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Dynamic polarizabilities and related properties of clock states of the ytterbium atom

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Abstract

We carry out relativistic many-body calculations of the static and dynamic dipole polarizabilities of the ground $6s^{2} {}^{1}S_{0}$ and the first excited $6s6p {}^{3}P_{0}^{0}$ states of Yb. With these polarizabilities, we compute several properties of Yb relevant to optical lattice clocks operating on the $6s^{2} {}^{1}S_{0}$ - $6s6p {}^{3}P_{0}^{0}$ transition. We determine (i) the first four *magic* wavelengths of the laser field for which the frequency of the clock transition is insensitive to the laser intensity. While the first magic wavelength is known, we predict the second, the third and the fourth magic wavelengths to be 551 nm, 465 nm and 413 nm. (ii) We re-evaluate the effect of black-body radiation on the frequency of the clock transition, the resulting clock shift at T = 300 K being -1.41(17) Hz. (iii) We compute long-range interatomic van der Waals coefficients (in a.u.) $C_{6}(6s^{2} {}^{1}S_{0} + 6s^{2} {}^{1}S_{0}) = 1909(160)$, $C_{6}(6s^{2} {}^{1}S_{0} + 6s6p {}^{3}P_{0}) = 2709(338)$ and $C_{6}(6s6p {}^{3}P_{0} + 6s6p {}^{3}P_{0}) = 3.886(360)$. Finally, we determine the atom-wall interaction coefficients (in a.u.), $C_{3}(6s^{2} {}^{1}S_{0}) = 3.34$ and $C_{3}(6s6p {}^{3}P_{0}) = 3.68$. We also address and resolve a disagreement between previous calculations of the static polarizability of the ground state.

1. Introduction

The ytterbium atom is employed in a number of projects aimed at studying fundamental problems of modern physics by means of atomic physics. A large parity-violating signal on the $6s^{2} {}^{1}S_{0}$ -6s5d ${}^{3}D_{1}$ 408 nm transition of Yb has been recently observed at Berkeley [1]. A search for the CP-violating (T-, Podd) permanent electric dipole moment of Yb was initiated [2]. A Bose–Einstein condensate in a dilute gas of the ground-state Yb atoms has been attained [3] and a number of experiments with ultracold Yb atoms have been carried out (see, e.g., [4–6]).

Here we focus on the optical lattice clock applications of Yb. In optical lattice clocks, neutral atoms are trapped in a standing wave of laser light (optical lattice) operated at a certain 'magic' wavelength [7]. At this wavelength a differential light perturbation of the two clock levels vanishes exactly. Using Yb atom in lattice clocks was proposed in [8] and experimentally demonstrated in Boulder [9, 10]. Apart from numerous technical applications, such atomic clocks may be used for searching for a variation of fundamental constants [11], mapping out atom–wall interaction [12] and serve as a basis for quantum information processing (see, e.g., [13, 14]).

In the present paper, we evaluate dynamic electric-dipole polarizabilities of the clock states, $6s^{2} {}^{1}S_{0}$ and $6s6p {}^{3}P_{0}^{0}$. Dynamic polarizabilities of real and imaginary frequencies are linked to a variety of important atomic and molecular properties. In particular, we compute magic wavelengths, the effect of black-body radiation on the frequency of the clock transition, atom–wall interaction constants C_{3} and finally, long-range molecular van der Waals coefficients C_{6} for the clock states. We take advantage of a close link between these properties and improve our theoretical predictions by referencing to measured properties, such as the first magic wavelength [9, 15] and the van der Waals coefficient for two ground-state atoms [5].

Ytterbium has been studied both experimentally (see, e.g., [5, 6, 9, 10, 15–18]) and theoretically (see, e.g., [8, 19–26]. Trapping and cooling schemes were discussed and implemented, the frequency of the clock transition was measured to a high accuracy [10], the first magic wavelength

[9, 15] and the van der Waals coefficient for the ground state of Yb were also measured [5].

Theoretical studies are not free from controversy. The calculated polarizability of the Yb ground state reported in [20] is in disagreement with most other *ab initio* or semi-empirical analyses. The source of this disagreement is explained below and the discrepancy is resolved. The resolution is closely related to the role of the excitations from the core which are discussed in detail. The corresponding statement can be formulated as a general theorem: *Consider a second-order transition matrix element, involving summation over a complete set of intermediate states. Then, a contribution from a subspace spanned by degenerate states does not depend on mixing of these states.* This statement has implications not only for computing polarizabilities, but also for the two-photon transition amplitudes, parity-violating amplitudes, and other properties in Yb and other atoms.

2. Method of calculation

In this section, we start by recapitulating expressions for dynamic polarizability and its relations to magic wavelengths, and atom-atom and atom-wall interaction coefficients. Next we discuss our computational relativistic many-body method.

The ac Stark shift of an energy of a spherically symmetric, J = 0, atomic state v in electro-magnetic wave of amplitude ε and frequency ω is given by

$$\Delta E_{v}(\omega) = -\alpha_{v}(\omega) \left(\frac{\varepsilon}{2}\right)^{2}, \qquad (1)$$

where the electric-dipole dynamic polarizability reads (we use atomic units in which $\hbar = 1$, |e| = 1 and $m_e = 1$)

$$\alpha_{v}(\omega) = \frac{2}{3} \sum_{k} \frac{E_{k} - E_{v}}{(E_{k} - E_{v})^{2} - \omega^{2}} \langle v \| \mathbf{d} \| k \rangle^{2} \,. \tag{2}$$

Here **d** is an electric dipole operator and summation is over a complete set of many-body states.

