High-Accuracy Calculation of the Blackbody Radiation Shift in the $^{133}$Cs Primary Frequency Standard

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The blackbody radiation (BBR) shift is an important systematic correction for the atomic frequency standards realizing the SI unit of time. Presently, there is controversy over the value of the BBR shift for the primary $^{133}$Cs standard. At room temperatures, the values from various groups differ at the $3 \times 10^{-15}$ level, while modern clocks are aiming at $10^{-16}$ accuracies. We carry out high-precision relativistic many-body calculations of the BBR shift. For the BBR coefficient $\beta$ at $T = 300$ K, we obtain $\beta = -(1.710 \pm 0.006) \times 10^{-14}$, implying $6 \times 10^{-17}$ fractional uncertainty. While in accord with the most accurate measurement, our 0.35% accurate value is in a substantial (10%) disagreement with recent semiempirical calculations. We identify an oversight in those calculations.

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First, let us review the underlying theory of the Cs BBR shift. BBR causes a weak oscillating perturbation of atomic energy levels. Conventionally, the BBR contribution is parametrized as the fractional correction to the unperturbed clock frequency $\nu_0 = 9192631770$ Hz

$$\delta \nu^{\text{BBR}} / \nu_0 = \beta \times (T/T_0)^4 (1 + \varepsilon \times (T/T_0)^2),$$

where $T_0 = 300$ K; $\varepsilon$ is a small, relatively well-known 1.4% correction [2], while the overall factor $\beta$ requires careful evaluation and is the subject of the present work. $\beta$ can be related to the scalar differential polarizability for the hyperfine manifold of the $6s_{1/2}$ Cs ground state. Indeed, the characteristic thermal photon energy at room temperatures is much smaller than the atomic energies, so that the perturbation can be well described in the static limit. Moreover, contributions of electromagnetic BBR multipoles beyond electric dipoles, as well as retardation corrections, are highly suppressed [10]. Then the BBR shift of the energy level is given by (atomic units are used throughout; $\alpha$ is the fine-structure constant)

$$\delta E^{\text{BBR}}_F = \frac{4}{15} (\alpha \pi) F^4 \alpha_F(0),$$

where $\alpha_F(0)$ is the static scalar electric-dipole polarizability of the hyperfine level $F$. The vector and tensor parts of the polarizability average out due to the isotropic nature of the BBR.

The relation of the BBR shift to polarizability has been exploited in the most accurate measurement of the differential Stark shift [3]. However, a recent direct temperature-dependent measurement [5] of the BBR shift turned out to be differing by about 2 standard deviations from the indirect measurement. Namely, this difference has stimulated the recent interest in the Cs BBR shift.

The overall BBR shift of the clock frequency is the difference of the individual shifts for the two hyperfine states ($F = 4$ and $F = 3$) involved in the transition. While taking the difference, the traditional lowest-order polar-
izability of the $6s_{1/2}$ level cancels out, and one needs to evaluate the third-order $F$-dependent polarizability $\alpha^{(3)}(0)$. This contribution involves two $E1$ interactions $V = -\mathbf{D} \cdot \mathbf{E}$ with the external electric field ($\mathbf{D}$ is the dipole operator) and one hyperfine $H_{\text{hfs}}$ coupling [11]. We parametrize the general correction to the energy as a sum of four diagrams:

$$\delta E^{(3)}_F = \langle \mathcal{F}|H_{\text{hfs}}RVRV|\mathcal{F}\rangle + \langle \mathcal{F}|VRH_{\text{hfs}}RV|\mathcal{F}\rangle + \langle \mathcal{F}|VRVRH_{\text{hfs}}|\mathcal{F}\rangle - \langle \mathcal{F}|H_{\text{hfs}}|\mathcal{F}\rangle\langle \mathcal{F}|VRVRV|\mathcal{F}\rangle,$$

where $R = (H - E_v)^{-1}$ is the resolvent operator, with $H$ and $E_v$ being the atomic Hamiltonian and the ground state energy, respectively. The four contributions will give rise to top, center, bottom, and residual contributions discussed below. (The naming convention reflects the relative position of the hyperfine operator in the string of the three interactions.) The top and bottom diagrams are equal due to Hermiticity considerations. Because of angular selection rules, only magnetic dipolar interaction remains; we write $H_{\text{hfs}} = \mu \cdot \mathcal{T}^{(1)}$, where $\mu$ is the nuclear magnetic moment and $\mathcal{T}^{(1)} = -i\sqrt{2}(\alpha C_{1A}^{(0)}(i))/(c r^2)$ is the relevant relativistic coupling tensor [12].

