Relativistic corrections for the ground electronic state of molecular hydrogen

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We recalculate the leading relativistic corrections for the ground electronic state of the hydrogen molecule using the variational method with explicitly correlated functions that satisfy the interelectronic cusp condition. The new computational approach allowed for the control of the numerical precision which reached about eight significant digits. More importantly, the updated theoretical energies became discrepant with the known experimental values and we conclude that the yet unknown relativistic recoil corrections might be larger than previously anticipated.

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I. INTRODUCTION

Theoretical studies of the hydrogen molecule are the cornerstone of the molecular quantum mechanics. Due to its simplicity, the achieved precision is the highest among all molecules and still has the potential of significant enhancement. This high precision of theoretical predictions for H₂ leads to improved tests of quantum electrodynamics and improved bounds on hypothetical interactions [1]. Moreover, at the 10^{-7} cm⁻¹ precision level the dissociation energy is sensitive to the proton charge radius, which may help to resolve the so-called proton radius conundrum [2]. This requires high accuracy calculations of not only nonrelativistic energies, but also leading relativistic $O(\alpha^2)$, QED $O(\alpha^3)$, as well as the higher order corrections $O(\alpha^4)$ and $O(\alpha^5)$. In fact, the nonrelativistic energies can already be calculated with the precision of 10^{-7} cm⁻¹, as demonstrated in Ref. [3]. The $O(\alpha^4)$ contribution has very recently been calculated [4] using explicitly correlated Gaussian (ECG) functions with $1 + r_{12}/2$ prefactor (rECG) that causes the interelectronic cusp condition to be exactly satisfied. Here, we report the results for the leading $O(\alpha^2)$ relativistic correction using rECG functions and conclude that the compilation of previous results in Ref. [5] has underestimated numerical uncertainties. We improve the numerical precision by 3-4 orders of magnitude and present in detail our computational approach.

II. COMPUTATIONAL METHOD

In the Born-Oppenheimer (BO) approximation the total wave function is assumed to be a product of the electronic and nuclear functions. The Schrödinger equation for the electronic wave functions in the infinite nuclear mass limit (assuming atomic units) is

$$H\psi(\vec{r}_1, \vec{r}_2) = \mathcal{E}(R)\psi(\vec{r}_1, \vec{r}_2)$$
 (1)

where

$$H = \frac{1}{2} (\vec{p}_1^2 + \vec{p}_2^2) + V \tag{2}$$

and

$$V = \frac{1}{R} - \frac{1}{r_{1A}} - \frac{1}{r_{2A}} - \frac{1}{r_{1B}} - \frac{1}{r_{2B}} + \frac{1}{r}$$
 (3)

with $R=r_{AB}$ and $r=r_{12}$, and where indices 1 and 2 correspond to electrons, whereas A and B correspond to the nuclei. The leading relativistic correction in the BO approximation is $\mathcal{E}_{\mathrm{rel}}(R)$ to the nonrelativistic potential $\mathcal{E}(R)$. This correction can be expressed in terms of the expectation value

$$\mathcal{E}_{\rm rel}(R) = \langle \psi | H_{\rm rel} | \psi \rangle \tag{4}$$

of the Breit-Pauli Hamiltonian [6]

$$H_{\text{rel}} = -\frac{1}{8} \left(p_1^4 + p_2^4 \right) + \pi \, \delta^3(r) - \frac{1}{2} \, p_1^i \left(\frac{\delta^{ij}}{r} + \frac{r^i \, r^j}{r^3} \right) p_2^j + \frac{\pi}{2} \left(\delta^3(r_{1A}) + \delta^3(r_{2A}) + \delta^3(r_{1B}) + \delta^3(r_{2B}) \right), (5)$$

where we neglected spin dependent terms vanishing for the ground electronic state of $^{1}\Sigma_{g}^{+}$ symmetry. The accurate calculation of the above expectation value is the principal goal of this work. We assume that every rECG basis function contains the 1 + r/2 factor and perform special transformation (regularization) of matrix elements, including those with the Dirac- δ function [7], to improve the numerical convergence. We demonstrate a significant enhancement in numerical precision and indicate that previous numerical results [5] were not as accurate as claimed. In order to be more convincing, we provide results obtained in three approaches: i) direct (no regularization) with ECG, ii) standard regularization with ECG, and iii) modified regularization with rECG functions. To test the convergence of these three different approaches, first we perform calculations for R=0, namely for the helium atom, for which highly accurate reference results can be obtained using explicitly correlated exponential functions. Next, the ECG calculations are performed for molecular hydrogen. Comparison of individual operators from different approaches is presented for the equilibrium internuclear distance, namely for R = 1.4 a.u. The most accurate predictions were obtained for the regularization with rECG functions at 53 points in the range R = 0.0 - 10 a.u.

Except for our recent paper [4], there has been no similar study of regularization techniques in the literature due to difficulties with two-center integrals involving inverse powers of interparticle distances. In relation to this, we have introduced a novel algorithm for numerical quadrature of non-standard ECG two-center integrals [4], which enables very efficient calculation of all complicated matrix elements.

