Self-consistent Theory of Finite Fermi Systems vs Skyrme–Hartree–Fock method. Spherical nuclei.

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Abstract. Recent results of the Fayans energy density functional (EDF) for spherical nuclei are reviewed. A comparison is made with predictions of several Skyrme EDFs. The charge radii and characteristics of the first $2^+$ excitations in semi-magic nuclei are briefly discussed. The single-particle spectra of doubly magic nuclei are considered in more detail. The phonon-particle coupling effects are analyzed including the tadpole term.

1 Introduction
In this century, the Hartree–Fock (HF) method with Skyrme forces\textsuperscript{(1)} dominates in the theoretical description of ground states of heavy nuclei. This method is often considered as a version of the Energy Density Functional (EDF) method by Kohn and Sham [2], which is based on the Hohenberg–Kohn theorem [3], stating that the ground state energy $E_0$ of a Fermi system is a functional of its density $\rho$. The Skyrme HF ansatz leads to a rather simple form of the nuclear EDF. The same is true for the HF method with Gogny force [4] which becomes again popular last decade. For completeness, we mention also the relativistic mean field (RMF) model, see recent Ref. 5 and links there, and a newly developed self-consistent approach, known as a method BCPM (Barcelona - Catania - Paris - Madrid) [6, 7].

We use an alternative EDF method developed by Fayans with coauthors\textsuperscript{8,11} with more sophisticated density dependence. The main, in-volume term of the Fayans EDF can be symbolically written as

$$\varepsilon_\rho(r) = \frac{a_\rho^2}{2} \frac{1 + \alpha_\rho^2 r^2}{1 + \gamma_\rho^2 r^2}, \quad (1)$$

where $\rho(r)$ is the total nuclear density, $\alpha, \alpha, \sigma, \gamma$ being parameters. The corresponding term of the Skyrme EDF corresponds to $\gamma=0$ in this relation. The use of the bare mass $m^*=m$ is another peculiarity of the Fayans EDF. Both the features of the Fayans method are closely related to the self-consistent Theory of Finite Fermi Systems (TFFS) and represent, in a hidden form, energy dependence effects inherent to this approach. The latter is based on the general principles of the TFFS [12] with inclusion of the TFFS self-consistency relations [13]. The final version of this approach [14, 15] was formulated in terms of the quasiparticle Lagrangian $L_q$, which is constructed to produce the quasiparticle mass operator $\Sigma_q(r, k^2; \varepsilon)$. By definition, the latter coincides with the exact mass operator $\Sigma(r, k^2; \varepsilon)$ at the Fermi surface. In the mixed coordinate-momentum representation it depends linearly on the momentum squared $k^2$ and the energy $\varepsilon$ as well [12].

In the TFFS, the effective mass is a product $m^*=m^k_1\cdot m^E_1$ of the “k-mass” and the “E-mass”. The two effects compensate each other almost exactly [15] resulting in $m^* \approx 1$ which justifies the Fayans choice of the bare mass instead of the effective one. The EDF of the self-consistent TFFS is found from the Lagrangian $L_q$ according the canonical rules. It includes implicitly the Z-factor

$$Z(r) = \left(1 - \frac{\partial \Sigma}{\partial \varepsilon} \right)_{\varepsilon=0}^{-1}, \quad (2)$$

where the index 0 means that the energy and momentum variables are taken at the Fermi surface. Its density dependence can be found explicitly [15]:

$$Z(r) = 2 \left(1 + \frac{1 - 4C_0L_0(r)/\varepsilon_0^2}{\varepsilon_0^2 - 1} \right)^{-1}, \quad (3)$$

where $C_0 = (dn/d\varepsilon_0^2)^{-1} = \pi^2/m^2_0$ is the usual TFFS normalization factor, inverse density of states at the Fermi surface, and the dimensionless parameter $\lambda_0$ determines the Z-factor of nuclear matter $Z_0$. In the result, the density dependence of the EDF becomes rather complicated [15]. Fayans with coauthors found [6] that, in a wide density region, it can be approximated with high accuracy by a more simple expression [1]. Thus, the Fayans EDF method can be interpreted as a version of the self-consistent TFFS.

