of imaginary time τ , based on the NV2+3-Ia Hamiltonian. Results with the one-body current at LO and currents beyond LO are indicated with 1b and 2b, respectively. Dashed and solid lines represent central values and associated error bars. Black dashed and solid lines denote the VMC results. See text for further explanations.

teractions [4]. Moreover, we base our calculations on tree-level corrections only, and disregard the (problematic) N4LO loop contributions discussed above. This choice is advantageous also because it allows for a clearer comparison with the no-core shell model and coupled-cluster calculations of Ref. [9], which are also based on tree-level axial currents alone (albeit derived in a Δ -less χ EFT). Corrections at N4LO in the present formulation are, in practice, subsumed in the LECs of the theory, which have been determined by fits to experimental data.

Before moving on to a (brief) discussion of these axial currents, it is worthwhile pointing out that manybody corrections to leading one-body transition operators have been shown to be crucial for providing a quantitatively successful description of many nuclear electroweak observables [57], such as nuclear electromagnetic form factors [58–61], low-energy electroweak transitions [6, 52, 53, 62–65], and electroweak scattering [66]. They have also been used in studies of double beta decay matrix elements [39, 67–69].

The N3LO axial currents used in this work are repre-

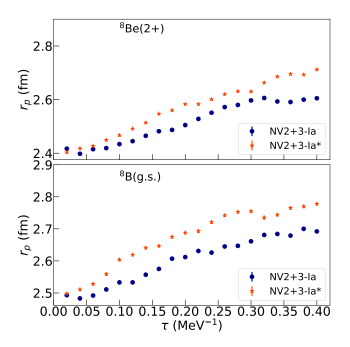


FIG. 3: GFMC propagation of the point-proton radius of the first excited state of ⁸Be (upper panel) and ⁸B (lower panel) based on the NV2+3-Ia and NV2+3-Ia* Hamiltonians.

sented diagrammatically Fig. 1. We refer the interested reader to Refs. [4, 5] for additional details and explicit expressions of the operators; here, we only note that we do not show diagrams that lead to vanishing contributions as well as pion-pole terms which give negligible corrections to the matrix elements under study.

The LO term, which scales as Q^{-3} in the power counting (Q denotes generically a low-momentum scale), is shown in panel (a) and reads

$$\mathbf{j}_{5,a}^{\mathrm{LO}}(\mathbf{q}) = -\frac{g_A}{2} \, \tau_{i,a} \, \boldsymbol{\sigma}_i \, \mathrm{e}^{i\mathbf{q} \cdot \mathbf{r}_i} \,, \tag{10}$$

where g_A is the nucleon axial coupling constant $(g_A = 1.2723 \ [31])$, f_{π} is the pion-decay amplitude $(f_{\pi} = 92.4 \ \text{MeV})$, σ_i and τ_i are the spin and isospin Pauli matrices of nucleon i, \mathbf{q} is the external field momentum, \mathbf{r}_i is the position of nucleon i, and the subscript a specifies the isospin component (a = x, y, z).

At N2LO there are two contributions (scaling as Q^{-1}). The first one is a relativistic correction to the single-nucleon operator at LO and is diagrammatically illustrated in panel (b), while the second involves the excitation of a nucleon into a Δ by pion exchange, as illustrated in panel (c). In the tables and figures below, we will denote these two contributions with N2LO-RC and N2LO- Δ , respectively. We use the same notation introduced in Ref. [4] and write the cumulative N2LO contribution as

$$\mathbf{j}_{5,a}^{\text{N2LO}}(\mathbf{q}) = \mathbf{j}_{5,a}^{\text{N2LO}}(\mathbf{q}; \text{RC}) + \mathbf{j}_{5,a}^{\text{N2LO}}(\mathbf{q}; \Delta)$$
 (11)

At N3LO (or Q^0 in the chiral expansion), there is a term of one-pion range illustrated in panel (d), and a

contact term shown in panel (e), which together give the following N3LO correction

$$\mathbf{j}_{5,a}^{\text{N3LO}}(\mathbf{q}) = \mathbf{j}_{5,a}^{\text{N3LO}}(\mathbf{q}; \text{OPE}) + \mathbf{j}_{5,a}^{\text{N3LO}}(\mathbf{q}; \text{CT})$$
. (12)

We will denote the individual terms with N3LO-OPE and N3LO-CT, respectively.

The configuration-space expressions of these currents are given in Eqs. (2.7)–(2.10) of Ref. [4]. Here, we limit ourselves to report the expression of the N3LO contact term used in this work to explicitly show the relation between the LECs entering this axial current and the LEC c_D in the three-nucleon interaction. In r-space the N3LO-CT current reads

$$\mathbf{j}_{5,a}^{\text{N3LO}}(\mathbf{q}; \text{CT}) = z_0 e^{i \mathbf{q} \cdot \mathbf{R}_{ij}} \frac{e^{-z_{ij}^2}}{\pi^{3/2}} (\boldsymbol{\tau}_i \times \boldsymbol{\tau}_j)_a (\boldsymbol{\sigma}_i \times \boldsymbol{\sigma}_j) ,$$
(13)

where

$$\mathbf{R}_{ij} = (\mathbf{r}_i + \mathbf{r}_j)/2 , \qquad z_{ij} = r_{ij}/R_{\mathrm{S}} , \qquad (14)$$

and r_{ij} is the interparticle distance. The δ -function in the contact axial current has been smeared by replacing it with a Gaussian cutoff of range R_S ,

