Constraints on extra dimensions from precision molecular spectroscopy

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Abstract

Accurate investigations of quantum-level energies in molecular systems are shown to provide a testing ground to constrain the size of compactified extra dimensions. This is made possible by recent progress in precision metrology with ultrastable lasers on energy levels in neutral molecular hydrogen (H2, HD, and D2) and molecular hydrogen ions (H2+, HD+, and D2+). Comparisons between experiment and quantum electrodynamics calculations for these molecular systems can be interpreted in terms of probing large extra dimensions, under which conditions gravity will become much stronger. Molecules are a probe of spacetime geometry at typical distances where chemical bonds are effective (i.e., at length scales of an Å). Constraints on compactification radii for extra dimensions are derived within the Arkani-Hamed-Dimopoulos-Dvali framework, while constraints for curvature or brane separation are derived within the Randall-Sundrum framework. Based on the molecular spectroscopy of D2 molecules and HD+ ions, the compactification size for seven extra dimensions (in connection to M-theory defined in 11 dimensions) of equal size is shown to be limited to \( R_7 < 0.6 \mu m \). While limits on compactification sizes of extra dimensions based on other branches of physics are compared, the prospect of further tightening constraints from the molecular method is discussed.

1. Introduction

A standard description of the world is usually presented in terms of the observable 3+1 spatiotemporal dimensions. However, string theories have been developed, seeking to produce a consistent description of the Standard Model of physics including the phenomenon of gravity, which appears to be most consistent if large numbers of dimensions are postulated. A 26-dimensional space time was deemed necessary for bosonic strings [1], and a ten-dimensional spacetime was deemed necessary for type-II [2, 3] and heterotic strings [4]. The latter theories are closely related to a mysterious theory called M-theory, which lives in 11 dimensions [5]. In contrast, classical physics requires three spatial dimensions (e.g., to accommodate Newton’s inverse square law, as argued already by Immanuel Kant). Ehrenfest has shown that atoms only exhibit stable orbits in a three-dimensional space [6]. These contradictions between requirements from classical and quantum physics for a three-dimensional space and the possibility of a theory involving higher dimensions were resolved in 1926 by Klein, who invoked the concept of compactification [7].

In the present study the accurate results from precision measurements on molecules are exploited to constrain existing theories of higher dimensions. For molecular systems, state-of-the-art quantum-level calculations of the molecular ions H2+, HD+, and D2+, which are all fundamental three-particle Coulomb systems, have reached such precision that the uncertainty becomes limited by the precision at which values of the fundamental mass ratios, \( m_p m_e \) and \( m_p m_e \), are known [8–10]. However, the recently improved determination of \( m_p m_e \) [11] demonstrates active progress on the experimental side. While experiments on the re-vibrational
spectrum of the H₂⁺ isotopomer are still under way [12], the small dipole moment of the HD⁺ isotopomer has enabled the accurate study of electric-dipole-allowed transitions in various bands [13–15].

In recent years, great progress has also been made in the calculation of level energies in the neutral hydrogenic molecules. Accurate Born–Oppenheimer energies have been calculated for the electronic ground state of H₂, HD, and D₂ [16], as well as non adiabatic interactions [17, 18] and relativistic and quantum electrodynamical (QED) corrections [19, 20]. Now a full set of ro-vibrational level energies of all quantum states up to the dissociation limit is available for all three isotopomers [21, 22]. These calculations on the ground electronic quantum levels were tested in experiments measuring the dissociation limits of H₂ [23], D₂ [24], and HD [25]. Further, they were compared to the experimental values for the fundamental vibrational splitting in both H₂ and hydrogen isotopomers [26, 27], to a measurement of the first overtone in H₂ [28, 30] and D₂ [29], a measurement of the second overtone in H₂ [31, 32], and measurements of highly excited rotational levels in H₂ [33]. The results from a variety of experimental precision measurements on both the ionic and neutral hydrogenic molecules are generally in excellent agreement with the QED calculations, within combined uncertainty limits from theory and experiment.

The agreement between experiment and first-principles calculations on the quantum level energies of molecules has inspired an interpretation of these data that goes beyond molecular physics. Since weak, strong, and (Newtonian) gravitational forces have negligible contributions to their quantum-level structure, electromagnetism is the sole force acting between the charged particles within light molecules, and QED is the fully-encompassing framework to perform the calculations. This makes it possible to derive bounds on possible fifth forces between hadrons from molecular precision experiments compared with QED calculations [34, 35].

