



Ab initio description of monopole resonances in light- and medium-mass nuclei

III. Moments evaluation in ab initio PGCM calculations

A. Porro^{1,2,3,a} , T. Duguet^{3,4}, J.-P. Ebran^{5,6}, M. Frosini⁷, R. Roth^{1,8}, V. Somà³

¹ Department of Physics, Technische Universität Darmstadt, 64289 Darmstadt, Germany

² ExtreMe Matter Institute EMMI, GSI Helmholtzzentrum für Schwerionenforschung GmbH, 64291 Darmstadt, Germany

³ IRFU, CEA, Université Paris-Saclay, 91191 Gif-sur-Yvette, France

⁴ KU Leuven, Department of Physics and Astronomy, Instituut voor Kern- en Stralingsfysica, 3001 Leuven, Belgium

⁵ CEA, DAM, DIF, 91297 Arpajon, France

⁶ Laboratoire Matière en Conditions Extrêmes, Université Paris-Saclay, CEA, 91680 Bruyères-le-Châtel, France

⁷ CEA, DES, IRESNE, DER, SPRC, 13108 Saint-Paul-lès-Durance, France

⁸ Helmholtz Forschungsakademie Hessen für FAIR, GSI Helmholtzzentrum, 64289 Darmstadt, Germany

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Abstract The paper is the third of a series dedicated to the ab initio description of monopole giant resonances in mid-mass closed- and open-shell nuclei via the so-called projected generator coordinate method. The present focus is on the computation of the moments m_k of the monopole strength distribution, which are used to quantify its centroid energy and dispersion. First, the capacity to compute low-order moments via two different methods is developed and benchmarked for the m_1 moment. Second, the impact of the angular momentum projection on the centroid energy and dispersion of the monopole strength is analysed before comparing the results to those obtained from consistent quasi-particle random phase approximation calculations. Next, the so-called energy weighted sum rule (EWSR) is investigated. First, the appropriate EWSR in the center-of-mass frame is derived analytically. Second, the intrinsic EWSR is tested in order to quantify the (unwanted) local-gauge symmetry breaking of the presently employed chiral effective field theory (χ EFT) interactions. Finally, the infinite nuclear matter incompressibility associated with the employed χ EFT interactions is extracted by extrapolating the finite-nucleus incompressibility computed from the monopole centroid energy.

1 Introduction

The study of giant resonances provides valuable insights into the structural and dynamical properties of atomic nuclei [1–

4]. In particular, the characteristics of the isoscalar giant monopole resonance (ISGMR or GMR for brevity here) and of the isovector giant dipole resonance (IVGDR) not only deepen our comprehension of nuclear structure but also have implications for the modelization of several astrophysical systems. This is the case, for instance, of the description of core-collapse supernovae explosions [5,6] and neutron stars mergers [7–10], both phenomena being associated to the nucleosynthesis of heavy elements and the behavior of nuclear matter under extreme conditions [11].

This article is the third (Paper III) of a series of four addressing the properties of the GMR in closed- and open-shell nuclei from an *ab initio* standpoint using the so-called projected generator coordinate method (PGCM). While the first paper (Paper I) [12] detailed the uncertainty budget associated to PGCM calculations of monopole and quadrupole responses, the second paper (Paper II) [13] focused on the GMR properties of ^{16}O , ^{24}Mg , ^{28}Si and ^{46}Ti . Two-dimensional PGCM calculations were shown to account well for the fragmented monopole response of (rather) light doubly open-shell nuclei thanks to their capacity (i) to capture the impact of the intrinsic static quadrupole deformation and of its fluctuations on the position of the breathing mode (typically at play in spherical nuclei), (ii) to describe in a refined way the coupling between the GMR and the giant quadrupole resonance (GQR) mechanism responsible for the appearance of an additional component in the GMR of intrinsically-deformed nuclei and (iii) to seize anharmonic effects that were shown to be significant in light systems.

^a e-mail: aporro@theorie.ikp.physik.tu-darmstadt.de (corresponding author)

The present paper focuses on the computation of the moments m_k of the monopole strength distribution in order to quantify its main characteristics such as its centroid energy and dispersion. Furthermore, the first moment m_1 leads to the so-called energy-weighted sum rule (EWSR) that is used to extract experimental strength functions. Also, the inverse-energy weighted sum rule (IEWSR) associated with the moment m_{-1} delivers, when applied to the dipole response, the so-called dipole polarizability that is relevant to the computation of radiative capture cross sections. Finally, the centroid energy of the monopole strength distribution gives access to the nucleus-dependent nuclear compressibility K_A that can eventually be linked to the nuclear matter incompressibility K_∞ [14, 15]. The latter quantity is a key characteristic of the nuclear equation of state and, as such, has a clear interest for several astrophysical applications.

The moments of a strength function can be computed in two ways. The first one involves an explicit sum over excited states and matrix elements of the simple one-body excitation operator F . The second one does involve the expectation value of a complicated many-body operator, but in the sole ground state. The first approach is presently denoted as the *sum over excited states* (SOES) method whereas the second one is referred to as the *ground-state expectation value* (GSEV) method. For a given many-body method, the agreement between the two approaches constitutes an internal-consistency test to pass¹ in order to correctly describe the excitation mode defined by the operator F .

In this context, the formal capacity to compute low-order moments via the GSEV approach is developed in Sects. 2 and Appendix D. Based on such an advancement, and after briefly introducing the numerical setting in Sect. 3, the SOES and GSEV approaches to m_1 are compared in Sect. 4 using the PGCM monopole responses of ^{16}O , ^{24}Mg , ^{28}Si and ^{46}Ti . Section 5 further discusses the impact of angular-momentum projection in the SOES approach. Next, PGCM moments are compared in Sect. 6 to those obtained via the quasi-particle random phase approximation (QRPA), the goal being to complement the study of Paper II [13] dedicated to the monopole strength function by focusing on its global characteristics. Section 7 focuses on the EWSR. First, it is demonstrated that the textbook expression of the EWSR must be corrected for the fact that nuclear excitations of interest are *intrinsic* excitations in the center-of-mass frame. Second, the intrinsic EWSR is tested in order to quantify the (unwanted) local-gauge symmetry breaking of the presently employed chiral Hamiltonian. Eventually, Sect. 8 is dedicated to accessing K_A in ^{16}O , ^{24}Mg , ^{28}Si and ^{46}Ti . The computed values are then employed to extract K_∞ and verify if the result thus

obtained is consistent with empirical expectations. This constitutes an important test for the chiral Hamiltonian under present use. The main findings of this work are summarised in Sect. 9 whereas a set of technical appendices complement the main body of the paper.

2 Moments of the strength function

In many respects, the present section follows Ref. [16]. By convention, all operators at play are redefined in such a way that their expectation value in the ground state is subtracted, i.e. for a given operator O its rescaled companion is introduced as²

$$\mathbf{O} \equiv O - \langle \Psi_0^{\sigma^0} | O | \Psi_0^{\sigma^0} \rangle, \quad (1)$$

where $|\Psi_0^{\sigma^0}\rangle$ denotes the ground state and the superscript σ^0 characterises its symmetry quantum numbers, as specified in the next paragraph. Equation (1) has the effect of removing the elastic part from the nuclear response function.

2.1 Definition

The strength function associated with a generic transition operator F reads as

$$S(E) \equiv \sum_{\nu\sigma} |\langle \Psi_\nu^\sigma | \mathbf{F} | \Psi_0^{\sigma^0} \rangle|^2 \delta(E_\nu^\sigma - E_0^{\sigma^0} - E), \quad (2)$$

where $E_0^{\sigma^0}$ denotes the ground-state energy whereas $\{|\Psi_\nu^\sigma\rangle(E_\nu^\sigma), \nu = 1, \dots, \nu_{\max}\}$ designates the set of excited states (energies). The superscript $\sigma \equiv (JM\Pi NZ)$ characterises the symmetry quantum numbers carried by the eigenstates, i.e. the angular momentum J and its projection M , the parity $\Pi = \pm 1$ as well as neutron N and proton Z numbers.

The k -th moment of the strength distribution³ associated with the operator \mathbf{F} is defined as⁴

$$m_k \equiv \int_0^{+\infty} E^k S(E) dE. \quad (3)$$

² Notice that Eq. (1) is no longer relevant as soon as one deals with commutators, since it is immediate to show that $[A - \langle A \rangle, B - \langle B \rangle] = [A, B]$.

³ From a mathematical standpoint, the moment m_k constitutes the k -th moment of a discretized probability distribution associated with the transition generated by F . Moments of a physical strength function are not guaranteed to be finite. The fact that it is indeed the case or not depends on mathematical characteristics of the Hamiltonian, e.g. of inter-nucleon interactions [16].

⁴ Except for m_0 , and as it will become evident below, there is in fact no difference in using \mathbf{F} or F in Eq. (8).

¹ It is a necessary but not sufficient condition for the response function associated with the operator F to be a good approximation of the exact one.

2.2 Mean energy(ies) and dispersion

Two sets of quantities having the dimension of an energy are introduced according to

$$\bar{E}_k \equiv \frac{m_k}{m_{k-1}}, \tag{4a}$$

$$\tilde{E}_k \equiv \left(\frac{m_k}{m_{k-2}}\right)^{1/2}. \tag{4b}$$

They coincide for all k 's if the strength distribution is concentrated in a single peak. The degree to which they differ reflects the fragmentation of the distribution. By definition, the average value of the energy distribution is given by

$$\bar{E}_1 = \frac{m_1}{m_0}. \tag{5a}$$

In this work the following energy averages are also employed

$$\tilde{E}_1 = \sqrt{\frac{m_1}{m_{-1}}}, \tag{5b}$$

$$\tilde{E}_3 = \sqrt{\frac{m_3}{m_1}}. \tag{5c}$$

Compared to the centroid energy \bar{E}_1 , the scaled (constrained) energy \tilde{E}_3 (\tilde{E}_1) is more sensitive to the high (low) energy part of the strength.

As shown in [Appendix A](#), the moments entertain the set of inequalities

$$\dots \geq \frac{m_{k+2}}{m_{k+1}} \geq \sqrt{\frac{m_{k+2}}{m_k}} \geq \frac{m_{k+1}}{m_k} \geq \sqrt{\frac{m_{k+1}}{m_{k-1}}} \geq \dots, \tag{6}$$

providing a practical tool to set boundaries on a specific moment in case it cannot be easily computed.⁵ Thanks to these inequalities, the variance of the strength distribution is shown to satisfy

$$\sigma^2 = \frac{m_2}{m_0} - \left(\frac{m_1}{m_0}\right)^2 \geq 0. \tag{7}$$

2.3 SOES formulation

Inserting Eq. (2) into Eq. (3) delivers the expression

$$m_k \equiv \sum_{\nu\sigma} (E_\nu^\sigma - E_0^{\sigma_0})^k |\langle \Psi_\nu^\sigma | \mathbf{F} | \Psi_0^{\sigma_0} \rangle|^2, \tag{8}$$

requiring the knowledge of excited states of the system. Equation (8) constitutes the SOES approach to the moments computation.

⁵ From a practical standpoint, Eq. (6) holds if the involved moments are all computed within the same approximation scheme.