III. REGULARIZATION OF THE RELATIVISTIC CORRECTION

In this section we provide regularization formulas for matrix elements with Dirac- δ and p^4 operators in $H_{\rm rel}$. The latter operator can be regularized according to two schemes: the standard one, already employed in the past in quantum molecular computations, [8] and the modified scheme, valid in the case of the wave function obeying Kato's cusp condition.

According to the standard scheme, the relativistic operators are transformed into the regular form by the following relations

$$4\pi \,\delta^3(r_{1A}) = 4\pi \,[\delta^3(r_{1A})]_r + \left\{\frac{2}{r_{1A}}, H - \mathcal{E}\right\},\tag{6}$$

$$4\pi \,\delta^3(r) = 4\pi \,[\delta^3(r)]_r + \left\{\frac{1}{r}, H - \mathcal{E}\right\},$$
 (7)

$$p_1^4 + p_2^4 = [p_1^4 + p_2^4]_r + 4\left\{\mathcal{E} - V, H - \mathcal{E}\right\} + 4\left(H - \mathcal{E}\right)^2,$$
(8)

where

$$4\pi \left[\delta^{3}(r_{1A})\right]_{r} = \frac{4}{r_{1A}}(\mathcal{E} - V) - \vec{p}_{1} \frac{2}{r_{1A}} \vec{p}_{1} - \vec{p}_{2} \frac{2}{r_{1A}} \vec{p}_{2},$$
(9)

$$4\pi \left[\delta^{3}(r)\right]_{r} = \frac{2}{r}(\mathcal{E} - V) - \vec{p}_{1} \frac{1}{r} \vec{p}_{1} - \vec{p}_{2} \frac{1}{r} \vec{p}_{2}, \qquad (10)$$

$$[p_1^4 + p_2^4]_r = 4(\mathcal{E} - V)^2 - 2p_1^2p_2^2.$$
(11)

For the exact wave function ψ , which fulfills the electronic Schrödinger equation $(\mathcal{E}-H)\psi=0$, the expectation value identity holds $\langle\psi|\dots|\psi\rangle=\langle\psi|[\dots]_r|\psi\rangle$, since for an arbitrary operator Q, $\langle\psi|\{Q,H-\mathcal{E}\}|\psi\rangle=0$. For an approximate function $\tilde{\psi}$, such expectation values do not vanish, but converge to zero at the limit $\tilde{\psi}\to\psi$. In practice, the numerical convergence of the regularized form is much faster, so the leading relativistic correction shall be evaluated as

$$\mathcal{E}_{\rm rel}(R) = \langle \psi | [H_{\rm rel}]_r | \psi \rangle \tag{12}$$

with

$$[H_{\rm rel}]_r = -\frac{1}{8} [p_1^4 + p_2^4]_r + \frac{\pi}{2} ([\delta^3(r_{1A})]_r + [\delta^3(r_{2A})]_r + [\delta^3(r_{1B})]_r + [\delta^3(r_{2B})]_r) + \pi [\delta^3(r)]_r - \frac{1}{2} p_1^i \left(\frac{\delta^{ij}}{r} + \frac{r^i r^j}{r^3}\right) p_2^j.$$
(13)

The modified regularization is applied when the wave function $\tilde{\psi}$ exactly satisfies the interelectronic cusp condition, as

for example the *r*ECG function does. In this case, the action of $[p_1^4 + p_2^4]_r$ on such a function can be represented as

$$[p_1^4 + p_2^4]_r |\tilde{\psi}\rangle = \left\{ 4 (\mathcal{E} - V)^2 - 2 \,\tilde{p}_1^2 \,\tilde{p}_2^2 + 8\pi \,\delta^3(r) \right\} |\tilde{\psi}\rangle. \tag{14}$$

The new term $\tilde{p}_1^2 \, \tilde{p}_2^2$ (in contrast to $p_1^2 \, p_2^2$) is understood as the differentiation $\nabla_1^2 \, \nabla_2^2$ of $\tilde{\psi}$ as a function, and what is omitted

$$-2\nabla_1^2 \nabla_2^2 (1+r/2) = 8\pi \delta^3(r). \tag{15}$$

coincides with the last term in Eq. (14). Now, if we are interested in determination of the $\langle \tilde{\psi} | p_1^4 + p_2^4 | \tilde{\psi} \rangle$ alone, we can additionally replace the Dirac- δ operator by its regularized form and obtain the fully-regularized expectation value

$$\langle \psi | p_1^4 + p_2^4 | \psi \rangle = \langle \psi | 4 (\mathcal{E} - V)^2 - 2 \, \tilde{p}_1^2 \, \tilde{p}_2^2 + 8\pi \, [\delta^3(r)]_r | \psi \rangle. \tag{16}$$

