Nuclear excitations as coupled one and two random–phase–approximation modes

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We present an extension of the random–phase approximation (RPA) where the RPA phonons are used as building blocks to construct the excited states. In our model, that we call double RPA (DRPA), we include up to two RPA phonons. This is an approximate and simplified way, with respect to the full second random–phase approximation (SRPA), to extend the RPA by including two particle–two hole configurations. Some limitations of the standard SRPA model, related to the violation of the stability condition, are not encountered in the DRPA. We also verify in this work that the energy–weighted sum rules are satisfied. The DRPA is applied to low–energy modes and giant resonances in the nucleus $^{16}$O. We show that the model (i) produces a global downwards shift of the energies with respect to the RPA spectra; (ii) provides a shift that is however strongly reduced compared to that generated by the standard SRPA. This model represents an alternative way of correcting for the SRPA anomalous energy shift, compared to a recently developed extension of the SRPA, where a subtraction procedure is applied. The DRPA provides results in good agreement with the experimental energies, with the exception of those low–lying states that have a dominant two particle–two hole nature. For describing such states, higher–order calculations are needed.

I. INTRODUCTION

A common feature of many-body systems is the presence of collective modes in their excitation spectra. Particularly interesting are those which can be interpreted in terms of vibrations. In nuclei, they have been known for a long time [1] and are observed both in the low–lying and in the higher–energy region (giant resonances (GR)) [1,2]. Another example of collective vibrations is the dipole plasmon resonance in metallic clusters [3,4], a very collective mode which is the analogue of the nuclear dipole GR and is due to the vibration of the center of mass of the delocalized valence electrons against that of the positive ions.

In all these cases, several characteristic properties are rather well reproduced within the random–phase approximation (RPA), which is for this reason considered a good microscopic theory for the study of collective states in many–body systems [1,2]. Among its merits, we mention that the RPA preserves the energy–weighted sum rules (EWSR), in the sense of the Thouless theorem [2]. This feature is very important because it guarantees that spurious states associated with broken symmetries are exactly separated in RPA from the physical states of the system.

On the other hand, the RPA model has some limits. Among them we recall that, by construction, it predicts a perfectly harmonic spectrum and cannot reproduce the spreading width of the excited modes. As underlined in several studies, the coupling of one particle–one hole (1p-1h) configurations with more complex states is very important to overcome these limitations and to describe the spreading widths and the anharmonicities in the excitation spectra [see, for example Refs. [3] and references therein]. A natural extension of the RPA is the so–called second RPA (SRPA), which amounts to enlarge the space of basic elementary excitations by including two particle–two hole (2p-2h) configurations and by coupling them with the 1p-1h ones and among themselves. It was shown [1,4,13] that the EWSR are exactly the same in SRPA and in RPA.

Recent applications of the SRPA model were performed without any cut and approximations in the matrices. Such applications provided a very strong modification (a large downwards energy shift) of the RPA excitation spectra, also for those collective states whose description within the RPA is rather good. This was obtained for collective nuclear excitations with Skyrme and Gogny forces [14,15], as well as with an interaction derived from the Argonne V18 nucleon–nucleon potential with the Unitary Correlation Operator Method [16]. In the case of metallic clusters, within the jellium model [4], the modifications found when passing from RPA to SRPA are very large [17] and completely spoil the quality of the RPA results concerning the dipole strength distribution. In this application, the limiting case of highly ionized clusters is very striking. The RPA is able to reproduce the exact theoretical prediction on the position of the dipole plasmon excitation (at the Mie energy) and the fact that the dipole strength is totally concentrated at this energy [18]. These very nice features are completely lost in SRPA [17]. In the nuclear case, one might associate such unpleasant result to some inadequacy of the used nuclear interaction. This could be especially true for the Skyrme– and Gogny–based SRPA calculations because some matrix elements appearing in SRPA are not
Also affected by the cutoff dependence and the double–
condition [7, 13] encountered in SRPA. They are however
mode, do not suffer from the violation of the stability
hole configuration is coupled to a correlated collective
vibration coupling [30], where one uncorrelated particle–
atomic nuclei is hardly feasible.