Till this year, Ref. [16], all self-consistent calculations with Fayans functionals were carried out for spherical nuclei only. It proved out successful in systematic description of nuclear magnetic [17, 18] and quadrupole [13, 21] moments and nuclear radii [22] as well. In the latter case, agreement with the data is better than that in all SHF calculations we know. There are two examples more where we may compare the two approaches directly. The energies and $B(E2)$ values for the first $2^+$ excitations in semi-
The single-particle energies (SPEs) $\varepsilon_i$ in seven doubly magic nuclei, for which the experimental spectra are known [25], is another example [26]. The PC corrections to $\varepsilon_i$ were found self-consistently with account for the tadpole diagram. A systematic comparison to the SHF predictions with a popular HFB-17 EDF [27] was carried out. Even at the level of the mean field theory, the Fayans EDF results are significantly better. Inclusion of the PC corrections makes the agreement better yet.

The article is organized as follows. Sect. 2 contains a brief comparison with several Skyrme EDFs of predictions for the charge radii and characteristics of the first $2^+$ excitations in semi-magic nuclei. Sect. 3 is devoted to the description of SPEs in doubly magic nuclei. Sect. 4 contains conclusions.

2 Charge radii and characteristics of $2^+_1$ levels in semi-magic nuclei

In Ref. [22], systematic self-consistent calculations of charge radii $R_{cb}$ were made on the base of the DF3-a EDF. Deformed nuclei were also included into analysis, with an approximate taking into account the deformation effect. Agreement with the data on the level of 0.01 fm was achieved, noticeably better than for SLy4 and HFB-17 Skyrme EDFs taken for comparison. The HFB-17 predictions were taken from [28], whereas for the SLy4 EDF calculations were made in [22]. As an example, the lead charge radii are displayed in Fig. 1. We see that the Fayans EDF, indeed, describes the radii perfectly well. The HFB-17 one reproduces the data reasonably for heavy Pb isotopes but fails systematically for $A<190$, the disagreement reaching 0.1 fm. The reason for this is that this EDF erroneously predicts rather strong stable deformation for the light Pb isotopes which leads to a significant increase of $R_{cb}$ values. This problem is discussed in detail in [16]. The SLy4 $R_{cb}$ values are systematically higher than the experimental ones at approximately 0.03 fm. This is a typical scale of accuracy in describing nuclear radii for different Skyrme EDFs.

![Figure 1](image1.png)

**Figure 1.** Charge radii in lead isotopes. Solid line shows the Liquid Drop model predictions.

A high accuracy of reproducing the charge radii may be linked to more adequate density dependence of the Fayans EDF compared to the Skyrme one. Indeed, if we denote the average error in describing the binding energies as $\delta E$ and that for the charge radii as $\delta R_{cb}$, these quantities should be, due to the Hohenberg–Kohn theorem [3], proportional to each other,

$$\delta R_{cb} = \alpha \delta E,$$

where the coefficient $\alpha$ depends on the functional we use. This almost obvious relation can be proved with such simple consideration. Let we have an exact EDF $E_0(\rho)$ and add to it a small addendum $\delta E_0 = \lambda f(\rho)$, where the coefficient is small, $\lambda \ll 1$. Obviously, we get $\delta E = \lambda \int dr f(\rho(r)) = \lambda a$; then the change of the mean field $\delta U = \delta f/\delta \rho$ is proportional to $\lambda$. Let us find the change of the density $\rho$ in the first order of the perturbation theory $\delta U$. It can be easily seen that $\delta \rho$ and the corresponding change of the radius $\delta R_{cb}$ are also proportional to $\lambda$, $\delta R_{cb} = \lambda b$. In the result, we obtain (4) with $\alpha = b/a$. As a rule, a fine tuning of the EDF parameters is performed by focusing mainly on reproduction of the nuclear masses within a minimal value of $\delta E$. In this case, the accuracy of reproducing the charge radii is proportional to the coefficient $\alpha$. As the analysis of [22] showed, for the Fayans EDF this coefficient is less than those of the HFB-17 and SLy4 functionals. This observation may be linked to more sophisticated density dependence of Fayans functional, which allows to incorporate implicitly the energy dependence effects.