We emphasize that here, unlike $p_1^2 p_2^2$, the $\tilde{p}_1^2 \tilde{p}_2^2$ term differentiates the right-hand-side wave function only. The specific relationship (14) can also be further employed to simplify the expectation value of the overall Breit-Pauli Hamiltonian (5) by complete elimination of the $\pi \delta^3(r)$ term

$$\mathcal{E}_{\rm rel}(R) = \langle \psi | [H_{\rm rel}]'_r | \psi \rangle, \tag{17}$$

$$[H_{\rm rel}]'_r = -\frac{1}{2} (\mathcal{E} - V) \left(\mathcal{E} - \frac{1}{R} - \frac{1}{r} \right)$$

$$+ \frac{1}{4} \left(\tilde{p}_1^2 \, \tilde{p}_2^2 + \vec{p}_1 \, \tilde{V} \, \vec{p}_1 + \vec{p}_2 \, \tilde{V} \, \vec{p}_2 \right)$$

$$- \frac{1}{2} \, p_1^i \left(\frac{\delta^{ij}}{r} + \frac{r^i \, r^j}{r^3} \right) p_2^j \tag{18}$$

with $\tilde{V}=-1/r_{1A}-1/r_{1B}-1/r_{2A}-1/r_{2B}$. Apart from its compactness, this formula has an additional important advantage which is not readily noticeable. Due to the above cancellations, all the time-consuming integrals with three odd powers of interparticle distances do not appear in the matrix elements with *r*ECG functions. This non-trivial cancellation has remarkable impact on calculations of the relativistic correction.

IV. INTEGRALS WITH ECG FUNCTIONS

The variational wave function

$$\psi = \sum_{i} c_i \psi_i(\vec{r}_1, \vec{r}_2), \qquad (19)$$

$$\psi_i = (1 + \hat{i}) (1 + P_{1 \leftrightarrow 2}) \phi_i(\vec{r_1}, \vec{r_2}), \tag{20}$$

where \hat{i} and $P_{1\leftrightarrow 2}$ are the inversion and the electron exchange operators, can be accurately represented in the basis of ECG functions of the form

$$\phi_{\Sigma^{+}} = e^{-a_{1A} r_{1A}^2 - a_{1B} r_{1B}^2 - a_{2A} r_{2A}^2 - a_{2B} r_{2B}^2 - a_{12} r^2}$$
 (21)

or in the basis of the modified rECG functions

$$\phi_{\Sigma^{+}} = \left(1 + \frac{r}{2}\right) e^{-a_{1A} r_{1A}^{2} - a_{1B} r_{1B}^{2} - a_{2A} r_{2A}^{2} - a_{2B} r_{2B}^{2} - a_{12} r^{2}}.$$
(22)

Nonlinear a-parameters are determined variationally for every ECG or rECG basis function, and linear c-parameters come from the solution of the general eigenvalue problem. The primary advantage of the ECG type of functions is that all integrals necessary for the calculations of nonrelativistic and relativistic operators can be evaluated very effectively as described below.

Each matrix element can be expressed as a linear combination of the following ECG integrals

$$f(n_1, n_2, n_3, n_4, n_5) = \frac{1}{\pi^3} \int d^3r_1 \int d^3r_2 \, r_{1A}^{n_1} r_{1B}^{n_2} r_{2A}^{n_3} r_{2B}^{n_4} r_{12}^{n_5}$$
$$\times e^{-a_{1A} \, r_{1A}^2 - a_{1B} \, r_{1B}^2 - a_{2A} \, r_{2A}^2 - a_{2B} \, r_{2B}^2 - a_{12} \, r_{12}^2} (23)$$

with integers n_i and real parameters a. Among all the integrals represented by the above formula, we can distinguish two subsets that can be evaluated analytically. The first subset contains the regular ECG integrals with the non-negative even integers n_i such that $\sum_i n_i \leq \Omega_1$, where the shell parameter $\Omega_1 = 0, 2, 4, \ldots$ These integrals can be generated by differentiation over a-parameters of the following master integral

$$f(0,0,0,0,0) = X^{-3/2}e^{-R^2\frac{Y}{X}},$$
(24)

where

$$X = (a_{1A} + a_{1B} + a_{12})(a_{2A} + a_{2B} + a_{12}) - a_{12}^{2}$$
(25)

$$Y = (a_{1B} + a_{1A}) a_{2A} a_{2B} + a_{1A} a_{1B} (a_{2A} + a_{2B})$$

$$+ a_{12} (a_{1A} + a_{2A})(a_{1B} + a_{2B}).$$
(26)

Each differentiation raises one of the n_i exponents by two. The second subset of integrals permits a single odd index $n_i \geq -1$ for which $\sum_i n_i \leq \Omega_2$ ($\Omega_2 = -1, 1, 3, \ldots$). These so-called *Coulomb* ECG integrals can also be obtained analytically by differentiation of another master integral. For instance, when $n_1 = -1$ the master integral reads

$$f(-1,0,0,0,0) = \frac{1}{X\sqrt{X_1}} e^{-R^2 \frac{Y}{X}} F\left[R^2 \left(\frac{Y_1}{X_1} - \frac{Y}{X}\right)\right],\tag{27}$$

where $X_1 = \partial_{a_{1A}}X$, $Y_1 = \partial_{a_{1A}}Y$, and F(x) = erf(x)/x.