However, the problem of the large downwards energy
shift exists also in the case of metallic clusters [17] where
there is no cutoff dependence and the Coulomb interaction
does not contain any parameter [12]. The origin of the
problem must then be different.

In deriving the equations of motion in SRPA [11, 20],
use is made, as in RPA, of the quasi–boson approxima-
tion (QBA) and it has been argued [21, 22] that its use
in SRPA is even more questionable than in RPA. Re-
cently, this was more precisely understood as related to
the replacement of the correlated ground state with the
Hartree–Fock (HF) one, that is generated by the use of
the QBA. This replacement produces a violation of the
stability condition at the SRPA level [13]. A careful anal-
ysis of the merits and limits of the SRPA was presented
in Ref. [13]. In particular, the violation of the stability
condition in SRPA [21] is illustrated and a generalization
of the Thouless theorem [7] is proven in the case where
a correlated ground state is used. Very recently, a sub-
traction procedure able to remove the double counting
in beyond–mean–field theories based on effective interac-
tions [24, 25] was applied to the SRPA case [26]. Refer-
ce [25] showed explicitly that this subtraction method
is also able to ensure the stability condition in extended
RPA models such as the SRPA. In Ref. [26], very encour-
aging results were found for both low–lying and higher–
energy excitations. In particular, the strong SRPA down-
ward shift is corrected for those states having a dominant
1p-1h nature such as the giant resonances, and the found
results are in much better agreement with the experi-
mental ones, compared to the standard SRPA results.

In Ref. [28], an extension of the SRPA including corre-
lations in the ground state was tested by applying it to
a solvable model and by comparing the results with the
exact ones. It was also applied to metallic clusters [29],
finding much better results than in SRPA. However, such
an extended SRPA is very demanding from a computa-
tional point of view and its application to the study of
atomic nuclei is hardly feasible.

On the other hand, approaches like the particle–
vibration coupling [30], where one uncorrelated particle–
hole configuration is coupled to a correlated collective
mode, do not suffer from the violation of the stability
condition [7, 13] encountered in SRPA. They are however
also affected by the cutoff dependence and the double–
counting problems if phenomenological zero–range inter-
actions are employed.

Extensions of RPA including multiphonon excitations
have been applied in several contexts. They make use
of mapping procedures to replace the fermion particle–
hole operators by their boson images [32]. The limits
of such procedure were tested in Ref. [33] within a solv-
able model, by comparing the exact results with those
obtained by truncating the mapping at different levels.
It was shown that in order to correct for the violations
of the Pauli principle, higher–order terms are required in
the boson expansion. This makes not feasible the applica-
tion of the approach to realistic systems. The procedure
presented in Refs. [34, 35] allows one to enlarge iter-
atively the space in which the Hamiltonian is diagonalized
by including multiphonon states built by Tamm–Dancoff
phonon operators remaining in the fermion space.

The approach that we are going to present consists in
considering the RPA collective fermion operators as
building blocks to construct the space of states by acting
repeatedly on the RPA ground state (that is, the vacuum
of the RPA operators). The Hamiltonian is then diagno-
salized in such space. Since the fermionic structure of
the collective elementary excitations is fully accounted for,
one may expect to get satisfactory enough results with
a basis of states containing up to two such elementary
excitations. Thus, one considers, as in SRPA, a space
containing 1p-1h and 2p-2h configurations, the latter ap-
pearing only through the two RPA phonon states. Since
the 2p-2h sector of the basis space is built with those
collective excitations which already contain some effects
of the residual interaction, one might envisage to reduce
the dimensions of the space by including only some con-
fugations selected, for example, on the basis of their
collectivity. When studying the excitation spectrum one
may introduce an energy cutoff and consider only those
basis vectors whose RPA energy is lower. This would al-
low one to reduce the heavy computational effort often
required within the SRPA. This approach is very much
related to that of Refs. [34, 35]. It is however to be noted
that in such references Tamm Dancoff phonons were used
to construct the basis states by acting on the uncorre-
lated HF ground state. We consider here RPA operators
acting on the correlated RPA ground state. Therefore,
even considering the mixing of configurations containing
up to two phonons, we take into account higher–order
excitations. We call our model double RPA and use in
what follows the acronym DRPA to denote it.

