Dynamics of the Vacuum and Casimir Analogs to the Hydrogen Atom

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This paper will discuss the current viewpoint of the vacuum state and explore the idea of a “natural” vacuum as opposed to immutable, non-degradable vacuum. This concept will be explored for all primary quantum numbers to show consistency with observation at the level of Bohr theory. A comparison with the Casimir force per unit area will be made, and an explicit function for the spatial variation of the vacuum density around the atomic nucleus will be derived. This explicit function will be numerically modeled using the industry multi-physics tool, COMSOL®, and the eigenfrequencies for the \( n = 1 \) to \( n = 7 \) states will be found and compared to expectation.

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I. INTRODUCTION

The current viewpoint of the quantum vacuum, or vacuum state, is that it is an immutable, non-degradable state for all observers and systems with no structure or variation. The concept of the vacuum state is typically introduced as a ground state of a harmonic oscillator, so the viewpoint that it is immutable is reasonable. How can the vacuum, being the ground state of a harmonic oscillator, be anything other than “zero” for all observers? What if, however, the vacuum could be posited to be a plenum that can be shown to be degradable, and has the capability to support particle-vacuum or particle-particle interactions that allow lower energy, ground states? It is known from experimental observation that the vacuum can exhibit characteristics that can best be associated with a degraded vacuum in the form of the Casimir force [1–10]. The Casimir force arises as a result of a geometric conducting boundary in the form of two plates being placed in close proximity to one another such that the geometry of the cavity can preclude the ability for certain wavelengths of light from being present in one direction. The integral of the spectrum of vacuum fluctuation frequencies in between the two plates has a different starting point than the integral of the spectrum of vacuum fluctuations frequencies outside of the plates. The difference between these two integrals is what is considered to be responsible for the manifestation of this force. In this scenario, the vacuum state between the two plates is considered to be at a lower state than the vacuum state on the outside of the plates, or the toy vacuum model harmonic oscillator is at a lower state than the “zero” state outside of the plates. Perhaps this is illustrating a deficiency in the way in which the vacuum state is described and understood. The Casimir force strongly indicates that the vacuum is degradable, however, this concept is at odds with the idea of a zero state of a harmonic oscillator, so perhaps a prudent path to explore is to consider the concept that the quantum vacuum is a bit more “natural” than a toy harmonic oscillator and can have spatial and temporal variations, and to see if this mutable quantum vacuum identifies with any inconsistencies with observation.

II. VACUUM “DENSITY”

The idea of a vacuum “density” expectation value will be explored by first starting with the Bohr formula that relates allowed energy levels to the primary quantum numbers. The energy for the \( n^{th} \) primary quantum number level of the hydrogen atom is given by the Bohr formula in Equation (1).

\[
E_n = -\frac{m}{2\hbar} \left( \frac{\epsilon^2}{4\pi\epsilon_0} \right)^2 \frac{1}{n^2}
\]

\[
= -\frac{1}{n^2} 13.6 eV, \ n = 1, 2, 3, ...
\]  

(1)

The allowed “orbit” radius for this energy level is shown in Equation (2).

\[
r_n = \frac{4\pi\epsilon_0 n^2 \hbar^2}{\epsilon^2 m_e}
\]

\[
= n^2 5.29 \times 10^{-11} \text{ meters}, \ n = 1, 2, 3, ...
\]  

(2)

Equation (3) shows the energy relationship with the primary quantum number for the \( n^{th} \) level associated with a hydrogen-like atom with the atomic number \( Z \).

\[
E_{Z,n} = -\frac{Z^2}{n^2} 13.6 eV, \ n = 1, 2, 3, ...
\]  

(3)
The allowed “orbit” radius for the energy level for any atom with atomic number \( Z \) is:

\[
r_{Z,n} = \frac{n^2}{Z} \times 5.29 \times 10^{-11} \text{meters}, \quad n = 1, 2, 3, \ldots
\]  

(4)

The historic perspective used in the development of the above relationships was that of the electron being in “orbit” around the nucleus in a quasi-classical sense. It is appropriate to think of these energy states as a wave function [11], so one can calculate an average “density” of this energy state by smearing the energy over a spherical region encapsulated by the allowed radius for that state (see Equation 5).

\[
\langle \rho \rangle = \frac{E_{Z,n}}{c^2 \frac{4}{3} \pi r_{Z,n}^3}.
\]  

(5)

The average density for hydrogen is calculated for the \( n = 1 \) to \( n = 7 \) states in TABLE I.