2.4 GSEV formulation

By means of the identity resolution on the A -body Hilbert space \mathcal{H}_A

$$\mathbb{1} = \sum_{\nu\sigma} |\Psi_\nu^\sigma\rangle \langle \Psi_\nu^\sigma|, \tag{9}$$

Equation (8) can be rewritten as a ground-state expectation value

$$\begin{aligned} m_k &= \sum_{\nu\sigma} (E_\nu^\sigma - E_0^{\sigma_0})^k \langle \Psi_0^{\sigma_0} | \mathbf{F} | \Psi_\nu^\sigma \rangle \langle \Psi_\nu^\sigma | \mathbf{F} | \Psi_0^{\sigma_0} \rangle \\ &= \sum_{\nu\sigma} \langle \Psi_0^{\sigma_0} | \mathbf{F} (H - E_0^{\sigma_0})^k | \Psi_\nu^\sigma \rangle \langle \Psi_\nu^\sigma | \mathbf{F} | \Psi_0^{\sigma_0} \rangle \\ &= \langle \Psi_0^{\sigma_0} | \mathbf{F} (H - E_0^{\sigma_0})^k \mathbf{F} | \Psi_0^{\sigma_0} \rangle. \end{aligned} \tag{10}$$

Computing moments via Eq. (10) constitutes the GSEV method based on the expectation value of a complicated operator in the sole ground state.

Clearly, the complexity of the operator at play in Eq. (10) increases with $|k|$. For $k \geq 0$ the many-body rank increases with k whereas for $k < 0$ it further involves a non-trivial inversion. Examples of calculations employing the GSEV formulation can be found in Refs. [17, 18].

2.5 Moment operators

Positive moments can be re-expressed in more convenient forms by invoking the appropriate definition of moment operators. As shown in [Appendix B](#), moments with $k \geq 0$ can be further rewritten as

$$m_k = (-1)^i \langle \Psi_0^{\sigma_0} | C_i C_j | \Psi_0^{\sigma_0} \rangle \tag{11}$$

with

$$\begin{aligned} C_l &\equiv \{H^l, F\} \\ &\equiv \underbrace{[H, [H, \dots, [H, [H, F]] \dots]]}_{l \text{ times}} \end{aligned} \tag{12}$$

and where i and j are any pair of integers fulfilling $i + j = k$. By definition $C_0 \equiv F$.

For odd moments, Eq. (11) can be further expressed in terms of a commutator

$$m_k = \frac{1}{2} (-1)^i \langle \Psi_0^{\sigma_0} | [C_i, C_j] | \Psi_0^{\sigma_0} \rangle. \tag{13}$$

The last step provides a useful simplification to the structure of the operator whose ground-state expectation value is to be computed. Indeed, taking F to be a one-body operator, while the product $C_i C_j$ contains up to $[(n - 1)k + 2]$ -body operators, n being the highest-rank component of H , the commutator contains only up to $[(n - 1)k + 1]$ -body operators. Because even moments can only be written in terms of

anti-commutators that have the same many-body rank as the product $C_i C_j$, this simplification does not occur in this case.

Eventually, two sets of *moment operators* are introduced according to

$$\check{M}_k(i, j) \equiv (-1)^i C_i C_j \quad \forall k \geq 0, \quad (14a)$$

$$M_k(i, j) \equiv \frac{1}{2}(-1)^i [C_i, C_j] \quad \forall \text{ odd } k > 0, \quad (14b)$$

whose expectation value in $|\Psi_0^{\sigma_0}\rangle$ delivers m_k and where i and j are any pair of integers fulfilling $i + j = k$.

Based on a Hamiltonian H containing up to three-body operators, the algebraic expressions of the tensors defining $M_1(1, 0)$ are explicitly derived in [Appendix D](#). The result is used to numerically compute the PGCM m_1 moment associated with the monopole operator $F = r^2$ via the GSEV approach in [Sect. 4.2](#).

2.6 Alternative formulation

It is possible to access the operator $M_k(j + 1, j)$ associated with the odd positive moment m_k in an alternative way. To do so, the similarity-transformed Hamiltonian

$$\begin{aligned} H_k(\eta) &\equiv e^{-\eta G_k} H e^{\eta G_k} \\ &= H + \eta[H, G_k] + \frac{1}{2!}\eta^2[[H, G_k], G_k] + \mathcal{O}(\eta^3), \end{aligned} \quad (15)$$

is introduced, where the expansion in powers of the parameter η results from the application of Baker-Campbell-Hausdorff's identity. To match the expression given in [Eq. \(14b\)](#) one takes $i = j + 1$ and

$$G_k \equiv C_j \quad \text{with } k \equiv 2j + 1, \quad (16)$$

such that

$$M_k(j + 1, j) = \frac{1}{2}(-1)^{j+1} \left. \frac{\partial^2}{\partial \eta^2} H_k(\eta) \right|_{\eta=0}. \quad (17)$$

Based on a Hamiltonian H containing up to three-body operators, the algebraic expressions of the tensors defining $M_1(1, 0)$ are also derived in [Appendix D](#) as a way to validate the correctness of the expressions obtained via the more direct commutator approach laid down in [Sect. 2.5](#).

2.7 Practical merits and limitations

The great practical advantage of the GSEV approach is to access strength function's moments based on the sole knowledge of the nuclear ground state. This indeed is a tremendous simplification given that accessing a complete-enough set of

excited states constitutes a challenge within any state-of-the-art *ab initio* many-body method⁶

Such a benefit however comes at the price of evaluating the ground-state expectation value of operators [[Eq. \(14\)](#)] whose many-body complexity increases with the moment order. The set of moment operators indeed involve the hierarchy of operators

$$\begin{aligned} C_1 &= [H, F], \\ C_2 &= [H, C_1], \\ &\vdots \\ C_{j+1} &= [H, C_j], \end{aligned}$$

whose many-body rank increases with j due to the new commutator involved at each step. With F a one-body operator and H containing up to three-body operators, C_1 contains up to three-body operators, C_2 up to five-body operators, C_3 up to seven-body operators, i.e. C_j contains up to $(2j + 1)$ -body operators. As a result, $\check{M}_k(i, j)$ contains up to $(2k + 2)$ -body operators and $M_k(i, j)$ up to $(2k + 1)$ operators. For example, $\check{M}_0(0, 0)$ contains up to two-body operators and $M_1(1, 0)$ contains up to three-body operators. Knowing that dealing with three-body operators constitutes the current computational limit, it makes possible to compute both m_0 and m_1 exactly via the GSEV approach in PGCM calculations.⁷ Moving further, $\check{M}_2(1, 1)$ contains up to six-body operators and $M_3(2, 1)$ contains up to seven-body operators, which makes them beyond reach.⁸ While it is in principle possible to design approximations to $\check{M}_2(1, 1)$ and $M_3(2, 1)$ based on rank-reduction techniques [[19](#)], this avenue is not pursued in the present work and PGCM moments such as m_2 and m_3 are accessed via the SOES approach.

3 Numerical setting

All calculations presented here use the same setting as in [Paper II \[13\]](#). The PGCM formalism and the characteristics of the numerical applications were detailed in [Paper I \[12\]](#). In particular, the definitions of mean-square-radius-like operators under use all throughout the present paper read

⁶ Interestingly, once the investment to set up a moment operator has been done, it can be employed with any method delivering the many-body ground-state.

⁷ Because in practice the Hamiltonian is first rank-reduced to be an effective two-body operator [[19](#)], $M_1(1, 0)$ only contains up to two-body operators in present calculations.

⁸ Because the Hamiltonian is first rank-reduced to be an effective two-body operator [[19](#)], $\check{M}_2(1, 1)$ would only contain up to four-body operators whereas $M_3(2, 1)$ would only contain up to five-body operators in the present calculations. It still makes them beyond the reach of current capabilities.

$$r^2 \equiv \sum_{i=1}^A r_i^2 \tag{18}$$

for the monopole operator,

$$r_{\text{lab}}^2 \equiv \frac{1}{A} \sum_{i=1}^A r_i^2 \tag{19}$$

for the radius in the laboratory frame and

$$r_{\text{int}}^2 \equiv \frac{1}{A} \left(1 - \frac{1}{A}\right) \sum_{i=1}^A r_i^2 - \frac{1}{A^2} \sum_{i,j=1}^A \vec{r}_i \cdot \vec{r}_j, \tag{20}$$

for the radius in the center-of-mass or intrinsic frame.

A one-body spherical harmonic oscillator basis characterised by the optimal frequency $\hbar\omega = 12$ MeV is employed. All states up to $e_{\text{max}} \equiv \max(2n + l) = 10$ are included, with n the principal quantum number and l the orbital angular momentum. The representation of three-body operators is further restricted by only employing three-body states up to $e_{3\text{max}} = 14$.

A Hamiltonian based on chiral effective field theory (χ EFT) and built at next-to-next-to-next-to-leading-order (N^3 LO) [20] is employed. It contains consistent two- (2N) and three-nucleon (3N) interactions and is further evolved via similarity renormalization group (SRG) transformations [21] to the low-momentum scale $\lambda = 1.88 \text{ fm}^{-1}$ (i.e. flow parameter $\alpha=0.08 \text{ fm}^4$) and truncated at the three-body operator level. The resulting three-body force is approximated via the rank-reduction method developed in Ref. [19].

Two-dimensional (2D) PGCM calculations mix a set of constrained HFB states with axial symmetry using the root-mean-square radius $r \equiv \sqrt{\langle r_{\text{lab}}^2 \rangle}$ and the axial mass quadrupole deformation parameter β_2 as generator coordinates. The QRPA calculations are performed at the HFB minimum via the quasi-particle finite amplitude method (QFAM) [22]. The QFAM monopole moments are computed via the contour integration of the response function in the complex energy plane [23] employing a 100 MeV contour radius.

Eventually, the present analysis is based on the (P)GCM and QFAM monopoles responses of ^{16}O , ^{24}Mg , ^{28}Si and ^{46}Ti . For ^{28}Si , both the oblate ground state and the prolate isomeric state have been analyzed. In the following, thus, the notation ^{28}Si is used in reference to the ground state while $^{28}\text{Si}_{\text{iso}}$ is employed for the isomeric state. Given that the present paper only focuses on spectral moments, the reader is referred to Paper II [13] for a detailed analysis of the corresponding strength functions.