Theories of higher dimensions were developed with the goal of resolving the hierarchy problem (i.e., the vast difference of scales between that of electro weak unification (1 TeV) and that of the Planck scale (10¹⁶ TeV), where gravity becomes strong). By permitting the leakage of gravity into higher dimensions while keeping the particles and the three forces of the Standard Model in 3+1 dimensions and invoking a compactification range for the extra dimensions exceeding 3+1, two different testable theories were phrased by Arkani-Hamed, Dimopoulos, and Dvali [36] and by Randall and Sundrum [37, 38]. The mathematical formalisms of these theories can be applied to molecular physics test bodies, from which constraints on the compactification distances can be deduced for the former theory, while constraints on the brane separation or curvature can be derived for the latter theory. That is the subject of the present paper.

2. The ADD model

The theory formulated by Arkani-Hamed, Dimopoulos, and Dvali [36], referred to as ADD theory (see also Antoniadis et al [39]) intends to establish an effective Planck scale to coincide with the electroweak scale by allowing gravity to propagate in extra dimensions. The three forces of the Standard Model, tested at very short distances in particle and atomic physics experiments, are considered to act locally within a three-brane (three spatial dimensions and a time dimension) embedded in a higher dimensional bulk, where gravity may act and allow gravitons to escape. By this process in ADD, the hierarchy problem is nullified, and the so-called desert range between the electroweak scale (Mₚₜ) of 1 TeV and the Planck scale (Mₚ) of 10¹⁶ TeV is avoided. The extension of the extra dimensions is necessarily limited in the case of the flat metrics considered in ADD, since Cavendish-type experiments have proven that gravity obeys the Newtonian 1/r² potential beyond the range of 1 cm [40]. Hence, the extra dimensions are considered to be compactified within a range parameter, Rₙ. While in principle the extra dimensions could exhibit differing range parameters, in the ADD formalism and in the present analysis, this difference is not made.

The Newtonian gravitational potential may be written as:

\[ V_{\text{SI}}(r) = -\frac{m_1 m_2}{r} = -\frac{m_1 m_2}{M_{\text{Planck}}^2} \frac{1}{r} \]

with the Planck mass defined as \( M_{\text{Planck}}^2 = \hbar c / G \) in SI units. In the following discussions, we adopt the natural units, \( \hbar = c = 1 \), and drop the \( (\hbar c) \)-factor in the potentials. The extra \( n \) spatial dimensions proposed in the ADD theory result in a modification of Newtonian gravity for distances shorter than the compactification length range, which is consistent with Gauss law:

\[ V_{\text{ADD}} = -\frac{m_1 m_2}{M_{(4+n)}^{n+2}} \frac{1}{r^{n+1}} \]

where the subscript 4 represents the known (3 + 1) spacetime dimensions, and \( M_{(4+n)} \) is the full higher-dimensional Planck mass. For separations larger than the compactification length, \( r > R_n \), the ADD potential should correspond to the Newtonian \( 1/r \) form.
\[ V_{ADD} = -\frac{m_1 m_2}{M_{n+2}(R_n)^n} r. \]  

(3)

To be more precise, \((R_n)^n\) should be the compactified volume of the extra dimensions, \(V_{n}\); thus a factor of order unity might be included for a specific compactification geometry.

The Planck mass, \(M_{Pl}\), is then related to the higher-dimensional mass \(M_{(n+0)}\), via:

\[ M_{Pl}^n = M_{(n+0)}^n. \]  

(4)

Thus the fundamental mass, \(M_{(n+0)}\), may still be small, and \(M_{Pl}\) becomes large due to the compactified volume of extra dimensions. Arkani-Hamed et al. have shown that if the fundamental mass is taken as \(M_{EW}\), one extra dimension would have a range of order of \(10^{10}\) km to account for the weakness of gravity. This is incompatible with the experimental evidence. But for two extra dimensions, \(R_n\) would be of submillimeter size [36], and therefore at a range where Newtonian gravity is not firmly tested. In our present study we will not set a certain energy scale, and in particular we do not assume that \(M_{4+n} \sim M_{EW}\). Our goal is to constrain \(R_n\) from molecular physics experiments without theoretical prejudice regarding the fundamental mass scale.

