Multiconfiguration Dirac-Hartree-Fock calculations of the electric dipole moment of radium induced by the nuclear Schiff moment

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(Rceived 23 March 2009; published 23 July 2009)

The multiconfiguration Dirac-Hartree-Fock theory has been employed to calculate the electric dipole moment of the $7s6d^3D_2$ state of radium induced by the nuclear Schiff moment. The results are dominated by valence and core-valence electron correlation effects. We show that the correlation effects can be evaluated in a converged series of multiconfiguration expansions.

DOI: 10.1103/PhysRevA.80.012513  PACS number(s): 31.30.jg, 11.30.Er, 21.10.Ky, 24.80.+y

I. INTRODUCTION

A nonzero permanent electric dipole moment (EDM) of an atom, molecule, or any other composite or elementary particle is one of the possible manifestations of parity ($P$) and time ($T$) reversal symmetry violations. In the absence of any external electromagnetic field an atom can have a permanent EDM either due to an intrinsic EDM of one of its constituent particles or due to $P$- and $T$-violating ($P$-odd and $T$-odd) interactions between these particles [1,2]. When compared with the intrinsic EDM of the constituent particles, the net induced EDM of an atom or molecule is often expected to be larger by several orders of magnitude due to various nuclear and atomic enhancement mechanisms. Therefore, atoms and molecules are considered to be very attractive for carrying out EDM experiments and in the search for "new physics" beyond the standard model since, in the latter case, the induced EDM is greatly suppressed when compared to the anticipated values from the "new" theories [3]. In atomic physics, in particular, the experimental search for a permanent EDM is gaining momentum due to recent advancements in trapping free neutral atoms [4–6], including various radioactive species [7,8].

During the last decade several atoms were considered as candidates for such experiments [3,9]. These involved (i) diamagnetic atoms (i.e., total angular momentum $J=0$) in their respective ground states, (ii) the alkali metals, and (iii) atoms with a single $p$ electron outside closed shells, which were investigated in laser traps for their prospectives to perform EDM experiments. Presently, radium appears to be the most promising candidate, and experiments on this element are under way at the Argonne National Laboratory [7,10,11] as well as at the Kernfysich Versneller Instituut [12–14]. The main advantages of radium lay in (i) large nuclear charge $Z$, (ii) simple electronic structure and closed-shell [Kr]$4d^{10}4f^{14}5s^25p^6d^6$ ground state, (iii) octupole deformations of the radium nuclei for several isotopes [15,16], as well as in (iv) coincidental proximity of two atomic levels of opposite parity, $7s7p$ $^3P_1$ and $7s6d$ $^3D_2$, which are separated by a very small energy interval $5.41\text{ cm}^{-1}$. In particular, the latter two advantages give rise to a relatively large enhancement factor, which is one of the largest among the atoms considered so far [17,18].

However, extraction of fundamental $P$- and $T$-violating parameters or coupling constants from experimentally measured atomic EDM requires atomic form factors which can be provided only by an ab initio atomic theory. Several of these form factors have been previously calculated by the group of Flambaum [9,17]. In practice, there are essentially four different form factors related to four mechanisms which can induce an atomic EDM. An atom can acquire a permanent EDM due to (1) $P$- and $T$-odd electron-nucleon interaction or due to the electromagnetic interaction of atomic electrons with nuclear $P$- and $T$-odd moments, of which the leading ones are (2) the Schiff moment, (3) the magnetic quadrupole moment, and (4) the electric octupole moment. The latter two moments may exist only in nuclei with spins larger than $I=1/2$, while the Schiff moment may exist also in isotopes with nuclear spin $I=1/2$. Nuclei with spins larger than $I=1/2$ produce electric quadrupole shifts which are difficult to account for in an EDM measurement. Therefore an EDM induced by the Schiff moment in $I=1/2$ isotopes
seems to be the property of choice among (most of) the experimenters [13,19,20]. The nuclear Schiff moment is a $P$-odd and $T$-odd (electric-dipole) moment that occurs due to $P$- and $T$-violating interactions at the nuclear scale. The Schiff moment mixes atomic states of opposite parity and may induce static EDM in atoms if magnetic and finite-size effects are taken into account in the electron-nucleus interaction [3].

In the present paper, we present calculations for the atomic EDM in radium as induced by the Schiff moment. The atomic wave functions were obtained within the framework of the multiconfiguration Dirac-Hartree-Fock (MCDHF) theory. The wave functions were separately optimized for the 7s6d $^3D_2$ and 7s7p $^3P_1$ states (similar calculations were carried out recently for the scalar-pseudoscalar contribution to the EDM in cesium [21]). The main purpose of this paper is to provide a systematic evaluation of the effects of electron correlation on the calculated EDM of radium. We demonstrate the saturation of the core-valence correlations, the dominant electron correlation effect beyond the Dirac-Fock approximation.

II. THEORY

The Hartree-Fock and Dirac-Hartree-Fock theories based either on the finite-grid, basis-set, or some other numerical methods provide a natural point of departure in describing the electronic structure of atoms and molecules. For medium and heavy elements, these methods are often combined with Breit-Pauli or Dirac-Coulomb-Breit Hamiltonians in order to account for relativistic and retardation effects on the wave functions and the level structure of complex atoms. However, the main hindrance in applying modern computational techniques arises from the electron-electron correlation, i.e., the residual interaction among the electrons beyond the atomic mean field, and this is especially true for systems with many electrons. In neutral or nearly neutral systems, missing electron-correlation effects are indeed often the main reason for the discrepancies between the observed and calculated properties of atoms.