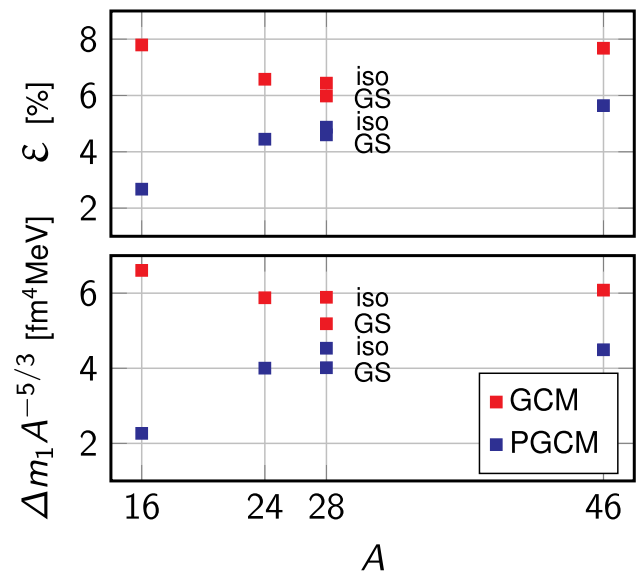


Fig. 1 Difference between monopole m_1 values obtained via the GSEV and SOES approaches in (P)GCM calculations as a function of A for ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti . Upper panel: difference in percentage. Lower panel: absolute difference multiplied by $A^{-5/3}$ to remove the expected trivial A dependence [see Eq. (25)]

4 SOES and GSEV approaches to m_1

4.1 Rationale

From a formal standpoint, the equivalence between the SOES and GSEV approaches relies on the completeness assumption from Eq. (9) allowing one to use the identity resolution. While the GSEV value of a given moment can be considered to be the formal value of reference, the SOES value is the one corresponding to the strength function actually computed in practice on the basis of a necessarily incomplete set of excited states.⁹ In this context, the agreement between the two values constitutes an internal-consistency test for the employed many-body method *relative to* the excitation operator F of interest. The agreement tests whether the vector $\mathbf{F}|\Psi_0^\sigma\rangle$ belongs to the subspace S spanned by the set of computed eigenstates $\{|\Psi_\nu^\sigma\rangle, \nu = 0, \dots, \nu_{\text{max}}\}$ explicitly at play in the SOES approach.

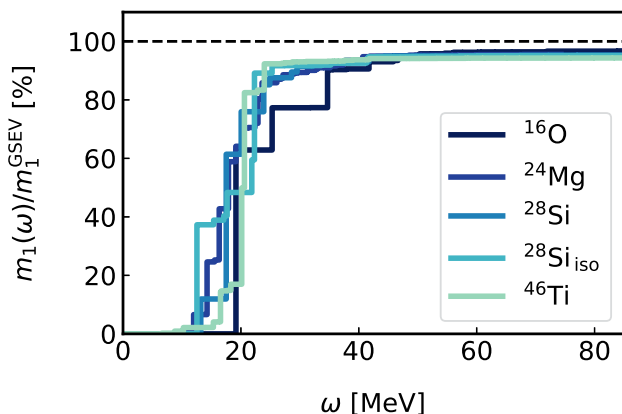
Presently employed (P)GCM eigenstates are linear combinations of the non-orthogonal (projected) Bogoliubov vacua defining the set $\{(P^\sigma)|\Phi(r^2, \beta_2)\}$ with $r \in [r_{\text{min}}, r_{\text{max}}]$ and $\beta_2 \in [\beta_{\text{min}}, \beta_{\text{max}}]$ ($\sigma \in \text{IRREPs}$). Consequently, the (P)GCM subspace $S^{(\sigma)}$ is nothing but the span of that set. In Appendix C of Ref. [24], the monopole and quadrupole operators were shown to be indeed exhausted for a GCM calculation based on Slater determinants built out of the

⁹ The moment obtained via the GSEV approach constitutes an upper bound of the value computed via the SOES approach.

Table 1 GCM and PGCM m_1 monopole moments computed via the SOES and GSEV approaches for ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

	^{16}O		^{24}Mg		^{28}Si		$^{28}\text{Si}_{\text{iso}}$		^{46}Ti	
	SOES	GSEV	SOES	GSEV	SOES	GSEV	SOES	GSEV	SOES	GSEV
GCM	7940	8611	16,676	17,850	21,046	22,384	22,104	23,625	43,185	46,776
PGCM	8386	8617	17,178	17,978	21,490	22,526	22,846	24,016	44,392	47,046

All quantities are in fm^4MeV

**Fig. 2** Integral $m_1(\omega)$ moment, as defined in Eq. (21), as a function of the maximum excitation energy and normalised by the GSEV value of m_1 from PGCM monopole calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

lowest eigenstates of axially deformed harmonic oscillators, the two generator coordinates being the corresponding axial and perpendicular oscillator frequencies. While realistic (P)GCM calculations rely on more general Bogoliubov vacua (and include particle-number and angular-momentum projections), such a proof gives some confidence that the monopole operator might be well exhausted in present 2D (P)GCM calculations using r^2 and β_2 as generator coordinates. It is the goal of the present section to test quantitatively to which extent this is indeed the case for m_1 .

4.2 Results

The (P)GCM m_1 values obtained from both evaluation methods are reported in Table 1. Furthermore, their difference [rescaled according to their expected $A^{5/3}$ scaling; see Eq. (25)] is displayed in Fig. 1 along with the difference in percentage.

Results obtained via the SOES approach are about 6–7% smaller than their GSEV counterpart across the five cases under consideration. The underestimation of the SOES approach is stable from $A = 16$ to $A = 46$ once the $A^{5/3}$ scaling has been removed. The small but systematic improvement of the PGCM over the GCM is attributed to the benefit of the symmetry restoration, i.e. symmetry contaminants are

removed by the angular momentum projection on $J = 0$ such that the operator r^2 is better exhausted by the corresponding subspace S_P . For PGCM calculations the SOES m_1 moment as a function of the maximum excitation energy, reading

$$m_1(\omega) \equiv \int_0^\omega E S(E) dE, \quad (21)$$

is displayed in Fig. 2 normalised by the corresponding GSEV value. The excited states included in the SOES evaluation reach a maximum energy of 97 MeV for ^{16}O , 74 MeV for ^{24}Mg , 94 and 98 MeV for the ground and isomeric state of ^{28}Si and 102 MeV for ^{46}Ti .

Eventually, the operator r^2 is exhausted, within a few percents, by the (P)GCM subspace $S_{(P)}$. This translates into the fact that the SOES approach to m_1 can be safely used within a few percent uncertainty.¹⁰ Differences between the GSEV and SOES approaches signal the necessity of improving the determination of an optimal (P)GCM subspace $S_{(P)}$. While this topic is of current interest, it goes beyond the scope of the present article.

5 Angular-momentum projection

The effect of angular momentum projection on the monopole moments m_k , $k = -1, 0, 1, 2, 3$, evaluated via the SOES approach is presently quantified by comparing results from GCM and PGCM calculations. As seen in Table 2, the angular momentum projection systematically enlarges m_k in a way that increases with k . In fact, while the increase with the moment order is rather marked in ^{16}O , it is limited in ^{24}Mg and has entirely disappeared in ^{46}Ti . Thus, and even though the range of nuclei presently tested is too limited to draw general conclusions, the impact of the angular momentum projection seems to decrease with A .

¹⁰ The resulting uncertainty for a moment m_k can be conjectured to increase with k . Indeed, the energy weight E^k entering m_k accentuates the importance of higher-energy states as k increases while the truncation of the completeness relation in the SOES approach probably affects more this higher-energy domain. Given that m_1 is the highest moment that can be computed exactly within the GSEV approach, this conjecture cannot be presently tested.

Table 2 Monopole moments computed using the SOES approach for GCM and PGCM calculations of ^{16}O , ^{24}Mg and ^{46}Ti

SOES	^{16}O			^{24}Mg			^{46}Ti		
	$(\beta_2 = 0.00)$			$(\beta_2 = 0.56)$			$(\beta_2 = 0.27)$		
	GCM	ε [%]	PGCM	GCM	ε [%]	PGCM	GCM	ε [%]	PGCM
m_{-1} [$\text{fm}^4\text{MeV}^{-1}$]	17.83	0.4	17.90	57.95	0.4	58.16	120.75	0.1	120.86
m_0 [fm^4]	369	2.1	377	955	1.4	969	2226	1.7	2264
m_1 [fm^4MeV]	7940	5.2	8386	16,676	2.9	17,178	43,185	2.7	44,392
m_2 [fm^4MeV^2]	182,667	11.0	205,277	315,053	4.4	329,629	871,376	3.2	900,105
m_3 [fm^4MeV^3]	4,718,706	19.8	5,887,075	6,661,187	5.3	7,031,885	18,469,397	2.7	18,976,050

Numbers in between GCM and PGCM results indicate the variation between the former and the latter in percentage

Table 3 Average energies and dispersion computed using the SOES for GCM and PGCM calculations of ^{16}O , ^{24}Mg and ^{46}Ti

SOES	^{16}O			^{24}Mg			^{46}Ti		
	$(\beta_2 = 0.00)$			$(\beta_2 = 0.56)$			$(\beta_2 = 0.27)$		
	GCM	ε [%]	PGCM	GCM	ε [%]	PGCM	GCM	ε [%]	PGCM
\bar{E}_1	21.10	2.5	21.64	16.96	1.3	17.19	18.91	1.3	19.17
\bar{E}_1	21.51	3.4	22.26	17.46	1.5	17.72	19.40	1.1	19.60
σ	5.67	19.3	7.02	5.01	1.7	5.09	3.89	7.1	3.63

All results are expressed in MeV units. Numbers in between GCM and PGCM results indicate the variation between the former and the latter in percentage

While the behavior of specific moments is interesting, it is more pertinent to investigate how this translates into the modification of physically-relevant quantities, e.g. the mean value and the dispersion of the monopole strength function. As visible from Table 3, the impact of the angular momentum projection on the centroid energy \bar{E}_1 decreases from 3.4% in ^{16}O to 1.5% in ^{24}Mg , and eventually down to 1.1% in ^{46}Ti . Except in ^{16}O , where it amounts to 750 keV, the GMR energy shift due to angular momentum projection is thus essentially negligible. The situation is similar for \bar{E}_1 that is used as an alternative to evaluate the GMR energy.

The impact on the dispersion is typically more significant. Again, the set of nuclei is too limited to draw general conclusions. Still, the dispersion varies by as much as 19.3% in ^{16}O and 7.1% in ^{46}Ti . In ^{24}Mg , the impact of angular momentum projection on σ is small but the strongly fragmented monopole strength is in fact significantly modified as can be seen in Paper II [13], which reflects the fact that \bar{E}_1 and σ are anyway insufficient to characterize the behavior of the strength in such a case.

6 Comparison to QRPA

In Paper II [13], the QFAM and GCM monopole strengths of ^{16}O , ^{24}Mg , ^{28}Si and ^{46}Ti were discussed at length. In comparing GCM to QFAM calculations, the objective is to analyse the impact of additional ground-state correlations in

the evaluation of moments, excluding the effects associated with symmetry restoration, which were addressed in Sect. 5. The GCM and QFAM m_1 values calculated within the GSEV approach are compared in Table 4 and shown to agree to better than 3% across the four nuclei under consideration, GCM values being systematically larger than QFAM ones.

The QRPA is known to fully exhaust any one-body excitation operator in odd- k moments, i.e. it can be shown analytically that, within the quasi-boson approximation, odd- k QRPA moments computed with the GSEV and the SOES approaches are strictly identical, the state at play in the GSEV being the HFB ground state [25–27]. This result demonstrates the internal consistency of the QRPA as far as strength functions are concerned. While the GCM does not strictly share this property as discussed above, the GCM ground-state is *necessarily* a better approximation of the exact ground state than the HFB state, such that GCM moments based on the GSEV approach are necessarily better than QRPA ones¹¹. This is testified by the larger values of the GCM m_1 moment reflecting the beneficial impact of (static) correlations associated with fluctuations of r^2 and β_2 leading to slightly larger GCM mean-square radii compared to HFB ones.¹²

¹¹ This recalls that the capacity of a method to fully exhaust the strength of a given excitation operator is *not* a sufficient condition to deliver a better approximation of the exact moments than a method that does not fully exhaust it.