While dealing with molecules, the unit attraction of gravity can be chosen as that between two protons, and a dimensionless gravitational coupling strength is defined as:

\[ \alpha_G = \frac{G m_p^2}{\hbar c}. \]  

(5)

Note that this particular choice of the gravitational coupling constant is equivalent to specifying \(\alpha_G = \left(\frac{m_p}{M_{Pl}}\right)^2 = 5.9 \times 10^{-39}\). Then the Newtonian attraction between two particles consisting of \(N_1\) and \(N_2\) protons or neutrons \((m_n \simeq m_p\) is adopted) can be written as:

\[ V_N(r) = -\alpha_G N_1 N_2 \frac{1}{r}. \]  

(6)

From equation (4), the ADD potential of equation (2) within the compactification radius, \(r < R_n\), may be rewritten as:

\[ V_{ADD}(r) = -\alpha_G N_1 N_2 R_n^n \frac{1}{r^{n+1}}, \]  

(7)

while this potential reduces to normal Newtonian gravity, \(V_N\), for the range outside the compactification length range, \(r > R_n\).

For molecules, this gravitational potential has an effect on the level energy of a molecular quantum state with wave function \(\Psi(r)\), to be written as an expectation value:

\[ \left\langle \Delta V_{ADD}(n, R_n) \right\rangle = -\alpha_G N_1 N_2 R_n^n \left[ \int_{R_n}^{\infty} \Psi^*(r) \frac{1}{r} \Psi(r) r^2 dr + R_n^n \int_{0}^{R_n} \Psi^*(r) \frac{1}{r^{n+1}} \Psi(r) r^2 dr \right] \]  

(8)

Note that the wave functions are given along a single coordinate, \(r\) (i.e., the vibrational coordinate, which probes the gravitational forces between nucleons). Here the nuclear displacement is separated from electronic motion and the wave function, \(\Psi(r)\), represents the probability that the nuclei in the molecule are at internuclear separation, \(r\). The first integral term represents the ordinary gravitational attraction, which for protons is \(8 \times 10^{-37}\) times weaker than the electrostatic repulsion, and can therefore be neglected. The second integral represents the effect of modified gravity and is evaluated using accurate wave functions for \(H_2\). The wave functions of the \(H_2\) ground electronic state for the \(v = 0\) and 1 vibrational levels are shown in figure 1. In practice, the integration is performed up to \(r = 10\ Å\) since the wave function amplitude is negligible beyond that. Also, at shorter distances of \(r \leq 0.1\ Å\) the wave function amplitude becomes negligible, which is why the second integral in equation (8) converges without additional assumptions. The HD+ \(v = 0, J = 2\) ground electronic state wave function is also displayed in figure 1, showing the larger internuclear distance of the ion with respect to the neutral.

For transitions between quantum states \(\Psi_1\) and \(\Psi_2\), as in spectroscopic transitions in molecules, a differential effect must be calculated:

\[ \left\langle \Delta V_{ADD}(n, R_n) \right\rangle = -\alpha_G N_1 N_2 R_n^n \left[ \left\langle \Psi_1 \left| \frac{1}{r^{n+1}} \right| \Psi_2 \right\rangle - \left\langle \Psi_2 \left| \frac{1}{r^{n+1}} \right| \Psi_2 \right\rangle \right]. \]  

(9)

This equation represents the expectation value for a high-dimensional gravity contribution to transitions in molecules. Here the ADD expectation value is written explicitly as a function of the two relevant parameters: the number, \(n\), of extra spatial dimensions and the compactification scale, \(R_n\). From equation (9), it is clear that a stronger effect can be expected if the difference in wave functions of the two states, \(\Psi_1\) and \(\Psi_2\), is greater. For this
reason, measurements on the dissociation limit in molecules, where $\Psi_1$ is lowest energy bound state and $\Psi_2$ is the noninteracting two-atom limit at $r = \infty$, are the most sensitive probes.

3. The Randall-Sundrum models

Let us now consider the Randall-Sundrum (RS) scenarios, RS-I and RS-II, to approach the physical description of extra dimensions in an alternative manner [37, 38]. In these scenarios, the particles and interactions of the Standard Model are confined in the SM-brane, separated by some distance, $y_c$, from another (hidden) three-brane along one extra dimension, $y$. The branes and the bulk are sources of gravity that were shown to produce an anti de Sitter metric:

$$ds^2 = e^{-2k_y} \eta_{\mu \nu} dx^\mu dx^\nu + dy^2,$$

where $\exp (-k_y y)$ is a so-called warp factor and $k$ is the bulk curvature [37]. The warped metric differentiates the RS models from the ADD model with a flat metric where $k = 0$. Thus, the exponential warp factor in the RS scenarios solves the hierarchy problem alternatively, without requiring large extra dimensions as assumed in the ADD model.