In the standard use of ECG functions the *regular* integrals with $\Omega_1=2$ and *Coulomb* with $\Omega_2=-1$ are sufficient to evaluate matrix elements of the electronic Schrödinger equation (1) and thus to perform calculations of the nonrelativistic energy of the ground state in molecular hydrogen. If additionally analytic gradient minimization is employed, the integrals with $\Omega_1=4$ and $\Omega_2=1$ are required. Such nonrelativistic calculations have been widely used for many atomic and molecular systems [9].

The molecular ECG integrals, as opposed to the atomic ones, have no known analytic form when two or more n_i are odd. Such *extended* integrals originate from regularization of the relativistic operators, for example from V^2 in Eq. (11), or from matrix elements of the nonrelativistic Hamiltonian with rECG basis. The algorithm for numerical evaluation of this *extended* type of integrals relies on the following relation,

which decreases one of the indices by one

$$f(n_1 - 1, n_2, n_3, n_4, n_5) = \frac{2}{\sqrt{\pi}} \int_0^\infty dy f(n_1, n_2, n_3, n_4, n_5) \Big|_{a_{1A} \to a_{1A} + y^2}.$$
(28)

The right-hand-side f is understood as the integral $f(n_1,n_2,n_3,n_4,n_5)$ evaluated with the a_{1A} parameter replaced by $a_{1A}+y^2$. The transformation y=-1+1/x converts the infinite integration domain to the finite interval (0,1) for which an m-point generalized Gaussian quadrature with logarithmic end-point singularity [10] is applied

$$\int_0^1 dx \left[W_1(x) + \ln(x) W_2(x) \right]$$

$$= \sum_{i=1}^m w_i \left[W_1(x_i) + \ln(x_i) W_2(x_i) \right]. \tag{29}$$

The $W_{1,2}$ are arbitrary polynomials of maximal degree m-1, w_i are weights, and x_i are nodes. In terms of this quadrature the integral (28) can be approximated by the formula

$$f(n_1 - 1, n_2, n_3, n_4, n_5) = \frac{2}{\sqrt{\pi}} \sum_{i=1}^{m} w_i (y_i + 1)^2 \quad (30)$$
$$\times f(n_1, n_2, n_3, n_4, n_5) \Big|_{a_{1A} \to a_{1A} + y_i^2}.$$

This quadrature is very efficient for extended integrals with two odd indices, for which typically only m=30 nodes allows about 16 significant digits to be obtained. These extended integrals are sufficient not only for all the relativistic operators with the regularization applied to ECG wave function, but also for the modified regularization (17)-(18) of $H_{\rm rel}$ with rECG wave function.

Nevertheless, calculations with rECG wave function of the expectation values of the individual relativistic operators involve *extended* integrals with three odd indices. They can be obtained by the double numerical integration of *Coulomb* ECG integrals over 30^2 nodes to achieve numerical precision of about 16 significant digits. This two dimensional integration is numerically stable but time consuming.

V. CALCULATIONS OF RELATIVISTIC CORRECTIONS

Relativistic corrections to the BO potential were calculated according to Eqs. (4), (12), and (17). In order to demonstrate the convergence of these three different approaches with ECG functions we compared results at R=0, i.e. for the helium atom, to the results obtained with explicitly correlated exponential (ECE) functions. Calculations with ECE functions are well known in literature (see e.g. Refs. [11] and [12]) and may serve as an excellent reference point and a rigorous test of the convergence of ECG results. The numerical values presented in Table I were obtained with 128, 256, 512, and 1024 ECG basis functions. Direct and standard regularization methods

were used with ECG functions, whereas the modified regularization methods were used with rECG functions. We observe a significant enhancement of numerical convergence of relativistic operators obtained with rECG basis. The total relativistic correction with N=1024 is accurate to 9 digits in the rECG basis, and to 5-6 digits in the ECG basis. A similar enhancement is observed for H_2 at R = 1.4 in Table II, where we compared our results for Dirac- δ functions with those obtained in the ECE basis. The accuracy of the extrapolated value for the total relativistic correction is estimated to have at least eight significant digits after the decimal point. In Table III we provide results for the nonrelativistic energy \mathcal{E} , for the relativistic correction $\mathcal{E}_{\mathrm{rel}}$, and for all four individual components of the relativistic correction evaluated at $R \in (0, 10)$ a.u. with the 1024-term basis of rECG functions. Our results for the overall relativistic correction $\mathcal{E}_{\mathrm{rel}}$ for H_2 are estimated to have eight significant decimal digits.

VI. VIBRATIONAL AVERAGING

In order to obtain the final value of the α^2 relativistic component of the dissociation energy, we solved the radial Schrödinger equation for two potentials. The first potential, used as a reference, is the nonrelativistic (BO) potential $\mathcal{E}(R)$ [14], which yielded the nonrelativistic (BO) energy level E. The second was the potential augmented by the relativistic correction $\mathcal{E}_{\mathrm{rel}}(R)$, which gave the eigenvalue corresponding to the relativistic energy level $E+E^{(2)}$. The difference between both the eigenvalues $E^{(2)}$ is the relativistic correction to molecular levels.