Today, many-body perturbation theory (MBPT) [22–25] and various variational methods, often referred to as the multiconfiguration Hartree-Fock (MCHF) theory (or its relativistic counterpart, the Dirac-Hartree-Fock theory [26–28]) are the two dominant pillars in performing atomic structure calculations. These methods are designed to evaluate the electron correlation effects in a systematic manner. For MBPT and most related methods, a nearby closed-shell configuration of the atom or ion is typically a convenient starting point. During the past decades, therefore, MBPT techniques were mainly applied to systems with either no or just a single electron outside closed shells. Systems with several electrons in open shells are, in contrast, much more difficult to deal with if benefit is to be taken from the theory of angular momentum and spherical tensor operators, i.e., by using a restricted representation of the one-electron orbitals. Difficulties occur then due to the large departure from a closed-shell $V^0$ potential and the rapid increase in the complexity of all perturbation expansions with any additional electron outside closed shells. The difficulties with the open-shell systems in MBPT approach can, in principle, be treated by applying a $V^{N-M}$ approximation [23] or by employing one of several hybrid approaches in which the configuration-interaction method is combined with the many-body perturbation theory [29,30] or with the coupled cluster approach [31]. Although in the variational multiconfiguration methods the algebraic complexity also depends on the shell structure of the atom, these methods can be applied more easily to systems with an arbitrary number of electrons outside closed shells. Apart from the number of open shells, the accuracy of multiconfiguration Dirac-Hartree-Fock calculations depends crucially also on the occupation of the valence shell(s) but this occurs rather indirectly through the limiting number of configuration state functions that can be included in a particular wave function expansion. In this sense, the limitations are less conceptual but arise from the available computer resources. These limitations are typically related to the structure of the valence shell(s), i.e., to the angular properties of the valence electrons and their couplings.

A. MCDHF theory

We used a slightly modified version of the General Relativistic Atomic Structure Package (GRASP) [28] to generate the electronic wave functions. In the multiconfiguration Dirac-Hartree-Fock method, the wave function for a particular atomic state $\Psi(\gamma PJM_j)$ is obtained as a linear combination of configuration state functions (CSFs) which are eigenfunctions of the parity $P$ and the total angular momentum operators $J^2$ and $J_z$,

$$\Psi(\gamma PJM_j) = \sum_r c^*_r \Phi(\gamma PJM_j).$$

In the present computations, the wave functions were separately generated for the 7s7p $^3P_1$ and 7s6d $^3D_2$ states of radium. Each wave function was obtained as self-consistent solution of the Dirac-Hartree-Fock equations [26] using a systematically increased multiconfiguration bases (of size $N_{\text{CSF}}$) of symmetry-adapted configuration state functions $\Phi(\gamma PJM_j)$. Configuration mixing coefficients $c_r$ were obtained through the diagonalization of the Dirac-Coulomb Hamiltonian

$$\hat{H}_{\text{DC}} = \sum_{j=1}^N \left[ c \alpha_j \cdot p_j + (\beta_j - 1)c^2 + V(r_j) \right] + \sum_{j>k} 1/r_{jk},$$

where $V(r)$ is the monopole part of the electron-nucleus interaction. A more detailed description of the theory [27,32] and method of calculation [33,34] can be found elsewhere.

B. Nuclear shape

All self-consistent-field calculations were done with the nucleus modeled as a spherical ball by applying a two-parameter Fermi distribution.
in order to approximate the radial dependence of the nuclear charge density \( \rho(r) \). The parameter \( \rho_0 \) is derived from the normalization condition \( \int \rho(r) d^3r = Z_e \). Figure 1 shows the nuclear charge density \( \rho(r) \) inside the \( ^{223}\text{Ra} \) nucleus calculated with the parameters \( c = 6.85 \) fm and \( a = 0.523 \) fm (see [35,36] for details). For the other two isotopes the “half-charge-density” parameter \( c \) was set to \( c = 6.83 \) fm (\( ^{223}\text{Ra} \)) and \( c = 6.73 \) fm (\( ^{225}\text{Ra} \)), with the “nuclear-skin-thickness” \( a \) value unchanged. These nuclear charge distributions were also used in the subsequent calculations of the Schiff moment expectation values [Eq. (14)].

In the GRASP code [28] all electronic (radial) orbitals are represented on a numerical grid which increases exponentially in order to ensure an accurate representation of the atomic wave functions near the nucleus. The grid is generated from the formula \( r_i = r_0 \exp[(i-1)h] \), with \( r_0 = 2.0 \times 10^{-8} \times a_0 \), \( a_0 = 1.0 \times 10^{-3} \), and \( i = 1, \ldots , 4000 \). With the above parameters, there were 1255 grid points within the half-charge-density nuclear radius \( r_{nc} = 6.85 \) fm, while the numerical representation of the full extent of all radial (core and valence) electronic orbitals required more than 3000 points. With the above choice of the radial grid, all necessary one- and two-particle matrix elements can be calculated with a (relative) accuracy of the order of \( 10^{-8} \) or better. Note however that larger uncertainties may arise from the radial matrix elements in Eq. (14) due to the approximate nature of the nuclear charge distribution in Eq. (3) and due to deviations from radial symmetry of those isotopes, for which nuclear deformations are significant for model dependence see [37].

C. Atomic EDM

Neglecting the contributions from the off-diagonal hyperfine interaction the coupled wave function of the total system “electrons+ nucleus” is given by the Clebsch-Gordan expansion [38]

\[
\Psi(\gamma\nu PJFM_F) = \sum_{M_J M_F} \langle J M_J M_F | PJFM_F \rangle \Psi(\gamma PJM_J) \Psi(\nu M_F),
\]

where \( \Psi(\nu M_F) \) represents the ground state of the nucleus and where the standard notation is used for the Clebsch-Gordan coefficients. For high-Z elements with closed levels of opposite parity, such as radium in the \( ^3P_1 \) and \( ^3D_2 \) levels, one of the most important P and T symmetry violating interactions is caused by the nuclear Schiff moment \( S \), which gives rise to the electron-nucleus interaction:

\[
\hat{H}_{SM} = 4\pi \sum_{j=1}^{N} (S \cdot \nabla_j) \rho(r_j).
\]