In the RS scenarios the modified gravitational potential between two masses separated by a distance, $r$, in the SM-brane can be expressed as:

$$V_{RS}(r) = -G m_1 m_2 \left( 1 + \Delta_{RS} \right),$$

where $\Delta_{RS}$ is the correction to the Newtonian potential. Callin [41] computed the potential in the framework of the RS-I scenario, obtaining for short distances:

$$\Delta_{RS-1}(r) \approx \frac{4}{3 \pi k r} \frac{1 - e^{-2k_y r}}{1 + \frac{1}{3} e^{-2k_y r}}, \quad kr \ll 1. \quad (12)$$

Here one can distinguish two regimes, $k \chi \ll 1$ and $k \chi \gg 1$, with the result up at leading orders:

$$\Delta_{RS-1}(r) \approx \begin{cases} \frac{2 \chi}{\pi r} + \ldots, & k \chi \ll 1, \\ \frac{4}{3 \pi k r} + \ldots, & k \chi \gg 1. \end{cases} \quad (13)$$

It turns out that the RS potential for long distances ($kr \gg 1$) is not applicable to molecules and is not considered further.

In the RS-II scenario, the hidden three-brane is chosen to be infinitely far ($y_c \to \infty$) from the SM-brane, resulting in an effective model with a single three-brane (SM-brane) in the bulk. In contrast to the ADD model, this solution thus offers the existence of extra dimensions that do not require compactification. For short distances in the RS-II scenario, Callin and Ravndall [42] obtained...
Δ π = + ≪ − r kr kr (14) RS II

for the RS correction. Note the correspondence of equation (14) with that of equation (13) for ky ≫ 1, which is expected since the latter RS-I condition implies the transition to RS-II at infinite brane separation.

From these RS potential corrections, the expectation values of the leading-order shifts of transitions in molecules in the short distance separation (kr ≪ 1) regime are therefore:

$$\Delta \alpha \Psi \Psi \Psi \Psi = - \frac{V k N N k r r}{4} 3^1, \text{ (15) }$$

where $\Delta E_{\text{exp}}$ and $\Delta E_{\text{theory}}$ signify uncertainties of theory and experiment. In all but two cases the values for $\Delta E$ were found to be larger than the discrepancies between theory and experiment, denoted by $\Delta E = E_{\text{exp}} - E_{\text{theory}}$, while the $H_2 \nu = 0 \rightarrow 1$ is within two standard deviations ($\Delta E < 2 \delta E$). From these results it is concluded that QED theory for these molecular systems is in very good agreement with observations. Recent calculations by Korobov et al [43] result in an increased discrepancy with the experimental results of Bressel et al [14] at the level of 2.6 standard deviations, and we do not include the HD+ values in the comparisons.

The agreement between theory and experiment for molecular systems is now translated into a constraining relation for higher dimensions in the ADD framework:

$$\Delta V_{\text{ADD}}(n, R_n) < \delta E.$$  (17)
As a first example we take the measurement on the fundamental vibration in the H$_2$ molecule. This is one of the most accurately measured numbers in neutral molecules, while the QED calculations for this fundamental rotationless transition are also more accurate by an order of magnitude with respect to the absolute binding energies because of the cancellation of errors for nonrotating molecules [26]. Constraints on $R_n$ can be derived via:

$$\left( R_n \right)^n < \frac{\delta E}{\alpha_G N_1 N_2 \Delta}$$

with $\Delta$ being the difference in expectation values over the wave function densities between $v = 0$ and $v = 1$ vibrational states in the molecule:

$$\Delta = \left[ \left\langle \frac{1}{r^{n+1}} \right\rangle_{\Psi_1} - \left\langle \frac{1}{r^{n+1}} \right\rangle_{\Psi_2} \right].$$

Figure 1 plots the wave functions for the lowest vibrational states, in the case of H$_2$ and for $J = 0$, as obtained from ab initio calculations [21, 22]. Since the wave functions are located in the same region of space, the fundamental vibrational transition in the hydrogen molecule ($v = 0 \rightarrow v = 1$) probes only a differential effect. Figure 2 presents the resulting constraints on $R_n$ from the measurement of the fundamental vibration in the H$_2$ molecule for the range of extra dimensions, $n = 1 - 8$. The sloping lines in figure 2 represent calculated $V_{\text{ADD}}/V_N$ for different $n$ and $R_n$ values. The horizontal dashed line, $\delta E/V_N$, indicates limits from molecular spectroscopy. Hence, for certain numbers of extra dimensions, $n$, $R_n$ is constrained to be less than the value where the $V_{\text{ADD}}/V_N$ and $\delta E/V_N$ intersect in the graph. Constraints on $R_n$ obtained from a comparison with the fundamental vibrational transition of H$_2$, are presented in table 2.