¹² The argument qualitatively relates to the EWSR expressing m_1 in terms of the ground-state mean-square matter radius.

Table 4 Monopole m_1 moment computed via the GSEV approach for GCM and QFAM calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

GSEV	^{16}O		^{24}Mg		^{46}Ti		^{28}Si		$^{28}\text{Si}_{\text{iso}}$	
	QFAM	GCM	QFAM	GCM	QFAM	GCM	QFAM	GCM	QFAM	GCM
m_1 [fm^4MeV]	8356	8611	17,478	17,850	46,387	46,776	22,080	22,384	23,075	23,625

The trend with A of the difference between GCM and QRPA m_1 values is better inferred from Fig. 3. Given the hypothesis at the heart of the QRPA, such a difference is expected to increase with the degree of *anharmonicity* of the system. As expected, and as discussed in Sect. 6 of Paper II [13], larger systems are more harmonic than lighter ones. This is indeed consistent with the fact that the difference with GCM values decreases with A . This interpretation is further supported by Fig. 4 where the difference is shown to grow with the size of the cubic coefficient¹³ a_3 extracted in Sect. 6 of Paper II [13].

Figure 3 also displays the deviation of QFAM m_1 values from GCM results based on the SOES approach. In this case, QFAM values are systematically a few percent *above* GCM ones; i.e. they are located in between the two sets of GCM values. Eventually, the disagreement between QRPA and GCM is smaller than the uncertainty in the evaluation of the GCM values. Contrary to values based on the GSEV approach, GCM values obtained from the SOES approach do not converge towards QRPA as A increases and thus do not scale as expected with the harmonic character of the system.

Eventually, the centroid and the dispersion of the QRPA monopole strength function are compared to GCM values based on the SOES approach in Table 5. The GCM centroid energy is typically 4–6% below the QRPA one¹⁴, which amounts to less than 1 MeV difference in the studied nuclei. The QRPA and GCM dispersions are also very consistent, especially in view of the remaining many-body uncertainty.

7 Energy weighted sum rule

7.1 Definition

The EWSR is a standard quantity in studies of giant resonances, and its exhaustion by a specific nuclear excitation serves as a good indicator of its degree of collectivity [16,28,29]. Furthermore, the EWSR is used to extract

¹³ The cubic coefficient is rescaled by $A^{-3/2}$ to remove its trivial A dependence due to the use of the rms radius as the variable in the fitted function; see Paper II [13] for details.

¹⁴ While it is true for the centroid of the actually computed GCM strength function (SOES value), the formal (GSEV) value not computed here is probably *higher* in view of the behavior of m_1 studied above.

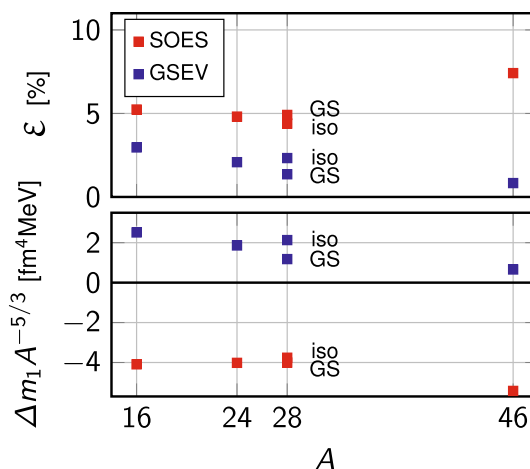


Fig. 3 Difference between the GCM and the QRPA monopole m_1 values as a function of A for ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti . The GCM moment is evaluated both through the SOES and GSEV approaches. Upper panel: absolute difference in percentage. Lower panel: difference multiplied by $A^{-5/3}$ to remove the expected trivial A dependence [see Eq. (25)]

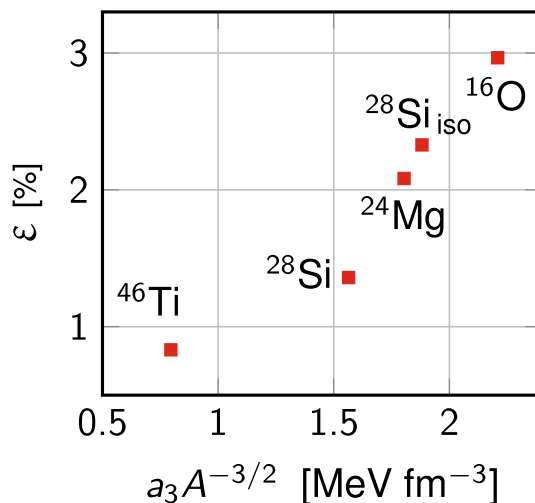


Fig. 4 Relative difference between the QFAM and the GSEV GCM monopole m_1 values as a function of the cubic coefficient a_3 (see Sect. 6 of Paper II [13]). The factor $A^{-3/2}$ is included to remove the trivial A dependence of a_3 . See text for details

strength functions from experimental data as briefly recalled in Appendix E.

The ESWR relies on an analytical evaluation of m_1 via Eq. (14), i.e. using the GSEV approach. Targeting the first

Table 5 Centroid energy and dispersion from QFAM and GCM(SOES) calculations of ^{16}O , ^{24}Mg and ^{46}Ti

	^{16}O			^{24}Mg			^{46}Ti		
	$(\beta_2 = 0.00)$			$(\beta_2 = 0.56)$			$(\beta_2 = 0.27)$		
	QFAM	ε [%]	GCM	QFAM	ε [%]	GCM	QFAM	ε [%]	GCM
\bar{E}_1	22.33	3.8	21.51	18.48	5.9	17.46	20.15	3.9	19.40
σ	5.55	2.1	5.67	5.58	11.5	5.01	3.96	1.8	3.89

All results are expressed in MeV units. Numbers in between QFAM and GCM results indicate the variation between the former and the latter in percentage

moment of the isoscalar monopole strength function, the similarity transformation of the Hamiltonian in Eq. (15) computed with the isoscalar local operator $G_1 = F = r^2$ is nothing but a local-gauge transformation. In case the inter-nucleon interaction V is local-gauge invariant, Eq. (15) reads

$$\begin{aligned} H_1(\eta) &= e^{-\eta r^2} H e^{\eta r^2} \\ &= e^{-\eta r^2} T e^{\eta r^2} + e^{-\eta r^2} V e^{\eta r^2} \\ &= e^{-\eta r^2} T e^{\eta r^2} + V \\ &= T(\eta) + V, \end{aligned} \tag{22}$$

such that V does not contribute to the quadratic term in η entering Eq. (15). Under such an assumption, thus, only the kinetic-energy operator T contributes to $M_1(1, 0)$. The kinetic term is then divided into its laboratory- and center-of-mass component according to

$$\begin{aligned} H &= T + V \\ &\equiv T_{\text{lab}} - T_{\text{cm}} + V. \end{aligned} \tag{23}$$

If the sole laboratory-frame component

$$T_{\text{lab}} \equiv \frac{1}{2m} \sum_{i=1}^A p_i^2 \tag{24}$$

is considered in the evaluation of the EWSR starting from $M_1(1, 0)$, m_1 is then obtained analytically under the form [16,30]

$$\text{EWSR}_{\text{lab}}(r^2) = \frac{2\hbar^2 A}{m} \langle \Psi_0^{\sigma_0} | r_{\text{lab}}^2 | \Psi_0^{\sigma_0} \rangle, \tag{25}$$

which constitutes the textbook EWSR formula for the isoscalar monopole mode. Interestingly, $\text{EWSR}_{\text{lab}}(r^2)$ is proportional to the laboratory-frame ground-state mean-square matter radius from Eq. (19). Thus, accessing it only requires the computation of that mean-square radius from the many-body method of interest, e.g. using the Bogoliubov state at the HFB minimum in QFAM calculations or the (P)GCM ground state in (P)GCM calculations.

However, it happens that nuclear excitations of interest are *intrinsic* excitations. Consequently, present many-body calculations employ the intrinsic Hamiltonian H from

Eq. (23) containing the intrinsic kinetic-energy operator $T_{\text{int}} \equiv T_{\text{lab}} - T_{\text{cm}}$, with the subtracted center-of-mass kinetic-energy operator reading as

$$T_{\text{cm}} \equiv \frac{1}{2mA} \sum_{i,j=1}^A \vec{p}_i \cdot \vec{p}_j. \tag{26}$$

In this context, the monopole m_1 moment reads as

$$\begin{aligned} m_1 &= \frac{1}{2} \langle \Psi_0^{\sigma_0} | [r^2, [H, r^2]] | \Psi_0^{\sigma_0} \rangle \\ &= \text{EWSR}_{\text{lab}}(r^2) - \frac{1}{2} \langle \Psi_0^{\sigma_0} | [r^2, [T_{\text{cm}}, r^2]] | \Psi_0^{\sigma_0} \rangle \\ &\equiv \text{EWSR}_{\text{lab}}(r^2) - \frac{2\hbar^2 A}{m} \langle \Psi_0^{\sigma_0} | R_{\text{cm}}^2 | \Psi_0^{\sigma_0} \rangle, \end{aligned} \tag{27}$$

with \vec{R}_{cm} the center-of-mass position vector. The derivation of the correction term from T_{cm} is provided in Appendix F. Eventually, the two terms can be combined such that the appropriate, i.e. *intrinsic*, ESWR is given by

$$\text{EWSR}_{\text{int}}(r^2) = \frac{2\hbar^2 A}{m} \langle \Psi_0^{\sigma_0} | r_{\text{int}}^2 | \Psi_0^{\sigma_0} \rangle, \tag{28}$$

and thus amounts to using the intrinsic mean-square radius from Eq. (20) rather than the laboratory-frame one.

The $\text{EWSR}_{\text{int}}(r^2)$ must in principle be fulfilled in *ab initio* calculations given that χ EFT-based 2N and 3N interactions are meant to be local-gauge invariant, which is a necessary condition to achieve a consistent coupling to the electromagnetic field [31]. However, enforcing the local-gauge invariance is not straightforward in practice. First, it cannot be exactly fulfilled if the same EFT truncation level is applied to both nuclear interactions and currents, even in the case of dimensional regularization. Second, the use of (nonlocal) cutoff regulators makes its fulfillment even more challenging [32]. Eventually, existing χ EFT-based 2N and 3N interactions are not strictly local-gauge invariant and it is our goal to quantify such a feature by testing the departure of $\text{EWSR}_{\text{int}}(r^2)$ from the computed m_1 .