The approach is described in the next section. The
energy–weighted sum rules are shown to be satisfied in
Sec. III, and applications to the nucleus $^{16}$O are pre-
sented in Sec. IV. Finally, in Sec. V we draw some
conclusions.
II. THE DRPA MODEL

In RPA one defines the operators

$$Q^\dagger_\nu = \sum_{ph} (X^\nu_{ph}b^\dagger_{ph} - Y^\nu_{ph}b_{ph})$$

and $Q_\nu$, such that

$$Q_\nu |\Psi_0(RPA)\rangle = 0$$

and

$$|\Psi_\nu(RPA)\rangle = Q^\dagger_\nu|\Psi_0(RPA)\rangle,$$

where $|\Psi_0(RPA)\rangle$ and $|\Psi_\nu(RPA)\rangle$ are the RPA ground and excited states of the system, respectively. The operators $b^\dagger_{ph}$ and $b_{ph}$ create and annihilate, respectively, a particle-hole pair ($b^\dagger_{ph} = a^\dagger_{ph}a_{ph}$). Here and in the following, coupling to total quantum numbers is understood. The $X$ and $Y$ amplitudes appearing in Eq. (1) are solutions of the equations of motion

$$\begin{pmatrix} A^{11} & B^{11} \\ A^{*11} & B^{*11} \end{pmatrix} \begin{pmatrix} X^\nu \\ Y^\nu \end{pmatrix} = E_\nu \begin{pmatrix} G^{11} & 0 \\ 0 & -G^{*11} \end{pmatrix} \begin{pmatrix} X^\nu \\ Y^\nu \end{pmatrix},$$

where

$$A^{11}_{ph,p'h'} = \langle \Psi_0(RPA)|[b_{ph}, [H, b^\dagger_{p'h'}]]|\Psi_0(RPA)\rangle \approx HF|[b_{ph}, [H, b^\dagger_{p'h'}]]|HF\rangle,$$

and

$$B^{11}_{ph,p'h'} = -\langle \Psi_0(RPA)|[b_{ph}, [H, b^\dagger_{p'h'}]]|\Psi_0(RPA)\rangle \approx -HF|[b_{ph}, [H, b^\dagger_{p'h'}]]|HF\rangle.$$

As explicitly shown, in order to calculate the matrix elements appearing in Eq. (4), use is made of the QBA, replacing the RPA ground state ($|\Psi_0(RPA)\rangle$) with the HF one ($|HF\rangle$).

In SRPA, the excitation phonon operators are generalized to

$$Q^\dagger_\rho = \sum_{ph} (X^\rho_{ph}b^\dagger_{ph} - Y^\rho_{ph}b_{ph}) + \sum_{p_1h_1} (X^\rho_{p_1h_1p_2h_2}b^\dagger_{p_1h_1p_2h_2}b^\dagger_{p_2h_2}b_{p_1h_1}),$$

which contain 2p-2h components in addition to the 1p-1h ones. This extension allows one to go beyond the harmonic approximation of the RPA and to describe the spreading width of the excited states through the coupling of the 1p-1h configurations with the 2p-2h ones. As in RPA, one assumes

$$|\Psi_\rho(SRPA)\rangle = Q^\dagger_\rho|\Psi_0(SRPA)\rangle$$

and

$$Q_\rho|\Psi_0(SRPA)\rangle = 0,$$

where $|\Psi_\rho(SRPA)\rangle$ and $|\Psi_0(SRPA)\rangle$ are the SRPA excited and ground states, respectively. The amplitudes appearing in Eq. (3) and the energies $E_\rho$ of the excited states are solutions of

$$\begin{pmatrix} A & B \\ B^* & A^* \end{pmatrix} \begin{pmatrix} X^\rho \\ Y^\rho \end{pmatrix} = E_\rho \begin{pmatrix} G & 0 \\ 0 & -G^* \end{pmatrix} \begin{pmatrix} X^\rho \\ Y^\rho \end{pmatrix},$$

where

$$X_\rho = \begin{pmatrix} X^\rho_{ph} \\ X^\rho_{p_1h_1p_2h_2} \end{pmatrix}, Y_\rho = \begin{pmatrix} Y^\rho_{ph} \\ Y^\rho_{p_1h_1p_2h_2} \end{pmatrix}$$