In this Hamiltonian, \( \rho(r) \) is the normalized to unity nuclear density function from Eq. (3), and the Schiff moment \( S \) is directed along the nuclear spin \( \hat{S} = \hat{S}_I/\ell \). The interaction in Eq. (5) mixes states of different parity and may also induce a static electric dipole moment of the atom. Since the Schiff moment interaction is quite weak, we can express the wave function of the (mixed-parity) hyperfine state \( |F, M_F\rangle \) of the level \( J^P \) as [21]

\[
\tilde{\Psi}(\gamma\nu PJFM_F) = a \Psi(\gamma\nu PJFM_F) + \sum_{i=1}^{m} b_i \Psi(\alpha_i \nu(-P)JIFM_F),
\]

where the coefficient \( a \) of the given hyperfine state can be set to 1. The expansion coefficients of the other (hyperfine) states of opposite parity can be perturbatively approximated by

\[
b_i = \frac{\langle \Psi(\alpha_i \nu(-P)JIFM_F) | \hat{H}_{SM} | \tilde{\Psi}(\gamma\nu PJFM_F) \rangle}{E(\gamma PJ) - E(\alpha_i (-P)J_i)}. \tag{7}
\]

The mixed-parity wave function in Eq. (6) for the hyperfine state \( |F, M_F\rangle \) of a particular atomic level \( ^{2S+1}L_J \) induces a static EDM of an atom:

\[
D_A = \langle \tilde{\Psi}(\gamma\nu PJFM_F) | \hat{D}_z | \tilde{\Psi}(\gamma\nu PJFM_F) \rangle = 2 \sum_{i=1}^{m} b_i \langle \Psi(\gamma\nu PJFM_F) | \hat{D}_z | \Psi(\alpha_i \nu(-P)JIFM_F) \rangle, \tag{8}
\]

where \( \hat{D}_z \) denotes the z projection of the electric-dipole moment operator. For this electric-dipole operator, the matrix element between (hyperfine) states of different parity can be expressed as

\[
\langle \Psi(\gamma\nu PJFM_F) | \hat{D}_z | \Psi(\alpha_i \nu(-P)JIFM_F) \rangle = (-1)^{J_z + J_z + F + 1} (2J + 1) \begin{pmatrix} F & 1 & F \\ -M_F & 0 & M_F \end{pmatrix} \times \begin{pmatrix} J & F & \ell \\ F & J_i & 1 \end{pmatrix} \langle \Psi(\gamma PJ) | \hat{D}_z | \Psi(\alpha_i (-P)J_i) \rangle, \tag{9}
\]

while the matrix element of the (scalar) electron-nucleus interaction in Eq. (5), induced by the Schiff moment, is written as
\[
\langle \Psi(\gamma \nu P J I M F) | \hat{H}_{SM} | \Psi(\alpha, (\nu-P) J I M F) \rangle 
= (-1)^{F+J} \sqrt{2J+1} \frac{1}{\varpi} \left\{ 2(\nu-J+1) \frac{1}{\varpi} \left[ \mathbf{I} \mathbf{I} \mathbf{J} \mathbf{J} \mathbf{F} \right] \right\} \]
\]

and is independent of \(M_F\). In the following, we shall refer to the last term on the right-hand side of Eq. (10) as the reduced matrix element of the Schiff operator for the two (fine-structure) levels \(J\) and \(J_f\) of different parity and shall assume \(M_F = F\), in line with optical pumping schemes of hyperfine levels with circularly polarized light. For the \(|F, M_F\rangle\) hyperfine state of the \(^3D_2\) level, the static EDM in Eq. (8), induced by the \(P\)-odd and \(T\)-odd nuclear Schiff moment, becomes

\[
D_N(^3D_2, FM_F) = 2\frac{\langle \Psi(^3D_2, FM_F) | \hat{D}_2 | \Psi(^3\nu P J I M F) \rangle \langle \Psi(^3\nu P J I M F) | \hat{H}_{SM} | \Psi(^3D_2, FM_F) \rangle}{E(^3D_2) - E(^3\nu P)}
\]

For the calculations of the matrix elements we extended the GRASP [28] and MDFGME [40] relativistic atomic structure packages. The extension, presented in this work, includes programs for both Schiff moment interaction and electric-dipole moment matrix elements. Experimental energy differences were used in the calculations of all expectation values. The energy values for the two levels of interest \((E_{^3\nu P} - E_{^3D_2}) = 13 \text{ 993.97 cm}^{-1}\) and \((E_{^3\nu P} - E_{^3D_2}) = 13 \text{ 999.38 cm}^{-1}\) were taken from the tables of Moore [41]. The nuclear spin and magnetic moment data were taken from the tables of Raghavan [42].

D. Handling nonorthogonalities

The electronic wave functions were optimized separately for the two levels of interest. All expectation values were evaluated with the biorthogonal technique developed by Malmquist [43]. For two atomic state functions

\[
\Psi(\gamma \nu P J) = \sum_r c_r \Phi(\gamma \nu P J)
\]

and

\[
\Psi(\alpha(-P) J) = \sum_s c_s \Phi(\alpha(-P) J)
\]

the reduction in a general matrix element

\[
\langle \Psi(\gamma \nu P J) | \hat{D}_n | \Psi(\alpha(-P) J) \rangle
\]

into a sum of radial integral \(\times\) angular coefficient terms is based on tensor algebra techniques. In the decomposition [Eq. (13)], it is usually assumed that the (many-electron) configuration states on both sides of the matrix element are built from a common set of spin orbitals. This is a very severe restriction since a high-quality wave function demands orbitals optimized for the specific electronic state. Instead of the standard decomposition [Eq. (13)] based on tensor algebra techniques, Malmquist showed [43] that for very general expansions, where the two atomic states are described by different orbital sets, it is possible to transform the wave function representations of the two states in such a way that standard techniques can be used for the reduction of the matrix elements in the new representation. This procedure
has been implemented in the modules that compute Schiff moments, and it can be summarized as follows:

1. Perform MCDHF or configuration interaction (CI) calculations for the two states where the orbital sets of the two wave functions are not required to be identical.

2. Change the wave function representations by transforming the two orbital sets to a biorthogonal basis. This is followed by a countertransformation of the expansion coefficients \( c_i \) and \( c_j \), so as to leave the resultant wave functions invariant.