The *magic* laser frequency ω^* is found from the condition that the ac Stark shifts of both clock states v and w are identical so that the frequency of the transition does not depend on the laser intensity,

$$\Delta E_{v}(\omega) - \Delta E_{w}(\omega) = -(\alpha_{v}(\omega) - \alpha_{w}(\omega)) \left(\frac{\varepsilon}{2}\right)^{2} = 0. \quad (3)$$

Simply, at the magic frequencies, $\alpha_v(\omega^*) = \alpha_w(\omega^*)$.

The clock frequency shift due to black-body radiation is expressed in terms of the difference of static polarizabilities of the two clock levels (see detailed discussion in [21]).

We also require atomic polarizabilities evaluated at purely imaginary frequencies

$$\alpha_{v}(i\omega) = \frac{2}{3} \sum_{k} \frac{E_{k} - E_{v}}{(E_{k} - E_{v})^{2} + \omega^{2}} \langle v \| \mathbf{d} \| k \rangle^{2}.$$
(4)

These are used in evaluating the atom–wall interaction constant C_3 [27]

$$C_3 = \frac{1}{4\pi} \int_0^\infty \alpha(i\omega) \,\mathrm{d}\omega,\tag{5}$$

and the van der Waals coefficients C_6 . For two separated atoms in states w and v,

$$C_6(w+v) = \frac{3}{\pi} \int_0^\infty \alpha_w(i\omega) \alpha_v(i\omega) \,\mathrm{d}\omega. \tag{6}$$

The C_6 for two atoms interacting in identical states is subsumed by the above formula. Equation (6) is derived under assumption that there are no downwards dipole transitions from either w or v state, which holds in our case.

It is clear that for computing the polarizabilities, we need to determine electric dipoles and energies, and to perform the summation over a complete set of atomic states. To this end, we use a combination of the configuration-interaction (CI) method and the many-body perturbation theory (MBPT) to construct an effective Hamiltonian for two valence electrons (the CI+MBPT method [28, 29]). Further, we employ the Dalgarno–Lewis method [30] to reduce the summation over a complete set of many-electron states to solving an inhomogeneous system of linear equations.

2.1. CI+MBPT method

The ground-state configuration of Yb is $4f^{14}6s^2$ and most of the low-energy excited states correspond to configurations with one of the 6s electrons being excited. All these states can be represented to a relatively high accuracy as states with two valence electrons above a closed-shell core. This is the starting point of our calculations: the computational (CI) model space is spanned by all possible excitations of the two valence electrons. Implications arising from the states with excitations from the outer 4f core subshell will be discussed below.

The effective CI+MBPT Hamiltonian for two valence electrons has the form

$$\hat{H}^{\text{eff}} = \hat{h}_1(r_1) + \hat{h}_1(r_2) + \hat{h}_2(r_1, r_2),$$
(7)

where \hat{h}_1 is the single-electron part of the relativistic Hamiltonian

$$\hat{h}_1 = c\hat{\boldsymbol{\alpha}} \cdot \mathbf{p} + (\hat{\boldsymbol{\beta}} - 1)m_e c^2 - \frac{Ze^2}{r} + V^{N-2} + \hat{\boldsymbol{\Sigma}}_1 \qquad (8)$$

and \hat{h}_2 is the two-electron part of the Hamiltonian

$$\hat{h}_2(r_1, r_2) = \frac{e^2}{|\mathbf{r}_1 - \mathbf{r}_2|} + \hat{\Sigma}_2(r_1, r_2).$$
(9)

In these equations, $\hat{\alpha}$ and $\hat{\beta}$ are the conventional Dirac matrices, V^{N-2} is the Dirac–Hartree–Fock (DHF) potential of the closed-shell atomic core (N-2=68, Z=70) and $\hat{\Sigma}$ is the correlation operator. It represents terms in the Hamiltonian arising due to virtual excitations from atomic core (see [28, 29] for details). $\hat{\Sigma} \equiv 0$ corresponds to the standard CI method. $\hat{\Sigma}_1$ is a single-electron operator. It represents a correlation interaction (core-polarization) of a particular valence electron with the atomic core. $\hat{\Sigma}_2$ is a two-electron operator. It represents screening of the Coulomb interaction between the two valence electrons by the core electrons. We calculate $\hat{\Sigma}$ in the second order of the MBPT. We use a B-spline technique [31] to construct a pseudospectrum. We use 40 B-splines in a cavity of radius $R = 40 a_B$ and calculate the eigenstates

Table 1. Energy levels of Yb (cm⁻¹). The 4f¹³5d6s²(7/2, 5/2)^o₁ level lies outside of the computational model space.