After angular and many-body reduction, the scalar polarizability may be expressed as

$$\alpha^{(3)}(0) = \frac{1}{3}\sqrt{2(2I)(2I + 1)(2I + 2)} \left[ \begin{array}{ccc} j_v & I & F \\ 1 & j_v & 1 \end{array} \right] \times g_1 \mu_n (-1)^{I+j_v+j_s}(2T + C + R),$$

where $g_1$ is the nuclear gyromagnetic ratio, $\mu_n$ is the nuclear magneton, $I = 7/2$ is the nuclear spin, and $j_v = 1/2$ is the total (electron) angular momentum of the ground state. The $F$-independent sums are ($|\nu\rangle \equiv |6s_{1/2}\rangle$)

$$T = \sum_{m,n,\sigma} \frac{(-1)^{I_m+j_s}}{2j_v + 1} \times \frac{\langle \nu | D | m \rangle \langle m | D | n \rangle \langle n | \mathcal{T}^{(1)} | \nu \rangle}{(E_m - E_v)(E_n - E_v)} \delta_{j_m,j_s},$$

$$C = \sum_{m,n,\sigma} \frac{(-1)^{I_m-j_v}}{2j_m + 1} \times \frac{\langle \nu | D | m \rangle \langle m | \mathcal{T}^{(1)} | n \rangle \langle n | D | \nu \rangle}{(E_m - E_v)(E_n - E_v)},$$

$$R = \frac{\langle \nu | \mathcal{T}^{(1)} | \nu \rangle}{2j_v + 1} \sum_{m \in \text{val}} \sum_{m \in \text{core}} \frac{\langle \nu | D | m \rangle^2}{(E_m - E_v)^2},$$

The summation indices $m$ and $n$ range over valence bound and continuum many-body states and also over single-particle core orbitals. With this convention, the above expressions subsume contributions from intermediate valence and core-excited states and they also take into account so-called core-valence counterterms [13]. Selection rules impose the following angular symmetries on the intermediate states: $s_{1/2}$ for $|n\rangle$, $p_{1/2,3/2}$ for $|m\rangle$ in the top diagram, $p_{1/2,3/2}$ for both $|m\rangle$ and $|n\rangle$ in the center diagram, and $p_{1/2,3/2}$ for the $|m\rangle$ in the residual term.

We will tabulate our results in terms of the conventional scalar Stark shift coefficient $k_s = 1/2(\alpha^{(3)}(0) - \alpha^{(3)}_{F-3}(0))$. For $^{133}$Cs, this coefficient can be written more explicitly in terms of the $F$-independent diagrams as $k_s = -3(2/3)^{1/2}g_1 \mu_n (2T + C + R)$, with the BBR coefficient $\beta = -4/15(\alpha \pi)^2 T_0^6 / \nu_0 \times k_s$.

Numerical evaluation of the diagrams $T, C$, and $R$ can be carried out either using the Dalgarno-Lewis method or by directly summing over individual intermediate states. Here we use the direct summation approach. This treatment is similar to the one used in high-accuracy calculations of atomic parity violation in $^{133}$Cs [14]. The main advantage of this method is that one could explicitly exploit high-accuracy experimental data for energies, dipole-matrix elements, and hyperfine constants. When the accurate values are unknown, we use ab initio data of proper accuracy. In addition, this approach facilitates comparison with recent calculations [6,7], which also use the direct summation approach.

The central technical issue arising in direct summation over a complete set of states is representation of the innumerable spectrum of atomic states. For example, even without the continuum, the bound spectrum contains an infinite number of states. A powerful numerical method for reducing the infinite summations or integrations to a finite number of contributions is the basis-set technique. In particular, we employ the $B$-spline technique [15]. In this approach, an atom is placed in a large spherical cavity and the single-particle Dirac-Hartree-Fock (DHF) orbitals are expanded in terms of a finite set of $B$ splines. The expansion coefficients are obtained by invoke the variational Galerkin principle. The resulting set of the single-particle orbitals is numerically complete and finite. The technique has a high numerical accuracy, and we refer the reader to a review [16] on numerous applications of $B$ splines for details.

We use a $B$-spline set with 70 splines of order 7 for each partial wave and constrain the orbitals to a cavity of radius $R_{\text{cav}} = 220$ a.u. This particular choice of $R_{\text{cav}}$ ensures that the lowest-energy atomic orbitals are not perturbed by the cavity. In particular, all core and valence DHF orbitals with radial quantum numbers 1–12 from the basis set produce energies and matrix elements in close numerical agreement with the data from the traditional finite-difference DHF code. These low-energy orbitals will produce true many-body states for a cavity-unconstrained atom.