and

$$A = \begin{pmatrix} A^{11}_{ph,p'h'} & A^{12}_{p_1h_1p_2h_2,p'h'} \\ A^{21}_{p_1h_1p_2h_2,p'h'} & A^{22}_{p_1h_1p_2h_2,p'h'} \end{pmatrix}.$$
is made, the vacuum condition, Eq. (16), follows immediately from Eq. (2), satisfied by the elementary operators $Q_{\nu}$ at the RPA level. By following the same procedure used to derive the RPA equations of motion we get

$$H|\Psi_\alpha> = E_\alpha|\Psi_\alpha> \Rightarrow$$

$$[H, \Lambda^\dagger_\alpha]|\Psi_0> = (E_\alpha - E_0)\Lambda^\dagger_\alpha|\Psi_0> .$$

From Eq. (15) and from the vacuum condition, Eq. (2), it follows

$$\sum_\nu c_\nu^\alpha <Q_{\nu}|[H,Q^\dagger_{\nu}]|Q_{\nu}> + \sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} <\Psi_0|[Q_{\nu},[H,Q^\dagger_{\nu_1}]]|\Psi_0> = (E_\alpha - E_0)\sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} <\Psi_0|[Q_{\nu},Q^\dagger_{\nu_1}]|\Psi_0> .$$

and

$$\sum_\nu c_\nu^\alpha <Q_{\nu}|[Q_{\nu}Q^\dagger_{\nu_1}]|\Psi_0> + \sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} <\Psi_0|[Q_{\nu},Q^\dagger_{\nu_1}]Q_{\nu_2}]|\Psi_0> = (E_\alpha - E_0)\sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} <\Psi_0|[Q_{\nu},Q^\dagger_{\nu_1},Q^\dagger_{\nu_2}]|\Psi_0> .$$

By using Eq. (15) and by employing the QBA at the RPA level, the matrix elements appearing in Eqs. (15) and (19) can be expressed in terms of the $X$ and $Y$ RPA amplitudes and of matrix elements similar to the SRPA matrix elements $A^{ij}$ and $G^{ij}$, with the HF ground state replacing $|\Psi_0(SRPA)>$. We can rewrite Eqs. (18) and (19) as

$$\sum_\nu c_\nu^\alpha A_{\nu\nu_1\nu_2} + \sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} A_{\nu\nu_1\nu_2} = (E_\alpha - E_0)c_\nu^\alpha$$

and

$$\sum_\nu d_{\nu\nu_1\nu_2} A^\dagger_{\nu\nu_1\nu_2} + \sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} A_{\nu\nu_1\nu_2} = (E_\alpha - E_0)\sum_{\nu\nu_1}\nu_2 d_{\nu\nu_1\nu_2} G_{\nu\nu_1\nu_2}.\tag{20}$$

with

$$A_{\nu\nu_1\nu_2} \equiv <HF|[Q_{\nu},[H,Q^\dagger_{\nu_1}]]|HF> = E_{\nu\nu_1\nu_2} \delta_{\nu,\nu_1} \delta_{\nu_2,\nu_2},\tag{22}$$

$$A^\dagger_{\nu\nu_1\nu_2} \equiv <HF|[Q_{\nu},[H,Q^\dagger_{\nu_1}]]|HF> = \sum_{p_1 h_1 p_2 h_2} p_1 h_1 p_2 h_2 \times (X_{p_1 h_1}^{\nu\nu_1} X_{p_2 h_2}^{\nu_2} - Y_{p_1 h_1}^{\nu\nu_1} Y_{p_2 h_2}^{\nu_2}) \times A^\dagger_{p_1 h_1\nu_2 p_2 h_2},$$

$$\times A^\dagger_{p_1 h_1\nu_2 p_2 h_2},\tag{23}$$

$$A_{\nu\nu_1\nu_2} \equiv <HF|[Q_{\nu},[H,Q^\dagger_{\nu_1}]Q^\dagger_{\nu_2}]|HF> = \sum_{p_1 h_1 p_2 h_2} p_1 h_1 p_2 h_2 \times (X_{p_1 h_1}^{\nu\nu_1} X_{p_2 h_2}^{\nu_2} - Y_{p_1 h_1}^{\nu\nu_1} Y_{p_2 h_2}^{\nu_2}) \times A_{\nu\nu_1\nu_2} \delta_{\nu,\nu_1} \delta_{\nu_2,\nu_2} + O(B^{11}),$$