$$C_{R_{\rm S}}(r_{ij}) = \frac{1}{\pi^{3/2} R_{\rm S}^3} e^{-(r_{ij}/R_{\rm S})^2},$$
 (15)

as previously done for the contact-like terms of the NV2 interactions. The adimensional LEC z_0 (reported in Table III) is given by

$$z_{0} = \frac{g_{A}}{2} \frac{m_{\pi}^{2}}{f_{\pi}^{2}} \frac{1}{(m_{\pi} R_{S})^{3}} \left[-\frac{m_{\pi}}{4 g_{A} \Lambda_{\chi}} c_{D} + \frac{m_{\pi}}{3} (c_{3} + 2 c_{4}) + \frac{m_{\pi}}{6 m} \right], \qquad (16)$$

where c_D is the LEC multiplying one of the contact terms in the three-nucleon interaction [33] given in Table III, $\Lambda_{\chi} = 1$ GeV is the chiral symmetry breaking scale, c_3 and c_4 are given in Table I, and m and m_{π} are the average nucleon and pion masses. It has recently been realized [13] that the relation between z_0 and c_D had been given erroneously in the original reference [12], a – sign and a factor 1/4 were missing in the term proportional to c_D .

V. RESULTS

The GFMC energies of the nuclei of interest calculated using the NV2+3-Ia and NV2+3-Ia* models are listed in Table IV along with the dominant spatial symmetry (s.s.) of the variational wave functions [77]. The energies are obtained using $\approx 80,000$ walkers, and are all well converged by 30 unconstrained steps [78]. All the GFMC results presented in this article (but for the two cases

TABLE IV: GFMC predictions for $A \leq 10$ nuclear states studied in this work, compared to experimental values [70, 71]. Numbers in parentheses are statistical errors for the GFMC calculations; experimental errors, being negligible, are not indicated. The dominant spatial symmetry (s.s.) of the nuclear wave function is given in the second column.

$^{A}Z(J^{\pi};T)$	s.s.	E [N		
		Ia	Ia*	Expt.
$^{6}\text{Li}(1^{+};0)$	[42]	-31.97(6)	-31.06(8)	-31.99
$^{6}\text{He}(0^{+};0)$ $^{7}\text{Li}(\frac{3}{2}^{-};\frac{1}{2})$	[42] [43]	-29.32(4) -39.25(15)	-28.46(5) -38.27(14)	-29.27 -39.25
${}^{7}\text{Li}(\frac{1}{2}^{-}; \frac{1}{2})$ ${}^{7}\text{Be}(\frac{3}{2}^{-}; \frac{1}{2})$	[43] [43]	-39.18(15) -37.75(8)	-37.66(15) -36.56(10)	-38.76 -37.60
⁸ He(0 ⁺ ;1) ⁸ Li(2 ⁺ ;1)	[422] [431]	-31.33(7) $-41.59(10)$	-28.53(6) $-38.89(7)$	-31.40 -41.28
${}^{8}\text{Li}(1^{+};1)$ ${}^{8}\text{B}(2^{+};1)$ ${}^{8}\text{Be}(2^{+};0)$	[431] [431] [44]	-40.59(7) $-37.87(8)$ $-54.07(7)$	-37.78(7) $-35.63(8)$ $-53.16(11)$	-40.30 -37.74 -53.47
$^{10}B(1^+;0)$ $^{10}C(0^+;1)$	[442] [442]	-64.61(41) $-61.01(50)$	-53.16(11) $-60.46(30)$ $-56.65(22)$	-64.03 -60.32

discussed below) are averages over the imaginary time τ from 0.2 to 0.82 MeV⁻¹. Results obtained with the NV2+3-Ia interaction are in statistical agreement with those published in Ref. [37] based on the same nuclear Hamiltonian. Model NV2+3-Ia leads to predictions that are in excellent agreement with the data. We also report for the first time results based on the second generation of NV2+3 interactions, specifically model NV2+3-Ia*, whose three-nucleon interaction has been constrained by fits to the experimental trinucleon binding energies and tritium GT matrix element. Results obtained with the NV2+3-Ia* Hamiltonian display a somewhat less satisfactory agreement with the experimental data, but still less than 4% away from them.

A typical imaginary-time evolution of the GFMC transition matrix elements is shown in Fig. 2. As can be seen, there is a rapid drop of 3\% from the initial VMC estimate at $\tau = 0$ that reaches a stable value around 0.2 MeV⁻¹. The results for all transitions presented in this article are averages over τ from 0.2 to 0.82 MeV⁻¹, as indicated by the dashed lines, with statistical errors denoted by the solid lines. The calculations of weak transitions involving the $(J^{\pi}, T) = (2^+, 0)$ state of ⁸Be and the ground state of ⁸B are treated differently. For these two states, we observe that the binding energy, magnitude of the quadrupole moment, and point-proton radius all increase monotonically as the imaginary time increases. This can be appreciated in Fig. 3 where we show the point-proton radii of the two nuclear states. We interpret this behavior as an indication that the resonant excited state of ⁸Be is dissolving into two separated α 's, while ⁸B is breaking into p+7Be. In the case of 8 Be, this issue has been addressed already in Refs. [64, 65, 79]. Here, we use similar techniques to treat these systems and extract matrix elements from the GFMC data. In particular,

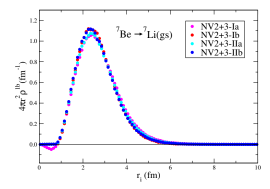


FIG. 4: (Color online) One-body density–defined in Eq. (18)–of the $^7\mathrm{Be}$ to $^7\mathrm{Li}(gs)$ GT RME obtained with models NV2+3-Ia/b and NV2+3-IIa/b.(see text for explanation).

