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Relationship Between the Fine-Structure Constant Betwixt Golden Ratio, Euler Number, Pi & Fibonacci Sequence

Shuvro Sarwar

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ABSTRACT : Since the inception of the Fine-Structure Constant (α), for the last hundred years, scientists have been trying to explain its origin using other mathematical constants, as it is considered as one of the fundamental constants of nature as well as it is dimensionless. The value is very nearly equal to 1/(137.036) and for reasons of convenience, the reciprocal value of the fine-structure constant is often specified. The 2018 CODATA recommended value is given by $\alpha^{-1} = 137.035999084$ [1]. In this article, a simpler way was devised to obtain the inverse of this cosmological constant, with no more care given to physical dimensional analysis by using Golden Ratio ($\Phi = 1.618034$), Euler's number (e = 2.718282), Pi ($\pi = 3.1416$) and Fibonacci sequence. Upon implementation of fractal dimensional analysis, with only 0.00022% error it can be claimed that,



However, a more elegant yet less accurate relationship with 0.00077% error can be formed by introducing a new constant K = $(\pi + \pi.\Phi + \Phi)$. This introduction of K = 9.84283033... also helps to avoid the Fibonacci sequence.



Again, by using Stirling's formula to figure out factorial it has been observed that, $(\pi \cdot \phi)! - (e/\phi)!! = 137.04$. And

$$\sqrt{\left[(\pi + e + \Phi) + \frac{1 + \sqrt{2}}{2(\pi + e + \Phi)}\right]!} = 137.022$$

$$3(e^{\pi} + \pi^{e}) + \frac{e^{\pi}}{2(e^{\pi} + \pi^{e} + \phi e)} = 137.031$$

Last but not the least, $[(1 + \gamma) + (2 + \gamma) + (3 + \gamma) + \dots + (16 + \gamma)] = 137.04$, here, $\gamma = \frac{\sqrt{2}}{\sqrt{3}} \times \frac{1.1}{e\pi\phi} = 0.065$

INTRODUCTION I.

The fine structure constant got its name from Arnold Sommerfield, who introduced it in 1916 [2]. It is noted that when an electron orbits the nuclei in different energy shells, the energy levels of each individual shell split into much finer ones. And the gaps between the fine layers of these energy levels are directly proportional to the square of number of protons in the nucleus multiplied by α [3]. And thus it got its name.

The value of fine structure constant as well as its reciprocal $\left(\frac{1}{\alpha} = 10^2 + \frac{10^3 - 10^{-3}}{3^3} - 10^{-3}\right)$ can be derived from other constants like: G (Newton's constant), c (Einstein's constant), h (reduced Planck's constant), K_B (Boltzmann's constant), K_E (Coulomb's constant), and e (Charge of an electron).

Sign	Name	Formula	Value	Dimension
G	The universal gravitational constant	$G = gR_E^2/M_E$	$6.67408 \times 10^{-11} \text{ m}^3 \text{ kg}^{-1} \text{ s}^{-2}$	$M^{-1}L^{3}T^{-2}$
с	The speed of light in a free space	$c = 1/\sqrt{\mu_o \epsilon_o}$	$299792458 \approx 3 \times 10^8 \text{ m s}^{-1}$	LT^{-1}
ħ	The reduced Planck's constant	$\hbar = h/2\pi$	$1.0545718 \times 10^{-34} \text{ m}^2 \text{ kg s}^{-1} \text{ (J.s)}$	ML^2T^{-1}
K _B	The Boltzmann's constant	$K_B = R/N_A$	$1.3806 \times 10^{-23} \text{ m}^2 \text{ kg s}^{-2} \text{ K}^{-1} (\text{J/K})$	$ML^2T^{-2}\theta^{-1}$
K _E	The Coulomb's constant	$K_E = 1/4\pi\epsilon_0$	$8.9875517923 \times 10^9 \text{ m}^3 \cdot \text{kg.s}^{-2} \cdot \text{C}^{-2}$	$ML^{3}T^{-2}Q^{-2}$
e	Charge of an electron	96500/N _A	1.60217662 ×10 ⁻¹⁹ C	Q

 $N_A = Avogadro's number = 6.023 \times 10^{23} mole^{-1}$. Faraday's constant = 96 485.3321 A.s / mol \approx 96500 C / mol.