The potential breaking of the local gauge invariance can be straightforwardly formulated by schematically expressing

Table 6 Isoscalar monopole $EWSR_{\text{int}}$ from Eq. (28) for QFAM and PGCM calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

$EWSR_{\text{int}}$	QFAM	PGCM
^{16}O	8462	8475
^{24}Mg	17,480	17,658
^{28}Si	22,004	22,129
$^{28}\text{Si}_{\text{iso}}$	22,883	23,352
^{46}Ti	45,600	45,844

All results are expressed in fm^4MeV

Table 7 Isoscalar monopole $EWSR_{\text{lab}}$ from Eq. (25) for QFAM and PGCM calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

$EWSR_{\text{lab}}$	QFAM	PGCM
^{16}O	8832	8851
^{24}Mg	17,885	18,064
^{28}Si	22,415	22,540
$^{28}\text{Si}_{\text{iso}}$	23,287	23,749
^{46}Ti	46,053	46,299

All results are expressed in fm^4MeV

the intrinsic Hamiltonian as

$$\begin{aligned}
 H &= T_{\text{lab}} - T_{\text{cm}} + V \\
 &\equiv T_{\text{lab}} - T_{\text{cm}} + V_{\text{lgf}} + V_{\text{lgf}}, \quad (29)
 \end{aligned}$$

where $V_{\text{lgf}} \equiv V - V_{\text{lgf}}$ formally defines the departure of the nuclear interactions from their local-gauge invariant formulation. Given Eq. (29), the monopole m_1 moment effectively reads in practice as

$$\begin{aligned}
 m_1 &= \frac{1}{2} \langle [r^2, [H, r^2]] \rangle_{\text{gs}} \\
 &= EWSR_{\text{int}}(r^2) + \frac{1}{2} \langle [r^2, [V_{\text{lgf}}, r^2]] \rangle_{\text{gs}} \\
 &\equiv EWSR_{\text{int}}(r^2) + \delta m_1^{\text{lgf}}, \quad (30)
 \end{aligned}$$

where δm_1^{lgf} quantifies the effective breaking of $EWSR_{\text{int}}(r^2)$.

7.2 $EWSR_{\text{lab}}$ versus $EWSR_{\text{int}}$

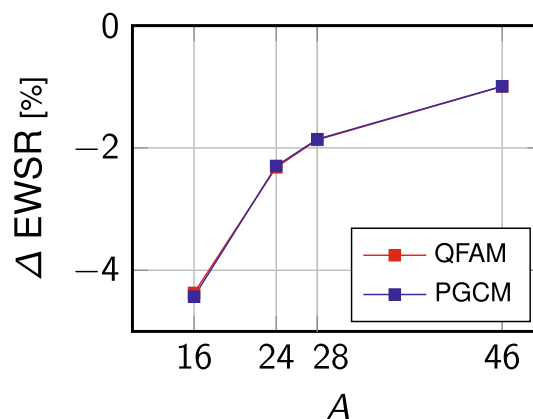
Values of $EWSR_{\text{int}}(r^2)$ ($EWSR_{\text{lab}}(r^2)$) from QFAM and PGCM calculations are reported in Table 6 (Table 7). While PGCM values are systematically larger, the difference is eventually very small. These features reflect the behavior of the point-matter radii reported in Table 8.¹⁵

¹⁵ The difference between HFB and PGCM radii relates to the impact of so-called static correlations beyond the mean-field included into

Table 8 Ground-state expectation value of point-matter nuclear radii in the laboratory and intrinsic frames for HFB and PGCM calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

	$\sqrt{\langle r_{\text{lab}}^2 \rangle_{\text{gs}}}$		$\sqrt{\langle r_{\text{int}}^2 \rangle_{\text{gs}}}$	
	HFB	PGCM	HFB	PGCM
^{16}O	2.580	2.583	2.525	2.527
^{24}Mg	2.998	3.013	2.963	2.979
^{28}Si	3.107	3.115	3.078	3.087
$^{28}\text{Si}_{\text{iso}}$	3.167	3.198	3.139	3.171
^{46}Ti	3.474	3.484	3.457	3.467

All results are expressed in fm

**Fig. 5** Relative difference between the monopole $EWSR_{\text{int}}$ and $EWSR_{\text{lab}}$ as a function of A for QFAM and PGCM calculations

The relative difference between $EWSR_{\text{int}}$ and $EWSR_{\text{lab}}$ is plotted as a function of A in Fig. 5. Results are identical for QFAM and PGCM calculations. The A dependence of the difference relates to the $-A^{-1}$ scaling driven by the center-of-mass-correction entering $EWSR_{\text{int}}$, as is analytically demonstrated in Appendix F for its one-body component. Again, the trend reflects directly how the difference of the mean-square point matter radii computed in the intrinsic frame and in the laboratory frame decreases with A .

the PGCM ansatz. In general, static correlations have little impact on radii, the exceptions being light spherical nuclei in which they can non-negligibly increase radii and transitional nuclei in which they can strongly reduce them. In nuclei displaying a sharp total energy surface around the deformed HFB minimum, as is the case here, the impact of static correlations on the mean-square radius is typically very small [33]. The presently employed χ EFT Hamiltonian typically delivers good radii such that the further addition of missing dynamical correlations to the PGCM ansatz [19] is expected to enlarge radii.

Table 9 Percent agreement between GSEV-based m_1 monopole moment and $EWSR_{int}$ in PGCM and QFAM calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti

% of $EWSR_{int}$	QFAM	PGCM
^{16}O	98.74	101.67
^{24}Mg	99.99	101.81
^{28}Si	100.35	101.80
$^{28}\text{Si}_{iso}$	100.84	102.85
^{46}Ti	101.73	102.62

7.3 Departure from sum rules

The actual deviation from the values provided in Table 6 is now tested. Introducing the discrepancy in percentage

$$\varepsilon \equiv \left(\frac{m_1}{EWSR_{int}} - 1 \right) \times 100, \quad (31)$$

the percentage $(100 + \varepsilon)$ of $EWSR_{int}(r^2)$ is reported in Table 9 for QFAM and PGCM calculations. It is observed that PGCM results overshoot the $EWSR_{int}(r^2)$ by [1.7, 2.6]% whereas QFAM results are slightly lower and fulfill it within $[-1.3, +0.8]\%$.

Overall, the violation of $EWSR_{int}(r^2)$ due to the breaking $\delta m_1^{\chi EFT}$ of local-gauge invariance by the presently employed χEFT interactions is small and remains below 3% in the present calculations. Still, it manifests slightly differently depending on the (approximate) many-body method, the nucleus or the eigenstate under consideration.

To illustrate this point more transparently, ε is plotted in Fig. 6 as a function of A . The difference to $EWSR_{int}(r^2)$ evolves with A for the ground states under consideration and is systematically larger for PGCM than for QRPA. One observes that the trend with A is flatter for the PGCM and that QFAM results seem to approach PGCM ones as the mass increases. One may conjecture that this is a sign of better convergence of the PGCM ground-state.

Eventually, a thorough investigation of the violation of $EWSR_{int}(r^2)$ requires a larger set of nuclei and excited states as well as to employ an expansion many-body method at various truncation orders. Furthermore, $\delta m_1^{\chi EFT}$ must be studied as a function of the chiral order and for various regularizations of the employed χEFT interactions. Such a systematic study is left to a future work.

8 Nuclear incompressibility

The monopole breathing mode probes the compressibility of nuclear matter. Consequently, the infinite matter incompressibility modulus K_∞ has been extracted based on micro-

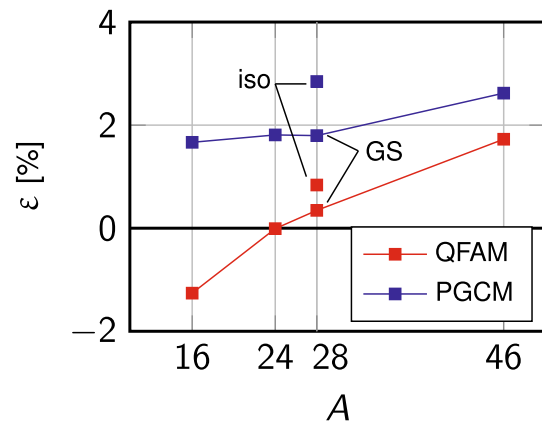


Fig. 6 Percent variation of the computed monopole m_1 compared to the corresponding $EWSR_{int}$ as a function of A for QFAM and PGCM calculations of ^{16}O , ^{24}Mg , ^{28}Si (ground-state and prolate isomer) and ^{46}Ti . The PGCM values are based on the GSEV approach

scopic calculations of E_{GMR} , typically within the frame of the nuclear energy density functional (EDF) method [3]. As a matter of fact, the extraction procedure is not unambiguous in itself. Furthermore, while originally applying it to a couple of doubly closed-shell nuclei (^{208}Pb and ^{92}Zr) led to consistent values of K_∞ , the more recent use of open-shell nuclei produced conflicting results [3, 34–40].

The goal is to extract the value of K_∞ associated with χEFT -based interactions via ab initio calculations. In EDF calculations, it has become customary to extract K_∞ by computing directly the symmetric nuclear matter equation of state, while checking that E_{GMR} is well reproduced in a selected set of finite nuclei on the basis of the same EDF parameterization. Another approach, presently in use, consists of extracting K_∞ from the leptodermous expansion of the finite-nucleus compressibility modulus computed microscopically [14]. While the former approach typically carries smaller uncertainties, the latter bypasses the need to compute the infinite matter equation of state.

The second approach was recently employed to extract K_∞ for NNLO_{sat} [41] and NNLO_{opt} [42] χEFT -based Hamiltonians via symmetry-adapted no core shell model (SA-NCSM) calculations of ^4He , ^{16}O , ^{20}Ne and ^{40}Ca [43]. The extracted result for NNLO_{sat} ($K_\infty = 297(37)$ MeV) was shown to be consistent, within the rather large extrapolation uncertainties, with the value ($K_\infty = 253$ MeV) based on the computation of the equation of state with the same Hamiltonian [41].

Following the same protocol but only relying on a set of intrinsically-deformed nuclei, i.e. ^{24}Mg , ^{28}Si and ^{46}Ti , the compressibility modulus K_∞ associated with the $N^3\text{LO}$ Hamiltonian under use [20] is presently estimated based on PGCM and QRPA calculations.