3. Calculate the matrix elements with the transformed wave functions for which now standard techniques can be used [27].

The transformation of wave functions is very fast since it relies only on angular coefficients for a one-electron operator of rank zero which appears in the evaluation of the kinetic-energy term in the MCDHF or CI step. The details of the transformations are discussed in [44].

III. METHOD OF CALCULATION

In the first phase of the computations, all spectroscopic orbitals required to form a reference wave function were obtained with a minimal configuration expansion with full relaxation. These orbitals were determined from a symmetry-adapted Dirac-Fock calculation with only those configurations which arise in \( j \)-\( j \) coupling for a particular state of interest. The spectroscopic orbitals were kept frozen in all later steps.

In the second phase the virtual orbitals were generated in five consecutive steps. At each step the virtual set has been extended by one layer of virtual orbitals. A layer is defined as a set of virtual orbitals with different angular symmetries. In the present paper five layers of virtual orbitals were generated, each layer comprising orbitals with symmetries \( s, p, d, f, g, \) and \( h \). At each step the configuration expansions were limited to single and double substitutions from valence shells to all new orbitals as well as to all (virtual) orbitals of the previously generated layers. These substitutions were augmented by a small subset of dominant single and double substitutions from core and valence shells, with the further restriction that at most one electron may be promoted from the core shells (i.e., in a double substitution at least one electron is promoted from a valence shell). All configurations from earlier steps were retained, with all previously generated orbitals fixed, and all new orbitals made orthogonal to all others of the same symmetry. The initial shapes of radial orbitals were obtained by means of a Thomas-Fermi potential and then driven to convergence with the self-consistency threshold set to \( 10^{-10} \) for spectroscopic orbitals and \( 10^{-8} \) for virtual orbitals, respectively. All radial orbitals were separately optimized for each atomic state. The Optimal Level form of the variational expression [35] was applied in all variational calculations.

B. Configuration-interaction calculations

In the third phase of the computations, the configuration-interaction calculations (i.e., without changing the radial shapes of the one-electron spin orbitals) were performed, with multiconfiguration expansions tailored in such a way as to capture the dominant electron correlation contributions to the expectation values. All single and double substitutions were allowed from several core shells and from both valence shells (i.e., 7s7p or 7s6d, depending on the state) to all virtual shells, with the same restriction as above, i.e., that at most one electron may be promoted from core shells. The virtual set was systematically increased from one to five layers. In a similar manner, several core subshells were systematically opened for electron substitutions — from the outermost 6p up to the 5s5p5d6s6p subshells.

The convergence of the calculations can be observed by monitoring the dependence of the matrix elements on the size of the virtual set as well as on the number of core subshells that are opened for electron substitutions. The effects of substitutions from 4s4p4d4f and still deeper shells were estimated in our previous papers and turned out to be below 1% in the case of hyperfine structures [45,46] and a fraction of a percent in the case of transition rates [47,48].

IV. RESULTS

Table I shows the values of the reduced matrix element of the Schiff operators \( 4\sqrt{3}[P_i\|\hat{V}\|P_j]\|D_2 \) from Eq. (10). The expectation values were calculated for the multiconfiguration expansions obtained from all possible combinations of virtual sets, and from opening sequentially the core subshells, as described in Sec. III B.

| Table I | presents the values of the reduced matrix element of the EDM operator \( 3\sqrt{3}[P_i\|-e\|P_j]\|D_2 \) from Eq. (9). As in Table I, the EDM matrix elements were calculated for the multiconfiguration expansions resulting from all possible combinations of virtual sets and from opening up the core subshells (cf. Sec. III B).

The data from Tables I and II are collected also in Fig. 2, where they are presented as functions of the size \( N_{SSF} \) [see Eq. (1)] of the multiconfiguration expansion. Figure 2(a) shows the reduced matrix element of the Schiff operator, while Fig. 2(b) shows the value of reduced matrix element of EDM operator. The resultant EDM induced by the nuclear

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A. Electron correlation effects

The calculations appear to be converged within the multiconfiguration approximation employed in the present paper. A comparison of the “DF” value with the final result in Table III shows that correlation effects are dominant in these calculations. An inspection of Tables I and II reveals that the Schiff moment matrix element is mainly responsible for the correlation correction to the atomic EDM. Overall, the correlated Schiff value is more than an order of magnitude related to the uncorrelated Schiff value is more than an order of magnitude related value of the EDM matrix element is about 30% achieved if all two-body (double substitutions) and possibly also three- and four-body (triple and quadruple substitutions) correlation effects are fully taken into consideration in these calculations in order to capture the bulk of the electron correlation effects:

(1) Schiff interaction is localized inside nucleus [see Eq. (14)], therefore only the (one-electron) matrix elements \[ \langle s_{1/2} | V^\dagger \rho(r) | p_{1/2} \rangle \] and \[ \langle s_{1/2} | V^\dagger \rho(r) | p_{3/2} \rangle \] contribute appreciably in Eq. (13) (see also Table II in Ref. [17]). All other spin orbitals may contribute through the indirect electron correlation effects [49].

(2) Only the inner regions of the spin-orbitals contribute to the Schiff moment matrix elements. Therefore it is essential to accurately represent core polarization in the atomic wave functions.

(3) For the electric dipole interaction, the outer parts of the spin orbitals are important. Therefore the valence correlation effects have to be accounted for. The Babushkin and Coulomb gauge values differ by 5 orders of magnitude in the calculation of the very weak \( ^3P_{1/2} \rightarrow ^3D_{2} \) transition [48].

Transition energy is difficult to reproduce accurately in variational calculations when the wave functions for the two levels are generated separately and when their one-electron orbitals are optimized independently. It is necessary to obtain a balanced description of both states [50], which can only be achieved if all two-body (double substitutions) and possibly also three- and four-body (triple and quadruple substitutions) correlation effects are fully taken into account. In general, the two gauges exhibit different energy dependence: the Coulomb gauge is strongly energy dependent, therefore the Babushkin gauge value is usually adopted in unsaturated calculations since it is less dependent on calculated transition energy value. The calculations of energy-level differences require well balanced orbital sets and typically demand highly extensive multiconfiguration expansions. The results are usually in better agreement with experiment if a common set of orbitals is used for both states.