$$G_{\nu\nu_1\nu_2,\nu_1\nu_2} = <HF|[Q_{\nu},Q^\dagger_{\nu_1}]Q^\dagger_{\nu_2}]|HF>$$

$$= \sum_{p_1 h_1 p_2 h_2} p_1 h_1 p_2 h_2 \times (X_{p_1 h_1}^{\nu\nu_1} X_{p_2 h_2}^{\nu_2} - Y_{p_1 h_1}^{\nu\nu_1} Y_{p_2 h_2}^{\nu_2}) \times G_{\nu\nu_1\nu_2,\nu_1\nu_2}.\tag{26}$$

The matrices in the right-hand sides of Eqs. (23)- (26) are denoted and have the same expressions as the SRPA $A^{ij}$ and $G^{ij}$ matrices, where the HF ground state is used instead of $|\Psi_0(SRPA)>$. In Eq. (25), $O(B^{11})$ denotes several terms containing the RPA matrix $B^{11}$ of Eq. (6). In the calculations that we present in this work we neglect for simplicity the terms $O(B^{11})$ appearing in Eq. (26). We have checked within an exactly solvable 3-level Lipkin model that their contribution to the DRPA energies is negligible.

### III. ENERGY WEIGHTED SUM RULES

It is very well known that, if $|\Psi_0>$ and $|\Psi_\nu>$ are the exact eigenstates of a system with Hamiltonian $H$, the following equality holds

$$\sum_\nu E_{\nu} <\Psi_\nu |T| \Psi_0>^2 = \frac{1}{2} <\Psi_0|[T,[H,T]]|\Psi_0>$$

with $T$ any transition operator. The above equality denotes the EWSR. The Thouless theorem states that in RPA the EWSR are valid if the sum in the l.h.s. is calculated with the RPA energies and transition probabilities, whereas the uncorrelated ground state $|HF>$ is used in the r.h.s. It was shown that the EWSR are valid also in SRPA. Following step by step the demonstration of Ref. 11, one easily shows their validity also in the present approach. For any self-adjoint transition operator $T$ connecting the ground state to the excited ones we can write

$$T = \sum_\alpha <\Psi_\alpha |T| \Psi_0> \Lambda^\dagger_\alpha + h.c.$$ \tag{28}

and

$$<\Psi_\alpha |T| \Psi_0> = <\Psi_\alpha |[\Lambda_\alpha, T]| \Psi_0> = \sum_\nu c_\nu^\alpha <\Psi_\alpha |[Q_{\nu}, T]| \Psi_0>$$

$$+ \sum_\nu d_{\nu\nu_1\nu_2} <\Psi_\alpha|[Q_{\nu}, Q^\dagger_{\nu_1}, T]| \Psi_0> .\tag{29}$$

In the spirit of the Thouless theorem the r.h.s. of Eq. (27) is approximated as

$$<\Psi_0|[T,[H,T]]| \Psi_0> \approx <HF|[T,[H,T]]|HF> .\tag{30}$$

Since, as can be easily verified

$$<HF|[\Lambda^\dagger_\alpha, [H, \Lambda^\dagger_\alpha]]|HF> = 0,$$ \tag{31}
one gets

\[
\frac{1}{2} < HF | [T, [H,T]] | HF > \\
= \sum_{\alpha,\alpha'} T_{0\alpha} T_{0\alpha'} ^* < HF | [\Lambda_{\alpha}, [H, \Lambda_{\alpha}]] | HF > \\
= \sum_{\alpha,\alpha'} T_{0\alpha} T_{0\alpha'} ^* \left[ \sum_{\nu,\nu'} c_{\nu}^{\alpha'} c_{\nu'}^{\alpha} A_{\nu,\nu'}^{11} + \sum_{\nu,\nu'} c_{\nu}^{\alpha'} d_{\nu,\nu'}^{0} A_{\nu,\nu'}^{12} + \sum_{\nu,\nu'} c_{\nu}^{\alpha'} c_{\nu'}^{\alpha} d_{\nu,\nu'}^{2} A_{\nu,\nu'}^{22} \right].
\] (32)