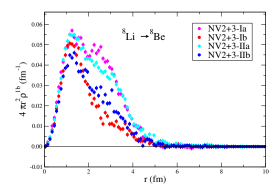


FIG. 5: (Color online) Same as Fig. 4 but for the $^8{\rm Li}$ to $^8{\rm Be}$ GT RME.

we note that the ground-state energy of ⁸Be drops very quickly as the imaginary time increases and reaches stability around $\tau \sim 0.1 \text{ MeV}^{-1}$. The transition matrix elements involving the two dissolving states have been determined assuming that, also for these states, $\tau \approx 0.1$ MeV^{-1} is the point at which spurious contaminations in the nuclear wave functions have been removed by the GFMC propagation. We then average in a small interval around this point, typically between 0.06 and 0.14 MeV^{-1} . Calculations involving these two systems are clearly affected by a systematic error. To have a rather rough estimate of this error, we study the sensitivity of the extracted matrix elements with respect to variations in the imaginary time interval selected for the averaging. We find that such a procedure generates an additional uncertainty of $\approx 5\%$ which is added in quadrature to the statistical one, and quoted in Table VI below of GFMC results.

TABLE V: Gamow-Teller RMEs in A=6, 7, 8, and 10 nuclei obtained with chiral axial currents [4] and VMC wave functions corresponding to the NV2+3-Ia/b and NV2+3-Ia/b (NV2+3-Ia/b* and NV2+3-IIa/b*) Hamiltonian models [1, 2, 4, 37]. Columns labeled with LO, N2LO-(RC+ Δ), N3LO-OPE, and N3LO-CT refer to the contributions given by the diagrams illustrated in panel (a), panels (b) plus (c), panel (d), and panel (e) of Fig. 1, respectively. The cumulative results are reported in the column labeled "Total", while results including only corrections beyond LO are listed under "Total-LO". Experimental values from Refs. [72–76] are given in the last column. The dominant spatial symmetries of the VMC wave functions are reported in the first column. Statistical errors associated with the Monte Carlo integrations are not shown, but are below 1%.

Transition	Model	LO	N2LO-(RC+ Δ)	N3LO-OPE	N3LO-CT	Total-LO	Total	Expt.
6 He(0 ⁺ ;1)→ 6 Li(1 ⁺ ;0) [42]→[42]	IIa (IIb) Ia* (Ib*)	2.207 (2.212) 2.192 (2.256)	0.022 (0.056) 0.027 (0.043) 0.021 (0.056) 0.027 (0.044)	0.043 (0.055) 0.038 (0.063)	-0.005 (-0.068) -0.034 (-0.082) -0.054 (-0.097) -0.057 (-0.090)	0.036 (0.016) 0.005 (0.022)	2.256 (2.306) 2.243 (2.228) 2.197 (2.279) 2.216 (2.228)	,
${}^{7}\text{Be}(\frac{3}{2}^{-};\frac{1}{2}) \rightarrow {}^{7}\text{Li}(\frac{3}{2}^{-};\frac{1}{2})$ $[43] \rightarrow [43]$	IIa (IIb) Ia* (Ib*)	2.293 (2.309) 2.327 (2.307)	0.099 (0.162) 0.102 (0.153) 0.098 (0.161) 0.103 (0.154)	0.078 (0.113) 0.076 (0.117)	-0.010 (-0.148) -0.070 (-0.190) -0.121 (-0.212) -0.120 (-0.210)	0.110 (0.076) 0.053 (0.066)	2.482 (2.427) 2.403 (2.385) 2.380 (2.373) 2.357 (2.374)	,
${}^{7}\text{Be}(\frac{3}{2}^{-};\frac{1}{2}) \rightarrow {}^{7}\text{Li}(\frac{1}{2}^{-};\frac{1}{2})$ $[43] \rightarrow [43]$	Ia^* (Ib^*)	2.128 (2.145) 2.158 (2.124)	0.066 (0.122) 0.069 (0.111) 0.065 (0.119) 0.067 (0.111)	0.065 (0.095) 0.063 (0.099)	-0.009 (-0.125) -0.059 (-0.162) -0.103 (-0.180) -0.101 (-0.178)	0.074 (0.044) 0.025 (0.038)	2.278 (2.215) 2.202 (2.189) 2.183 (2.162) 2.161 (2.176)	,
$^{8}\text{Li}(2^{+};1) \rightarrow ^{8}\text{Be}(2^{+};0)$ $[431] \rightarrow [44]$	Ia (Ib) IIa (IIb) Ia* (Ib*) IIa* (IIb*)	0.144 (0.101) 0.148 (0.099)	0.032 (0.028) 0.031 (0.033) 0.032 (0.033) 0.032 (0.037)	0.010 (0.011) 0.010 (0.012)	-0.001 (-0.014) -0.008 (-0.019) -0.016 (-0.020) -0.014 (-0.023)	0.033 (0.025) 0.026 (0.025)	(,	
$^{8}B(2^{+};1)\rightarrow^{8}Be(2^{+};0)$ [431] \rightarrow [44]	Ia (Ib) IIa (IIb) Ia* (Ib*) IIa* (IIb*)	0.144 (0.102) 0.148 (0.098)	0.032 (0.032) 0.031 (0.033) 0.032 (0.034) 0.032 (0.037)	0.010 (0.011) 0.010 (0.012)	-0.001 (-0.014) -0.008 (-0.019) -0.016 (-0.020) -0.014 (-0.022)	0.033 (0.026) 0.026 (0.025)	0.188 (0.122) 0.177 (0.128) 0.174 (0.123) 0.154 (0.145)	,
$^{8}\text{He}(0^{+};1)\rightarrow^{8}\text{Li}(1^{+};1)$ [422] \rightarrow [431]	Ia (Ib) IIa (IIb) Ia* (Ib*) IIa* (IIb*)	0.465 (0.370) 0.362 (0.377)	0.030 (0.034) 0.032 (0.034) 0.031 (0.034) 0.033 (0.035)	0.012 (0.011) 0.009 (0.014)	-0.001 (-0.014) -0.009 (-0.017) -0.010 (-0.022) -0.017 (-0.019)	0.035 (0.028) 0.029 (0.026)	0.424 (0.396) 0.500 (0.398) 0.391 (0.402) 0.510 (0.391)	· · /
$^{10}C(0^+;1) \rightarrow ^{10}B(1^+;0)$ $[442] \rightarrow [442]$	IIa (IIb) Ia* (Ib*)	2.157 (2.176) 2.015 (2.123)	0.003 (0.026) -0.002 (0.026) 0.024 (0.022) 0.015 (0.031)	0.046 (0.072) 0.059 (0.069)	-0.006 (-0.081) -0.042 (-0.120) -0.119 (-0.117) -0.126 (-0.123)	0.002 (-0.022) -0.037 (-0.028)	2.159 (2.154) 1.978 (2.095)	,