To understand the relative role of various contributions, we start by computing the Stark shift at the DHF level. We obtain the Stark coefficient of $R^{\text{DHF}} = -2.799 \times 10^{-10}$ Hz/(V/m)$^2$ [17], with fractional contributions $T^{\text{DHF}} = 0.418$, $C^{\text{DHF}} = 0.003$, and $R^{\text{DHF}} = 0.580$. It is clear that the top and residual
terms dominate over the center diagram. The bulk (99.8%) of the value of the residual term is accumulated due to the principal $6s - 6p_{1/2,3/2}$ transitions. For the top term, the saturation of the sum is not as rapid, but still the dominant contributions come from the lowest-energy excitations: Limiting the summations to the first four excited states recovers only 68% of the total value for the top diagram. Additionally, core-excited states contribute only 0.1% to the final value.

The above observations determine our strategy for more accurate calculations. We group the entire set of atomic states into the “main” low-lying-energy states (principal quantum numbers $n \leq 12$) and remaining “tail” states. We will account for the contribution from the main states using high-accuracy experimental and ab initio values. The contribution from the tail will be obtained using either the DHF or the mixed approach.

First, we describe the high-accuracy data used in our calculations. We need dipole and hyperfine matrix elements and energies. Experimental values for the dipole-matrix elements for the following six transitions were taken from the literature (see compilations in Refs. [6,18]): $6s_{1/2} - 6p_{1/2,3/2}$, $7s_{1/2} - 6p_{1/2,3/2}$, $7s_{1/2} - 7p_{1/2,3/2}$. Crucial to the accuracy of the present analysis were the matrix elements for the principal $6s_{1/2} - 6p_{1/2,3/2}$ transitions. We have used a 0.005%-accurate value for $\langle 6s_{1/2} | D | 6p_{1/2}\rangle$ from Ref. [19]. The value for $\langle 6s_{1/2} | D | 6p_{1/2}\rangle$ was obtained by using the above $6s_{1/2} - 6p_{1/2}$ matrix element and a 0.03%-accurate measured ratio [20] of these matrix elements. These six experimental matrix elements were supplemented by 92 values ($ns_{1/2} - np_{1/2,3/2}$ values for $n, n' = 6-12$) from high-accuracy ab initio calculations. We employ the relativistic linearized coupled-cluster singles-doubles (LCCSD) method. The underlying formalism, implementation, and results for alkali atoms are described in Ref. [18]. For dipole-matrix elements, the accuracy of the ab initio LCCSD method is a few tenths of a percent.

As to the high-accuracy values of the matrix elements of the hyperfine coupling, the diagonal matrix elements of the $T^{(1)}$ tensor are directly related to the conventional hyperfine constants: $A = g_s \mu_s / j_v (2j_v + 1)/(2j_v + 2))^{1/2} \langle \langle T^{(1)} | \nu \rangle \rangle$. We have used the compilation of hyperfine constants from Ref. [21] for the main $n = 6-12$ states. Off-diagonal matrix elements between the $s$ states were evaluated using the geometric-mean formula

$$\langle n_{1/2} | T^{(1)} | n'_{1/2}\rangle = \{(\langle n_{1/2} | T^{(1)} | n_{1/2}\rangle \times \langle n'_{1/2} | T^{(1)} | n'_{1/2}\rangle)\}^{1/2}. $$

This expression has been shown to hold to about $10^{-3}$ in Ref. [22] (notice that the radiative corrections would start playing a role at a few tenths of a percent as well). If we had $6 \leq n \leq 12$ in the above expression, then the experimental value is used for its corresponding diagonal element on the right. If we also had $6 \leq n' \leq 12$, then that experimental value is also used for the corresponding diagonal element on the right; otherwise, the DHF value is taken. ($n$ and $n'$ can be obviously interchanged in this prescription.) This mixed approach has allowed us to uniformly improve the accuracy of the calculations. Indeed, in the numerically important top term, the hyperfine matrix elements come in the combination $\langle ns_{1/2} | T^{(1)} | 6s_{1/2}\rangle$. As $n$ grows, the correlations become less important, so the dominant correlation correction comes from the $6s_{1/2}$ state. Using the described mixed approach allows us to account for this dominant correlation. The geometric-mean formula holds only for the $s$ states. For the off-diagonal matrix elements between various combinations of $6p_{1/2,3/2}$ and $np_{1/2,3/2}$ ($n = 6-9$) states, we employed a modification of the LCCSD method augmented by a perturbative treatment of the valence triple excitations (LCCSDpV method) [18]. The accuracy of these matrix elements is a few percent. As these matrix elements enter the relatively small center term, the effect on the overall theoretical error is negligible.