To establish reliable uncertainties for the final results we studied two sources of error: the convergence of single point calculations and the polynomial interpolation. As mentioned above, the relativistic correction was evaluated using basis sets of increasing size, which permitted a detailed analysis of the convergence at each internuclear distance (see Tab. II). From this analysis we estimated that in the vicinity of the equilibrium distance the $\mathcal{E}_{\mathrm{rel}}(R)$ bears an uncertainty of $7 \cdot 10^{-9}$ a.u. equivalent to $2 \cdot 10^{-8}$ cm⁻¹. The influence of the density of the points, at which $\mathcal{E}_{\mathrm{rel}}(R)$ was evaluated, on the accuracy of the final result was assessed by doubling the number of points, which however were calculated only with 512-term basis. As a consequence $E^{(2)}$ was shifted by $5 \cdot 10^{-7}$ cm⁻¹. The related uncertainty due to the selection of the degree of the interpolation polynomial was also investigated. By changing the degree in the range 5-12, we observed changes in the relativistic correction at the level of 10^{-7} cm⁻¹. To summarize, the largest contribution to the uncertainty of the relativistic correction $E^{(2)}$ comes from the limited number of points (and the necessity of interpolation) at which the relativistic potential was evaluated. The final relative uncertainty is assumed to be smaller than 10^{-6} cm⁻¹.

VII. RESULTS AND SUMMARY

Results of our calculations for the dissociation energy and the two selected most accurately measured transitions in H_2 and D_2 are presented in Tables IV and V. The nonrelativistic energy E for H_2 was calculated by solving the full nonadiabatic Schrödinger equation in the exponential basis [3], whereas for D_2 using the NAPT [16] expansion with the neglect of $O(1/\mu)^3$ terms. All the corrections were obtained within the adiabatic approximation. The relativistic correction $E^{(2)}$ was evaluated and reported in this work. The leading QED correction $E^{(3)}$ was obtained in Ref. [17], while the higher-order QED, namely $E^{(4)}$, in Ref. [4]. $E^{(5)}$ was estimated from the correction analogous to that of atomic hydrogen with the assumption that it is proportional to the electronnucleus Dirac- δ , and the related uncertainty was assigned to be 50%.

Our results for the leading relativistic corrections significantly differ from those by Piszczatowski *et al.* [5], whose compilation partially relies on the former calculations by Wolniewicz [8]. For example, our relativistic correction to D_0 of H_2 is $-0.533\,121(1)\,\mathrm{cm}^{-1}$ whereas Piszczatowski et al. reported $-0.531\,9(3)\,\mathrm{cm}^{-1}$. Interestingly, our result is closer to that obtained by Wolniewicz $-0.533\,0\,\mathrm{cm}^{-1}$, despite the differences at the level of individual operators. In our opinion, these differences come from the much more accurate calculation of relativistic matrix elements performed here.

Most importantly, our final theoretical predictions for D_0 are now in disagreement with experimental values, in contrast to the previous theoretical results [5]. This disagreement probably comes from the underestimation of the relativistic nuclear recoil correction. We have previously assumed that these corrections are of the order of the ratio of the electron mass to the reduced mass of the nuclei, which for H_2 is $\sim 10^{-3}$. This might be incorrect because the nonrecoil relativistic correction is anomalously small. This assertion is supported by the example of the helium atom. The nonrecoil relativistic correction to the ⁴He ionization energy is 16 904.024 MHz, while the nuclear recoil is -103.724 MHz, so the ratio is $6 \cdot 10^{-3}$, which is an order of magnitude higher than the estimate based on the helium mass ratio 10^{-4} . On the other hand, in the separated atoms limit the relativistic recoil correction exactly vanishes. Therefore, at present this correction cannot be reliably estimated.

If the relativistic nuclear recoil correction in $\rm H_2$ is underestimated, the difference between our predictions and the experimental values for $\rm D_2$ should be smaller than that for $\rm H_2$ and this is really the case (*vide* Tables IV and V). We emphasize that our theoretical predictions should be treated as preliminary until the relativistic nuclear recoil corrections are reliably calculated. In fact, such corrections have already been obtained by Stanke and Adamowicz [19] for purely vibrational states. However, their result for the total relativistic dissociation energy of $\rm H_2$ (with $E^{(2)} = -0.5691\,\rm cm^{-1}$), when augmented by missing higher order corrections, yields $\rm D_0 = 36\,118.0318\,cm^{-1}$, which differs from the experimental value by as much as $\rm 0.038\,cm^{-1}$, so its numerical uncertainty is out of control. We plan to calculate these nonadia-

batic corrections using the fully nonadiabatic wave function in an exponential basis as in Ref. [3] or by using nonadiabatic perturbation theory (NAPT) [13]. Certainly, this calculation has to be performed to resolve discrepancies with H_2 experiments.

In conclusion, the former excellent agreement of theoretical predictions with experimental D_0 values was accidental and the improved calculations of the leading relativistic corrections result in a few σ disagreements with experimental values for dissociation energies and transition energies, which is probably caused by the unknown relativistic nuclear recoil (nonadiabatic) effects in the relativistic corrections.

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TABLE I. Convergence of matrix elements of relativistic operators at ${\cal R}=0$ a.u. (helium atom limit).