A. Weak Transitions in A=6-10 Nuclei

In this section, we present results for the GT reduced matrix element (RME) defined as

$$RME = \frac{\sqrt{2J_f + 1}}{g_A} \frac{\langle J_f M | j_{5,\pm}^z | J_i M \rangle}{\langle J_i M, 10 | J_f M \rangle} , \qquad (17)$$

where $j_{5,\pm}^z$ is the z-component (at vanishing momentum transfer) of the charge-raising/lowering current with $\mathbf{j}_{5,\pm} = \mathbf{j}_{5,x} \pm i \mathbf{j}_{5,y}$, and $\langle J_i M, 10 | J_f M \rangle$ is a Clebsch-Gordan coefficient.

Results for the GT RMEs in $A\!=\!6\!-\!10$ nuclei based on variational wave functions are reported in Table V for the eight different NV2+3 interactions discussed above, namely the NV2+3-Ia/b, NV2+3-IIa/b and corresponding starred models. The one-body axial current at LO,

illustrated in panel (a) of Fig. 1, leads to contributions to the matrix elements reported in the third column of Table V. One-body relativistic corrections (N2LO-RC) and two-body currents of one-pion range (N2LO- Δ) at N2LO, displayed in panels (b) and (c) of Fig. 1, are added up and given in the fourth column of Table V labeled with N2LO-(RC+ Δ). A rough estimate of the size of the RC corrections can be obtained suppressing each LO term by a factor of $(Q/m)^2 \approx 0.01$, where we used a "typical" nucleon's low-momentum $Q \approx 100$ MeV. Contributions at N3LO are given in the columns labeled by N3LO-OPE and N3LO-CT, corresponding to the one-pion range and contact currents (and displayed in panels (d) and (e) of Fig. 1). The cumulative contributions are given in the next to last column, while the contributions beyond LO only in the column labeled "Total-LO". Experimental

TABLE VI: Gamow-Teller RMEs in A=6, 7, 8, and 10 nuclei obtained with chiral axial currents [4] and GFMC (VMC) wave functions corresponding to the NV2+3-Ia and NV2+3-Ia* Hamiltonian models [1, 2, 4, 37]. Results corresponding to the one-body current at LO (column labeled "LO"), and to the sum of all the corrections beyond LO (column labeled "Total-LO") are given, along with the cumulative contributions (column labeled 'Total') to be compared with the experimental data [72–76] reported in the last row. Results from Ref. [6] based on the AV18 and IL7 nuclear Hamiltonian are also shown where available. Statistical errors associated with the Monte Carlo integrations are not shown, but are below 1%. Transitions for the A=8 systems are affected by an additional systematic error of $\approx 5\%$, see text for explanation.