Schiff moment in the \( ^3D_2 \) state of the isotope Ra-223 is presented in Table III.

### Table I. The values of the reduced matrix element of the Schiff operator \( 4\sqrt{3}|P_e||r||D_2| \) from Eq. (10) (a.u.). Electron substitutions from different sets of spectroscopic orbitals: \( 6p|7s6d|7s7p \) (second column), \( 6p|7s6d|7s7p \) (third column), ... and \( 5spd6sp|7s6d|7s7p \) (sixth column). Electron substitutions to different sets of virtual orbitals: (1v) one layer of virtual orbitals, (2v) two layers of virtual orbitals, ..., and (5v) five layers of virtual orbitals. ‘DF’ = uncorrelated Dirac-Fock value.

| Virtual set | 6p | 6sp | 5d6sp | 5pd6sp | 5spd6sp |
|-------------|----|-----|-------|--------|---------|
| 1v          | 1931 | 1892 | 1862  | 1749   | 1647    |
| 2v          | 4597 | 7544 | 7553  | 7598   | 7754    |
| 3v          | 5141 | 8073 | 8112  | 8125   | 8327    |
| 4v          | 4760 | 7947 | 7889  | 8117   |         |
| 5v          | 4642 | 7834 | 7743  | 8217   |         |

### Table II. The values of the reduced matrix element of the EDM operator \( 4\sqrt{3}|P_e||r||D_2| \) from Eq. (9) (a.u.). The notations are the same as in Table I.

| Virtual set | 6p | 6sp | 5d6sp | 5pd6sp | 5spd6sp |
|-------------|----|-----|-------|--------|---------|
| 1v          | 4.818 | 4.811 | 4.810  | 4.810  | 4.810   |
| 2v          | 4.514 | 4.467 | 4.460  | 4.460  | 4.460   |
| 3v          | 4.270 | 4.192 | 4.157  | 4.155  | 4.155   |
| 4v          | 4.251 | 4.168 | 4.114  | 4.109  | 4.108   |
| 5v          | 4.240 | 4.153 | 4.083  | 4.076  | 4.075   |
(4) On the other hand, the wave functions for the two levels have to be generated separately, if the effects of non-orthogonality are to be taken into account correctly. Non-orthogonality between the spin orbitals is essential in the evaluation of nondiagonal matrix elements (see Ref. [48]).

**B. Breit and QED corrections**

The calculations described in the previous sections as well as the data in Tables I–III were obtained with only the Coulomb interaction included in the differential equations which are iteratively solved during the self-consistent-field process (SCF). The effects of magnetic and retardation corrections can be evaluated by introducing the full Breit operator in the self-consistent-field process. Such modified differential equations lead to change in the final wave function shape, which in turn modifies the matrix elements evaluated in the calculation of the Schiff moment. Quantum electrodynamics (QED) correction is dominated by two contributions. The self-energy (SE) part cannot be easily evaluated. Its contribution to the Cs parity-violation amplitude has been calculated in Ref. [51] and turned out to be of the order of −0.7%. On the other hand, the vacuum polarization (VP) can be easily calculated by introducing the Uehling potential into the SCF, as was done, for example, in calculations of Li-like ions hyperfine matrix elements [52] or in Cs parity-violation amplitude [53]. In the latter case, it leads to a 0.4% increase in the amplitude. If both are evaluated, the SE and VP contributions partly compensate. We have used the MDFGME code [40] in its 2008 version, which now includes a Schiff moment option, to evaluate the effect of the Breit interaction and vacuum polarization on the Schiff moment. The results are presented in Table IV. The inclusion of the Breit interaction in the SCF reduced the value of the Schiff moment by a factor of 1.7%, while the vacuum polarization increased it by roughly 0.6%. The total correction is a reduction in the order of 1.1%. From Ref. [51] one should expect a partial mutual compensation of the VP and SE contributions. The total correction would then be dominated by the Breit contribution.

The results presented in Table IV were obtained in the single configuration approximation, i.e., by neglecting the electron correlation effects. In the single configuration approximation the expectation values of the Schiff moment operator and of the electric dipole operator arise from the 7p–6d one-electron matrix element in Eq. (13). However, the 7p–6d amplitude contributes roughly 10% of the total value of the atomic EDM (see Tables I and III), while the dominant part comes from s–p amplitudes, through the configuration interaction mixing [17]. We made several attempts to calculate the QED corrections to the 7s–7p amplitudes, but we were unable to obtain a convergent solution for a correlated atomic wave function when Breit interaction was included in the self-consistent-field process. Our experience indicates that a fully correlated calculation, i.e., with the multiconfiguration expansion (1) described in Sec. III B and with the

| Shells opened for substitutions | 6p | 6sp | 5d6sp | 5pd6sp | 5spd6sp |
|-------------------------------|----|-----|-------|--------|--------|
| Virtual set                  |    |     |       |        |        |
| 1v                            | 0.1812 | 0.1181 | 0.1163 | 0.1092 | 0.1028 |
| 2v                            | 0.2694 | 0.4378 | 0.4374 | 0.4399 | 0.4489 |
| 3v                            | 0.2850 | 0.4393 | 0.4378 | 0.4383 | 0.4492 |
| 4v                            | 0.2627 | 0.4293 | 0.4244 | 0.4208 | 0.4329 |
| 5v                            | 0.2555 | 0.4224 | 0.4144 | 0.4097 | 0.4346 |
TABLE IV. Effect of the Breit and vacuum polarization on the reduced matrix element of the Schiff operator (Schiff), reduced matrix element of the EDM operator (Stark), and the atomic EDM \((D_{a}[10^{9}S/I])\) of the \(F,M_F=3/2,3/2\) hyperfine state of the \(^3D_2\) level induced by the nuclear Schiff moment in the isotope \(^{225}\text{Ra}\) \((I=\frac{5}{2})\) (a.u.). VP—vacuum polarization. DF—Coulomb interaction only.