Fundamental Entity	Planck's Expression	Value in SI unit	Stoney's Expression	Value in SI unit
Length	$\sqrt{\frac{\hbar G}{c^3}}$	1.616255×10 ⁻³⁵ m	$\sqrt{\frac{GK_e e^2}{c^4}}$	1.3807×10 ⁻³⁶ m
Mass	G	2.176434×10 ⁻⁸ kg	$\sqrt{\frac{k_e e^2}{G}}$	1.8592×10 ⁻⁹ kg
Time	$\sqrt{\frac{\hbar G}{c^5}}$	5.391247×10 ⁻⁴⁴ s	$\sqrt{\frac{GK_ee^2}{c^6}}$	4.6054×10 ⁻⁴⁵ s
Temperature (Absolute Hot)	$\sqrt{\frac{\hbar c^5}{GK_B^2}}$	1.416784×10 ³² K	$\sqrt{\frac{K_e e^2 c^4}{GK_B^2}}$	1.2119522×10 ³¹ K
Charge	$\sqrt{\frac{\hbar c}{K_e}}$	1.875546×10 ⁻¹⁸ C	е	1.6021766×10 ⁻¹⁹ C

Now from the table above it can be seen that the ratio between the corresponding values of Planck's units & Stoney's units are also a constant, which is **11.706237481** = $\sqrt{137.036}$. Means, the ratio has the value of $\alpha^{-1/2}$. It can be observed that if we take square of each expressions and after that divide the Stoney's unit with corresponding Planck's unit then the ratio will always be $\frac{K_e e^2}{\hbar c}$. Hence, this is the fundamental expression for α .

	K _e e ²	μ ₀ св ²	e ²	µ _o ce ²	μος	e^2Z_o	e²Zo	ve_	$\left(\frac{e}{e}\right)^2$	λε	λ.	2256	r_0
<i>a</i> –	ħc -	4π ħ	2ɛohc	2h	$2R_{K}$	2h	4πħ	c	$\left(\overline{q_p}\right)$	$2\pi r_e$	λ	N _{Edd}	re

	Dimensional Analysis of Stoney's Units	Dimensional Analysis of Planck's Units
L	$\sqrt{\frac{GK_{e}e^{2}}{c^{4}}} = [M^{-1}L^{3}T^{-2}, ML^{3}T^{-2}Q^{-2}, Q^{2}, L^{-4}T^{4}]^{(0.5)} = \sqrt{L^{2}}$	$\sqrt{\frac{hG}{c^3}} = [ML^2T^{-1} \cdot M^{-1}L^3T^{-2} \cdot L^{-3}T^3]^{(0.5)} = \sqrt{L^2}$
М	$\sqrt{\frac{k_{d}e^{2}}{G}} = [ML^{3}T^{-2}Q^{-2}.Q^{2}.ML^{-3}T^{2}]^{(0.5)} = \sqrt{M^{2}}$	$\sqrt{\frac{hc}{g}} = [ML^2T^{-1} \cdot LT^{-1} \cdot ML^{-3}T^2]^{(0.5)} = \sqrt{M^2}$
Т	$\sqrt{\frac{GK_{e}e^{2}}{c^{6}}} = [M^{-1}L^{3}T^{-2}, ML^{3}T^{-2}Q^{-2}, Q^{2}, L^{-6}T^{6}]^{(0.5)} = \sqrt{T^{2}}$	$\sqrt{\frac{hG}{c^5}} = [ML^2T^{-1} \cdot M^{-1}L^3T^{-2} \cdot L^{-5}T^5]^{(0.5)} = \sqrt{T^2}$
θ	$\sqrt{\frac{K_{e}e^{2}c^{4}}{GK_{B}^{2}}} = [ML^{3}T^{2}Q^{-2}, Q^{2}, L^{4}T^{-4}, ML^{-3}T^{2}, M^{-2}L^{-4}T^{4}\theta^{2}]^{(0.5)}$	$\sqrt{\frac{\hbar c^5}{G K_{\theta}^2}} = [ML^2 T^{-1} \cdot L^5 T^{-5} \cdot ML^{-3} T^2 \cdot M^{-2} L^{-4} T^4 \theta^2]^{(0.5)}$
Q	e = Charge of a single electron or a single proton	$\sqrt{\frac{hc}{\kappa_{e}}} = [ML^{2}T^{-1}, LT^{-1}, M^{-1}L^{-3}T^{2}Q^{2}]^{(0.5)} = \sqrt{Q^{2}}$