In Ref. [19], excitation energies $\omega_2$ and transition probabilities $B(E2)$ of the first $2^+$ excitations in even tin and lead isotopes as well as the quadrupole moments of odd neighbors of these isotopes were calculated within the self-consistent TFFS with the use of the DF3-a EDF. The effect of the density dependence of the effective pairing interaction was analyzed by comparing results obtained by solving the QRPA equations with volume and surface pairing. The effect was found to be noticeable giving evidences in favor of the surface pairing. For example, the $2^+$-energies are systematically higher at 200–400 keV for the volume pairing as compared with the surface pairing case, the latter being on average better. Obtained results
were compared with predictions of [24] for the Skyrme EDFs SkM* and SLy4. For tin isotopes, the comparison of $\epsilon_{g_1}$ and $B(E2)$ values is given in Figs. 2 and 3, correspondingly. On the average, predictions of the Fayans EDF with both models for pairing reasonably agrees with the data perfectly well, significantly better than those for both the Skyrme EDFs. Among the latter, the SkM* EDF turns out to be more successful. For SLy4 EDF, the $^{112}$Sn nucleus and its neighbors are very close to the point of the quadrupole instability, in contradiction with the data.

3 Single-particle spectra of magic nuclei

The major part of modern calculations with the Fayans functional in spherical nuclei are made with the EDF DF3-a. This is a version [29] of the EDF DF3 [9, 11] with modified spin-dependent parameters, the spin-orbit $\kappa$, $\kappa'$ and the first spin harmonics $g_1$, $g'_1$ which play the role of the effective tensor forces. The DF3 and DF3-a EDFs both contain 3 non-zero spin-dependent parameters, specifically the value of $g_1=0$ is putted. The bulk of the data [25] for SPEs contains 65 spin-orbit differences which permits to try to find an optimal set of the spin-dependent parameters of the EDF. Such a set DF3-b, with $g_1\neq0$, was found in [26], but the overall description of these differences turned out only a bit better than for DF3 or DF3-a sets. Moreover, the average accuracy in reproducing SPEs $\epsilon_{i,j}$, see Table 1, for DF3-b EDF is the same as for DF3 and only a little better than for the DF3-a EDF. The latter was chosen for systematic calculations as far as it previously proved to be successful in description of different nuclear phenomena [19, 21, 29]. For comparison, we found the SPEs of all nuclei under consideration for the Skyrme EDF HFB-17 [27] which is a record-holder in reproducing nuclear masses.

Figs. 4 and 5 show the comparison to the data [25] of predictions of all the EDFs under discussion for the SPEs in $^{208}$Pb for neutrons and protons, correspondingly. We see that both HFB-17 spectra, especially the neutron one, are too expanded. This is the result of influence of the effective mass $m^* < m$. The analysis in Ref. [30] confirmed a preference of the choice of the bare mass for describing SPEs with Skyrme EDFs without PC corrections.

**Table 1.** Average deviations $\langle \delta\epsilon_{i,j}\rangle_{\text{rms}}$ (MeV) of the theory predictions for the single-particle energies from the experimental values for magic nuclei.

| Nucleus | $N$ | DF3-b | DF3-a | DF3 | HFB17 |
|---------|-----|-------|-------|-----|-------|
| $^{40}$Ca | 14 | 1.08 | 1.25 | 1.35 | 1.64 |
| $^{48}$Ca | 12 | 0.89 | 1.01 | 1.00 | 1.70 |
| $^{56}$Ni | 14 | 1.00 | 0.97 | 0.85 | 1.40 |
| $^{78}$Ni | 11 | 1.24 | 1.41 | 1.09 | 1.32 |
| $^{100}$Sn | 13 | 1.09 | 1.17 | 1.01 | 1.56 |
| $^{132}$Sn | 17 | 0.58 | 0.66 | 0.55 | 1.15 |
| $^{208}$Pb | 24 | 0.44 | 0.51 | 0.43 | 1.15 |
| Total | 105 | 0.89 | 0.98 | 0.89 | 1.40 |