Transition	Model	s.s.	LO	Total-LO	Total	Expt.
$^{6}\text{He}(0^{+};1) \rightarrow ^{6}\text{Li}(1^{+};0)$	Ia Ia* Ref. [6]	$[42] \rightarrow [42]$	2.130(2.200) 2.107(2.192) 2.168(2.174)	0.070(0.056) 0.011(0.005) 0.037(0.030)	2.201(2.256) 2.118(2.197) 2.205(2.211)	2.1609(40)
$^{7}\text{Be}(\frac{3}{2}^{-};\frac{1}{2}) \rightarrow ^{7}\text{Li}(\frac{3}{2}^{-};\frac{1}{2})$	Ia Ia* Ref. [6]	[43]→[43]	2.273(2.317) 2.286(2.327) 2.294(2.334)	0.164(0.165) 0.052(0.053) 0.061(0.050)	2.440(2.482) 2.338(2.380) 2.355(2.384)	2.3556(47)
$^{7}\text{Be}(\frac{3}{2}^{-};\frac{1}{2}) \rightarrow ^{7}\text{Li}(\frac{1}{2}^{-};\frac{1}{2})$	Ia Ia* Ref. [6]	[43]→[43]	2.065(2.157) 2.061(2.158) 2.083(2.150)	0.103(0.121) 0.009(0.025) 0.046(0.046)	2.168(2.278) 2.070(2.183) 2.129(2.196)	2.1116(57)
$^{8}\text{Li}(2^{+};1) \rightarrow ^{8}\text{Be}(2^{+};0)$	Ia Ia*	$[431] \rightarrow [44]$	0.074(0.147) 0.096(0.148)	0.029(0.041) 0.025(0.026)	0.103(0.188) 0.120(0.174)	0.284 Ref. [75] 0.190 Ref. [76]
$^{8}B(2^{+};1) \rightarrow ^{8}Be(2^{+};0)$	Ia Ia*	$[431] \rightarrow [44]$	0.091(0.146) 0.102(0.148)	0.035(0.042) 0.024(0.026)	0.125(0.188) 0.126(0.174)	0.269(20)
$^{8}\text{He}(0^{+};1) \rightarrow ^{8}\text{Li}(1^{+};1)$	Ia Ia*	[422]→[431]	0.262(0.386) 0.297(0.362)	0.040(0.038) 0.025(0.029)	0.302(0.424) 0.322(0.391)	0.512(6)
$^{10}\mathrm{C}(0^+;1) \rightarrow ^{10}\mathrm{B}(1^+;0)$	Ia Ia* Ref. [6]	[442]→[442]	1.928(1.940) 2.086(2.015) 2.032(2.062)	0.050(0.041) -0.031(-0.037) 0.016(0.015)	1.978(1.981) 2.055(1.978) 2.048(2.077)	1.8331(34)

data from Refs. [72–76] are reported in the last column of Table V.

All the calculations use axial currents at tree-level which are consistent with the specific NV2+3 model used to generate the VMC wave functions. VMC results based on different nuclear Hamiltonians are qualitatively in agreement. In particular, for the A=3,6,7 and 10 systems the LO contribution provides about 97% of the total matrix elements with currents beyond LO giving the remaining $\lesssim 3\%$ correction. This correction adds up constructively to the LO contribution for all nuclei being considered, but for the A = 10 transition. For this last transition, we find that the contributions beyond LO give a correction that quenches the LO results obtained with all the starred models, and with the un-starred NV2+3-IIb interaction. More details about this calculation will be given in the following section. We emphasize that the "Total-LO" column includes, in addition to two-body contributions, also a small correction resulting from the one-body N2LO-RC current.

Transitions involving A = 8 nuclei exhibit a large sup-

pression at LO. This behavior is attributable to the fact that the initial and final VMC wave functions are characterized by different dominant spatial symmetries, which make their overlap small compared to cases in which both the initial and final states display the same dominant spatial symmetry. As a consequence, in these cases the LO term is only about $\approx 40-50\%$ of the total matrix element, with two-body currents providing a large correction. Two-body currents, while improving the agreement with the experimental values, are insufficient to fully explain them. Because of the reduced overlap between dominant components in the wave functions, these matrix elements are particularly sensitive to small components, which are poorly constrained and model dependent. This can be appreciated by looking at the one-body transition densities, $\rho^{1b}(r_i)$, defined as

RME(1b) = RME(LO) =
$$4 \pi \int dr_i r_i^2 \rho^{1b}(r_i)$$
, (18)

where r_i is the distance of nucleon i from the center-of-mass of the system.

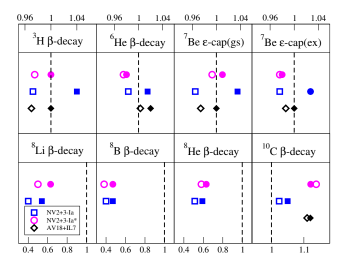


FIG. 6: (Color online) Ratios of GFMC to experimental values of the GT RMEs in the 3 H, 6 He, 7 Be, 8 B, 8 Be, 8 He and 10 C weak transitions. Theory predictions correspond to the χ EFT axial current at LO (empty symbols) and up to N3LO (filled symbols) obtained with the NV2+3-Ia and NV2+3-Ia* models. Results from Ref. [6] based on the AV18+IL7 nuclear Hamiltonian and N4LO currents from Ref. [5] are also shown. Results for the 3 H weak transition were reported in Ref. [4].

In Figs. 4 and 5, we show one-body densities for two transitions, namely the ϵ -capture of the ⁷Be ground state to the ⁷Li ground state and the ⁸Li β -decay. The former, involves initial and final states with the same [43] (dominant) spatial symmetry, while for the latter the initial state is in a [431] spatial symmetry configuration and the final state is in a [44] one. The densities are calculated using the NV2+3-Ia/b and NV2+3-IIa/b interactions. From the figures we can see that the A=7 one-body densities are well constrained and essentially model independent, while the A=8 ones are particularly sensitive to the nuclear Hamiltonian used to generate the wave functions. Of course, these considerations are based on VMC results. A GFMC propagation might mitigate the observed model dependence.

In all the cases we studied, the contact-like current at N3LO provides a correction that quenches the LO terms, while the currents of one-pion range at N2LO and N3LO add up constructively to the LO contributions (see Table V). The main difference between the un-starred and starred calculations is observed in the size of the contact contribution. In particular, starred models are characterized by a larger value of z_0 (see Table III), which in turns leads to a larger (in magnitude) N3LO-CT correction.