Both the experimental and theoretical results for the fundamental vibration in the hydrogen molecule are known to the $10^{-4}$ cm$^{-1}$ level, which is an order of magnitude more accurate than the values for the binding energies [26, 27]. However, for a comparison of dissociation limits, it is no longer a small difference along the internuclear coordinate axis that is probed, but the difference between the 1 Å molecular scale and infinite atomic separation. The expectation value for the ADD contribution to the binding energy of the lowest bound state in the H$_2$ molecule, or the $D_0$ binding energy, is:

$$\left\langle \Delta V_{\text{ADD}} \left( n, R_n \right) \right\rangle = \alpha_G N_1 N_2 R_n \left( \frac{1}{r^{n+1}} \right)_{\Psi_1}.$$ 

By comparing to the experimental findings on $D_0$(H$_2$) [23], this leads to another set of constraints on $R_n$ for $n$ extra dimensions, which are also listed in table 2.

The method was further applied to the fundamental vibration of HD and D$_2$, where the experimental and theoretical uncertainties are similar to those in H$_2$. Although the heavier masses of the isotopomers improve the constraints obtained from H$_2$, as expected from equation (18), the HD and D$_2$ fundamental vibration constraints are still less stringent compared to those from the H$_2$ dissociation limit. The results obtained for D$_2$ dissociation energy [24] lead to the tightest constraints on $R_n$ from the neutrals as listed in table 2, which scale by a factor $(\frac{1}{4})^{1/n}$ relative to H$_2$ due to the mass difference.
The experimental accuracy for the HD⁺ molecular ion transitions is an order of magnitude better than the corresponding neutral molecule system that stems mostly from the possibility of trapping the ionic species. The theoretical calculation for the three-body HD⁺ level energies is also more accurate than those of the neutral molecular hydrogen. However, the internuclear separation of HD⁺ (~1.1 Å) is greater than that of neutral hydrogen molecules (~0.76 Å), as shown in figure 1. Thus, the neutrals are inherently more sensitive as the wave functions probe shorter internuclear distances compared to their ionic counterparts. The constraints for \( R_n \) derived from the HD⁺ (\( v = 0, J = 2 \rightarrow v = 4, J = 3 \)) ro-vibrational transition from Koelemeij et al [13] are listed in table 2. In the table, the \( R_n \) constraints from \( D_2 D_0 \) are the most stringent for \( n = 4, 5, 6 \) extra dimensions, while the constraints from HD⁺ are the most constraining for \( n = 2, 3, 7 \). The higher-dimensional Planck mass, \( M_{4+n} \), and corresponding Planck length, \( R_{PL(4+n)} \), derived from the tightest \( R_n \) constraints obtained in this study are also listed in table 2.

Similarly, we derive constraints pertaining to corrections in the RS scenario with one extra dimension, and the combined uncertainty, \( \delta E \), for a specific molecular transition

\[
\left\{ \Delta V_{RS}(k) \right\} < \delta E .
\]

Using the combined uncertainties for the \( D_0(D_2) \) study, we present constraints for the RS schemes. For the RS-I scenario in the short distance (\( kr \ll 1 \)) regime, we obtain constraints for the brane separation of \( \chi < 1 \times 10^{18} \) m in the limit \( k \chi \ll 1 \) using equation (13). In the limit \( k \chi \gg 1 \) in equation (13), a constraint for the inverse of the curvature of \( 1/k < 2 \times 10^{18} \) m is obtained. For the RS-II model, we obtain constraints for the inverse of the curvature \( 1/k < 2 \times 10^{18} \) m for \( kr \ll 1 \) from equation (15).