**Figure 3.** $B(E2,\text{up})$ values for tin isotopes. Predictions for the SkM* and SLy4 force are taken from [24].

**Figure 4.** Neutron single-particle levels in $^{208}$Pb. Experimental data from [25].

**Figure 5.** Proton single-particle levels in $^{208}$Pb. Experimental data from [25].

**Figure 6.** PC corrections to the mass operator. The gray circle denotes the “tadpole” term.
Figure 7. The vertex $g_L$ for the $3^+_1$ state in $^{208}\text{Pb}$.

Let us go to the PC contributions to SPEs. Accounting for PC effects, the equation for SPEs and corresponding wave functions can be written as

$$ (\varepsilon - H_0 - \delta \Sigma^{PC}(\varepsilon)) \phi = 0, \quad (5) $$

where $H_0$ is the quasiparticle Hamiltonian with the spectrum $\varepsilon^{(0)}_q$ and $\delta \Sigma^{PC}$ is the PC correction to the quasiparticle mass operator. After expanding this term in the vicinity of $\varepsilon = \varepsilon^{(0)}_q$ one finds

$$ \varepsilon_q = \varepsilon^{(0)}_q + Z^{PC}_A \delta \Sigma^{PC}(\varepsilon^{(0)}_q), \quad (6) $$

with obvious notation. Here $Z^{PC}$ denotes the $Z$-factor due to the PC effects, i.e. that found from Eq. (2) with substitution of $\delta \Sigma^{PC}(\varepsilon)$ instead of the main mass operator $\Sigma(\varepsilon)$. Remember that in the TFFS the corresponding $Z$-factor is included in the quasiparticle Hamiltonian $H_0$.

The PC correction to the mass operator is displayed in Fig. 6, where $g_L$ is the vertex for creating the $L$-phonon. In magic nuclei, it obeys the equation [12]

$$ g_L(\omega) = \mathcal{F} A(\omega) g_L(\omega), \quad (7) $$

where $A(\omega) = \int G(\varepsilon + \omega/2) G(\varepsilon - \omega/2) d\varepsilon/(2\pi)$ is the particle-hole propagator, $G(\varepsilon)$ being the one-particle Green function. In obvious symbolic notation, the pole diagram corresponds to $\delta \Sigma^{pole} = (g_L, D_L g_L)$, where $D_L(\omega)$ is the phonon $D$-function.

All the low-lying phonons we consider have natural parity. In this case, the vertex $g_L$ possesses even $T$-parity. It is a sum of two components with spins $S = 0$ and $S = 1$, respectively,

$$ g_L = g_{13}T_{13}(n, a) + g_{11}T_{11}(n, a), \quad (8) $$

where $T_{13}$ and $T_{11}$ stand for the usual spin-angular tensor operators. The operators $T_{13}$ and $T_{11}$ have opposite $T$-parities, hence the spin component should be the odd function of the excitation energy, $g_{13} \propto \omega_L$. For the ghost dipole, $L = 1$ and $\omega_1 = 0$, Eq. (7), due to the TFFS self-consistency relation [13], has the exact solution

$$ g_L(r) = \alpha_1 (dU/dr) |_{\Omega} Y_{11}(n), \quad (9) $$

where $\alpha_1 = 1/\sqrt{2m\omega_B}$, $B_1 = 3mA/4\pi$ is the Bohr-Mottelson (BM) mass coefficient [13] and $U(r)$ is the central part of the mean-field potential generated by the energy functional.