Basis	Direct Standard regularization		rECG (+modified regularization)	
		\mathcal{E}		
128	-2.903724368357561		-2.903724366011805	
256	-2.903724376781020		-2.903724376765067	
512	-2.903724377031170		-2.903724377030040	
1024	-2.903724377034103		-2.903724377034089	
∞ -Slater a			-2.903724377034119598(1)	
		$p_1^4 + p_2^4$		
128	108.103812847	108.178260879	108.175893984	
256	108.149717136	108.176705311	108.176119036	
512	108.171069063	108.176261126	108.176133296	
1024	108.174593664	108.176173934	108.176134411	
∞ -Slater a			108.17613445(1)	
		$\delta^3(r_1) + \delta^3(r_2)$		
128	3.61807292223	3.62085592707	3.62085250490	
256	3.61983231429	3.62085832771	3.62085826322	
512	3.62066249308	3.62085862304	3.62085861086	
1024	3.62079894559	3.62085863628	3.62085863616	
∞ -Slater a			3.62085863700(1)	
		$\delta^3(r)$		
128	0.106521423626	0.106345075042	0.106345517181	
256	0.106391759156	0.106345347318	0.106345416874	
512	0.106355477797	0.106345369617	0.106345375554	
1024	0.106348511028	0.106345370530	0.106345370708	
∞ -Slater a			0.106345370634(1)	
		$p_1^i \left(\frac{\delta^{ij}}{r} + \frac{r^i r^j}{r^3} \right) p_2^j$		
128	0.278 191	140 60	0.27818821108	
256	0.278 189	53641	0.27818896140	
512	0.278 189	388 13	0.27818933907	
1024	0.278 189	381 79	0.27818938036	
∞ -Slater a			0.27818938108(1)	
		$\mathcal{E}_{ ext{rel}}$		
128	-1.95091394168	-1.95203089354	-1.95174292889	
256	-1.95153123551	-1.95182724821	-1.95175366041	
512	-1.95170604993	-1.95177065311	-1.95175469640	
1024	-1.95173983039	-1.95175970900	-1.95175476523	
∞ -Slater a			-1.951754768(1)	

 $[^]a$ The reference values were evaluated with the atomic ECE basis functions $\phi = \exp(-\alpha r_1 - \beta r_2 - \gamma r)$.

TABLE II. Convergence of matrix elements of relativistic operators at the equilibrium distance R=1.4 a.u.

Basis	Direct	Standard regularization	rECG (+modified regularization)
		\mathcal{E}	
128	-1.174 475 6	-1.174475640130736	
256	-1.174 475 7	711 731 700	-1.174475711200533
512	-1.174 475 7	714 117 150	-1.174475714015654
1024	-1.174 475 7	714 217 171	-1.174475714203071
∞ -J ${\sf C}^a$			-1.1744757142204434(5)
		$p_1^4+p_2^4$	
128	13.214563	13.2387714	13.237780507
256	13.231849	13.2383081	13.237929826
512	13.235568	13.2380459	13.237954576
1024	13.237266	13.2379814	13.237956021
∞	13.2387(10)	13.237960(16)	13.23795618(7)
		$\sum_{a,X} \delta^3(r_{aX})$	
128	0.9175507	0.91933127850	0.91933132106
256	0.9188789	0.91933592794	0.91933574110
512	0.9191535 0.91933617260		0.91933612742
1024	0.9192855 0.91933621005		0.91933619174
∞	0.91937(7)	0.9193362112(18)	0.919336206(7)
JC^b			0.919336211(2)
		$\delta^3(r)$	
128	0.016742915	0.016742915953	0.016743529776
256	0.016771639	0.016743229495	0.016743316316
512	0.016750461	0.016743274514	0.016743287361
1024	0.016745258 0.016743277598		0.016743278963
∞	0.0167435(4) $0.0167432781(7)$		0.0167432783(5)
JC^b			0.0167432783(3)
		$p_1^i \Big(rac{\delta^{ij}}{r} + rac{r^i r^j}{r^3}\Big) p_2^j$	
128	0.095 27	0.095 271 222 30	
256	0.095 26	9 308 16	0.09526858845
512	0.095 26	9 010 11	0.09526888381
1024	0.095 268 989 79		0.09526897683
∞	0.095 268 986(6)		0.095268987(4)
		$\mathcal{E}_{ ext{rel}}$. ,
128	-0.20534230	-0.20580042	-0.20567223422
256	-0.20555444	-0.20573326	-0.20568563769
512	-0.20565450	-0.20569982	-0.20568836337
1024	-0.20567565	-0.20569167	-0.20568851645
∞	-0.205682(8)	-0.205689(2)	-0.205688526(7)

 $[^]a$ Evaluated with James-Coolidge wave function, Ref. [14]. b Evaluated in this work.

TABLE III. The electronic energy \mathcal{E} , expectation values of individual relativistic operators, and the relativistic correction of Eq. (17) evaluated with 1024-term rECG basis for H₂ (all entries in a.u.).