				CI-	+MBPT
State		J	Exp. [32]	ab initio	Rescaled $\hat{\Sigma}_1$
$4f^{14}6s^2$	1 S	0	0.0	0.0	0.0
4f ¹⁴ 6s6p	${}^{3}P^{o}$	0	17 288	18246	17 289
		1	17 992	18946	17 996
		2	19710	20688	19759
4f ¹⁴ 5d6s	^{3}D	1	24 489	24 922	24 489
		2	24752	25 195	24 7 4 3
		3	25 27 1	25 765	25 276
4f ¹⁴ 6s6p	$^{1}\mathbf{P}^{o}$	1	25 068	26463	25611
$4f^{13}5d6s^2$	(7/2, 5/2)°	1	28 857		
4f ¹⁴ 5d6s	^{1}D	2	27 678	28 4 8 5	27 812
4f ¹⁴ 6s7s	³ S	1	32 695	33 262	32 668
4f ¹⁴ 6s7s	${}^{1}S$	0	34 351	34 871	34 280

of the V^{N-2} DHF Hamiltonian up to the maximum value of the angular momentum $l_{\text{max}} = 5$. The same basis is used in computing $\hat{\Sigma}$ and in constructing the two-electron states for the valence electrons. 30 out of 40 lowest-energy states for every *l* up to $l_{\text{max}} = 5$ are used to calculate $\hat{\Sigma}$ and 14 lowest states above the core are used for every *l* up to $l_{\text{max}} = 4$ to construct the two-electron states.

The two-electron valence states are found by solving the eigenvalue problem,

$$\hat{H}^{\rm eff}\Psi_v = E_v\Psi_v\,,\tag{10}$$

using the standard CI techniques. Calculated and experimental energies of a few lowest-energy states of Yb are presented in table 1. One can see that the pure *ab initio* energies are already close to the experimental values. However, for improving the accuracy further, we re-scale the correlation operator $\hat{\Sigma}_1$ by replacing $\hat{\Sigma}_1$ in the effective Hamiltonian (7) in each partial wave s, $p_{1/2}$, $p_{3/2}$, ... by $f_a \hat{\Sigma}_1$. The rescaling factors are $f_s = 0.8772$, $f_p = 1.03$, $f_d = 0.933$ and $f_f = 1$. These values are chosen to fit the experimental spectrum of Yb. The result of the fitting is shown in table 1. Note that the fitting is not exact for all the levels because the number of levels is larger than the number of fitting parameters.

Transition amplitudes are found with the random-phase approximation (RPA) [33, 34]

$$E1_{vw} = \langle \Psi_v \| d_z + \delta V^{N-2} \| \Psi_w \rangle, \tag{11}$$

where δV^{N-2} is the correction to the core potential due to core polarization by an external electric field. We compile our representative dipole matrix elements in table 2. The values were computed in the CI+MBPT approach. We also compare with a high-precision result [16] derived from fitting molecular long-range potentials to photoassociation spectra taken with ultracold samples. The 16% theory–experiment disagreement for the 6s² ¹S₀–6s6p ¹P₁^o amplitude is due to mixing of the 6s6p ¹P₁^o state with the core-excited state 4f¹³5d6s²(7/2, 5/2)₁^o, which lies outside the computational model space (see section 3.)

Table 2. Reduced matrix elements (a.u.) of the electric dipole operator for transitions in Yb. Values were computed in the CI+MBPT approach. The theory–experiment disagreement for the $6s^{2} \, {}^{1}S_{0}$ – $6s6p^{1}P_{1}^{o}$ is due to mixing of the $6s6p^{1}P_{1}^{o}$ state with the core-excited state $4f^{13}5d6s^{2}(7/2, 5/2)_{1}^{o}$, which lies outside the computational model space (see section 3.)

Transition	Calculated	Experimental
$6s^{2} {}^{1}S_{0} - 6s6p {}^{3}P_{1}^{0}$	0.587	
$6s^{2} {}^{1}S_{0} - 6s6p {}^{1}P_{1}^{0}$	4.825	$4.148(2)^{a}$
$6s6p^{3}P_{0}^{o}-5d6s^{3}D_{1}$	2.911	
$6s6p^{3}P_{0}^{\circ}-6s7s^{1}S_{0}$	1.952	

^a Reference [16].

2.2. Dalgarno-Lewis and RPA methods

Computing polarizability requires summing over a complete set of *many-body* states. The CI+MBPT subspace of two valence electrons covers only a part of this set. The other class of the intermediate state are core excitations, where the valence electrons serve as spectators. Note that even in the independent particle picture, the state of the valence electrons affects possible core excitations: core electrons cannot be promoted into excited orbitals occupied by the valence electrons due to the Pauli exclusion principle. Following this discussion, we divide the polarizability into three parts [27]:

$$\alpha(\omega) = \alpha_{\rm val}(\omega) + \alpha_{\rm core}(\omega) + \alpha_{\rm core-val}(\omega).$$

The valence contribution, $\alpha_{val}(\omega)$, is given by (2) where vand k are two-electron CI states (i.e., k lie entirely in the model space). We use the Dalgarno–Lewis method [30] for summing over a complete set of two-electron states. In this method, a correction $\delta \Psi_v$ to the two-electron wavefunction of the state v is introduced and the contribution of the valence electrons to the polarizability is expressed as (here we consider static polarizability)

$$\alpha_{\rm val}(0) = \frac{2}{3} \langle \delta \Psi_v \| \mathbf{d} \| \Psi_v \rangle. \tag{12}$$

The correction $\delta \Psi_v$ is found by solving the system of linear inhomogeneous equations

$$(\hat{H}^{\text{eff}} - E_v)\delta\Psi_v = -(d_z + \delta V^{N-2})\Psi_v.$$
(13)

In the case of dynamic polarizabilities we employ the identity

$$\frac{E_k - E_v}{(E_k - E_v)^2 - \omega^2} = \frac{1}{(E_k - E_v) - \omega} + \frac{1}{(E_k - E_v) + \omega}$$

i.e., equations (13) need to be solved twice at two different energies $E = E_v \pm \omega$. For polarizabilities of purely imaginary argument, $\omega \rightarrow i\omega$ in the above equations.