We performed GFMC propagations only for the NV2+3-Ia and NV2+3-Ia* models. GFMC results are reported in Table VI, where, for completeness, we also show the corresponding VMC values in parentheses along with the GFMC results from Ref. [6]. We summarize the GFMC results in Fig. 6 and compare them (where possible) to the results of Ref. [6] based on the AV18+IL7 nuclear Hamiltonian.

The effect of the GFMC propagation in imaginary time is to reduce the VMC results by $\lesssim 4\%$ in all selected transitions (but for the A=10 transition obtained with the NV2+3-Ia* model). The agreement with the data, after the inclusion of two-body currents, is at the $\approx 2\%$ ($\lesssim 2\%$) level for the A=6 transition with the NV2+3-Ia (NV2+3-Ia*) model; and at the $\lesssim 4\%$ ($\lesssim 1\%$) level for the A=7 cases with the NV2+3-Ia (NV2+3-Ia*) model. These results are in agreement with those obtained for the same transitions in the calculations of Ref. [6] which were based on the AV18+IL7 interactions. The NV2+3 models lead to a more satisfactory agreement with the data for the A=6 RME primarily because, with these interactions, the LO term is 2% smaller than obtained using AV18-IL7 model.

The largest discrepancy generated by the use of different nuclear Hamiltonians, including AV18+IL7, is observed in the A = 10 transition. This can be appreciated looking at both Table V and Table VI. From the former, we observe a rather large cutoff dependence (models a vs models b), and also a large sensitivity to the class (either I or II) used to generate the nuclear wave functions. From Table VI, we see that the results of Ref. [6], based on the AV18+IL7 Hamiltonian, lie between models Ia and Ia*. This large model and cutoff dependence can be traced back to the existence of two nearby $J^{\pi} = 1^+$ excited states in ¹⁰B, the lower one a predominantly ³S₁[442] state and the upper one a ${}^{3}D_{1}[442]$ state (in LS coupling), which are only 1 MeV apart. The transition from the ${}^{10}\mathrm{C}(0^+)$ state, which is predominantly ${}^{1}S_{0}[442]$, is large in the $S \to S$ components, but about five times smaller in the $S \to D$ components. This makes the GT matrix element particularly sensitive to the exact mixing of the ${}^{3}S_{1}$ and ${}^{3}\mathrm{D}_{1}$ components in the two ${}^{10}\mathrm{B}(1^{+})$ states produced by a given Hamiltonian. It would appear that none of the interactions models studied here gets quite the right mixing of these components. In particular, results based on the NV2+3-Ia and NV2+3-Ia* interactions over-predict the data by $\sim 7\%$ and 12%, respectively, which gives an indication of the spread of the theoretical estimates.

Predictions for the RMEs of A=8 transitions are the first QMC calculations for these systems that include corrections from two-body axial currents. As discussed above, RMEs are suppressed at leading order which gives only $\approx 50\%$ and $\approx 40\%$ of the experimental values for the $^8\mathrm{B} \to ^8\mathrm{Be}$ and $^8\mathrm{He} \to ^8\mathrm{Li}$ transitions. Two-body currents provide about 20–30% correction in the right direction which is, however, still insufficient to reach agreement with the experimental data. These transitions are challenging not only from the theoretical but also from the experimental point of view. For example, in Tables V and VI we quote two results for the RME of the $^8\mathrm{Li} \to ^8\mathrm{Be}$ decay obtained from the $\log(ft)$ -values of Ref. [75] and Ref. [76] via the following formula [74]

RME(EXPT) =
$$\frac{1}{g_A} \sqrt{2J_i + 1} \sqrt{\frac{6139 \pm 7}{ft}}$$
, (19)

where J_i is the angular momentum of the parent nucleus.

The Fermi transition strength is small enough in this case that it can be neglected in the above formula. We then obtain two values, namely, RME(Ref. [75])=0.284, and RME(Ref. [76])=0.190. We note that Refs. [73, 80], report a different overall factor of 6147 instead of 6139 in the formula given above. In our estimate we used $g_A=1.2723$. Despite this additional uncertainty in the deduced experimental values, our predictions still severely underestimate the data. For example, the calculated Ia* RME provides only $\approx 40\%$ and $\approx 60\%$ of the experimental values given in Ref. [75] and Ref. [76], respectively.

B. Two-Body Transition Densities

In order to have a better understanding of the twobody terms in the axial current and their contributions, it is helpful to study the associated two-body transition densities, which we define as [81]

RME(2b) =
$$4\pi \int_0^\infty dr \, r^2 \, \rho^{2b}(r)$$
, (20)

where r is the interparticle distance, and $2b = N2LO-\Delta$, N2LO-OPE, and N2LO-CT.

Two-body transition densities, calculated for selected nuclei with variational wave functions corresponding to the Hamiltonian models NV2+3-Ia (Ia) and NV2+3-Ia* (Ia*), are presented in Fig. 7. These models produce similar N2LO- Δ and N3LO-OPE densities, since they are based on the same underlying NV2 interaction and only differ in the NV3 interaction. Specifically, what differs is the strength z_0 of the contact current—linked to the (contact) three-nucleon interaction via the relation in Eq. (16). Because of the different methodologies adopted in constraining c_D and c_E (the LECs parameterizing the contact piece of the three-nucleon interaction), z_0 turns out to be much larger for Ia* than for Ia, see Table III. Since the contact current is proportional to z_0 , this also explains why the corresponding density for Ia* is much larger (in magnitude) than for Ia. Note that they are both negative. As a consequence, the total density (black symbols in Fig. 7) develops a node at around 1 fm in the case of model Ia*.