TABLE I: This table shows the derived “density” of a given energy state \( n \), with \( Z = 1 \)

<table>
<thead>
<tr>
<th>( n )</th>
<th>radius(m)</th>
<th>( E(eV) )</th>
<th>( E(J) )</th>
<th>( \rho ) (kg/m(^3))</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.29 \times 10^{-14}</td>
<td>13.60 \times 10^{-19}</td>
<td>2.176 \times 10^{-19}</td>
<td>3.905 \times 10^{-9}</td>
</tr>
<tr>
<td>2</td>
<td>2.11 \times 10^{-10}</td>
<td>3.40 \times 10^{-20}</td>
<td>5.440 \times 10^{-20}</td>
<td>1.525 \times 10^{-7}</td>
</tr>
<tr>
<td>3</td>
<td>4.76 \times 10^{-10}</td>
<td>1.51 \times 10^{-20}</td>
<td>2.418 \times 10^{-20}</td>
<td>5.952 \times 10^{-9}</td>
</tr>
<tr>
<td>4</td>
<td>8.46 \times 10^{-10}</td>
<td>0.85 \times 10^{-20}</td>
<td>1.360 \times 10^{-20}</td>
<td>5.959 \times 10^{-10}</td>
</tr>
<tr>
<td>5</td>
<td>1.32 \times 10^{-9}</td>
<td>0.54 \times 10^{-20}</td>
<td>8.704 \times 10^{-20}</td>
<td>9.997 \times 10^{-11}</td>
</tr>
<tr>
<td>6</td>
<td>1.90 \times 10^{-9}</td>
<td>0.38 \times 10^{-20}</td>
<td>6.044 \times 10^{-20}</td>
<td>2.325 \times 10^{-11}</td>
</tr>
<tr>
<td>7</td>
<td>2.59 \times 10^{-9}</td>
<td>0.28 \times 10^{-20}</td>
<td>4.441 \times 10^{-20}</td>
<td>6.774 \times 10^{-12}</td>
</tr>
</tbody>
</table>

The primary quantum number \( n \) is only varied from 1 to 7 here.

The question can be raised on what the calculated value for the average “density” really means —does it really indicate a perturbation (rarefaction or densification) of the quantum vacuum, or is it just a number that has no physical interpretation? To help consider this question, an equation can be fitted to the radius and density data presented in TABLE I and the result is shown in Equation (6).

\[
\rho (r) = \frac{2.98968 \times 10^{-46}}{r^4}.
\]  

(6)

A. Casimir Discussion

Equation (6) shows that the density value is dependent on \( 1/r^4 \), which is a similar dependency to the Casimir Force. At this point, the equation should not be viewed as a real continuous function; rather, it should be considered as a function that can only be used with the allowed, discrete values for the radius as input values. Recall that the Casimir force per unit area is \((1/c^2)\) added for unit consistency:

\[
\frac{1}{c^2} \frac{F}{A} = \frac{1 \ h c \pi^2}{c^2 240 d^4}
\]  

(7)

One can use Equation (7) to calculate a Casimir “density” value for the hydrogen primary quantum numbers \( 1 - 7 \) by equating the distance, \( d \), to twice the allowed orbit radius, \( 2r \). In a sense, the electron establishing a “boundary” at this radius could be envisioned as setting up some sort of boundary condition that mimics a Casimir cavity of sorts. These values are calculated and compared to the average density with a ratio provided in TABLE II (again Casimir force per unit area is multiplied by \( 1/c^2 \) for units consistency).

TABLE II: This table compares the derived “density” of a given energy state \( n \), with \( Z = 1 \) to the Casimir density for a cavity with a separation distance of \( 2r_n \).