The core polarizability is found using the RPA method [35] (linearized response theory). This method yields required eigen-energies of many-body particle-hole excitations from the core and relevant electric-dipole transition amplitudes. The dynamic core polarizability reads

$$\alpha_{\rm core}(i\omega) = \sum_{\omega_{\mu}>0} \frac{f_{\mu}}{(\omega_{\mu})^2 + \omega^2}.$$
 (14)

Here the summation is over particle–hole excitations from the ground state of the atomic core; ω_{μ} are excitation energies and f_{μ} are the corresponding electric-dipole oscillator strengths.

Technically, compared to the original, differential-equation method of [35], we use a B-spline basis set expansion. An important property of the RPA core polarizability is that it satisfies (non-relativistically) the Thomas–Reiche–Kuhn sum rule, $\lim_{\omega\to\infty} \omega^2 \alpha_{core}(i\omega) = N-2$. This property is especially important in evaluating the C_3 and C_6 coefficients for heavy atoms [27].

Finally, the core–valence counter term, $\alpha_{core-val}(\omega)$, is small. At the DHF level, we find that the relevant contributions to static polarizabilities are below 1% and we neglect it at our present level of accuracy.

3. The role of the $4f^{14}6s6p \ ^{1}P_{1}^{0}-4f^{13}5d6s^{2}(7/2, 5/2)_{1}^{0}$ mixing

There is a number of relatively low-lying states in the discrete spectrum of Yb which have an excitation from the outer 4f subshell of the core. These states certainly cannot be considered as states with two valence electrons above closed shells. One of such states is $4f^{13}5d6s^2(7/2, 5/2)_1^{\circ}$ which lies close to the $4f^{14}6s6p$ $^{1}P_1^{\circ}$ state (see table 1). Note that both states have the same total angular momentum and parity. Therefore the Coulomb interaction strongly mixes the two states. This mixing complicates the calculations. Our effective Hamiltonian (7) does take into account excitations from the core via second-order correlation operator $\hat{\Sigma}$. Yet, the second-order approximation is not adequate in this particular case, as the two states are separated by mere 0.0173 a.u. (3789 cm⁻¹).

The calculation of the ground-state polarizability is affected by the $4f^{14}6s6p \, {}^{1}P_{1}^{o}-4f^{13}5d6s^{2}6(7/2, 5/2)_{1}^{o}$ mixing. The $6s^{2} \, {}^{1}S_{0} \rightarrow 6s6p \, {}^{1}P_{1}^{o}$ electric dipole transition contributes about 90% to the polarizability of the ground state. Corresponding theoretical CI+MBPT transition amplitude, 4.825 a.u., deviates significantly from the experimental value 4.148 a.u. [16]. As was pointed out in [19] the reason for this disagreement is the above-mentioned mixing.

Calculation of the core polarizability displays a similar dependence on the mixing. Our RPA result for $\alpha_{core}(0) = 6.39$ a.u. At the same time, semi-empirical result [24] based on the lifetime measurements of low-lying core-excited states in Yb suggests a much higher value of 24(4) a.u. Again, the reason for this large difference lies in the strong mixing of the 4f¹³5d6s²(7/2, 5/2)^o₁ and 4f¹⁴6s6p ¹P^o₁ states. Indeed, in the RPA core polarizability calculations, we consider a different system, Yb²⁺, which is unaware of the presence of the two valence electrons.

Since the $6s^{2} {}^{1}S_{0} \rightarrow 6s6p {}^{1}P_{0}^{0}$ transition contributes about 90% to the polarizability of the ground state, the authors of [20] suggested to replace the CI+MBPT dipole matrix element by its experimental value. This was made to improve the accuracy of computing the polarizability. Below we argue that such a substitution cannot be justified unless a similar correction is done for the $4f_{7/2} \rightarrow 5d_{5/2}$ transition amplitude in the calculation of the *core* polarizability.

Let us use the following notation for the pure (unmixed) states:

$$\Phi_1 = |4f^{14}6s6p \ ^1P_1^o\rangle,$$

$$\Phi_2 = |4f^{13}5d6s^2 \ (7/2, 5/2)_1^o\rangle.$$

The mixed states can be written as

$$\Psi_1 = \Phi_1 \cos \phi - \Phi_2 \sin \phi,$$

$$\Psi_2 = \Phi_1 \sin \phi + \Phi_2 \cos \phi,$$

where ϕ is the mixing angle.

Let also use $\Psi_0 \equiv \Phi_0$ for the ground-state wavefunction.

If we neglect a small energy difference between the states Ψ_1 and Ψ_2 (the relevant energy denominators entering $\alpha(0)$ differ by about 10%) then their total contribution to the polarizability of the ground state is proportional to

$$\langle \Psi_0 \| \mathbf{d} \| \Psi_1 \rangle^2 + \langle \Psi_0 \| \mathbf{d} \| \Psi_2 \rangle^2 = \langle \Phi_0 \| \mathbf{d} \| \Phi_1 \rangle^2 + \langle \Phi_0 \| \mathbf{d} \| \Phi_2 \rangle^2$$
(15)

and does not depend on the mixing angle ϕ . This means that either bases $\{\Phi_0, \Phi_1, \Phi_2\}$ or $\{\Psi_0, \Psi_1, \Psi_2\}$ can be used in the calculations. The ab initio calculations within the CI+MBPT framework in which Yb is considered as a two-valenceelectrons atom use the non-mixed basis, while the analysis based on experimental data naturally corresponds to the use of the mixed basis. Both approaches produce close results as will be demonstrated in section 5. By contrast, the substitution of the experimental value for the $(6s^{2} {}^{1}S_{0}|\mathbf{D}|6s6p {}^{1}P_{1}^{o})$ transition amplitude is equivalent to using the mixed non-orthogonal basis { Ψ_0 , Ψ_1 , Φ_2 }. This seems to introduce a large error into the calculation of the polarizability. Indeed, the value reported in [20, 21] is 111.3 a.u. which is about 30% smaller than the results of most ab initio calculations and semi-empirical analyses, compiled in table 3. For example, the analysis [24] of the experimental data yielded $\alpha_{1S_0}(0) = 136.4(4.0)$ a.u.