Table 10 Average GMR energies in MeV computed from QFAM and PGCM calculations according to Eqs. (4)

E_{GMR}	QFAM		PGCM		
	\bar{E}_1	\bar{E}_3	\bar{E}_1	\bar{E}_1	\bar{E}_3
^{24}Mg	18.48	21.19	17.19	17.72	20.23
^{28}Si	18.88	20.91	18.04	18.45	20.60
^{46}Ti	20.15	21.33	19.17	19.60	20.68

Table 11 Finite-nucleus compression modulus K_A (in MeV) computed from QFAM and PGCM calculations

K_A	QFAM		PGCM		
	\bar{E}_1	\bar{E}_3	\bar{E}_1	\bar{E}_1	\bar{E}_3
^{24}Mg	74.0	97.3	64.6	68.7	89.5
^{28}Si	83.0	101.8	76.2	79.7	99.4
^{46}Ti	118.2	132.4	107.5	112.5	125.1

Values are categorised according to the definition of the GMR energy [see Eqs. (4)] employed to compute K_A via Eq. (32)

8.1 Finite-nucleus compression modulus

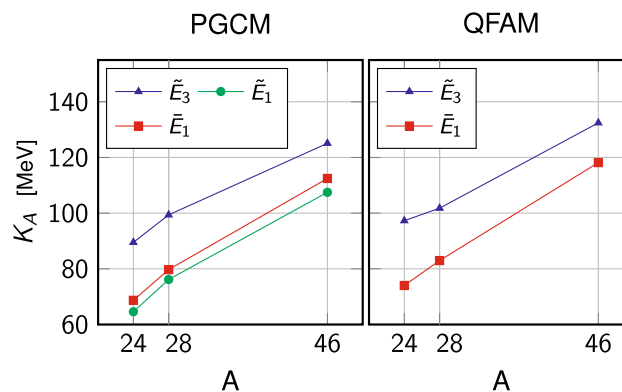
The first step consists of accessing the finite-system compression modulus given by [14]

$$K_A \equiv \frac{m}{\hbar^2} \langle \Psi_0^{\sigma_0} | r_{\text{lab}}^2 | \Psi_0^{\sigma_0} \rangle E_{\text{GMR}}^2, \quad (32)$$

which thus requires the ground-state mean-square matter radius and the GMR energy as inputs. In finite, especially light and deformed, nuclei the GMR strength is not concentrated into a single peak. Consequently, the choice of E_{GMR} to be used in Eq. (32) is neither unique nor obvious. Specific derivations support the use of \bar{E}_1 or \bar{E}_3 whereas general arguments also motivate the use of the centroid energy \bar{E}_1 [14]. In the following, all three cases are tested.¹⁶

Based on the GMR energies provided in Table 10, the set of K_A values are given in Table 11 and displayed in Fig. 7 as a function of A . The higher values of K_A in QRPA than in PGCM reflects the characteristics of the GMR energies pointed out earlier on whenever computing PGCM moments via the SOES approach as presently done. The spread of K_A values depending on the definition of E_{GMR} is the manifestation that \bar{E}_1 (\bar{E}_3) is more sensitive to the part of the strength located at lower (higher) energies than \bar{E}_1 . Eventually, K_A can typically vary by as much as 30% in ^{24}Mg depending on that choice. However, this variation quickly decreases with A to reach 14% in ^{46}Ti . Such a trend is encouraging in view of extracting K_∞ .

¹⁶ Whenever a single mode exhausts the complete monopole response, the three energies are the same and the choice is thus straightforward.

**Fig. 7** Finite-nuclei compression modulus K_A as a function of A for PGCM and QFAM calculations. Different definitions of the average GMR energy E_{GMR} entering Eq. (32) are used, see Eq. (5) for the notation

8.2 Extraction of K_∞

The method to extract K_∞ is based on the leptodermous expansion of K_A given by [14]

$$K_A = K_{\text{vol}} + K_{\text{surf}}A^{-1/3} + K_{\text{Coul}}Z^2A^{-4/3} + K_{\text{sym}}\beta^2, \quad (33)$$

where K_{vol} , K_{surf} , K_{Coul} and K_{sym} are the volume, surface, Coulomb and symmetry contributions to the compression modulus, respectively. The parameter β characterizes the isospin asymmetry

$$\beta \equiv \frac{N - Z}{N + Z}, \quad (34)$$

where N (Z) denotes the neutron (proton) number. Equation (33) is fitted based on the values of K_A given in Table 11 and K_{vol} is interpreted as the infinite nuclear matter incompressibility K_∞ . Given that the Coulomb and symmetry terms do not significantly impact the asymptotic behaviour of K_A for very large A [43], K_∞ can be obtained via a simple linear fit in the variable $x \equiv A^{-1/3}$

$$K_A = K_\infty + K_{\text{surf}}x. \quad (35)$$

While the linear fits are displayed in Fig. 8, the corresponding values of K_∞ and K_{surf} are reported in Table 12 along with the uncertainties associated with the fit. The extracted incompressibility is $K_\infty \approx 290$ MeV. While QRPA central values are a few MeV higher than PGCM ones, they only differ by about 3.3% and 4.2% when using $E_{\text{GMR}} \equiv \bar{E}_1$ and $E_{\text{GMR}} \equiv \bar{E}_3$, respectively. Eventually, QRPA and PGCM values are consistent within extrapolation uncertainties, which are significantly larger for QRPA than for PGCM results.¹⁷

¹⁷ The tiny extrapolation uncertainty of the PGCM results might be accidental, i.e. it may simply reflect the small number of points employed in the fit rather than a genuine behavior following strictly the

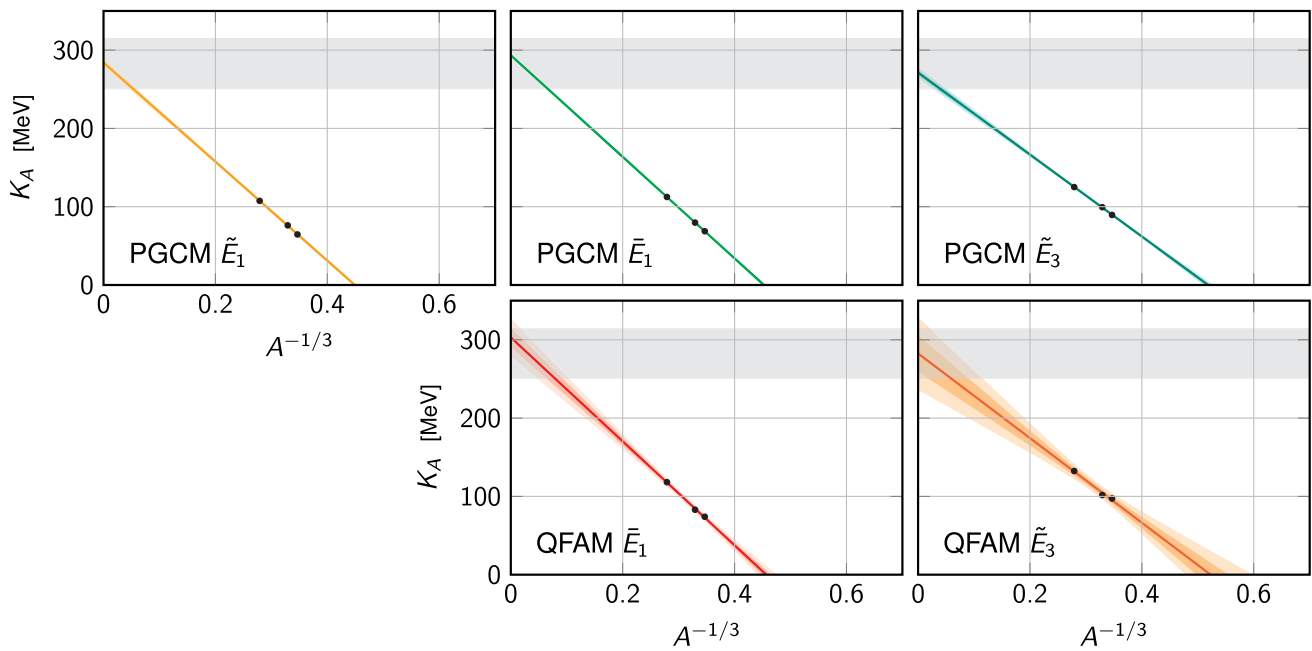


Fig. 8 Finite-nucleus compression modulus K_A as a function of $A^{-1/3}$ obtained from PGCM and QRPA calculations (black circles). The best fit is shown in all cases with the corresponding 1σ (darker shade)

and 2σ (lighter shade) bands of regression (see, e.g., Chap. 3 of Ref. [44]). The shaded gray area represents the empirically accepted range $250 < K_\infty < 315$ MeV [45]

Interestingly, while the hierarchy $K_A(\tilde{E}_1) < K_A(\bar{E}_1) < K_A(\tilde{E}_3)$ is systematically valid for all computed nuclei with $A \leq 46$, the trends are such that the extrapolation to very large A values leads to K_∞ being the smallest for $E_{\text{GMR}} \equiv \bar{E}_3$. Eventually, the nuclear matter incompressibility varies by 6.6% (7.5%) for QRPA (PGCM) between the two extreme values obtained for \tilde{E}_3 and \bar{E}_1 . This confirms the trend observed above for K_A as a function of A .

In Fig. 8, the shaded gray area accounts for the generally accepted range $250 < K_\infty < 315$ MeV [45]. All values of K_∞ fall, within extrapolation uncertainties, into this region.¹⁸

Uncertainties of the present theoretical predictions (partially) evaluated in Paper I [12] are not presently propagated to K_∞ . While they are not negligible, they are typically sub-leading compared to the uncertainties associated with the choice of E_{GMR} and with the extrapolation based on the leptodermous expansion. While the range of masses presently used in the fit allows one to make quantitative statement, the use of (much) heavier systems in the future will help reducing the extrapolation uncertainty and ensure the stability of the fit. In any case, and as already stipulated in Ref. [43], the present work demonstrates that extrapolating the finite-nuclei compressibility modulus for a large enough set of nuclei can

Table 12 Fitting parameters for the linear extrapolation given in Eq. (35)

	K_∞	K_{surf}
QFAM \bar{E}_1	303(12)	-664(38)
QFAM \tilde{E}_3	283(23)	-540(72)
PGCM \bar{E}_1	284(3)	-632(9)
PGCM \bar{E}_1	293(1)	-649(3)
PGCM \tilde{E}_3	271(4)	-523(12)

All quantities are expressed in MeV

be complementary to the computation of the equation of state in order to extract the nuclear matter compressibility.

9 Conclusions

The present paper focused on the ab initio computation of the monopole strength's moments m_k . As a first step, the formal capacity to compute low-order moments in PGCM calculations via the ground-state expectation value of moment operators was achieved. This development was then exploited to validate the use, within a few percent uncertainty, of the approach based on the explicit sum over excited states for the first moment m_1 in ^{16}O , ^{24}Mg , ^{28}Si and ^{46}Ti .

With this at hand, the angular momentum projection was shown to have little impact on the centroid but to affect significantly the dispersion of the monopole strength distribution. Next, the centroid energy obtained in GCM calculations

¹⁸ $A^{-1/3}$ law of Eq. (35). The exercise needs to be repeated in the future with a significantly larger number of points.

¹⁸ Values of K_{surf} are also in qualitative agreement with systematic studies [45].

was demonstrated to be typically 4–6% below QRPA results, which amounts to less than 1 MeV difference in the nuclei under study. The QRPA and GCM dispersions were seen to be also very consistent.

The next part of the study focused on the EWSR and first demonstrated that its textbook expression must be corrected for the fact that nuclear excitations of interest are *intrinsic* excitations in the center-of-mass frame. Having derived the appropriate *intrinsic* analytical EWSR, deviations were shown to be of 3% as a result of the (unwanted) local-gauge symmetry breaking of the employed χ EFT-based Hamiltonian [20].

Eventually, the finite-nucleus compressibility K_A was computed in ^{24}Mg , ^{28}Si and ^{46}Ti in order to extract the infinite matter nuclear incompressibility $K_\infty = 290(15)$ MeV that happens to be consistent, within uncertainties, with empirical expectations.

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Appendix A Schwartz’ inequalities

With $\rho(E)$ a positive definite function and, thus, $\rho(E)dE$ a positive measure, Schwartz’s inequality reads as

$$\int f^2(E)\rho(E)dE \int g^2(E)\rho(E)dE \geq \left(\int f(E)g(E)\rho(E)dE \right)^2, \tag{A.1}$$

with f and g two arbitrary functions. The strength function $S(E)$ is defined for positive values of E [see Eq. (2)], so that

$$\rho(E) \equiv E^k S(E) \tag{A.2}$$

is positive definite. For $f(E) = E$ and $g(E) = 1$ Eq. (A.1) reads

$$m_{k+2}m_k \geq m_{k+1}^2, \tag{A.3}$$

which provides the sequence of inequalities in Eq. (6).