<table>
<thead>
<tr>
<th>( n )</th>
<th>radius(m)</th>
<th>( \rho ) (kg/m(^3))</th>
<th>Casimir (kg/m(^3))</th>
<th>Ratio(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5.29 \times 10^{-10}</td>
<td>3.91 \times 10^{-9}</td>
<td>1.16 \times 10^{-4}</td>
<td>2.96</td>
</tr>
<tr>
<td>2</td>
<td>2.11 \times 10^{-10}</td>
<td>1.53 \times 10^{-7}</td>
<td>4.51 \times 10^{-7}</td>
<td>2.96</td>
</tr>
<tr>
<td>3</td>
<td>4.76 \times 10^{-10}</td>
<td>5.95 \times 10^{-9}</td>
<td>1.76 \times 10^{-8}</td>
<td>2.96</td>
</tr>
<tr>
<td>4</td>
<td>8.46 \times 10^{-10}</td>
<td>5.96 \times 10^{-10}</td>
<td>1.76 \times 10^{-9}</td>
<td>2.96</td>
</tr>
<tr>
<td>5</td>
<td>1.32 \times 10^{-9}</td>
<td>1.00 \times 10^{-10}</td>
<td>2.96 \times 10^{-10}</td>
<td>2.96</td>
</tr>
<tr>
<td>6</td>
<td>1.90 \times 10^{-9}</td>
<td>2.33 \times 10^{-11}</td>
<td>6.88 \times 10^{-11}</td>
<td>2.96</td>
</tr>
<tr>
<td>7</td>
<td>2.59 \times 10^{-9}</td>
<td>6.77 \times 10^{-12}</td>
<td>2.00 \times 10^{-11}</td>
<td>2.96</td>
</tr>
</tbody>
</table>

\( ^a \) This is the Casimir force per unit area multiplied by \( 1/c^2 \).

\( ^b \) This ratio is the Casimir column value divided by the \( \rho \) column value

TABLE II shows that the values will be equal if there is a factor of \( 1/3 \) added to the Casimir force per unit area \( \times 1/c^2 \). As a short side discussion to help possibly provide an explanation for the \( 1/3 \) factor, consider the Friedmann equation:

\[
\frac{\ddot{a}}{a} = -\frac{4\pi G}{3} \left( \rho c^2 + 3P \right)
\]

In order for there to be a universe that is accelerating as evidenced by observation, the equation of state for the vacuum, \( P = \omega \rho c^2 \) must be such that \( P < -\rho c^2 / 3 \), or \( \omega < -1/3 \) [12]. If the equation of state of the quantum vacuum is just such that \( \omega \sim -1/3 \), this might explain the need for the addition of a factor of \( 1/3 \) in TABLE II. It could be posited that the predicted altered ground state of the vacuum around the Hydrogen nucleus resulting from the presence of the “orbiting” electron may be responsible for helping establish the stable energy state and preclude the “orbiting” electron from radiating down to the nucleus. An approach similar to this was also successfully explored in [13]. Equipped with this insight, one can combine Equation (5) and Equation (7) to yield Equation (8)

\[
\langle \rho \rangle = \frac{E_{Z,n}}{c^2 \frac{4}{3} \pi r_{Z,n}^3} = \frac{1}{3} \frac{h c \pi^2}{240 d^4}
\]  

(8)

The significance of this equation is that it indicates that the calculated “density” expectation value using the Bohr relationships for the energy and radius may have physical meaning as opposed to just a calculated number.
Consider the Casimir force (and Casimir equation) as it has been explored in the lab to date by numerous experimentalists with the work done by Steven Lamoreaux in 1996 establishing convincing experimental evidence of the phenomenon [2]. The Casimir force equation is used in practice to assess the force developed for a given geometry associated with a real test article that will be studied in the laboratory. When the force per unit area equation is applied to the real test article being studied in the lab, the equation provides a number that really does have physical meaning as evidenced by the real force that is measured during the study of the test article. The state of the quantum vacuum between two plates that establish a Casimir cavity is perturbed (rarified) as a result of the plates precluding certain vacuum fluctuation modes from being manifested in the cavity. The pressure inside the cavity is less than the pressure outside of the cavity.