The result of [20, 21] is brought into an agreement with that of [24] by consistently correcting the core polarizability for the mixing. A DHF contribution from the 4f subshell to core polarizability is 1.9 a.u. If we replace this value by the value [24] based on experimental data (i.e., account for the mixing), 24(4) then the result of [20] increases to $\alpha_{1S_0}(0) = 133$ a.u. consistent with the value of 136.4(4.0) a.u. from [24].

It is useful to find out what happens if the difference in energies of the two mixed states is not neglected. We will use a shorthand notation $\Delta E_1 = E(\Psi_1) - E(\Psi_0)$, $\Delta E_2 = E(\Psi_2) - E(\Psi_0)$ and $\delta = 1/\Delta E_2 - 1/\Delta E_1$. Since the state $\Psi_2 \equiv 4f^{13}5d6s^26(7/2, 5/2)_1^{\circ}$ is above the state $\Psi_1 \equiv 4f^{14}6s6p^1P_1^{\circ}$ the parameter δ is negative. Further, $A_{01} = \langle \Phi_0 \| \mathbf{d} \| \Phi_1 \rangle$ and $A_{02} = \langle \Phi_0 \| \mathbf{d} \| \Phi_2 \rangle$.

A partial contribution of the two states into the polarizability reads

$$\delta \alpha = \frac{1}{\Delta E_1} \left(A_{01}^2 + A_{02}^2 \right) + \delta (A_{01} \sin \phi + A_{02} \cos \phi)^2.$$
(16)

We see that when the core-excited states lie above the states with which they are mixed, the correction to the polarizability due to this mixing is negative. This is the case for both polarizabilities of Yb considered in this work. We keep this in mind while analysing the accuracy of the calculations in section 5.

Table 3. Static electric-dipole polarizability $\alpha(0)$ (in a.u.) of the $6s^{2} {}^{1}S_{0}$ and $6s6p {}^{3}P_{0}^{\circ}$ clock states of a Yb atom in various approximations and comparison with literature values.

$\alpha(0), {}^{1}S_{0}$	$\alpha(0), {}^{3}P_{0}^{o}$	Comment
This work		
138.9	315.9	ab initio with no fitting
145.7	301.6	Energies are fitted by rescaling $\hat{\Sigma}_1$
139.1		With a correction to $\alpha({}^{1}S_{0})$ to fit the magic frequency $\omega = 0.0600$ a.u.
135.2		Using experimental data for dominant contributions
138.8		Rescaled to fit experimental data[5] for $C_6 = 1932(30)$ a.u.
141(6)	302(14)	Final
other		
131.6		Wang <i>et al</i> [37]; density functional theory, 1995
145.3		Wang and Dolg [38]; CCSD(T), 1998
141.7		Miller [39]; density functional theory, 2002
118(45)	252(25)	Porsev et al [19]; CI+MBPT, 1999
111.3(5)	266(15)	Porsev and Derevianko [20, 21]; CI+MBPT, 2006
154.7		Buchachenko <i>et al</i> [40]; CCSD(T), 2006
136.4(4.0)		Zhang and Dalgarno [24]; based on experimental data, 2007
157.3		Chu <i>et al</i> [23]; density functional theory, 2007
143		Recommended value by Zhang and Dalgarno [24], 2007
144.59		Sahoo and Das [26]; relativistic coupled-cluster theory, 2008

4. Mixing of degenerate states

In the previous section, we considered a particular case of mixing of the $4f^{14}6s6p$ ${}^{1}P_{1}^{o}$ and $4f^{13}5d6s^{2}6(7/2, 5/2)_{1}^{o}$ states of Yb and demonstrated that this mixing can be ignored in the *ab initio* calculations of the ground-state polarizability of Yb. Note that similar problems arise in computing the polarizability of the ${}^{3}P_{0}^{o}$ state and in many other cases not limited to calculations of the polarizabilities of Yb. Therefore, it may be useful to treat a general case and formulate the following theorem: *Consider a second-order transition amplitude, involving summation over a complete set of intermediate states. Then, a contribution from a subspace spanned by degenerate states does not depend on mixing of these states.*

A generic contribution to a second-order transition amplitude can be written as (here E_0 is some energy off-set)

$$A_{if} = \sum_{m} \frac{\langle F_{\rm f} | \hat{P} | F_m \rangle \langle F_m | \hat{Q} | F_{\rm i} \rangle}{E_0 - E_m} \tag{17}$$

Here F_i and F_f are some initial and final atomic states, and \hat{P} and \hat{Q} are some operators. Let us assume that the summation over the complete set of intermediate states F_m subsumes a summation over a group of degenerate states \mathcal{D} and rewrite the corresponding partial contribution as

$$\Delta A_{\rm if} = \frac{1}{\Delta E} \sum_{m \in \mathcal{D}} \langle F_{\rm f} | \hat{P} | F_m \rangle \langle F_m | \hat{Q} | F_{\rm i} \rangle.$$
(18)

Here the summation ranges only over degenerate states and ΔE is their common energy denominator.