5. Comparison with other constraints

The constraints obtained from molecular systems probe length scales in the order of Angstroms. This complements bounds that probe subatomic-to-astronomical length scales obtained from other studies using distinct methodologies. Length scales of several hundred nanometers to microns are probed in Casimir-force studies using cantilevers [44] or atomic-force microscopy [45]. The micrometer-to-millimeter range is probed in torsion-balance type experiments, with the tightest constraint obtained by Kapner et al [46] for a single extra dimension of \( R_1 < 4.4 \times 10^{-5} \) m. The centimeter-to-meter separations are accessed by Cavendish- or Eötvös-type investigations in the laboratory, while astronomical scales can be probed in satellite or planetary orbits that also serve to constrain the universality of free fall and deviations from the gravitational inverse-square law [40]. Constraints for the RS-theories are obtained by Iorio [47] using data from the orbital motions of satellites or astronomical objects, with the tightest constraint for the inverse of the curvature of \( 1/k < 5 \) m obtained from the motion of the GRACE satellite. The latter constraint is in the \( kr \gg 1 \) regime of equation (14) and probes a different distance range to that of molecules (\( kr \ll 1 \)).

Precision spectroscopies of hydrogen [48, 49] and muonic atoms [50, 51] have been interpreted along the same lines in terms of the ADD model [52], resulting in typical constraints of \( R_1 < 10^{-5} \) m. The interpretation is not straightforward because of the proton size puzzle [53]; in fact, the argument has been turned around, where the existence of extra dimensions are instead invoked as a possible solution to the puzzle [54]. In the treatment of atoms, some assumptions had to be made on the wave function density at \( r = 0 \), typical for the s-states involved, causing problems in calculating the second integral of equation (8) over the electronic wave function that has a
significant amplitude at \( r = 0 \) in atoms. Note that these difficulties are absent in molecules, as the molecular wave function probes the 0.1–5 Ångström distance range.

To probe length scales in the subatomic range, one is ultimately limited by the increasing contributions from nuclear structure and the strong interaction (e.g., in neutron scattering studies [55]). In contrast to QED calculations, the most accurate lattice-QCD calculation of light hadron masses only achieves relative accuracies, on the order of a few percent. Nevertheless, the smaller nucleon size presents higher sensitivity to the effects of ADD-type interactions, and constraints for the size of extra dimensions may be extracted. The general method for molecules presented here may be applied to a comparison of \textit{ab initio} lattice-QCD calculations with the measurements of light hadron masses. The corresponding QCD test probes length scales the size of a nucleon, at \( \sim 10^{-15} \) m. The \textit{ab initio} calculations of Dürr \textit{et al.} [56] for the nucleon mass are estimated to be accurate to around 50 MeV/c\(^2\), while the experimental mass values are accurate to 20 MeV/c\(^2\). The calculated nucleon mass, with 3\% relative accuracy, is the isospin average of proton, \( m_p \), and neutron, \( m_n \), masses, while \( m_p \) is known experimentally to be 0.1\% smaller than \( m_n \). Constraints based on experimental nucleon masses and QCD theory have not been explored, but we produce here a rough first estimate by assuming that the three constituent quarks each have an effective mass that is \( \frac{1}{3} \) of the nucleon mass, and have separation distances \( \sim r_p \). Analogous to equation (17) for QED interactions in molecules, the expectation value for an ADD contribution on the mass of the proton can be written as \( V_{\text{ADD}}(p)/c^2 < \delta m_p \), yielding a bound for the case of seven extra dimensions of \( R_7 < 2.4 \times 10^{-16} \) m.

In high-energy particle collisions, higher-dimensional gravitons may be produced that could escape into the bulk, leading to events with missing energy in (3+1)-dimensional spacetime [36, 57, 58]. Based on this premise of an energy-loss mechanism, the phenomenology of the SN 1987 A supernova was investigated, imposing limits on extra dimensions of \( R_2 < 3 \times 10^{-6} \) m, \( R_3 < 4 \times 10^{-7} \) m, and \( R_4 < 2 \times 10^{-8} \) m [59]. Similarly, from a missing energy analysis of proton-colliding events at the Large Hadron Collider (LHC), a constraint for \( R_2 < 3.2 \times 10^{-4} \) m can be extracted from the \( M_{\text{e+e}} = 1.93 \) TeV bound for \( n = 2 \) given in [60]. For comparison, the Planck energy scale in (4 + \( n \)) dimensions in table 2 turns out to be in the range between 1–1000 MeV, but it is derived from a completely independent methodology. Also for \( n > 2 \), the bounds derived from the LHC are nominally more stringent than those from molecules. However, additional assumptions beyond the ADD potential in equation (2) (e.g., the fundamental quantization of gravity, the existence and propagation of gravitons in (4 + \( n \)) dimensions, and postulating the existence of massive new particles), are necessary for an effective theory [57, 58] to interpret the LHC’s missing energy signals. Such assumptions are not needed for the molecular physics bounds, which are not sensitive to physics at very short distances.