Since the “density” using the Bohr relationships has been shown to make predictions of the energy density around the hydrogen nucleus that are identical to the modified Casimir force per unit area equation, this may indicate that these numerical values do have physical meaning and are not just a numerical calculation with no basis in nature. To be explicit, these values may indicate that the quantum vacuum around the hydrogen nucleus is not an immutable and non-degradable medium with no variation or structure, rather the vacuum appears to have a perturbation around the hydrogen nucleus that exhibits a strong dependency on $1/r^4$.

What if one considers the scenario when the atomic number $Z$ is varied? The course is similar to the above treatment for hydrogen, except the Bohr relationships used are the equations with the $Z$ dependency included. For this discussion, the primary quantum number $n$ will be spanned from 1 to 7, and the atomic number $Z$ will be spanned from 1 to 7, which corresponds to hydrogen, helium, lithium, beryllium, boron, carbon, and nitrogen respectively. The expectation value for the “density” is shown in FIG. 1 for each atomic number. The observed “density” is determined using Equation (5), and the Casimir “density” is determined using Equation (9):

$$\langle \rho \rangle = \frac{Z}{3} \frac{hc\pi^2}{240d^2}.$$  \hfill (9)

A trend line has been added to each series to visually link each set of primary quantum numbers together for a given atomic number $Z$, and help illustrate the $1/r^4$ dependency. A mark is plotted for each combination of $Z$ and $n$ for the “density” expectation value. The modified Casimir force per unit area value is also calculated for each point with the inclusion of $Z$ in the numerator of the equation as shown in Equation (9), and is plotted as an open circle. The figure shows that as the atomic number $Z$ increases, the curves trends upward, and the $n = 1$ state progresses to the left as it should. For higher atomic numbers, the perturbation of the quantum vacuum is slightly higher than for lower atomic numbers.

### B. Continuous Density Function

To this point, the discussion has been about the expectation value for the “density” of the quantum vacuum for a given primary quantum number with no consideration of substructure or variation within the given spherical region. The interpretation is that the predicted “density” is an isotropic state throughout the orbital defined by the corresponding allowed radius. Since it was just shown that the expectation value for the density at each allowed orbit radius is dependent on $1/r^4$, it is reasonable to expect that there is a more continuous representation of the density of the quantum vacuum that varies continuously over the radial distance from the nucleus. This continuous function can be found by establishing the requirement that the volumetric integral of the density (multiplied by $c^2$) is equal to the energy level allowed for each primary quantum number. The differential element to integrate is a thin spherical shell element that starts at the “radius” of the proton and continues to the allowed radius associated with the target quantum number:

$$dE = \rho(r)c^24\pi r^2 dr, \quad \rho(r) = \frac{A_0}{r^4}, \quad (10)$$

where the term $A_0$ is a proportionality constant that is to be determined to ensure that the continuous function will yield the energy level for each quantum number. The integral to solve is simple enough (neglecting the constant

![FIG. 1: Perturbation of the Vacuum - derived “density” plotted for $n = 1$ to $n = 7$ for $Z = 1$ through $Z = 7$. The bottom line labelled $Z = 1$ represents hydrogen, and labels for the points representing $n = 1$ and $n = 7$ have been added. The Casimir values are also plotted as hollow circles.](image-url)
of integration):

$$E = c^2 \int_{r_0}^{r_n} \frac{A_0}{r^2} 4\pi dr$$

$$= -4\pi A_0 c^2 \left( \frac{1}{r_n} - \frac{1}{r_0} \right).$$  \hspace{1cm} (11)

The term $r_0$ is the effective “radius” of the proton, and $r_n$ is the allowed radius of the quantum number $n$. There are three options to use for the proton radius in this calculation. The first option is to use the charge radius of the proton which has a measured value of 0.88fm [14], the second is to use the Compton wavelength of the proton which is 1.32fm, and the third option is to use the historical Rutherford empirical equation, $R = R_0 Z^{\frac{2}{3}}$ where $R_0 = 1.2 fm$, and $Z$ again is the atomic number yielding a radius of 1.2fm for the hydrogen nucleus. The third option of 1.2fm will be used for the remainder of this treatment. Using this yields a value for $A_0$ of $2.312 \times 10^{-31}$. The continuous density function for the quantum vacuum is shown in Equation (12) and includes the necessary dependency on the atomic number $Z$:

$$\rho (r) = \frac{Z^2}{n^2} \frac{2.312 \times 10^{-51}}{r^4}. \hspace{1cm} (12)$$