As in the previous section, we introduce the pure nonmixed basis states Φ_k and rewrite the real (mixed) states F_m as an expansion

$$F_m = \sum_k c_{mk} \Phi_k. \tag{19}$$

Both sets Φ and *F* can be made orthonormal, with the matrix $C = \{c_{mk}\}$ being a unitary matrix, i.e., $CC^{\dagger} = I$. Substitution of (19) into (18) leads to

$$\Delta A_{\rm if} = \frac{1}{\Delta E} \sum_{kl} \langle F_{\rm f} | \hat{P} | \Phi_k \rangle \langle \Phi_l | \hat{Q} | F_{\rm i} \rangle \sum_m c_{mk} c_{ml}^*$$
$$= \frac{1}{\Delta E} \sum_k \langle F_{\rm f} | \hat{P} | \Phi_k \rangle \langle \Phi_k | \hat{Q} | F_{\rm i} \rangle.$$
(20)

Here we used the ortho-normality condition, $\sum_{k} c_{km} c_{kl}^* = \delta_{ml}$. We see that the final expression does not depend on the mixing coefficients *c*.

We stress two conditions under which this theorem is valid. It is assumed that the unmixed states possess the same energy and that there is no mixing with the states outside the degenerate subspace. These two conditions are probably never fulfilled exactly. However, there is a large number of cases when they are fulfilled approximately. One of the criteria is that the average energy spread in the quasi-degenerate subspace $|\delta E_{\mathcal{D}}| \ll |E_0 - \bar{E}_{\mathcal{D}}|$, where $\bar{E}_{\mathcal{D}}$ is the average energy in the subspace.

Why is this practically important? In Yb, to rigorously account for the mixing, one has to extend the two-valenceelectron model space to include computationally expensive core-excited states. The theorem claims that while computing polarizabilities, a much smaller two-valence-electron space would suffice. In other words, at the CI stage, the excitations from the 4f subshell can be safely ignored (even) if there are nearby two-electron states of the same symmetry. (Of course, *virtual* core excitations are included in the self-energy MBPT operator Σ .) Excitations from the core will have to be included into calculation of the polarizability of the core. This corresponds to working with the non-mixed basis Φ_k .

The independence on mixing explains why pure twovalence *ab initio* calculations of polarizabilities give good results for Yb despite its complicated structure. This should be true not only for polarizabilities but also for two-photon transition amplitudes, parity non-conservation, etc, and not only for Yb but for some other atoms as well.

5. Results and discussion

5.1. Static polarizabilities

Our computed static polarizabilities of the clock states are presented in table 3. There we tabulate values obtained in various approximations and we also compile results from the literature. Below we analyse our results and estimate theoretical uncertainties.

The first line of table 3 lists the results of pure *ab initio* calculations, next line gives the results obtained when energies

are fitted as explained in section 2. Then, to gauge the accuracy of the calculations, we introduce various corrections to the polarizability of the ground state based on available experimental data. In the third line of the table we correct α_0 to reproduce the experimental value for the magic frequency [9, 15]. This correction will be explained in the following section. The fourth value (135.2) is obtained by replacing the dominant ab initio contribution by semi-empirical values. This corresponds to changing from the unmixed Φ_1 , Φ_2 bases to the mixed Ψ_1 , Ψ_2 bases (see section 3 for details). Specifically, we use the value 4.148 a.u. [16] for the $\langle 6s^{2} {}^{1}S_{0} \| d \| 6s6p {}^{1}P_{1} \rangle$ transition amplitude and $\delta \alpha = 24(4)$ [24] for the contribution of the excitations from the 4f core subshell. The resulting value is smaller than both ab initio results (with and without fitting) consistent with our observation made at the end of section 3. Finally, we rescale $\alpha(0)$ of the ground state to fit the experimental value for the van der Waals coefficient C_6 [5]. Since C_6 is simply obtained by integrating $\alpha(i\omega)^2$, equation (6), and both the integral and the static polarizability are dominated by the principal transition, the uncertainties of $\alpha(0)$ and C_6 are correlated (see a detailed discussion in [36]). The final value for the ground state and its uncertainty, 141(6), are chosen to cover the spread of our numbers in the table.

Now we turn to the polarizability of the ${}^{3}P_{0}^{o}$ state. The central point of our final value is chosen to coincide with the fitted value. Choosing the lower calculated value as the central point is justified by the fact that there is a correction due to excitations for 4f and this correction is negative (see the end of section 3 for details). The value of this correction must be smaller than that for the ground state because corresponding states with excitations from 4f are higher in the spectrum and closer to the neighbouring two-electron states of the same symmetry. Note that the polarizabilities of both clock states end up having a comparable fractional accuracy. Qualitatively, this may be explained that while the convergence with respect to increasing basis is slower for the ${}^{3}P_{0}^{o}$ state, calculations for the ${}^{1}S_{0}$ state are affected more strongly by the core-excited states.

Results of other calculations and semi-empirical analyses are presented in table 3 for comparison. For the ground state note a good agreement with the value of [24], $\alpha(0) =$ 136.4(4.0), derived using only experimental data. The value $\alpha(0) =$ 143 recommended by the same authors is also consistent with our result.