### 6. Conclusion and outlook

The alternative approaches for constraining compactification radii for extra dimensions, partially surveyed here, are all complementary as they probe different length and energy scales. Some approaches serve to produce tighter limits, often at the expense of additional assumptions. In the present study, constraints are derived on compactification scales of extra dimensions from precision measurements on molecules, leading to straightforward interpretations. Molecules, particularly the lightest ones such as neutral and ionic molecular hydrogen, exhibit wave functions representing the internuclear distances, with amplitudes confined to the range 0.1–5 Å. Current state-of-the-art experiments on neutral molecular hydrogen determine vibrational splittings of the order of 10^{-8} \text{ cm}^{-1}, or 3 MHz [26]. Since the lifetimes of ro-vibrational quantum states in H\(_2\) are of the order of 10^6 s [61], measurements of vibrational splittings of the order of 10^14 Hz could, in principle, be possible at more than 20-digit precision, which leaves room for improvement ‘at the bottom’ of over 10 orders of magnitude if experimental techniques can be developed accordingly. Similar improvements in theory would make these molecular systems an ideal testing ground for constraining or detecting higher dimensions, as well as fifth forces [34]. After having performed a 15-digit accuracy calculation on Born–Oppenheimer energies [16], calculations of strongly improved accuracy have just been published [62], while improved calculations of nonadiabatic corrections are underway [63]. Immediate improvements, based on existing technologies, the experimental accuracies of the dissociation limits in the neutral hydrogen and its isotopomers [64], and the spectroscopy of HD\(^+\) [12, 65, 66] were discussed recently.