A plot of this function for hydrogen is shown in FIG. 2. Rather than being a series of step functions stepping down from a large value for $n = 1$ to a small value when $n = 7$ as was previously the scenario in the isotropic view, this curve is now a continuous function with slight shifts at the allowed radii which could be construed as soft boundaries between the energy levels. So rather than the density of the quantum vacuum being a uniform $3.9 \times 10^{-5} kg/m^3$ over the entire spherical region defined by the Bohr radius for $n = 1$, it starts out considerably higher just outside the nucleus and tapers down to $2.95 \times 10^{-10} kg/m^3$ at the Bohr radius. On average, it is $3.9 \times 10^{-5} kg/m^3$, but it varies continuously over radial distance $r$.

If the vacuum is indeed not an immutable and non-degradable medium, but rather a medium that can vary, as first evidenced by direct observation of the Casimir force, what can be said about what has been developed in this discussion? A thing to note is that the integral of the perturbation of the quantum vacuum around the nucleus appears to have characteristics very similar to that of the Casimir force per unit area, and since the Casimir force per unit area is negative; the integral of the vacuum perturbation would also be negative. So for $n = 1$, $Z = 1$, the energy for the captured electron is $-13.6 eV$, and likewise, the integral of the vacuum perturbation is $-13.6 eV$.

### C. Longitudinal Vacuum Waves

If the quantum vacuum is indeed not a static immutable medium, can be locally perturbed as the above assessment indicates may be the case, and that this perturbed medium can be shown to be related to the binding energy of trapped electrons, what other characteristics might the medium have that should be considered? If the quantum vacuum is a sea of vacuum fluctuations consisting of virtual photons and virtual fermions (e.g. electron-positron pairs and others), then it may be useful to study the types of wave modes that are possible for this medium that has a $1/r^4$ density dependency. The discussion will be specific to acoustic wave modes, as solutions to acoustic wave modes in a spherical region with soft boundaries are determined using the Helmholtz equation, and will yield wave modes with mathematical forms that are identical to the electron probability wave functions. A couple of points need to be made before proceeding. First, acoustic modes in air are longitudinal waves that propagate as a result of collisional process between the neutrals that make up the air. Acoustic wave modes in a plasma medium are not communicated by means of collisional process, rather the longitudinal wave propagates as a result of the charged particle motion being relayed to other charged particles by means of the

![FIG. 2: Continuous Function of the Vacuum Perturbation for Hydrogen: this function was used in the COMSOL analysis discussed in Section II C.](image-url)
intermediary electric field. Second, consider an acoustic mode analog that looks like one of the three $2p$ orbitals as shown in FIG. 3 [15]. In the acoustic case, the mode has two areas of maximum and minimum pressure separated by a region of neutral pressure that is defined by the nodal surface. In an acoustic mode, particles oscillate from the high pressure region to the low pressure region where they will reflect back again as the wave cycle oscillates. The particles are at their slowest, minimal displacements from reflection, and spend the most time in these extreme pressure regions, whereas they are at their fastest and largest displacements when they cross the nodal surface (or surfaces depending on mode). If one were to “mark” a particle that is a member of the acoustic continuum medium and try and “find” that particle or observe that particle at a particular moment in time, the odds are higher that the particle will be found within the high pressure lobes and lower that the particle will be found at the nodal surface. This is a classical analogy to the probability function that determines the likelihood of observing an electron (in a particular state) at some point around the nucleus.

The vacuum density function for hydrogen defined by Equation (12) and plotted in FIG. 2 can be modelled and studied using the high fidelity industry tool, COMSOL Multiphysics. The vacuum medium can be modelled as a virtual plasma consisting of predominantly electron-positron ephemeral pairs. Although there will be a spectrum of other fermion-antifermion particle pairs, the dominant members of the medium at any given time will be the $e^- - e^+$ pairs. The speed of sound in a plasma can be calculated using Equation (13) [16].

\[ v_s = \sqrt{\frac{kT_e}{M}} \] (13)