Appendix B Commutator approach

The introduction of moment operators first relies on expressing the moments in terms of the commutators C_l . This step relies on rewriting Eq. (10) as

$$\begin{aligned} m_k &= \langle \Psi_0^{\sigma_0} | \mathbf{F} \mathbf{H}^k \mathbf{F} | \Psi_0^{\sigma_0} \rangle \\ &= \langle \Psi_0^{\sigma_0} | \mathbf{F} \mathbf{H}^i \mathbf{H}^j \mathbf{F} | \Psi_0^{\sigma_0} \rangle \\ &= \langle \Psi_0^{\sigma_0} | [\mathbf{F}, \mathbf{H}^i][\mathbf{H}^j, \mathbf{F}] | \Psi_0^{\sigma_0} \rangle, \end{aligned} \tag{B.4}$$

with $i + j = k$ and where the property $\mathbf{H} |\Psi_0^{\sigma_0}\rangle = 0$ has been used. Since $[\mathbf{H}^n, \mathbf{F}] = [H^n, F]$ for $n \in \mathbb{N}$, the bold notation can in fact be omitted. The needed commutators can be rewritten [46] as

$$[H^l, F] = \sum_{n=0}^{l-1} \binom{l}{n} C_{l-n} H^n, \tag{B.5a}$$

$$[F, H^l] = \sum_{n=0}^{l-1} \binom{l}{n} H^n \tilde{C}_{l-n}, \tag{B.5b}$$

with C_l introduced in Eq. (12) and \tilde{C}_l defined through

$$\tilde{C}_l \equiv \{F, H^l\} \equiv [\dots [[F, H], H] \dots, H], \tag{B.6}$$

l times

the two being equal up to a sign, i.e. $\tilde{C}_l = (-1)^l C_l$. The only non-vanishing contributions to Eq. (B.4) are obtained for $n = 0$ in Eq. (B.5) by virtue of $\mathbf{H} |\Psi_0^{\sigma_0}\rangle = 0$ (even though the bold notation could be omitted in the meantime). This finishes to prove Eq. (11).

Appendix C Second-quantized operators

Given an arbitrary orthonormal basis of the one-body Hilbert space \mathcal{H}_1 represented by the particle annihilation and creation operators $\{c_p, c_p^\dagger\}$, a generic (particle-number conserving) operator O containing up to three-body operators reads as

$$\begin{aligned} O &\equiv O^{[0]} + O^{[2]} + O^{[4]} + O^{[6]} \\ &\equiv O^{00} + O^{11} + O^{22} + O^{33} \\ &\equiv O^{00} + \frac{1}{(1!)^2} \sum_{pq} o_{pq}^{11} c_p^\dagger c_q \\ &\quad + \frac{1}{(2!)^2} \sum_{pqrs} o_{pqrs}^{22} c_p^\dagger c_q^\dagger c_s c_r \\ &\quad + \frac{1}{(3!)^2} \sum_{pqrst} o_{pqrst}^{33} c_p^\dagger c_q^\dagger c_r^\dagger c_u c_t c_s, \end{aligned} \tag{C.7}$$

where $O^{[0]} = O^{00}$ is a number. Given that O is presently taken to be particle-number conserving, the k -body class $O^{[2k]}$ contains a single operator O^{kk} characterized by the equal number k of particle-creation and annihilation operators. Such an operator is obviously in normal order with respect to the particle vacuum.

Matrix elements entering Eq. (C.7) are fully antisymmetric, i.e.

$$o_{p_1 \dots p_k p_{k+1} \dots p_{2k}}^{kk} = (-1)^{\sigma(P)} o_{P(p_1 \dots p_k | p_{k+1} \dots p_{2k})}^{kk} \tag{C.8}$$

where $\sigma(P)$ refers to the signature of the permutation P . The notation $P(\dots | \dots)$ denotes a separation into the k particle-creation operators and the k particle-annihilation operators such that permutations are only considered between members of the same group.

Appendix D Operator $M_1(1, 0)$

The algebraic expressions of the matrix elements defining the operator $M_1(1, 0)$ allowing to m_1 via the GSEV approach are presently derived. All notations are consistent with Appendix C for operators expressed in normal order with respect to the particle vacuum.

There are two equivalent ways to obtain the odd-moment operators, namely given by Eqs. (14b) and (17), respectively. They are explored separately below.

Appendix D.1 Similarity-transformed H

Using Eq. (17) for $k = 1$, the operator is given by

$$M_1(1, 0) = -\frac{1}{2} \frac{\partial^2}{\partial \eta^2} H_1(\eta) \Big|_{\eta=0} \tag{D.9}$$

with

$$H_1(\eta) = e^{-\eta F} H e^{\eta F}, \tag{D.10}$$

such that

$$\begin{aligned} H_1(\eta) &= H_1^{[0]}(\eta) + \sum_{ab} h_{ab}^{11} e^{-\eta F} c_a^\dagger c_b e^{\eta F} \\ &\quad + \frac{1}{(2!)^2} \sum_{abcd} h_{abcd}^{22} e^{-\eta F} c_a^\dagger c_b^\dagger c_d c_c e^{\eta F} \\ &\quad + \frac{1}{(3!)^2} \sum_{abcdef} h_{abcdef}^{33} e^{-\eta F} c_a^\dagger c_b^\dagger c_c^\dagger c_f c_e c_d e^{\eta F}. \end{aligned} \tag{D.11}$$

As shown below, the similarity transformation, F being a one-body operator, does not change the rank of the operator, such that $M_1(1, 0)$ has the same rank as H . Introducing the identity operator in-between each pair of creation and/or annihilation operators under the form

$$1 = e^{\eta F} e^{-\eta F}, \tag{D.12}$$

the similarity transformation is separately performed on each creation (annihilation) operator. The elementary commutator

$$\begin{aligned} [F, c_a^\dagger] &= \sum_{kl} f_{kl}^{11} c_k^\dagger c_l c_a^\dagger - c_a^\dagger F \\ &= \sum_{kl} f_{kl}^{11} c_k^\dagger (\delta_{la} - c_a^\dagger c_l) - c_a^\dagger F \\ &= \sum_k f_{ka}^{11} c_k^\dagger, \end{aligned} \tag{D.13}$$

together with Baker-Campbell-Hausdorff's formula allows one to obtain

$$\begin{aligned} e^{-\eta F} c_a^\dagger e^{\eta F} &= c_a^\dagger + \sum_{n=1}^{\infty} \frac{(-\eta)^n}{n!} \underbrace{[F, [F, \dots, [F, [F, c_a^\dagger]] \dots]]}_{n \text{ times}} \\ &= c_a^\dagger - \eta \sum_k f_{ka}^{11} c_k^\dagger + \eta^2 \frac{1}{2!} \sum_{kl} f_{lk}^{11} f_{ka}^{11} c_l^\dagger + \dots \\ &= c_a^\dagger - \eta \sum_k f_{ka}^{11} c_k^\dagger + \eta^2 \frac{1}{2!} \sum_k (f^{11})_{ka}^2 c_k^\dagger + \dots \\ &= \sum_k (e^{-\eta f^{11}})_{ka} c_k^\dagger. \end{aligned} \tag{D.14a}$$

Similarly, one has

$$e^{-\eta F} c_a e^{\eta F} = \sum_k (e^{\eta f^{11}})_{ka} c_k. \tag{D.14b}$$

Eventually, Eq. (D.11) is written as

$$H_1(\eta) = H_1^{[0]}(\eta) + \sum_{kl} h_{kl}^{11}(\eta) c_k^\dagger c_l$$

$$\begin{aligned}
 &+ \frac{1}{(2!)^2} \sum_{klmn} h_{klmn}^{22}(\eta) c_k^\dagger c_l^\dagger c_n c_m \\
 &+ \frac{1}{(3!)^2} \sum_{klmnop} h_{klmnop}^{33}(\eta) c_k^\dagger c_l^\dagger c_m^\dagger c_p c_o c_n, \quad (D.15)
 \end{aligned}$$

with the similarity-transformed matrix elements being defined as

$$\begin{aligned}
 h_{k_1 \dots k_n l_1 \dots l_n}^{nn}(\eta) &\equiv \sum_{a_1 \dots a_n b_1 \dots b_n} h_{a_1 \dots a_n b_1 \dots b_n}^{nn} \\
 &\times (e^{-\eta f^{11}})_{k_1 a_1} \dots (e^{-\eta f^{11}})_{k_n a_n} \\
 &\times (e^{\eta f^{11}})_{l_1 b_1} \dots (e^{\eta f^{11}})_{l_n b_n} \quad (D.16)
 \end{aligned}$$

and

$$H_1^{[0]}(\eta) \equiv H^{[0]}. \quad (D.17)$$

The similarity-transformed matrix elements on the left-hand side of Eq. (D.16) naturally inherit the antisymmetry of the original matrix elements, as it can be checked directly. The second derivative with respect to η can now be explicitly performed to derive the matrix elements of $M_1(1, 0)$, i.e.

$$M_1^{[0]} = 0, \quad (D.18a)$$

$$m_{1,kl}^{11}(1, 0) \equiv -\frac{1}{2} \frac{\partial^2}{\partial \eta^2} h_{kl}^{11}(\eta) \Big|_{\eta=0}, \quad (D.18b)$$

$$m_{1,klmn}^{22}(1, 0) \equiv -\frac{1}{2} \frac{\partial^2}{\partial \eta^2} h_{klmn}^{22}(\eta) \Big|_{\eta=0}, \quad (D.18c)$$

$$m_{1,klmnop}^{33}(1, 0) \equiv -\frac{1}{2} \frac{\partial^2}{\partial \eta^2} h_{klmnop}^{33}(\eta) \Big|_{\eta=0}. \quad (D.18d)$$

As for the matrix elements of the similarity-transformed Hamiltonian in Eq. (D.16), the matrix elements of $M_1(1, 0)$ are manifestly antisymmetric. The explicit writing of Eq. (D.18) can be found in Sect. 4.3.2 of Ref. [27]. The results are strictly equivalent to those obtained via the commutator-based formulation described below.