The second, tadpole, term in Fig. 6 is

$$ \delta \Sigma^{tad} = \int \frac{d\omega}{2\pi} \delta L g_L D_L(\omega), \quad (10) $$

where $\delta L g_L$ can be found [15] by variation of Eq. (7) in the field of the $L$-phonon:

$$ \delta L g_L = \delta L F A(\omega_L) g_L + \mathcal{F} \delta L A(\omega_L) g_L + \mathcal{F} A(\omega_L) \delta L g_L, \quad (11) $$

The quantity $\delta L F$ can be readily obtained by variation of each Green function in the particle-hole propagator $A$ in field $g_L$ induced by the $L$-phonon. The explicit expression for the variation $\delta L F$ can not be found within the TFFS as in this approach the Landau–Migdal amplitude $\mathcal{F}$ is introduced as a phenomenological quantity. In [15] the ansatz was proposed,

$$ \delta L F = \frac{\delta F(\rho)}{\delta \rho} \delta \rho_L, \quad (12) $$

where $\delta \rho_L = A_L g_L$ is the transition density for excitation of the $L$-phonon.

The $L$-phonon excitation energies $\omega_L$ and creation amplitudes $g_L(r)$ were found by solving Eq. (7) in a self-consistent way with the DF3-a functional. In more detail, the procedure is described in [19]. All the $L$-phonons we consider are the surface vibrations which belong to the Goldstone mode corresponding to the spontaneous breaking of the translation symmetry in nuclei [11]. The coordinate form of their creation amplitudes $g_L(r)$ is very close to that, Eq. (9), for the ghost 1$^-$ phonon, which is the lowest energy member of this mode:

$$ g_L(r) = \alpha_L (dU/dr) + \chi_L(r), \quad (14) $$

where the in-volume correction $\chi_L(r)$ is rather small.

The smallness of the in-volume component $\chi_L$ is demonstrated in Fig. 7 for the $3^+_1$ state in $^{208}\text{Pb}$, which is the most collective one among the surface vibrations and plays the main role in PC corrections for this nucleus. The small spin components $S = 1$ are also displayed. To make them distinguishable, they are multiplied by the factor of 10. The smallness of the spin components is typical for $L$-phonons with a high collectivity. The first, surface term on the right-hand sight of Eq. (14) corresponds to the BM model for the surface vibrations [11], the amplitude $\alpha_L$ being related to the dimensionless BM amplitude $\beta_L$ as follows: $\alpha_L = R\beta_L$, where $R = r_0 A^{1/3}$ is the nucleus radius, and $r_0 = 1.2$ fm.

If one neglects in-volume contributions, the tadpole PC term [10] can be reduced to a very simple form:

$$ \delta \Sigma^{tad}_L = \frac{\alpha_L^2}{2} \frac{2L + 1}{3} \Delta U(r). \quad (15) $$
Table 2. Pole and tadpole contributions (MeV) to PC corrections from $^3-$states to neutron SPEs in $^{208}\text{Pb}$.

| $\lambda$ | $\delta\varepsilon^{\text{pole}}_{\lambda}$ | $\delta\varepsilon^{\text{ind}}_{\lambda}$ | $\delta\varepsilon_{\lambda}$ |
|-----------|-----------------|-----------------|-----------------|
| $3d_{3/2}$ | -0.150           | 0.012           | -0.137          |
| $2g_{7/2}$ | -0.142           | 0.061           | -0.081          |
| $4s_{1/2}$ | -0.134           | 0.016           | -0.118          |
| $3d_{5/2}$ | -0.147           | 0.023           | -0.124          |
| $1f_{5/2}$ | -0.708           | 0.204           | -0.504          |
| $1i_{11/2}$ | -0.058          | 0.198           | 0.140           |
| $2h_{9/2}$ | -0.244           | 0.076           | -0.167          |
| $3p_{1/2}$ | -0.220           | 0.053           | -0.167          |
| $2f_{5/2}$ | -0.186           | 0.094           | -0.092          |
| $3p_{3/2}$ | -0.205           | 0.056           | -0.149          |
| $1i_{13/2}$ | 0.057           | 0.211           | 0.269           |
| $2f_{7/2}$ | 0.724            | 0.091           | 0.815           |
| $1h_{9/2}$ | -0.014           | 0.197           | 0.184           |

Table 3. Pole and tadpole contributions (MeV) to PC corrections from $^3-$states to proton SPEs in $^{208}\text{Pb}$.