The way this equation is applied to a classical plasma is that the $T_e$ is the temperature of the electrons, and $M$ is the mass of the ions. In the initial consideration of the hydrogen atom, the mass of the proton will be used for $M$. It should be noted that it is likely that $M$ will be a lower mass value that is representative of either relativistic $e^- - e^+$ pairs, or heavier charged fermion-antifermion pairs. The “average” temperature of the quantum vacuum modelled as a virtual plasma within a given energy state around the hydrogen nucleus will be set equal to the equivalent kinetic temperature of the orbiting electron at that state. In this view, the real electron at the $n = 1$ state has a thermal speed of $ac$. The temperature corresponding to this thermal speed is found from the equipartition theorem using the relationship $1/2m_e v^2 = 3kT_e$, or $kT_e = 1/3m(ac)^2$ for the $n = 1$ case.

A possible source of longitudinal waves is the hydrogen nucleus. Continuing with the quasi-classical viewpoint, the electron in the $n = 1$ state is “orbiting” around the proton at an average distance of the Bohr radius $(5.29 \times 10^{-11} m)$ with an orbital speed of $ac$. Similarly, the proton is oscillating around the electron-proton system center of mass at the same frequency and it acts as a longitudinal wave source for the surrounding quantum vacuum medium. The frequency of this oscillation can be considered the eigenfrequency of the longitudinal wave associated with this quantum state. When running a COMSOL acoustic analysis of this system, the objective is to find the COMSOL-determined acoustic resonance frequency for each of the seven primary quantum number states (spherical acoustic modes defined by their corresponding Bohr radii) and compare it to the target frequency determined using the electrons orbital frequency. The target frequencies for the primary quantum numbers are given in TABLE III.

![FIG. 3: Plot for the $Z = 1, 2p$ orbital from Orbital Viewer Software.](image)

<table>
<thead>
<tr>
<th>$n$</th>
<th>Thermal vel$^a$</th>
<th>Orbital freq$^b$</th>
<th>Sound speed$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$2.19 \times 10^6$</td>
<td>$6.5808 \times 10^{12}$</td>
<td>29,476</td>
</tr>
<tr>
<td>2</td>
<td>$1.09 \times 10^6$</td>
<td>$8.2260 \times 10^{14}$</td>
<td>14,738</td>
</tr>
<tr>
<td>3</td>
<td>$7.29 \times 10^5$</td>
<td>$2.4373 \times 10^{14}$</td>
<td>9,825</td>
</tr>
<tr>
<td>4</td>
<td>$5.47 \times 10^5$</td>
<td>$1.0283 \times 10^{14}$</td>
<td>7,369</td>
</tr>
<tr>
<td>5</td>
<td>$4.37 \times 10^5$</td>
<td>$5.2646 \times 10^{13}$</td>
<td>5,895</td>
</tr>
<tr>
<td>6</td>
<td>$3.65 \times 10^5$</td>
<td>$3.0467 \times 10^{13}$</td>
<td>4,912</td>
</tr>
<tr>
<td>7</td>
<td>$3.12 \times 10^5$</td>
<td>$1.9186 \times 10^{13}$</td>
<td>4,210</td>
</tr>
</tbody>
</table>

$^a$ Thermal velocity is in $m/s$.
$^b$ Orbital frequency is in $Hz$.
$^c$ Sound speed is in $m/s$.

Since this phase of the analysis is centered on finding the spherical acoustic modes for all of the primary quantum numbers, a 2D axisymmetric model was used. This allowed for very fine mesh size when studying the $n = 1$ mode. FIG. 4 shows a close-up of the innermost region roughly defined by the Bohr radius, $5.29 \times 10^{-11} m$. The mesh size for this region of the model was set to one picometer within the spherical region defined by the Bohr radius.