| Schiff | Stark | EDM | Correction |
|--------|-------|-----|------------|
| DF     | 674   | 5.849 | 0.0512    |
| Breit SCF | 664 | 5.835 | 0.0503 | −1.7% |
| Breit SCF+VP SCF | 668 | 5.837 | 0.0507 | −1.1% |

self-consistent Breit interaction would lead to a larger (absolute value of) total correction. However, the increase in the number of extra integrals involved (roughly by 2 orders of magnitude) and substantial convergence difficulties render such a calculation virtually impossible with today’s computers. The final correction from Table IV (i.e., the reduction by a factor of 0.989%) has been carried over to the final summary of our results, which is presented in Table V. It should be noted, as explained above, that this is a correction to the \(7p-6d\) amplitude only, leaving the dominant \(7s-7p\) amplitudes unchanged. Therefore, the QED correction obtained in this manner, for the uncorrelated wave function, should be treated only as an order of magnitude estimate. This issue should be investigated in more detail in the future.

C. Accuracy

Although we have full control of the core polarization effects, the overall accuracy of these results depends primarily on the electron-correlation effects which were not included in the calculations, i.e., unrestricted double substitutions and triple substitutions. These unrestricted substitutions had to be omitted due to software and hardware limitations. A similar approximation, based on single and restricted double substitutions, was employed also in our previous papers on radium \([45–48]\). The accuracy of the present calculations can be indirectly inferred from a comparison of the previously calculated transition rates and hyperfine constants with experiment. Reference \([46]\) showed hyperfine constants calculated in an approach similar to that employed in the present paper. As discussed there, the overall accuracy of the calculated magnetic dipole constants was 6%, while for the electric quadrupole constants the estimated accuracy was 3%. The interaction responsible for the hyperfine shifts takes place in the vicinity of the nucleus. The bulk of the hyperfine integral (matrix element) come from the first oscillation of the one-electron wave function \([54,55]\). Therefore, the calculated hyperfine constants depend heavily on the inner regions of the one-electron wave functions. The interaction of the electronic cloud with the nuclear Schiff moment also takes place in the vicinity of the nucleus. In fact, the bulk of the Schiff integral (matrix element) comes from the nuclear skin [see Eq. \((14)\)]. Therefore we might expect that the accuracy of the calculated Schiff moment matrix elements is comparable to the accuracy of the calculated hyperfine constants. In conclusion, we might expect the accuracy of the calculated Schiff moment matrix elements of the order of 10% or better.

The electric-dipole moment matrix element [see Eq. \((15)\)] has similar radial dependence as the electric-dipole transition elements; therefore their accuracy might also be expected to be of the same order. The calculated rate for the strong transition \(^1P_{1/2}^-S_0\) showed excellent agreement between the values calculated in Babushkin and Coulomb gauges \([47]\), which is an indication (but not a proof) of convergence. The accuracy of the calculated rate for the \(^3P_{1/2}^-S_0\) transition was also reasonably good \([48]\). Although there was a 20% difference between the values calculated in the two gauges, the Babushkin gauge value fell within the experimental limit \([10]\). The gauge difference was somewhat larger (35%) for the calculated rate for the \(^3D_z^-S_0\) electric quadrupole transition \([47]\) which is the only significant decay channel of the metastable state \(^3D_2\). However, the calculated rates of other weak transitions were less accurate. In particular, the calculated rate for the \(^3P_{1/2}^-D_2\) transition showed very large gauge dependence, which seems to indicate that the core-core correlation effects (which were omitted in present calculations) are important (see, e.g., \([56]\)). These correlation contributions are currently beyond the capacity of the computer resources available to us [i.e., a heterogeneous set of LINUX workstations with the total number of 32 nodes working together as a parallel message passing interface (MPI) cluster]. We were unable to include or even to estimate the contributions of the omitted electron correlation effects, i.e., the unrestricted double substitutions and triple substitutions, to the atomic EDM. Such calculation would require a multiprocessor cluster in excess of a hundred processors \([57]\). In order to estimate the uncertainty of the electric dipole moment calcu-

TABLE V. The atomic EDM induced by the nuclear Schiff moment in the \(^3D_2\) electronic state for three isotopes of radium: \(^{213}\text{Ra}\) \((I=\frac{5}{2}, F=\frac{3}{2})\), \(^{223}\text{Ra}\) \((I=\frac{3}{2}, F=\frac{3}{2})\), and \(^{225}\text{Ra}\) \((I=\frac{5}{2}, F=\frac{3}{2})\), respectively. Results are shown as functions of the size of the virtual orbital set, including electron substitutions from the spectroscopic \(5sp6dp7p\) orbitals to: (1v) one layer of virtual orbitals, (2v) two layers of virtual orbitals, …, and (5v) five layers of virtual orbitals. ‘DF’=uncorrelated Dirac-Fock value. Our values include the “Breit SCF + VP SCF” correction from Table IV. The “RHF + CI” results in the last line are quoted from Ref. \([17]\).

| Layer | \(^{213}\text{Ra}\) | \(^{223}\text{Ra}\) | \(^{225}\text{Ra}\) |
|-------|----------------|----------------|----------------|
| DF    | 0.0159         | 0.0507         | 0.0158         |
| 1v    | 0.0320         | 0.1017         | 0.0318         |
| 2v    | 0.1393         | 0.4441         | 0.1387         |
| 3v    | 0.1394         | 0.4444         | 0.1388         |
| 4v    | 0.1344         | 0.4283         | 0.1338         |
| 5v    | 0.1349         | 0.4300         | 0.1343         |
| Final | 0.13(4)        | 0.43(14)       | 0.13(4)        |
| RHF+CI| 0.094          | 0.30           | 0.094          |
lation, we had to resort to an upper limit resulting from the hierarchy of electron-electron interactions. Table II demonstrates the saturation of the core-valence correlation correction, which is the dominant electron correlation effect beyond the Dirac-Fock approximation. Therefore we might expect that the entire (core-valence) electron correlation contribution would be a rather conservative estimate of the uncertainty of the electric dipole moment calculations. In conclusion, we may assume the accuracy of the calculated electric dipole moment matrix elements of the order of 30%.