| $\lambda$ | $\delta\varepsilon^{\text{pole}}_{\lambda}$ | $\delta\varepsilon^{\text{ind}}_{\lambda}$ | $\delta\varepsilon_{\lambda}$ |
|-----------|-----------------|-----------------|-----------------|
| $3p_{1/2}$ | -0.375           | 0.153           | -0.222          |
| $3p_{3/2}$ | -0.371           | 0.152           | -0.219          |
| $2f_{5/2}$ | -0.278           | 0.168           | -0.110          |
| $1i_{13/2}$ | -0.534           | 0.266           | -0.268          |
| $2f_{7/2}$ | -0.409           | 0.168           | -0.240          |
| $1h_{9/2}$ | -0.054           | 0.222           | 0.168           |
| $3s_{1/2}$ | -0.310           | 0.143           | -0.167          |
| $2d_{5/2}$ | -0.241           | 0.146           | -0.095          |
| $1h_{11/2}$ | -0.017           | 0.246           | 0.229           |
| $2d_{5/2}$ | 0.435            | 0.147           | 0.582           |
| $1g_{7/2}$ | -0.271           | 0.197           | -0.074          |

The general consideration of the tadpole term within the not self-consistent TFFS, with solving Eq. (11), was carried out in [32]. It was found that the in-volume corrections to Eq. (15) are, indeed, small for heavy nuclei, e.g., for $^{208}\text{Pb}$. At the same time, for light nuclei, e.g., $^{40,48}\text{Ca}$, the accurate solution of Eq. (11) diminishes the approximate value (15) for the tadpole term by $\approx 30\%$

Following to [26], we neglect the in-volume corrections for all nuclei considered. To find the phonon amplitudes $\alpha_\lambda$, we use the definition $\alpha_\lambda = \frac{g_{\varepsilon_\lambda}^{\text{max}}}{dU/dr} \bar{r}_{\varepsilon_\lambda}^{\text{max}}$, with obvious notation. It should be noted that the values of $\alpha_\lambda$ and $\alpha_\lambda^p$ are always very close to each other and to that which follows from the BM model formula for $B(EL)$: $B(EL)_{\text{BM}} = (3Z/4\pi)^2 \beta_1^2 R^{EL}$ [31], where the dimensionless BM phonon creation amplitude $\beta_1$ is related to that used by us as $\alpha_\lambda = \beta_1 R/\bar{r}_{\varepsilon}^{\text{max}}$, $R = 1.2 A^{1/3}$. For example, for the $3\gamma_2$ state in $^{208}\text{Pb}$ we have: $\alpha_\lambda = 0.32 \text{ fm}$, $\alpha_\lambda^p = 0.33 \text{ fm}$, and $\alpha_\lambda^{BM} = 0.30 \text{ fm}$.

A comparison of the pole and tadpole PC corrections to neutron and proton SPEs, induced by the $3\gamma_2$ state in $^{208}\text{Pb}$, are given in tables 2 and 3, correspondingly. The tadpole term is always positive, whereas the pole one is, as a rule, negative, and the two contributions are of the opposite sign. The magnitude of the tadpole term is, as a rule, less than the pole one, but comparable with the latter. Especially it this true for protons. Therefore the sum is often essentially less than the pole term alone. A typical suppression of the pole contribution is of 30–50%, but there are cases of a stronger suppression, e.g. $2d_{5/2}$ and $1g_{7/2}$ proton states. Moreover, there are several cases than the tadpole term dominates: the neutron $1i_{11/2}$ and $1h_{9/2}$ states and the proton $1h_{9/2}$ and $1i_{11/2}$ ones. In these cases, the total correction is of the opposite sign as compared with the pole term. The general conclusion is that one overestimates the PC correction to SPEs neglecting the tadpole term, and often it is better to omit it completely than consider the pole term alone.