FIG. 5 shows the eigenfrequency associated with the spherical acoustic mode that falls within the $n = 1$ spherical region of the model. Note that the quantum vacuum density used in the COMSOL analysis is the con-
continuous curve defined by Equation (12), and there are no “hard” boundaries in the model. The picture on the left of FIG. 5 is the view from the outer extents of the model, and the picture on the right is a close-up of the mode with a cutaway depicting the intensity distribution of the mode. This analysis result shows that this eigenfrequency is $5.3944 \times 10^{14} \text{Hz}$, whereas the hope was to find that this frequency was at $6.5808 \times 10^{15} \text{Hz}$. In order to ensure that the eigenfrequency falls at the target value, the sound speed model will need to be scaled up by a factor of 11. This means that all of the sound speeds listed in Table III will be an order of magnitude higher. This also requires that the mass in the denominator of Equation (13) is not the proton mass, but an effective mass of $15.2 \times$ that of the electron. It should be noted that in [17], the authors determined that in order to accurately predict the permittivity, $\varepsilon_0$, and permeability, $\mu_0$, of the vacuum of space arising as a result of quantum vacuum fermion pair fluctuation interaction with propagating photons, the quantum vacuum would need to routinely produce fermion pairs in a relativistic condition with a mass factor of 31.9. A scaling factor of 11 was used for all subsequent COMSOL analysis to find eigenfrequencies representative of the seven primary quantum numbers for the hydrogen model.

FIG. 6 shows the expanded view of the COMSOL 2D axisymmetric model used for finding eigenfrequencies. The model mesh sizing is provided to the right of the model picture, and the dense mesh sizing is apparent as the elements become unresolvable at the resolution of the figure below the primary quantum number, $n = 6$.

TABLE IV presents the longitudinal mode eigenfrequencies for the $n = 1$ to $n = 7$ orbitals. For each row, the expected value is listed in the left column, and the COMSOL eigenfrequency value is provided to the right along with a % error. For a primary quantum number, the expected value is the frequency that the proton would oscillate about the system center of mass that consists of the proton and the orbiting electron. FIG. 7 shows the eigenfrequency solution plots that correspond to those recorded in TABLE IV.

III. CONCLUSIONS

This paper has explored the idea of the quantum vacuum not being an absolute immutable and non-degradable state, and studied the ramifications of the quantum vacuum being able to support non-trivial spatial variations in “density.” These considerations showed no predictions that were contrary to observation, and in fact duplicated predictions for energy states associated with the primary quantum number. An explicit function of vacuum density spatial variation was de-
TABLE IV: This table shows the predicted “orbital” frequency and the COMSOL analysis eigenfrequencies for the \( n = 1 \) to \( n = 7 \) states for hydrogen.

<table>
<thead>
<tr>
<th>( n )</th>
<th>Orbital freq(^a)</th>
<th>COMSOL freq(^b)</th>
<th>%error</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>( 6.58 \times 10^{13} )</td>
<td>( 6.25 \times 10^{13} )</td>
<td>-4.98</td>
</tr>
<tr>
<td>2</td>
<td>( 8.23 \times 10^{14} )</td>
<td>( 8.23 \times 10^{14} )</td>
<td>0.05</td>
</tr>
<tr>
<td>3</td>
<td>( 2.44 \times 10^{14} )</td>
<td>( 2.38 \times 10^{14} )</td>
<td>-2.48</td>
</tr>
<tr>
<td>4</td>
<td>( 1.03 \times 10^{14} )</td>
<td>( 1.01 \times 10^{14} )</td>
<td>-1.59</td>
</tr>
<tr>
<td>5</td>
<td>( 5.26 \times 10^{13} )</td>
<td>( 4.98 \times 10^{13} )</td>
<td>-5.36</td>
</tr>
<tr>
<td>6</td>
<td>( 3.05 \times 10^{13} )</td>
<td>( 3.48 \times 10^{13} )</td>
<td>14.28</td>
</tr>
<tr>
<td>7</td>
<td>( 1.92 \times 10^{13} )</td>
<td>( 2.13 \times 10^{13} )</td>
<td>11.16</td>
</tr>
</tbody>
</table>

\(^{a}\) Orbital frequency is in Hz.
\(^{b}\) COMSOL frequency is in Hz.

FIG. 7: COMSOL analysis results of the acoustic “natural” vacuum model. The orbital shells (dark lines) can be counted, but the \( n = 1 \) radius is quite small as seen from the top left thumbnail that depicts the COMSOL eigenfrequency solution for that orbital.