Another contribution that affects the overall accuracy of the results arises from the nuclear-density dependence of radial matrix elements in Eq. (14), i.e., from the distribution of the nuclear charge, as discussed in Sec. II B. However, the error of the calculated value of atomic EDM is dominated by the electric dipole moment calculation. Therefore we assumed 30% accuracy of the final calculated value of the Schiff moment enhancement factor, as shown in Table V.

A more stringent accuracy assessment for the present calculations would require more experimental data to compare with. Hereby, we would like to encourage the experimenters to provide them. In particular, hyperfine constants for other levels, as well as lifetime and transition rate measurements for weak transitions would be very valuable.

The results presented in the second and in the last column of Table V illustrate the isotope dependence of the atomic EDM. The two isotopes in question, 223Ra and 225Ra, have the same nuclear spin (I = 1/2), but they have slightly different nuclear shapes [see Eq. (3) and the subsequent discussion in Sec. II B]. The above-mentioned isotope dependence, of the order of 0.4%, arises primarily through the derivative dp/dr in Eq. (14). The isotope dependence of the atomic wave functions [Eq. (1)] has been neglected in the present calculations.

V. CONCLUSIONS

Radium is well suited for EDM experiments because of its large nuclear charge Z and the two low-lying levels 3P1 and 3D2 of opposite parity, which are separated in energy by only 5 cm−1 = 2 × 10−5 a.u. The 3D2 level is metastable with a very long lifetime of the order of 4 s [48] and therefore suitable for laser-ion traps. For radium, moreover, there are several isotopes with mass ranging from 209 up to 229, of which 213Ra and 225Ra have nuclear spin I = 1/2. For EDM experiments the I = 1/2 isotopes are preferable since these isotopes cannot be disturbed by higher order electromagnetic moments.

The paper presents systematic computations of the static EDM of atomic radium in the 3D2 level induced by the nuclear Schiff moment. Table V shows the atomic EDM induced by the nuclear Schiff moment in the 3D2 electronic state for three isotopes of radium:

\[ ^{213}_{88} \text{Ra} \ (I = \frac{1}{2}, F = \frac{3}{2}, \mu = 0.6133), \]

\[ ^{223}_{88} \text{Ra} \ (I = \frac{3}{2}, F = \frac{3}{2}, \mu = 0.2705), \]

and

\[ ^{225}_{88} \text{Ra} \ (I = \frac{1}{2}, F = \frac{3}{2}, \mu = -0.7338). \]

The wave functions for the two levels, 3P1 and 3D2, were generated separately, in order to correctly reproduce the effects of nonorthogonality between one-electron spin orbitals. We demonstrate that core-valence electron correlation, which is the dominant electron correlation effect beyond the Dirac-Fock approximation, contributes almost 90% of the total EDM value. Our final value is about 30% larger than the relativistic Hartree-Fock and configuration interaction (RHF+CI) result from Ref. [17]. The difference can be attributed to different methods employed to account for core-valence electron correlation effects. Neither of these two calculations included the core-core correlation effects, which we believe to be the dominant source of uncertainty in our calculated value of the Schiff moment enhancement factor.

ACKNOWLEDGMENTS

This work was supported by the Polish Ministry of Science and Higher Education (MNiSW) in the framework of the scientific Grant No. 1P03B11030. S.F. acknowledges the support by the DFG under the Project No. FR 1251/13. E.G. acknowledges support from the Lithuanian Science Council. P.I. acknowledges the support from the Swedish Research Council (Vetenskapsrådet). P.I. acknowledges the support of the Helmholz Alliance Program of the Helmholtz Association Contract No. HA-216, “Extremes of Density and Temperature: Cosmic Matter in the Laboratory.” Laboratoire Kastler Brossel is “Unité Mixte de Recherche du CNRS, de l’ENS et de l’UPMC no 8552.” We would like to thank Klaus Jungmann, Jeffrey Guest, William Trumble, and Jean-Paul Desclaux for helpful discussions. We thank the (anonymous) referee for pointing our attention to the fact that our calculated QED correction arisen from the 7p-6d amplitude and therefore represented less than 10% of the total correction.