R	E	$p_1^4 + p_2^4$	$\sum_{a,X} \delta^3(r_{aX})$	$\delta^3(r)$	$p_1^i \left(\frac{\delta^{ij}}{r} + \frac{r^i r^j}{r^3} \right) p_2^j$	$\mathcal{E}_{ ext{rel}}$
0.0	∞ 17.104.040.505.799.0	108.176 134 41	7.241 717 273 9	0.106 345 370 636	0.278 189 381 06	-1.951754765
0.05	17.104 840 595 733 8	97.883 437 11	6.489 043 177 5	0.105 003 823 908	0.276 266 361 42	-1.850717847
0.1	7.127 216 731 179 9	87.359 441 20	5.770 597 148 6	0.101 570 564 053	0.271 273 759 54	-1.672040901
0.2	2.197 803 295 242 6	69.106 095 40	4.568 931 840 2	0.091 368 192 666	0.255 825 956 08	-1.302271866
0.4	-0.1202303411732	44.92674848	3.007 974 262 0	0.068 873 279 242	0.218 115 094 21	-0.783614393
$0.6 \\ 0.8$	-0.7696354294740 -1.0200566663407	31.45554441 23.55141269	2.1355542202 1.6165590023	0.050763240175 0.037667878063	0.18303648908 0.15386526748	-0.509463149 -0.363237148
1.0	-1.0200300003407 -1.1245397195256	18.631 892 59	1.288 195 811 0	0.028 345 276 112	0.13020020748 0.13021059568	-0.303237148 -0.281549312
1.1	-1.1243397193230 -1.1500573677202	16.862 669 10	1.168 538 437 1	0.028345270112 0.024725634292	0.120 121 309 61	-0.254680537
$1.1 \\ 1.2$	-1.1649352434217	15.417 026 26	1.069 918 912 8	0.024 723 034 292	0.12012130901 0.11101250931	-0.234030337 -0.234013225
1.3	-1.1723471490151	14.225 963 31	0.9879494323	0.019 008 309 934	0.11101230931 0.10276409419	-0.218043755
1.4	-1.1723471490131 -1.1744757142023	13.23795602	0.919 336 191 7	0.016 743 278 963	0.095 268 976 86	-0.218049799 -0.205688516
1.4011		13.22806227	0.9186457757	0.016 720 183 220	0.095 190 366 90	-0.205569553
1.45	-1.1744703313700 -1.17440570714499	12.807 526 10	0.889 230 483 6	0.015 730 076 395	0.09177395700	-0.200610272
1.5	-1.1728550795518	12.414 016 28	0.8615729317	0.014 787 413 515	0.088 432 824 23	-0.196156821
1.6	-1.1685833733462	11.724 276 68	0.8127287476	0.013 090 471 717	0.082 173 146 96	-0.188864898
1.7	-1.1624587268749	11.145 591 08	0.771 298 695 7	0.011611319478	0.076 418 016 59	-0.183376700
1.8	-1.1550687375868	10.659 827 14	0.736 097 157 4	0.010 316 180 045	0.071 104 900 35	-0.179362896
1.9	-1.1468506970016	10.25263362	0.706 180 601 1	0.009 177 235 715	0.066 179 438 68	-0.176571891
2.0	-1.1381329571022	9.912 536 44	0.680 790 743 4	0.008 171 495 990	0.061 594 536 96	-0.174809213
2.1	-1.1291638360667	9.630 266 86	0.659 312 181 5	0.007 279 910 924	0.05730939671	-0.173922389
2.2	-1.1201321168151	9.39825583	0.641 240 366 5	0.006 486 644 310	0.053 288 924 14	-0.173790035
2.3	-1.1111817651695	9.21024640	0.6261570146	0.005778509183	0.04950304198	-0.174313461
2.4	-1.1024226059759	9.06099378	0.6137109920	0.005144501580	0.04592628003	-0.175410463
2.5	-1.0939381299201	8.94602998	0.6036032828	0.004575430210	0.04253733371	-0.177010457
2.6	-1.0857912373625	8.86147693	0.5955750005	0.004063603931	0.03931882634	-0.179050818
2.7	-1.0780284841479	8.80389842	0.5893978526	0.003602589090	0.03625688742	-0.181473897
2.8	-1.0706832334498	8.77018399	0.5848665704	0.003186984950	0.03334102116	-0.184225040
2.9	-1.0637780087717	8.75745814	0.5817929471	0.002812250937	0.03056370383	-0.187250948
3.0	-1.0573262688383	8.76301572	0.5800014505	0.002474548852	0.02792005906	-0.190498823
3.2	-1.0457996613902	8.81876665	0.5796085282	0.001897578330	0.02302537071	-0.197450152
3.4	-1.0360753951531	8.91801344	0.5824483317	0.001434600771	0.01865530993	-0.204664704
3.6	-1.0280463083390	9.04310919	0.5873998954	0.001068345909	0.01482089794	-0.211757192
3.8	-1.0215497953795	9.17885316	0.5934913579	0.000783739899	0.01152959176	-0.218405205
4.0	-1.0163902529178	9.31327923	0.5999421995	0.000566901578	0.00877312169	-0.224378488
4.2	-1.0123599596533	9.43806455	0.6061874544	0.000404933730	0.00652243889	-0.229550124
4.4	-1.0092565162188	9.54839599	0.6118724265	0.000286163107	0.00472959559	-0.233888328
4.6	-1.0068952237883	9.64234777	0.6168193211	0.000200467812	0.00333424931	-0.237433284
4.8	-1.0051160060122	9.72004626	0.620 980 427 9	0.000 139 467 088	0.00227142658	-0.240269571
5.0	-1.0037856585418	9.78286262	0.624 391 465 4	0.000 096 514 570	0.00147803156	-0.242501813
5.2	-1.0027968162807	9.83276296	0.627 132 969 7	0.000 066 524 898	0.000 897 184 78	-0.244236803
5.4	-1.0020650571894	9.871 870 41	0.629 303 290 7	0.000 045 719 642	0.000 480 298 37	-0.245573020
5.6	-1.0015252518178	9.902 201 97	0.631 001 580 7	0.000 031 354 765	0.000 187 505 16	-0.246595529
5.8	-1.0011278808276	9.92554176	0.632 318 772 1	0.000 021 470 798	-0.00001291240	-0.247374807
6.0	-1.0008357076028	9.943 392 93	0.633 333 420 5	0.000 014 686 948	-0.00014560665	-0.247967362
6.5	-1.000 400 547 946 1	9.971 478 81	0.634 946 087 8	0.000 005 668 125	-0.000 293 186 06	-0.248899471
7.0	-1.0001979144266	9.985 482 43	0.635 760 762 7	0.000 002 183 134	-0.000 307 901 17	-0.249373815
7.5	-1.0001021060380	9.99243990	0.636 169 798 9	0.000 000 839 980	-0.00027491015	-0.249621706
8.0	-1.0000556046110	9.995 922 78	0.636 376 259 1	0.000 000 323 028	-0.00023037686	-0.249756677
8.5	-1.0000321717017	9.997 700 01	0.636 482 218 0	0.000 000 124 123	-0.000 188 104 44	-0.249834155
9.0	-1.0000197816909	9.998 632 62	0.636 537 991 1	0.000 000 047 661	-0.00015228986	-0.249881211
9.5	-1.0000128559933	9.999 142 48	0.636 568 497 4	0.000 000 018 282	-0.00012334587	-0.249911507
10.0	-1.0000087556935	9.999 434 61	0.636 586 008 8	0.000 000 007 007	-0.00010039054	-0.249932046
<u> </u>	-1.0	10.0	0.636 619 772 3	0.0	0.0	-0.25