FIG. 8: Acoustic eigenmodes for air in a 9in. diameter spherical isotropic air cavity with soft wall boundary. A detailed analysis and experiment measuring acoustic frequencies of a ringing basketball was performed and documented in [18].

medium are shown in FIG. 8. In the process of searching for the \( n = 1 \) to \( n = 7 \) primary quantum number eigenfrequencies using the 2D COMSOL model, some solutions mapped to the non-spherical, but axisymmetric electron orbitals from the \( p \), \( d \), and \( f \) families. Three examples are provided in FIG. 9. These results suggest that the analytical methods used in this paper that were successful at the Bohr level, can be expanded to yield solutions that exhibit the full wave characteristics of QED associated with the Schrödinger wave equation:

\[
i\hbar \frac{\partial \psi}{\partial t} = -\frac{\hbar^2}{2m} \frac{\partial^2 \psi}{\partial x^2} + V \psi.
\]

There are a number of approaches detailed in the literature that seek to develop different interpretations or understandings of the origin of the wave equation, and we will only touch on few of the concepts in closing. The orthodox view is of course the Copenhagen interpretation which, in short, does not seek to assign any classical nature to the wave equation by itself, rather only considers the wave’s statistical impact on configuration space[19]. The de Broglie-Bohm theory [20] posits the existence of a guiding global wave equation (known as the pilot wave) in a purely classical world and probabilistic quantum nature only arises as a consequence of the influence of this pilot wave on the classical world. Stochastic interpretations can be viewed as essentially fully classical interpretations of the wave equation that arises as a result of interaction with a pervasive classical zero-point radiation field (vacuum fluctuations) [21, 22]. Most recently, a paper was put forth that articulates another formulation in the debate known as a Many Interacting Worlds[23] formulation. In the MIW view, the world we live in is completely classical, but the equations of motion include a repulsive quantum force term that originates as a result of the interaction between \( N \) number of classical worlds that are close in configuration space (\( N \) can be finite or
infinite), and the wave equation is an emergent characteristic in the limit of this fully mechanical theory. In our approach discussed in this paper, the ground state of the quantum vacuum has been shown to possibly exhibit structure and spatial variation that depends on the boundary conditions present, and this dynamic variation can be successfully modelled in a quasi-classical (if not fully classical) sense with COMSOL. Indeed, the wave nature of the electron orbitals within the hydrogen atom possibly have a quantum vacuum longitudinal wave mode origin explanation.

FIG. 9: 2D Axisymmetric model results that capture axisymmetric acoustic solutions like 2p, 3d, and 4f orbitals ($m = 0$).

The paper will close with the following thought experiment: If the vacuum around the nucleus can be considered more of a “natural” vacuum as opposed to an immutable ground state with absolutely no spatial variation, and if there are ephemeral fermion/antifermion pairs dominated by electron-positron pairs that create and annihilate with a density that increases significantly as one moves closer to the nucleus, what is so special about the orbiting electron that allows it to be a “real” electron out of this vacuum soup? Perhaps it is not a case of uniqueness, but a case of non-uniqueness. Consider the following: A room full of paired square dancers progresses through the dance moves smoothly as called by the caller, and they occasionally change partners when instructed. What if there were an additional solitary dance partner of a given gender introduced to the ranks of this evenly matched group, and the rule is established that when a trade call is issued, the free dancer will couple to the nearest available dance partner of the opposite gender, and the previously paired dancer that misses out is now the free dancer until the next trade call is issued. As the evening progresses, nearly every dance partner of the gender that had the extra dancer has had a period where they were the “unique” solitary dancer. In an analogous way, perhaps the “real” electron is also “unique”. In one instance, the “real” electron collides with a positron vacuum fluctuation elevating the now un-paired electron vacuum fluctuation to the “real” state. This real electron continues in its real state for a brief period until it too collides with a positron vacuum fluctuation, elevating the next un-paired electron vacuum fluctuation to the “real” state. This process continues ad infinitum, and the “real” electron is not unique, rather it is non-unique in that the “real” descriptor is associated with the state, not the individual electron. So if the “real” electron is simply a unique state of the underlying natural vacuum, an unmatched dance partner in the sea of dancers, then the probability wave functions for the electron states may be a dual representation of the longitudinal acoustic wave modes that arise as a result of the dynamics of this natural vacuum.

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