By acting with \( \sum_{\alpha} T_{0\alpha} \sum_{\nu,\nu} c_{\nu}^{\alpha} \sum_{\alpha} T_{0\alpha} \) on the l.h.s. and r.h.s. of Eq. (18), and with \( \sum_{\alpha} T_{0\alpha} \sum_{\nu,\nu} c_{\nu}^{\alpha} \sum_{\alpha} T_{0\alpha} \) on Eq. (19), and summing side by side the obtained equations we get the EWSR

\[
\frac{1}{2} < HF | [T, [H,T]] | HF > = \sum_{\alpha} E_{\alpha} | T_{0\alpha} |^2.
\] (33)

We also note that, since the transition amplitudes are evaluated, as in RPA, with reference to the HF ground state, one has

\[
T_{0\alpha} \approx < HF | [\Lambda_{\alpha}, T] | HF > = \sum_{\nu} c_{\nu}^{\alpha} \sum_{ph} \left( X_{ph}^{\nu} + Y_{ph}^{\nu} \right) |p| T |h\rangle.
\] (34)

In deriving the last equation it has been assumed that \( T \) is a one-body operator.

**IV. RESULTS FOR \(^{16}\text{O}\)**

We have applied the DRPA method to the study of the excitation spectrum in \(^{16}\text{O}\). In this section we present the results of these calculations and compare them with those obtained in Ref. [14] within the SRPA. The latter were shown to be very different from the RPA ones, also in the parts of the spectrum which are expected to be reasonably well described by RPA. As in Ref. [14], all calculations have been done by using the SGII Skyrme interaction [37]. Starting with a HF calculation, natural parity RPA elementary excitations are obtained and used as building blocks to construct a basis of one- and two-phonon states with RPA energies up to some given cutoffs, \( E1C \) and \( E2C \), respectively. In order to avoid any misunderstanding due to the use of the word 'phonon',

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**FIG. 1:** (Color online) Monopole states calculated with the RPA and the DRPA, with three different values of the cutoff \( E1C \), from 20 to 30 MeV and a value of 40 MeV for the \( E2C \) cutoff.

**FIG. 2:** (Color online) Same as in Fig. 1, but for the dipole states.

**FIG. 3:** (Color online) Same as in Fig. 1, but for the quadrupole states.
we stress again that we do not make any bosonic mapping of the RPA elementary excitations, that is, the fermionic internal structure is maintained and the Pauli principle is fully preserved. First of all, it is necessary to assess the stability of the results with respect to the values of the cutoffs $E1C$ and $E2C$. In Figs. 1–4, we show the energies obtained with a fixed value of $E2C = 40$ MeV, with values of $E1C$ running from 20 to 30 MeV, for the multipole states $0^+$, $1^-$, $2^+$ and $3^-$. We report also the RPA energies. In all cases, as expected, we see that the energies obtained by diagonalizing the Hamiltonian in the space of one and two RPA phonons are lower than the RPA results. In the cases of positive parity states this lowering is very pronounced. This happens because, for the nucleus we are dealing with, 1p-1h configurations of positive parity correspond to a two–major–shell jump and the corresponding unperturbed energies are quite large. The lower–energy states found with the DRPA in the $0^+$ and $2^+$ cases have mostly a 2p-2h nature, as was shown with genuine SRPA calculations [26] and cannot thus be found with the RPA. They can be predicted here by coupling low–lying negative parity RPA phonons. However, for the moment, we want to stress that for all multiplicities the results stabilize with increasing $E1C$ from 20 to 30 MeV. To complete this analysis on the stability of our calculations, in Figs. 5–8, we show the results obtained by fixing $E1C$ to the value of 30 MeV and varying $E2C$ from 40 to 50 MeV. Again, the results are remarkably stable.