TABLE IV. Contributions to the dissociation energy D_0 and two selected most accurate experimental transitions in H_2 (in cm⁻¹). There are additional 10^{-3} relative uncertainties on $E^{(2)}$, $E^{(3)}$, and $E^{(4)}$ terms due to the BO approximation, which are included in the final result only.

Contrib.	D_0	$S_3(3)$	$Q_{1}(0)$
\overline{E}	36 118.797 746 12(5)	12 559.749 918 95(8)	4 161.163 977 09(6)
$E^{(2)}$	$-0.533121(1)^a$	0.065366	0.023397
$E^{(3)}$	-0.1948(2)	-0.06573(6)	-0.02129(2)
$E^{(4)}$	-0.002067(6)	-0.000599	-0.000192
$E^{(5)}$	0.00012(6)	0.000037(19)	0.000012(6)
E_{FS}	-0.000031	-0.000010	-0.000003
Total	36 118.067 8(6)	12559.74898(8)	4161.16590(3)
Exp.	$36118.06962(37)^b$	$12559.74952(5)^c$	$4161.16636(15)^d$
Diff.	0.0018(6)(4)	0.00054(8)(5)	0.00046(3)(15)

 $[^]a$ For comparison, Wolniewicz [8] obtained -0.5330, Piszczatowski et al. [5] -0.5319, and Stanke et al. [19] $-0.5691 \, \mathrm{cm}^{-1}$ (the latter value comes from nonadiabatic calculations).

TABLE V. Contributions to the dissociation energy D_0 and two selected most accurate experimental transitions in D_2 (in cm⁻¹). There are additional $5 \cdot 10^{-4}$ relative uncertainties on $E^{(2)}$, $E^{(3)}$, and $E^{(4)}$ terms due to the BO approximation, which are included in the final result only.

Contrib.	D_0	$S_2(2)$	$Q_1(0)$
\overline{E}	36 749.090 98(8)	6 241.120 96(30)	2 993.614 88(15)
$E^{(2)}$	-0.529170(1)	0.040057	0.017677
$E^{(3)}$	-0.1982(2)	-0.03315(3)	-0.01539(2)
$E^{(4)}$	-0.002096(6)	-0.000299	-0.000139
$E^{(5)}$	0.00012(6)	0.000019(10)	0.00009(5)
E_{FS}	-0.000204	-0.000032	-0.000015
Total	36748.3614(4)	6241.12755(30)	2993.61702(15)
Exp.	$36748.36286(68)^a$	$6241.12764(2)^b$	$2993.61706(15)^c$
Diff.	0.0015(4)(7)	0.00009(30)(2)	0.00004(15)(15)

^a Ref. [22]; ^b Ref. [23]; ^c Ref. [21].

^b Ref. [18]; ^c Ref. [20]; ^d Ref. [21].