[1] A.-M. Mårtensson-Pendrill, in Methods in Computational Chemistry, edited by S. Wilson (Plenum Press, New York, 1992), Vol. 5, pp. 99–156.
[2] I. B. Khriplovich and S. K. Lamoreaux, CP Violation Without Strangeness (Springer, Berlin, 1997).
[3] J. S. M. Ginges and V. V. Flambaum, Phys. Rep. 397, 63 (2004).
[4] T. Legero, T. Wilk, M. Henrich, G. Rempe, and A. Kuhn, Phys. Rev. Lett. 93, 070503 (2004).
[5] S. Kuhn, W. Alt, D. Schrader, I. Dotsenko, Y. Miroshnichenko, A. Rauschenbeutel, and D. Meschede, Phys. Rev. A 72, 023406 (2005).
[6] J. Beugnon, M. P. A. Jones, I. Dingjan, B. Darquié, G. Messin, A. Browaeys, and P. Grangier, Nature (London) 440, 779.
(2006).
[7] J. R. Guest, N. D. Sciellzo, I. Ahmad, K. Bailey, J. P. Greene, R. J. Holt, Z.-T. Lu, T. P. O'Connor, and D. H. Potterveld, Phys. Rev. Lett. 98, 093001 (2007).
[8] U. Dammalapati, S. De, K. Jungmann, and L. Willmann, Phys. Rev. A 79, 041402 (2009).
[9] V. A. Dzuba, V. V. Flambaum, J. S. M. Ginges, and M. G. Kozlov, Phys. Rev. A 66, 012111 (2002).
[10] N. D. Sciellzo, J. R. Guest, E. C. Schulte, I. Ahmad, K. Bailey, D. L. Bowers, R. J. Holt, Z.-T. Lu, T. P. O'Connor, and D. H. Potterveld, Phys. Rev. A 73, 010501(R) (2006).
[11] www-mep.phy.anl.gov/atta/research/radiumedm.html
[12] K. Jungmann, Acta Phys. Pol. B 33, 2049 (2002).
[13] K. Jungmann (private communication).
[14] www.kvi.nl/~trimp/web/html/trimp.html
[15] J. Engel, M. Bender, J. Dobaczewski, J. H. de Jesus, and P. Olbratowski, Phys. Rev. C 68, 025501 (2003).
[16] J. Dobaczewski and J. Engel, Phys. Rev. Lett. 94, 232502 (2005).
[17] V. A. Dzuba, V. V. Flambaum, and J. S. M. Ginges, Phys. Rev. A 61, 062509 (2000).
[18] V. V. Flambaum, Phys. Rev. A 60, R2611 (1999).
[19] J. Guest (private communication).
[20] W. Trimble (private communication).
[21] G. Gaigalas, E. Gaidamauskas, and P. Jönsson, J. Phys.: Conf. Ser. 130, 012008 (2008).
[22] I. Lindgren and J. Morrison, Atomic Many-Body Theory (Springer, Berlin, 1996).
[23] V. A. Dzuba, Phys. Rev. A 71, 032512 (2005).
[24] V. A. Dzuba and V. V. Flambaum, Phys. Rev. A 75, 052504 (2007).
[25] W. R. Johnson, Atomic Structure Theory: Lectures on Atomic Physics (Springer, Berlin, 2007).
[26] I. P. Grant, Relativistic Atomic Structure Calculations, Methods in Computational Chemistry Vol. 2, edited by S. Wilson (Plenum, New York, 1988).
[27] I. P. Grant, Relativistic Quantum Theory of Atoms and Molecules: Theory and Computation (Springer, New York, 2007).
[28] P. Jönsson, X. He, C. Froese Fischer, and I. P. Grant, Comput. Phys. Commun. 177, 597 (2007).
[29] V. A. Dzuba, V. V. Flambaum, and M. G. Kozlov, Phys. Rev. A 54, 3948 (1996).
[30] M. G. Kozlov, S. G. Porsev, and W. R. Johnson, Phys. Rev. A 64, 052107 (2001).
[31] M. G. Kozlov, Int. J. Quantum Chem. 100, 336 (2004).
[32] I. P. Grant, Comput. Phys. Commun. 84, 59 (1994).
[33] J. Bieroń, P. Jönsson, and C. Froese Fischer, Phys. Rev. A 53, 2181 (1996).
[34] J. Bieroń, P. Jönsson, and C. Froese Fischer, Phys. Rev. A 60, 3547 (1999).
[35] K. G. Dyall, I. P. Grant, C. T. Johnson, F. A. Parpia, and E. P. Plummer, Comput. Phys. Commun. 55, 425 (1989).
[36] W. R. Johnson and G. Soff, At. Data Nucl. Data Tables 33, 405 (1985).
[37] http://dirac.chem.sdu.dk/doc/FiniteNuclei/Finite Nuclei.shtml
[38] This approach is well justified since each nondiagonal term in Eq. (4) leads to an additional summation in expansion (6) \( \Sigma_a c_a \Psi(\cdots P J \cdots) \), with coefficients of the order of \( 10^{-5} \cdots 10^{-10} \), owing to weakness of the hyperfine interaction. These coefficients are to be compared with \( a \approx 1 \) in Eq. (6), and, hence, they strongly suppress the nondiagonal terms of the hyperfine structure.
[39] G. Gaigalas, S. Fritzsche, and I. P. Grant, Comput. Phys. Commun. 139, 263 (2001).
[40] P. Indelicato and J.-P. Desclaux, MCDGFME, a MultiConfiguration Dirac-Fock and General Matrix Elements program (release 2005), URL http://dirac.spectro.jussieu.fr/mcfd
[41] C. E. Moore, Atomic Energy Levels, NSRDS-NBS No. 35 (US Government Printing Office, Washington, DC, 1971).
[42] P. Raghavan, At. Data Nucl. Data Tables 42, 189 (1989).
[43] P. Å. Malmqvist, Int. J. Quantum Chem. 30, 479 (1986).
[44] J. Olsen, M. R. Godfrey, P. Jönsson, P. Å. Malmqvist, and C. Froese Fischer, Phys. Rev. E 52, 4499 (1995).
[45] J. Bieroń and P. Pykkö, Phys. Rev. A 71, 032502 (2005).
[46] J. Bieroń, J. Phys. B 38, 2221 (2005).
[47] J. Bieroń, C. Froese Fischer, S. Fritzsche, and K. Pachucki, J. Phys. B 37, L305 (2004).
[48] J. Bieroń, P. Indelicato, and P. Jönsson, Eur. Phys. J. Spec. Top. 144, 75 (2007).
[49] J. Bieroń, C. Froese Fischer, P. Jönsson, and P. Pykkö, J. Phys. B 41, 115002 (2008).
[50] C. Froese Fischer, Phys. Scr. T83, 49 (1999).
[51] V. M. Shabaev, K. Pachucki, I. I. Tupitsyn, and V. A. Yerokhin, Phys. Rev. Lett. 94, 213002 (2005).
[52] S. Boucard and P. Indelicato, Eur. Phys. J. D 8, 59 (2000).
[53] W. R. Johnson, I. Bednyakov, and G. Soff, Phys. Rev. Lett. 87, 233001 (2001).
[54] P. Pykkö, J. Magn. Reson. (1969-1992) 8, 15 (1972).
[55] P. Pykkö, E. Pajanne, and M. Inokuti, Int. J. Quantum Chem. 7, 785 (1973).
[56] P. Jönsson, A. Ynnerman, C. Froese Fischer, M. R. Godfrey, and J. Olsen, Phys. Rev. A 53, 4021 (1996).
[57] J. Bieroń, C. Froese Fischer, P. Indelicato, P. Jönsson, and P. Pykkö, Phys. Rev. A 79, 052502 (2009).