We stress that, due to the Pauli principle, the set of two phonon states is redundant. This means that the associated generalized eigenvalue problem may present unphysical solutions if the norm matrix presents singularities. The solution of the eigenvalue problem is achieved by using a $QZ$ algorithm [31]. The employed algorithm provides the physical eigenvalue as the ratio of two quantities $\alpha$ and $\beta$, the latter approaching zero, in the case of singularities of the norm matrix. Analyzing the $\beta$ quantities, spurious states associated to the redundancy can be identified and isolated. Moreover, we notice that, in general, two different groups of eigenvalues are found. A first class, for which the corresponding $\beta$ value is quite stable with respect to the increasing of the model space (that is, the single–particle basis, and the $E1C$ and $E2C$ cutoff values), and a second one exhibiting instead a quite

FIG. 4: (Color online) Same as in Fig. 1, but for the octupole states.

FIG. 5: (Color online) Monopole states calculated with the RPA and the DRPA, with three different values of the cutoff $E2C$, from 40 to 50 MeV and a value of 30 MeV for the $E1C$ cutoff.

FIG. 6: (Color online) Same as in Fig. 5, but for the dipole states.
strong dependence on the model space. Such states have beta values that are typically much smaller than those of the states belonging to the first group. In practice, for each solution, by increasing the model space dimension (in particular with respect to the $E2C$ cutoff), we studied carefully the behavior of the $\beta$ values and only those eigenvalues characterized by stable values have been retained.

Let us now turn our attention to the multipole strength distributions. The results within the present approach, with $E1C = 30$ MeV and $E2C = 50$ MeV are presented in Figs. 9–12. The shown curves have been obtained by folding the discrete spectrum with a Lorentzian having a 1 MeV width. In the same figures the RPA and SRPA results are shown.

Due to the one–body nature of the transition operators, the states having a vanishingly small one–phonon component cannot be excited from the ground state. This is the reason why almost no strength is seen below $\sim 12$ MeV in the DRPA response for the $0^+\text{ and } 2^+$ cases. As a general trend, we see in all cases a shift of the DRPA strength distribution with respect to the RPA, but much less pronounced than in the SRPA. This is a quite satisfactory result because the positions of the main
FIG. 11: (Color online) Same as in Fig. 9, but for the isoscalar quadrupole response.

FIG. 12: (Color online) Same as in Fig. 9, but for the isoscalar octupole response.

peaks of the isovector and isoscalar GR remain not much lower than in the RPA, and it is known that these excitations are in general well described by the RPA model. Our present results present a shift, with respect to the RPA energies, that is similar to that found with the subtracted SRPA scheme of Ref. [26], that provided a satisfactory agreement with the corresponding experimental responses [39].

In particular, the lowest $3^-$ level is predicted at an energy very close to the experimental value [40]. In Fig. 12 we see that the SRPA response shows a wide peak centered around 27 MeV, which is not present in RPA and DRPA. This comes from some RPA high-lying energy strength that is shifted downward in SRPA.

In Fig. 13 we show some positive (upper panel) and negative (lower panel) parity low-lying states obtained in DRPA with the experimental ones. In the DRPA case, the values in brackets indicate the corresponding one–phonon percentage in the wave functions.

V. CONCLUSIONS

We have presented an extension of the RPA model, denoted here as DRPA, allowing us to include approximately 2p–2h configurations, in a less explicit way compared to the SRPA model. The DRPA uses RPA phonons
(up to two) as building blocks to construct the excited states.

We have presented an application to low–lying states and giant resonances in the nucleus $^{16}$O.

In the case of giant resonances, the comparison with the RPA and the SRPA results indicates that (i) there is a global downwards shift of the energies with respect to the RPA spectra; (ii) this shift is however strongly reduced with respect to that produced by the standard SRPA. The DRPA results are in reasonably good agreement with the experimental responses.

This method represents for these cases an alternative way to include multiparticle–multihole configurations and to correct, at the same time, for the anomalous shift generated by the SRPA. Also for the low–lying states there is a global downwards shift with respect to the RPA results. The $1^-$ and $3^-$ states are in good agreement with the experimental values. The $0^+$ and $2^+$ states, however, have larger energies with respect to the measured values, even if the correction with respect to the RPA goes in the good direction, lowering the energies. This discrepancy with the experimental values may be understood as due to the almost pure $2p–2h$ nature of the low–lying $0^+$ and $2^+$ states. The implicit inclusion of $2p–2h$ configurations induced by the DRPA is not enough to well describe these excitations, and higher–order calculations would be necessary.

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