The result for the ground-state polarizability is in a disagreement with previous calculations [19–21]. As discussed in section 3 the disagreement for ${}^{1}S_{0}$ comes from the inconsistent use of experimental matrix element for the principal transition in these works. As for the ${}^{3}P_{0}^{o}$ state, the present result and that of [21] differ by about two standard deviations and in experimental work this level of agreement between two values would be acceptable.

Finally, let us emphasize that evaluating accuracy of theoretical calculations is a non-trivial exercise and due to the lack of high-accuracy experimental data for the ${}^{3}P_{0}^{0}$ state, the uncertainty in [21] was estimated as a half of the difference between pure *ab initio* result and the result obtained with the fitting of the energy. This may be an unreliable estimate.

Table 4. Magic wavelengths for the $6s^2 {}^1S_0 \rightarrow 6s6p {}^3P_0^o$ transition in Yb and values of dynamic polarizabilities at corresponding frequencies (a.u.).

a	b initio	Corrected ^a		Experiment
λ (nm)	α (ω) (a.u.)	$\overline{\lambda}$ (nm)	$\alpha(\omega)$ (a.u.)	λ (nm)
749.0	193	759.37	186	759.355 ^b
552.2	46	551.5	60	
459.3	478	465.4	382	
413.2	1246	413.25	817	
359.7	-742	402.55 ^c	365°	

^a $\alpha_{1S_0}(\omega)$ is corrected to fit experimental magic frequency.

^b Reference [15].

^c Unreliable.

Suppose we carry out a similar analysis based on the present calculations for the ground state. Fitting energy in Σ_1 moves our *ab initio* result, 138.9, to 145.7, so using prescription of [21] would yield 145.7(3.4) which is displaced compared to and is as twice as accurate as our final value of 141(6).

5.2. Magic frequencies

Magic frequencies are frequencies of the laser field at which the ac Stark shifts of both clock levels are the same so that the frequency of the clock transition is insensitive to the laser intensity. Magic frequencies, ω^* , are found from the condition

$$\alpha_{^{1}S_{0}}(\omega^{*}) = \alpha_{^{3}P_{0}}(\omega^{*}). \tag{21}$$

The first five calculated magic frequencies for Yb are presented in table 4. The first frequency was measured to a high precision [9, 15]. Our *ab initio* calculations reproduce it with the 1.3% accuracy. The accuracy for the other frequencies is likely to be worse. This is because the energy denominators for the dynamic polarizabilities (2) are shifted by the frequency of the laser field and the assumption that the difference in energies of the $4f^{14}6s6p$ ${}^{1}P_{1}^{o}$ and $4f^{13}5d6s^{2}6(7/2, 5/2)_{1}^{o}$ states can be neglected becomes less accurate (see section 3 for details).

To improve the accuracy of predicting the magic frequencies and corresponding polarizabilities we modify formula (2) in two ways. First, we use experimental data for the dominant terms: $\langle 6s^{21}S_0 \| d \| 6s6p^{1}P_1 \rangle = 4.148$ [16] and $\langle 6s^{21}S_0 \| d \| 4f^{13}5d6s^2 (7/2, 3/2)_1^{\circ} \rangle = 2$ [41]. Second, we introduce a correction to simulate the effect of other excitations from the 4f core shell,

$$\Delta \alpha = \frac{A^2}{3} \left(\frac{1}{E - \omega} + \frac{1}{E + \omega} \right), \tag{22}$$

where $E = 37414.59 \text{ cm}^{-1}$ is the energy of the nearby state with the excitation from the core and A is a fitting parameter. Its value is chosen to fit experimental magic frequency. The rest of contributions are taken from the *ab initio* calculations.

The above empirical correction has been introduced for the ground state only. The polarizability of the upper clock state was computed within the CI+MBPT method for the valence contribution and the DHF method for core polarizability.

The values of magic frequencies and corresponding polarizabilities calculated with this empirical correction are

Table 5. Atom–wall interaction coefficients C_3 and van der Waals coefficients C_6 (a.u.) for the $6s^{2} {}^{1}S_0$ and $6s6p {}^{3}P_0^{0}$ states of Yb.

	Value		Comment	
		Atom-	wall interaction	
C_3	${}^{1}S_{0}$	3.34	Calculated	This work
C_3	${}^{3}P_{0}^{o}$	3.68	Calculated	This work
		Atom-a	tom interaction	
C_6	${}^{1}S_{0} + {}^{1}S_{0}$	1909(160)	Calculated	This work
		2300(250)	Experimental	Enomoto <i>et al</i> [17]
		1932(35)	Experimental	Kitagawa <i>et al</i> [5]
		2400-2800	Calculated	Buchachenko et al [40]
		2292	Calculated	Chu <i>et al</i> [23]
		2062	Calculated	Zhang and Dalgarno [24]
C_6	${}^{3}P_{0}^{o} + {}^{3}P_{0}^{o}$	3886(360)	Calculated	this work
$C_6 \\ C_6$	${}^{1}S_{0}^{o} + {}^{3}P_{0}^{o}$	2709(338)	Calculated	this work

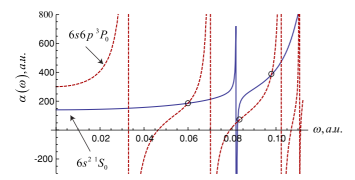


Figure 1. Dynamic polarizabilities of the two clock levels. 'Magic' conditions occur when the two polarizabilities intersect off resonance. These are marked by small circles on the plot. (This figure is in colour only in the electronic version)

listed in the third and fourth columns of table 4 and the ac polarizabilities are shown in figure 1. We see that the results for the first four magic frequencies change little, while those for the fifth magic frequency vary significantly. Still, while difficult to predict, the fifth magic frequency may be of practical interest for designing better clocks: by contrast to the first four frequencies, the ac polarizability here may be negative (that was the *ab initio* prediction, but the empirical correction flipped the sign of the polarizability). Because of that, the atoms are trapped at the minima of the laser intensity, which reduces perturbations of the clock frequency [42].