Appendix D.2 Commutator approach

The matrix elements of M_1 can also be derived from Eq. (14) for $k = 1$ (e.g. $i = 0$ and $j = 1$). This is achieved by applying Wick’s theorem with respect to the particle vacuum $|0\rangle$. In this case the only non-vanishing contraction at play is

$$\overline{c_a^\dagger c_b^\dagger} \equiv \langle 0 | c_a c_b^\dagger | 0 \rangle = \delta_{ab}. \quad (D.19)$$

The commutator $C_1 = [H, F]$ is computed separately for the various components of H . The operator F being a one-body operator, the commutator preserves the n -body nature of the component $H^{[n]}$ such that each n -body component of C_1 is

introduced as

$$[H^{[0]}, F] = 0, \quad (D.20a)$$

$$[H^{[1]}, F] \equiv \sum_{ab} c_{1,ab}^{11} c_a^\dagger c_b, \quad (D.20b)$$

$$[H^{[2]}, F] \equiv \frac{1}{(2!)^2} \sum_{abcd} c_{1,abcd}^{22} c_a^\dagger c_b^\dagger c_d c_c, \quad (D.20c)$$

$$[H^{[3]}, F] \equiv \frac{1}{(3!)^2} \sum_{abcdef} c_{1,abcdef}^{33} c_a^\dagger c_b^\dagger c_c^\dagger c_f c_e c_d. \quad (D.20d)$$

The derivation of the matrix elements from Eq. (D.20) relies on the tool developed in Ref. [47]. This tool allows one to compute the antisymmetrized matrix elements of the normal-ordered operator obtained via the commutator of any two normal-ordered operators. While the development was originally done with respect to a Bogoliubov vacuum $|\Phi_{\text{HFB}}\rangle$ and expressing normal-ordered operators in the associated quasi-particle basis, it can be readily exploited here by simply substituting quasi-particle operators β^\dagger (β) with particle operators c^\dagger (c) and by using the particle vacuum $|0\rangle$ instead of the Bogoliubov one. Naturally the particle formalism only needs to retain particle-number-conserving components.

Eventually, the matrix elements of the elementary commutator from Eq. (D.20) can be expressed as

$$c_{1,ab}^{11} = \sum_k h_{ak}^{11} f_{kb}^{11} - \sum_k f_{ak}^{11} h_{kb}^{11}, \quad (D.21a)$$

$$\begin{aligned}
 c_{1,abcd}^{22} &= P(c/d) \sum_k h_{abck}^{22} f_{kd}^{11} \\
 &- P(a/b) \sum_k f_{ak}^{11} h_{kbcd}^{22}, \quad (D.21b)
 \end{aligned}$$

$$\begin{aligned}
 c_{1,abcdef}^{33} &= P(de/f) \sum_k h_{abcdek}^{33} f_{kf}^{11} \\
 &- P(a/bc) \sum_k f_{ak}^{11} h_{kbcd}^{33}, \quad (D.21c)
 \end{aligned}$$

with

$$P(a/b) \equiv 1 - P_{ab}, \quad (D.22a)$$

$$P(a/bc) \equiv 1 - P_{ab} - P_{ac}, \quad (D.22b)$$

$$P(ab/c) \equiv 1 - P_{ac} - P_{bc}, \quad (D.22c)$$

and where P_{ab} denotes the transposition operator exchanging indices a and b . The extended writing of Eq. (D.21), i.e. with permutations explicitly carried out, can be found in Sect. 4.3.2 of Ref. [27].

The above result is exploited to readily compute the nested commutator needed to obtain the m_1 operator

$$M_1(1, 0) \equiv -\frac{1}{2} [[H, F], F] = -\frac{1}{2} [C_1, F], \quad (D.23)$$

by substituting H with C_1 in Eq. (D.21). Eventually, the matrix elements of $M_1(1, 0)$ are obtained as

$$M_1^{[0]} = 0, \tag{D.24a}$$

$$m_{1,ab}^{11} \equiv -\frac{1}{2} \sum_c \{c_{1,ac}^{11} f_{cb}^{11} - f_{ac}^{11} c_{1,cb}^{11}\}, \tag{D.24b}$$

$$m_{1,abcd}^{22} = \frac{1}{2} P(a/b) \sum_k f_{ak}^{11} c_{kbcd}^{22} - \frac{1}{2} P(c/d) \sum_k c_{abck}^{22} f_{kd}^{11}, \tag{D.24c}$$

$$m_{1,abcdef}^{33} = \frac{1}{2} P(a/bc) \sum_k f_{ak}^{11} c_{kbcdef}^{33} - \frac{1}{2} P(de/f) \sum_k c_{abcdek}^{33} f_{kf}^{11}. \tag{D.24d}$$

The extended writing of Eq. (D.24) is provided in Sect. 4.3.2 of Ref. [27] and is found to be identical to the similarity-evolved derivation from Appendix D.1.

Appendix E Strength function extraction

The actual relation of the strength function to scattering observables is hereby briefly discussed. At first order in perturbation theory, the transition rate $w_{0 \rightarrow \nu}$ from the ground state $|\Psi_0^{\sigma_0}\rangle$ to an excited state $|\Psi_\nu^\sigma\rangle$ mediated by the time-independent operator F is provided by Fermi’s golden rule

$$w_{0 \rightarrow \nu} = 2\pi |\langle \Psi_\nu^\sigma | F | \Psi_0^{\sigma_0} \rangle|^2 \delta(E_\nu^\sigma - E_0^{\sigma_0} - E). \tag{E.25}$$

The corresponding cross section $\sigma_{0 \rightarrow \nu}$ is obtained normalising the transition rate by the flux of incident particles and the number of scattering centers

$$\frac{d\sigma_{0 \rightarrow \nu}}{dE} = w_{0 \rightarrow \nu} \times \frac{1}{\text{flux}} \times \frac{1}{\# \text{ of sc. centers}}. \tag{E.26}$$

The total cross section is computed by summing over all possible final states ν so that it can be expressed as

$$\sigma = 2\pi \int_{-\infty}^{+\infty} S(E) dE = 2\pi m_0. \tag{E.27}$$

In practice, double-differential cross sections are experimentally measured to perform a *multipole-decomposition analysis* (MDA), allowing the extraction of the multipole strength distributions [3]. In the MDA process, the experimental cross-sections at each angle are binned into small (typically, ≤ 1 MeV) excitation energy intervals. The laboratory angular distributions for each excitation-energy bin are then converted into the centre-of-mass frame using standard Jacobian and relativistic kinematics. For each excitation energy bin, the experimental angular distributions are fitted

by means of the least-square method with the linear combination of the calculated double-differential cross sections associated to different multipoles:

$$\left. \frac{d^2\sigma^{\text{exp}}}{d\Omega dE} \right|_{E_x} = \sum_{L=0}^{\infty} a_L(E_x) \left. \frac{d^2\sigma_L^{\text{DWBA}}}{d\Omega dE} \right|_{E_x}, \tag{E.28}$$

where $a_L(E_x)$ is the m_1 sum rule fraction for the L -th component. The cross sections used for the fit procedure correspond to the 100% of m_1 for the L -th multipole at excitation energy E_x calculated using the distorted-wave Born approximation (DWBA). In such calculations an optical potential is used as the scattering potential. The fractions of m_1 , $a_L(E_x)$, for various multipole components are determined by minimising χ^2 error. Eventually, the strength distributions for different multipolarities are obtained by multiplying the extracted $a_L(E_x)$ ’s by the strength corresponding to 100% m_1 at the given energy E_x

$$S_L(E_x) = \frac{m_{L,1}}{E_x} a_L(E_x). \tag{E.29}$$

Traditionally, the energy-weighted sum rules $m_{L,1}$ employed in the above procedure for different L ’s are always the textbook $EWSR_{\text{lab}}$ rather than the appropriate intrinsic one discussed in Sect. 7.

Appendix F Intrinsic EWSR

The monopole EWSR from Eq. (25) is evaluated under the assumption that only the kinetic energy T_{lab} from Eq. (24) contributes to Eq. (14b), such that

$$EWSR_{\text{lab}}(r^2) = -\frac{1}{2} \langle [T_{\text{lab}}, r^2], r^2 \rangle_{\text{gs}} = \frac{2\hbar^2 A}{m} \langle r_{\text{lab}}^2 \rangle_{\text{gs}}. \tag{F.30}$$

The correction to $EWSR_{\text{lab}}(r^2)$ due to the subtraction of the center-of-mass kinetic energy T_{cm} [Eq. (26)] from the Hamiltonian is given by

$$\delta m_1^{\text{cm}}(r^2) \equiv \frac{1}{2} \langle [T_{\text{cm}}, r^2], r^2 \rangle_{\text{gs}} = \frac{1}{4mA} \sum_{ijkl=1}^A \langle [[\vec{p}_i \cdot \vec{p}_j, r_k^2], r_l^2] \rangle_{\text{gs}}. \tag{F.31}$$

The commutator relation

$$[\vec{p}_i, f(\vec{r}_k)] = -i\hbar \vec{\nabla}_i f(\vec{r}_k) \tag{F.32}$$

is employed to evaluate the elementary commutator

$$[\vec{p}_i, r_k^2] = -2i\hbar \vec{r}_i \delta_{ik}, \tag{F.33}$$

allowing to process Eq. (F.31) according to

$$\begin{aligned}\delta m_1^{\text{cm}}(r^2) &= \frac{1}{4mA} \sum_{ijkl} \langle [\vec{p}_i \cdot [\vec{p}_j, r_k^2], r_l^2] + [[\vec{p}_i, r_k^2] \cdot \vec{p}_j, r_l^2] \rangle_{\text{gs}} \\ &= \frac{1}{4mA} \sum_{ijkl} \langle [\vec{p}_i, r_l^2] \cdot [\vec{p}_j, r_k^2] + [\vec{p}_i, r_k^2] \cdot [\vec{p}_j, r_l^2] \rangle_{\text{gs}} \\ &= -\frac{2\hbar^2}{mA} \sum_{ij} \langle \vec{r}_i \cdot \vec{r}_j \rangle_{\text{gs}} \\ &= -\frac{2\hbar^2 A}{m} \langle R_{\text{cm}}^2 \rangle_{\text{gs}},\end{aligned}\quad (\text{F.34})$$

where the second equality follows from

$$[[\vec{p}_i, r_k^2], r_l^2] = -2i\hbar[\vec{r}_i, r_l^2]\delta_{ik} = 0, \quad (\text{F.35})$$

and where

$$\vec{R}_{\text{cm}} \equiv \frac{1}{A} \sum_{i=1}^A \vec{r}_i, \quad (\text{F.36})$$

denotes the center of mass coordinate.

Eventually, the EWSR associated with the intrinsic Hamiltonian can be written under the alternating forms

$$\begin{aligned}\text{EWSR}_{\text{int}}(r^2) &= \text{EWSR}_{\text{lab}}(r^2) + \delta m_1^{\text{cm}}(r^2) \\ &= \frac{2\hbar^2 A}{m} \langle r_{\text{lab}}^2 - R_{\text{cm}}^2 \rangle_{\text{gs}} \\ &= \frac{2\hbar^2 A}{m} \langle r_{\text{int}}^2 \rangle_{\text{gs}} \\ &= \text{EWSR}_{\text{lab}}(r^2) \left(1 - \frac{1}{A}\right) - \frac{2\hbar^2}{mA} \sum_{i \neq j} \langle \vec{r}_i \cdot \vec{r}_j \rangle_{\text{gs}}.\end{aligned}\quad (\text{F.37})$$

The above result demonstrates that the subtraction of T_{cm} in H leads to replacing the laboratory-frame mean-square radius $\langle r_{\text{lab}}^2 \rangle$ [Eq. (19)] by the intrinsic one $\langle r_{\text{int}}^2 \rangle$ [Eq. (20)]. The last line splits $\delta m_1^{\text{cm}}(r^2)$ into its one- and two-body contributions to demonstrate that the one-body part of T_{cm} leads to a simple A -dependent renormalization of $\text{EWSR}_{\text{lab}}(r^2)$ [48].

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