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References

[1] Lovelace C 1971 Phys. Lett. B 34 500
[2] Gliozzi F, Scherk L and Olive D I 1976 Phys. Lett. B 58 282
[3] Green M B and Schwarz J H 1981 Nucl. Phys. B 181 302
[4] Gross D J, Harvey J A, Martinez E and Rohl R 1985 Phys. Rev. Lett. 54 502
[5] Duff M J, Howe P S, Inami T and Stelle K S 1987 Phys. Lett. B 191 70
[6] Ehrenfest P 1920 Ann. Phys. Lpz. 61 140
[7] Klein O 1926 Z. Phys. A 37 995
[8] Korobov V I, Hilico L and Karr J P 2009 Phys. Rev. A 79 012501
[9] Karr J P, Hilico L and Korobov V I 2011 Can. J. Phys. 89 103
[10] Korobov V I and Zhong Z-X 2012 Phys. Rev. A 86 044501
[11] Sturm S, Kohler F, Zatorski I, Wagner A, Harman Z, Werth G, Quint W, Keitel C H and Blaum K 2014 Nature 506 467
[12] Karr J P 2014 J. Mol. Spectrosc. 308 37
[13] Koelmeijer J C J, Roth B, Wicht A, Ernsting I and Schiller S 2007 Phys. Rev. Lett. 98 173002
[14] Bressel U, Borodin A, Shen J, Hansen M, Ernsting I and Schiller S 2012 Phys. Rev. Lett. 108 183003
[15] Koelmeijer J C J, Nooij D W E, de Jong D, Haddad M A and Ubachs W 2012 Appl. Phys. B 107 1075
[16] Pachucki K 2010 Phys. Rev. A 82 032509
[17] Pachucki K and Komasa J 2008 J. Chem. Phys. 129 034102
[18] Pachucki K and Komasa J 2009 J. Chem. Phys. 130 164113
[19] Pachucki K 2005 Phys. Rev. A 71 012503
[20] Pachucki K 2007 Phys. Rev. A 76 022106
[21] Piszczatowski K, Lach G, Przybytek M, Komasa J, Pachucki K and Jeziorski B 2009 J. Chem. Theory-Comp. 5 3039
[22] Komasa J, Piszczatowski K, Lach G, Przybytek M, Jeziorski B and Pachucki K 2011 J. Chem. Theory-Comp. 7 3105
[23] Liu J, Salumbides E J, Hollenstein U, Koelmeijer J C J, Eikema K S E, Ubachs E and Merkt F 2009 J. Chem. Phys. 130 174306
[24] Liu J, Sprecher D, Jungen C, Ubachs W and Merkt F 2010 J. Chem. Phys. 132 154301
[25] Sprecher D, Liu J, Jungen C, Ubachs W and Merkt F 2010 J. Chem. Phys. 133 111102
[26] Dickenson G D, Niu M L, Salumbides E J, Komasa J, Eikema K S E, Pachucki K and Ubachs W 2013 Phys. Rev. Lett. 110 193601
[27] Niu M L, Salumbides E J, Dickenson G D, Eikema K S E and Ubachs W 2014 J. Mol. Spectrosc. 300 44
[28] Campargue A, Kassi S, Pachucki K and Komasa J 2012 Phys. Chem. Chem. Phys. 14 8602
[29] Kassi S, Campargue A, Pachucki K and Komasa J 2012 J. Chem. Phys. 136 184309
[30] Kassi S and Campargue A 2014 J. Mol. Spectrosc. 300 55
[31] Hu S-M, Pan H, Cheng C-F, Su Y R, Li X F, Wang J, Campargue A and Liu A W 2012 Astroph. J. 749 76
[32] Tan Y, Wang J, Cheng C-F, Zhao X-Q, Liu A-W and Hu S-M 2014 J. Mol. Spectrosc. 300 60
[33] Salumbides E J, Dickenson G, Ivanov T and Ubachs W 2011 Phys. Rev. Lett. 107 043005
[34] Salumbides E J, Koelmeijer J C J, Komasa J, Pachucki K, Eikema K S E and Ubachs W 2013 Phys. Rev. D 87 112008
[35] Salumbides E J, Ubachs W and Korobov V I 2014 J. Mol. Spectrosc. 300 65
[36] Arkani-Hamed N, Dimopoulos S and Dvali G 1988 Phys. Lett. B 249 263
[37] Randall L and Sundrum R 1999 Phys. Rev. Lett. 83 3370
[38] Randall L and Sundrum R 1999 Phys. Rev. Lett. 83 4690
[39] Antoniadis I, Arkani-Hamed N, Dimopoulos S and Dvali G 1998 Phys. Lett. B 436 257
[40] Adelberger E G, Heckel B R and Nelson A E 2003 Annu. Rev. Nucl. Part. Sci. 53 77
[41] Callin P 2004 arXiv:hep-ph/0407054
[42] Callin P and Ravndal F 2004 Phys. Rev. D 70 104009
[43] Korobov V I, Hilico L and Karr J P 2014 Phys. Rev. A 89 032511
[44] Zuuriber P, de Man S, Grucza G, Heeck K and Iannuzzi D 2011 New J. Phys. 13 023027
[45] Banishiv A A, Klimchitskaya G L, Mostepanenko V M and Mohideen U 2013 Phys. Rev. Lett. 110 132501
[46] Kapner D L, Cook T S, Adelberger E G, Gundlach J H, Heckel B R, Hoyle CD and Swanson H E 2007 Phys. Rev. Lett. 98 021101
[47] Iorio L 2012 Ann. Phys. (Berlin) 524 371
[48] Biraben F 2009 Eur. Phys. J. Spec. Top. 172 109
[49] Parthey C G et al 2010 Phys. Rev. Lett. 104 233001
[50] Pohl R et al 2010 Nature 466 213
[51] Antognini A et al 2013 Science 339 417
[52] Li Z-G, Ni W-T and Pulido Patón A 2007 Chin. Phys. B 17 70
[53] Pohl R, Gilman R, Miller G A and Pachucki K 2013 Annu. Rev. Nucl. Part. Sci. 63 175
[54] Wang L-B and NiW-T 2013 Mod. Phys. Lett. A 28 1350094
[55] Nesvizhetskii V V, Pignol G and Protasov K V 2008 Phys. Rev. D 77 034020
[56] Dürr S et al 2008 Science 322 1224
[57] Mirabeli E A, Perelstein M and Peskin M E 1999 Phys. Rev. Lett. 82 2236
[58] Giudice G E, Rattazzi R and Wells J D 1999 Nucl. Phys. B 544 3
[59] Cullen S and Perelstein M 1999 Phys. Rev. Lett. 83 268
[60] Aad G et al (Atlas Collaboration) 2013 Phys. Rev. Lett. 110 011802
[61] Black J H and Dalgaroon A 1976 Astroph. J. 203 132
[62] Pachucki K and Komasa J 2014 J. Chem. Phys. 141 224103
[63] Pachucki K 2015 private communication
[64] Sprecher D, Jungen C, Ubachs W and Merkt F 2011 Faraday Discuss. 150 51
[65] Tran V Q, Karr J-P, Douillet A, Koelmeijer J C J and Hilico L 2013 Phys. Rev. A 88 032511
[66] Schiller S, Bakalov D and Korobov V I 2014 Phys. Rev. Lett. 113 023004