Finally, we used experimental energy values from the NIST tabulation [23], for states with principle quantum number $n = 6-12$, and the DHF values otherwise.

With the described set, we report our final result for the scalar Stark coefficient and relative blackbody radiation shift at 300 K to be

$$k_s = -(2.271 \pm 0.008) \times 10^{-10} \text{ Hz/(V/m)},$$
$$\beta = -(1.710 \pm 0.006) \times 10^{-14}. $$

The fractional contributions of the individual diagrams are $2T_{C+R} = 0.442$, $2T_{C+R} = 0.002$, and $2T_{C+R} = 0.560$. When comparing with the DHF values, the most substantial modification due to correlations is in the center term, which changes the sign. Fortunately, this term is relatively small, and this extreme change does not substantially affect the final result.

The overall uncertainty of these results was determined from the uncertainties of the individual matrix elements and energy values used in their computation. Standard uncertainty analysis was done throughout all mathematical operations. For energy values taken from NIST, the uncertainty is assumed negligible. For all other experimental values, the reported uncertainty is used. For ab initio matrix elements (DHF, LCCSD, or LCCSDpV), we assigned an assumed uncertainty. These assumed uncertainties were based on comparison between calculated and high-accuracy experimental values. This resulted in a relative uncertainty for both the scalar Stark coefficient and the BBR shift of 0.35\%. We have performed several consistency checks, e.g., replacing experimental matrix elements and energies by ab initio LCCSD values or replacing the DHF values for states with $n = 13-26$ with the LCCSD values. The final result was stable to such modifications within the stated uncertainty in Eq. (2). These tests provide us with additional confidence with respect to our standard
error analysis based on errors of the used experimental values. It is also worth noting that the present calculation does not include radiative corrections, which may contribute at the level of a few tenths of a percent (some radiative corrections, e.g., vacuum polarization, are absorbed in our final value already as we use experimental hyperfine constants).

A comparison with recent theoretical and experimental work is presented in Table I. While agreeing with the most accurate measurement by Simon et al. [3], our results are in substantial disagreement with the recent calculations [6,7]. The principal differences between the present work and these calculations are (i) more sophisticated treatment of correlations and (ii) rigorous summation over the complete set of intermediate states in perturbative expressions. As discussed above, we used the numerically complete basis-set approach which approximates the Rydberg states and continuum with a quasispectrum. To illuminate the importance of these contributions, we truncate our summations at \( n = 12 \). The resulting value deviates from our final \( k_x \) result by 7%. The fact that the continuum needs to be included is hardly surprising, as, for example, about 20% of the textbook polarizability of the ground state of the hydrogen atom comes from the continuum states. Calculations [24] include the continuum via the Green’s function technique, and their result is in agreement with our value.

To conclude, here we have reported results of relativistic many-body calculations of the BBR shift, one of the leading systematic corrections in the \(^{133}\text{Cs} \) frequency standard and a subject of recent controversy. Our 0.35%-accurate result revalidates high-precision Stark shift measurements [3]. Our work also clarifies the origin of the reported discrepancy between that measurement and recent calculations [6,7].

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Note added.—While completing this manuscript, we have learned of another accurate many-body calculation of the BBR shift in \(^{133}\text{Cs} \) clock [25]. Their result \( k_x = 2.26 \times 10^{-10} \pm 1 \% \) Hz/(V/m)\(^2 \) is in agreement with our more accurate value.

Table I. Values of \( k_x \) in \( 10^{-10} \) Hz/(V/m)\(^2 \).

| Theory | -1.97 ± 0.09 | Ref. [6] |
| Theory | -2.06 ± 0.01 | Ref. [7] |
| Theory | -2.28 | Ref. [24] |
| Expt. | -2.05 ± 0.04 | Ref. [4] |
| Expt. | -2.271 ± 0.004 | Ref. [3] |
| Theory | -2.271 ± 0.008 | Present |

References

[12] This coupling is for a pointlike nucleus. In calculations, we include the effect of finite nuclear magnetization, modeling the nucleus as a uniformly magnetized ball of radius 5.6748 fm.
[17] To test the numerical convergence, we also used another basis set of 40 splines with \( R_{\text{av}} = 75 \) a.u. The resulting number was consistent with the value produced with the described larger set to one part in 40,000.