Finally, using the modified formula at $\omega = 0$ provides another accuracy test for our predicted static polarizability of the ground state (section 5.1). The resulting value, 139.1 a.u., was listed in table 3 and used in estimating the accuracy of the calculations.

5.3. Black-body radiation shift

The effect of black-body radiation (BBR) on the frequency of the Yb clock transition is expressed in terms of the static polarizabilities. The relevant clock shift reads (see detailed derivations in [21])

$$\delta \nu_{\rm BBR} \approx -\frac{2}{15} (\alpha_{\rm fs} \pi)^3 T^4 \big\{ \alpha_{^{3}\rm P_{0}}(0) - \alpha_{^{1}\rm S_{0}}(0) \big\}, \qquad (23)$$

where *T* is the temperature and $\alpha_{\rm fs}$ is the fine-structure constant. With our polarizabilities we find at T = 300 K, $\delta \nu_{\rm BBR} = -1.41(17)$ Hz which is in good agreement with the result, $\delta \nu_{\rm BBR} = -1.34(13)$ Hz, of [21] despite our significant differences in values of individual polarizabilities. The agreement is fortuitous, as both polarizabilities of [21] are smaller, most of the shift cancelling out while taking the difference in equation (23). Note, however, that a somewhat smaller uncertainty reported in [21] is mostly due to overestimating the theoretical accuracy for polarizabilities, especially for the ground state. See sections 3 and 5.1 for details.

It is worth emphasizing that at the time of writing, the BBR correction is the leading uncertainty in the error budget of the lattice clocks [10]. The present calculations demonstrate a difficulty of atomic theory in determining this important correction accurately. We hope that our analysis would motivate a further experimental work, for example, on cryogenically cooled clock chambers to reduce (and ultimately to accurately determine) the BBR shift.

5.4. Atom–wall interaction coefficients C_3 and van der Waals coefficients C_6

We calculate the atom-wall interaction constants C_3 and the van der Waals constant C_6 for both clock states using formulae (5) and (6). The results are presented in table 5 together with the results of other calculations and measurements for the C_6 coefficient for the ground state.

For two Yb atoms in their ground states we find $C_6({}^{1}S_0 + {}^{1}S_0) = 2041$ a.u. (CI+MBPT with rescaled Σ to fit the energies). However, from our calculations of the polarizabilities we know that the contributions from the excitations from the 4f core state are likely to make the results smaller. Therefore, we rescaled the central point for $C_6({}^{1}S_0 + {}^{1}S_0)$ at twice the rate as for the polarizability of the ground state (i.e., reduced it by 6.5%). This moves the central value to 1909 a.u. Since the C_6 is obtained as a quadrature of the square of polarizability, we assign a fractional uncertainty to C_6 as twice as large as that for the static polarizabilities. Note a good agreement of the ground-state C_6 with recent experimental determinations [5] and with calculations [24].

The results in the table for $C_6({}^{1}S_0 + {}^{3}P_0)$ and $C_6({}^{3}P_0 + {}^{3}P_0)$ were obtained in CI+MBPT with rescaled Σ to fit the energies.

Including core polarizability into computation of C_6 is important. We find that neglecting $\alpha_{core}(i\omega)$ reduces the final result for C_6 by about 10%. This is consistent with the earlier observation [27] for heavy alkali-metal atoms.

Atom–wall interaction coefficients C_3 for the clock states were obtained in CI+MBPT with rescaled Σ to fit the energies; these are also listed in table 5. Note that the difference in C_3 coefficients for the clock states was computed by us earlier in [12]. While forming the difference the contribution of core polarizability cancels out. For individual states, this contribution is substantial. For example, for the ground state the result is increased from 2.09 to 3.34 a.u. due to $\alpha_{core}(i\omega)$. As a large fraction of C_3 value is accumulated due to the core excitations, assigning error to C_3 is difficult. For example, to estimate the error in C_3 for simpler alkali-metal atoms in [27], the results of two methods of computing C_3 , (i) via the integral of $\alpha(i\omega)$ and (ii) by computing an expectation value of some two-body dipole-dipole interaction were compared. A highly technical evaluation of the expectation value of twobody operator is beyond the scope of the present paper and we do not assign uncertainties to C_3 . We expect that the errors in C_3 are unlikely to exceed 50%.

6. Conclusion

We calculated polarizabilities of the clock states of Yb with the accuracy of about 5%. The substantial disagreement with previous calculations for the ground-state polarizability was explained and resolved. We also computed the first four magic frequencies of the lattice laser field, the effect of blackbody radiation on the frequency of the clock transition, the C_3 atom-wall interaction constants and the C_6 van der Waals coefficients for both clock states. Polarizabilities, the first magic frequency and the C_6 coefficient for the ground state are in good agreement with the most accurate calculations and measurements. The presented data may be of interest for designing better clocks, for applications of the clocks in studying the atom-wall interaction and quantum information processing, and quantifying molecular potentials for ultracold collision studies.

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