Another interesting feature of Fig. 7 is the difference between the N2LO- Δ and N3LO-OPE densities at separations $r \gtrsim 2$ fm for the transitions in the larger systems, especially those involving the A=8 resonant states. In the limit of vanishing momentum transfer we are considering here, the corresponding currents have the same operator structure [4], up to a momentum dependent term, absent in the N2LO- Δ current, which, however, we have explicitly verified to give a numerically small contribution by direct calculation. Examination of Eqs. (2.9) and (2.10) of Ref. [4] shows that this common operator structure involves two independent correlations functions $I^{(1)}(\mu_{ij}; \alpha_p)$ and $I^{(2)}(\mu_{ij}; \alpha_p)$ with $\mu_{ij} = m_{\pi} r_{ij}$, proportional to different combinations of LECs, denoted by α_1

 (α_1^{Δ}) and α_2 (α_2^{Δ}) in the OPE (Δ) current, with

$$\frac{\alpha_1^{\Delta}}{\alpha_1} = \frac{c_4^{\Delta}}{c_4 + 1/(4m)} \approx 0.89 , \qquad \frac{\alpha_2^{\Delta}}{\alpha_2} = \frac{c_3^{\Delta}}{c_3} \approx 3.6 ,$$
$$\frac{\alpha_1^{\Delta}}{\alpha_2^{\Delta}} = -\frac{1}{4} , \qquad \qquad \frac{\alpha_1}{\alpha_2} \approx -1.0 \qquad (21)$$

using the values in Tables I and II (m) is the average nucleon mass). Here, $c_3^\Delta = -h_A^2/(9\,m_{\Delta N})$ and $c_4^\Delta = h_A^2/(18\,m_{\Delta N})$. Indeed, the N2LO- Δ current reads

$$\mathbf{j}_{\Delta}^{\text{N2LO}} = -\left(\boldsymbol{\tau}_{i} \times \boldsymbol{\tau}_{j}\right)_{a} \boldsymbol{\sigma}_{i} \times \mathbf{O}_{ij}^{(1)} - \boldsymbol{\tau}_{j,a} \mathbf{O}_{ij}^{(2)} + \left(i \rightleftharpoons j\right),$$
(22)

with

$$\mathbf{O}_{ij}^{(p)} = I^{(1)}(\mu_{ij}; \alpha_p^{\Delta}) \, \boldsymbol{\sigma}_j + I^{(2)}(\mu_{ij}; \alpha_p^{\Delta}) \, \hat{\mathbf{r}}_{ij} \, \boldsymbol{\sigma}_j \cdot \hat{\mathbf{r}}_{ij} . \tag{23}$$

A similar expression holds for $\mathbf{j}_{\pi}^{\text{N3LO}}$ with α_p replacing α_p^{Δ} . There are cancellations between the terms proportional to $\mathbf{O}_{ij}^{(1)}$ and $\mathbf{O}_{ij}^{(2)}$ in each of these currents, and these cancellations are sensitive to the values of the ratios $\alpha_1^{\Delta}/\alpha_2^{\Delta}$ and α_1/α_2 , and to the overlap between the wave functions of the states involved in the transition.

To gain insight into how short-range physics impacts these weak transitions across different nuclei, we display in Fig. 8 the densities corresponding to the individual two-body contributions, each normalized as $4\pi r^2 \rho^{2b}(r)/(4\pi r^2 \rho^{2b})_{\rm max}$, where $(4\pi r^2 \rho^{2b})_{\rm max}$ is the maximum attained value (in magnitude); so all curves peak at 1. We also display the total densities and note that, since both a positive peak and a negative valley are present in this case, each curve is normalized so that the value of the positive peak is 1.

The universal behavior exhibited by the N2LO- Δ , N3LO-OPE, N3LO-CT densities is quite striking, as the curves corresponding to different nuclei and different Hamiltonian models, essentially overlap for $r \lesssim 1/m_\pi$. (It is even more striking when the weighing r^2 factor is not included). Such behavior can be understood as follows.

In a charge-raising process the two-body weak transition operators primarily convert a pn pair with total spin/isospin S/T = 1/0 (nn pair with S/T = 0/1) to a pp pair with S/T = 0/1 (pn pair with S/T = 1/0) [81]—of course, similar considerations apply to a charge lowering process. These operators, at least in light systems, do not couple $TT_z = 10$ and 00 to $TT_z = 11$ in a significant way, since P-wave components are small in that case. At separations $\lesssim 1/m_{\pi}$, where these transitions operators are most effective, the pair wave functions with S/T = 1/0and 0/1 in different nuclei are similar in shape and only differ by a scale factor [82]—those corresponding to the Ia* model are illustrated in Fig. 9. This is the origin for the universal behavior observed in Fig. 8. Note that in Fig. 9 we also show the densities of pairs in S/T=0/0and 1/1, which do not scale. A complete analysis and interpretation of these results—in particular, of the role played by the tensor component of the nuclear interaction in shaping these densities—can be found in Ref. [82].