The total PC effect to SPEs of all doubly magic nuclei under consideration is shown in table 4. We see that it makes the agreement essentially better for $^{208}\text{Pb}$ but a bit less good in the light nuclei. The latter is explained mainly with the use the approximation (15) for the tadpole term. According to [32], the exact consideration for $^{40,48}\text{Ca}$ diminishes the tadpole term by approximately 30%. Such a correction should lead to a better agreement with the experimental SPEs. Let us stress once more a high accuracy of reproducing the experimental SPEs in $^{208}\text{Pb}$: $\langle \delta\varepsilon_1 \rangle_{\text{rms}} = 0.28 \text{ MeV}$. Indeed, the corresponding result [33], obtained within the RMF approach, is $\langle \delta\varepsilon_1 \rangle_{\text{rms}} = 0.85 \text{ MeV}$.

Table 4. PC effect on average deviations $\langle \delta\varepsilon_1 \rangle_{\text{rms}}$ (MeV) of the theory predictions for SPEs from the experimental values for the DF3-a functional.

| Nucleus | $N$ | $\text{DF3-a+PC}$ | $\text{DF3-a}$ |
|---------|-----|-------------------|----------------|
| $^{40}\text{Ca}$ | 14 | 1.30 | 1.25 |
| $^{48}\text{Ca}$ | 12 | 1.05 | 1.00 |
| $^{50}\text{Ni}$ | 14 | 0.98 | 0.97 |
| $^{78}\text{Ni}$ | 11 | 1.34 | 1.41 |
| $^{100}\text{Sn}$ | 13 | 1.21 | 1.17 |
| $^{132}\text{Sn}$ | 17 | 0.63 | 0.66 |
| $^{208}\text{Pb}$ | 24 | 0.38 | 0.51 |

4 Conclusions

Recent studies with the Fayans EDF for spherical nuclei are reviewed and compared to predictions of several Skyrme EDFs. The charge radii and the characteristics of the first $2^+$ excitations in semi-magic nuclei are briefly discussed. The accuracy of reproducing the charge radii is of the order of 0.01 fm, which is significantly better than that obtained with HFB-17 and SLy4 EDFs. The excitation energies and $B(E2)$ values of the first $2^+$ states in the lead and tin isotopic chains also agree with the data much better than the results [24] obtained with the SLy4 and SKM* EDFs.

The SPEs of doubly magic nuclei are considered in more detail. The PC effects in SPEs are analyzed particu-
larly. In addition to the usual pole diagram, the consideration includes the so-called tadpole term which is usually ignored \[^{13}[14]\]. The latter is considered approximately, with neglecting the in-volume components of the vertex \(g_\ell(r)\) of creating a surface \(L\)-phonon. This approximation works well in heavy nuclei, leading to a simple formula for the tadpole contribution which can be easily included into the calculation scheme. The tadpole contribution is often comparable with that of the pole diagram. As a rule, these two contributions to SPEs have different signs, their sum being often significantly less than that of the pole term alone. For the \(^{208}\)Pb nucleus, the calculation without PC corrections results in the average deviation \(\langle \delta \varepsilon_{L} \rangle_{\text{rms}}\) from the experimental values equal to 0.51 MeV for the Fayans EDF, whereas it is 1.15 MeV for the HFB-17 EDF. With the PC corrections, including the tadpole contribution, we obtained very high accuracy: \(\langle \delta \varepsilon_{L} \rangle_{\text{rms}}=0.34\) MeV. For a comparison, the corresponding quantity, found in \[^{13}\] within the RMF approach with PC corrections without the tadpole term, is \(\langle \delta \varepsilon_{L} \rangle_{\text{rms}}=0.85\) MeV.

To conclude, the Fayans EDF leads to a better agreement with the experimental data in all phenomena in spherical nuclei considered than several popular Skyrme EDFs chosen for a comparison. We relate it to the peculiarities of the Fayans EDF which reflect, in a hidden form, the energy dependence effects inherent to the self-consistent TFFS.

5 Acknowledgment
The work was partially supported by the Grant NSh-932.2014.2 of the Russian Ministry for Science and Education and by the RFBR Grants 13-02-00085-a, 13-02-12106_ofi-m, 14-02-00107-a, 14-22-03040_ofi-m.

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