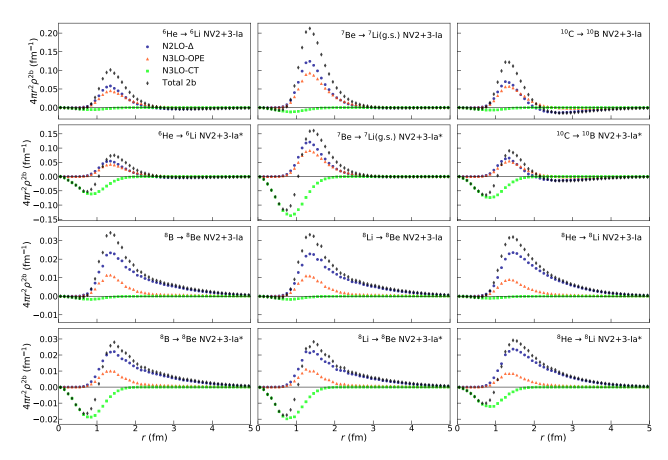


FIG. 7: (Color online) Two-body transition density—see Eq. (20)—for selected nuclei obtained with the NV2+3-Ia and NV2+3-Ia* models (see text for explanation).

However, at separations $r \gtrsim 1/m_{\pi}$, the N2LO- Δ and N3LO-OPE densities (especially the former) do not scale, particularly in the case of the heavier systems with $A \geq 8$, presumably due to delicate cancellations between the terms proportional to the operators $\mathbf{O}_{ij}^{(1)}$ and $\mathbf{O}_{ij}^{(2)}$ present in these currents (P-wave components in the wave functions of these systems may also play a significant role). As a matter of fact, because of the different longrange behavior present in the $A \geq 8$ transitions (note, e.g., how the N2LO- Δ density in the ¹⁰C transition assumes negative values at large separations, all the other densities being positive), and because of the rather large size of the contact contribution in the Ia* model, the total densities exhibit nodes when calculated with the starred interactions, which leads to non-trivial cancellations.

VI. CONCLUSIONS

In this work, we reported on a detailed study of weak matrix elements in $A\!=\!3\!-\!10$ systems based on chiral (two- and three-nucleon) interactions and associated (one- and two-body) axial currents at high orders in the chiral expansion. A summary of our results is displayed in Fig. 6. Agreement with the experimental data

is obtained when correlated nuclear wave functions are adopted. For these transitions the contribution of corrections beyond LO in the axial current is typically at the $\approx 2\%$ level of the value calculated with the LO Gamow-Teller operator. These findings are in line with those reported in the hybrid study of Ref. [6] for the same transitions. Here, we also present calculations of matrix elements entering the rates of the ⁸Li, ⁸B, and ⁸He beta decays. These matrix elements are found to be suppressed at LO, and N2LO and N3LO currents provide a large correction ($\approx 20-30\%$) which is, however, insufficient to explain the experimental data. We attribute the large suppression at LO to the fact that the Gamow-Teller operator is, in these transitions, connecting large to small components of the initial and final wave functions. Improving on these calculations will require the development of more sophisticated wave functions with better constrained small components.

Finally, we also reported on a careful analysis of oneand two-body transition densities shown in Figs. 4–8. The latter are especially interesting because they allow us to understand the spatial distributions of the various two-body current operators, that is their behavior as functions of interparticle distance. We have shown that, for each set of interactions and consistent currents (either NV2+3-Ia or NV2+3-Ia*), the two-body transition

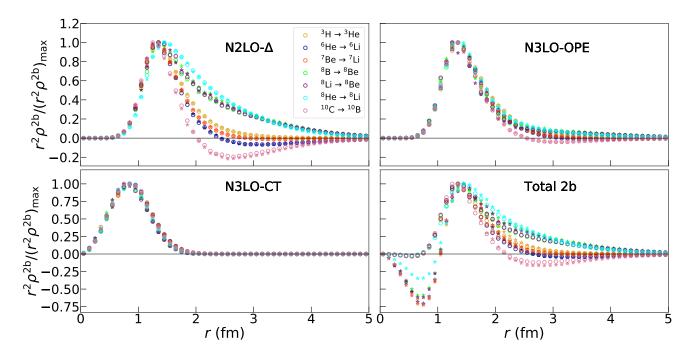


FIG. 8: (Color online) Two-body transition density for selected nuclei (see text for explanation). Different colors indicate different transitions. Results obtained from calculations using the NV2+3-Ia model are represented with open circles and results obtained from calculations employing the NV2+3-Ia* model results are represented by solid stars.

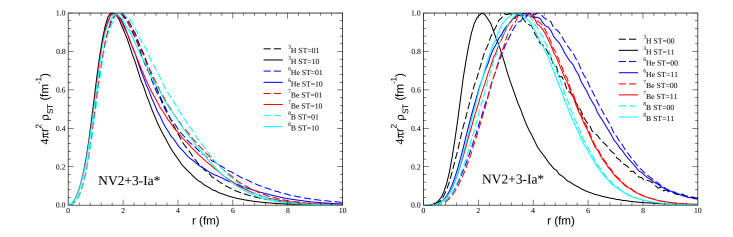


FIG. 9: (Color online) Pair densities in total spin/isospin S/T = 1/0 and 0/1 (left panel) and S/T = 0/0 and 1/1 (right panel) obtained with VMC wave functions corresponding to model Ia*. All curves have been rescaled so as peak at 1. Only central values are shown; statistical Monte Carlo errors are below a few %.

densities exhibit a universal behavior at short distance across all nuclei we have considered in the present study.

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