(Stoney Length ÷ Planck Length)	So (1) $E = mc^2$ (2) E	$=$ hf (3) c $=$ f λ	Let's say, $r_e = Bohr's$ radius $n = 1$
	(4) $\hbar = \frac{\hbar}{2\pi}$ (5) $2\pi r = r$	$h\lambda(6) c^2 = \frac{1}{100}$	$\lambda_e = (h/mc) = Compton wavelength$
$= \sqrt{\frac{GK_{e}e^{2}}{c^{4}}} \div \sqrt{\frac{\hbar G}{c^{3}}} = \sqrt{\frac{K_{e}e^{2}}{\hbar c}} =$	$E = mc^2 = hf$	From and the second	So, $2\pi r_e = n\lambda$
$\sqrt{\alpha}$	\Rightarrow mc.c = hf	Energy : $\mathbf{E} = \mathbf{m} \mathbf{e}^2$	$\Rightarrow 2\pi r_e = \lambda (For n = 1)$
$= 1.3807 \times 10^{-36} \div 1.61625 \times 10^{-35}$	\Rightarrow p.c = hf	$\Rightarrow E = mc.c$	$\Rightarrow 2\pi r_e = h/p$
= 1/√137.036	\Rightarrow p.f λ = hf	$\Rightarrow E = pc$	$\Rightarrow 2\pi r_e = h/mv_e$
(Stoney Mass + Dlanck Mass)	\Rightarrow p. λ = h	$\Rightarrow E = \frac{p}{\sqrt{p}}$	$\Rightarrow 2\pi r_e = h/mc\alpha$
(Stoney Mass + Flanck Mass)	BTW, $K = 2\pi/\lambda$	√µ0≈0	$\Rightarrow 2\pi r_e = \lambda_e / \alpha$
$=\sqrt{\frac{k_e e^2}{G}} \div \sqrt{\frac{\hbar c}{G}} = \sqrt{\frac{K_e e^2}{\hbar c}} = \sqrt{\alpha}$	\Rightarrow (mv). $\lambda = h$	び E = hf	$\therefore \alpha = \lambda_e / 2\pi r_e$
-1.8502×10^{-9} · 2.175424×10 ⁻⁸	\Rightarrow h = (mv). λ	$\Rightarrow E = h(c/\lambda)$	Velocity of the electron:
$= 1.6392 \times 10^{-2} \div 2.170434 \times 10^{-1}$	\Rightarrow h = (mv).(2 π r)	$\Rightarrow E = h/t$	$v_e = c\alpha = 3 \times 10^8 / 137.036 \text{ m/s}$
= 1/\(137.036	Considering n = 1	$\Rightarrow E = \frac{n}{2\pi} \cdot \frac{2\pi}{t}$	$\therefore v_{\rm e} = 2.2 \times 10^6 \text{ m/s}$
(Stoney Time ÷ Planck Time)	$h/2\pi = (mv).(r)$	\Rightarrow E = ħ.ω	Charge of the electron:
$\frac{GK_{\theta}e^2}{2} \cdot \frac{hG}{2} = \sqrt{\alpha}$	$\Rightarrow \hbar = mvr = L$	& P = ħ.K	$e = Faraday Const./Avogadro N_A.$
$= \sqrt{c^6} \cdot \sqrt{c^5} \cdot \sqrt{\hbar c} = \sqrt{\alpha}$			\Rightarrow e = [(96500)/(6.023×10 ²³)] C
$= 4.6054 \times 10^{-45} \div 5.39124 \times 10^{-44}$	Attractive force between electron & proton of a hydrogen atom $n = 1$.		$\therefore e = 1.6022 \times 10^{-18} C$
= 1/√137.036	$\Rightarrow F_1 = K_e \cdot e^2 / r_e^2, \text{ P.E.}$	$= (\mathbf{F}_1 \times \mathbf{r}_e).$	Mass of the electron:
	Centripetal force per same electron in the	rceived by the Bohr's model,	$\frac{\text{The Mass of a Proton } (\text{H}^+)}{\text{The Mass of an Electron } (m_c)} = 1836$
(Stoney Temp. ÷ Planck Temp.)	$\Rightarrow F_2 = m_e v_e^2 / r_e, \text{ K.E.}$	$= (\mathscr{V}_2 F_2 \times r_e).$	$m_e = 9.11 \times 10^{-28} \text{ gm} = 9.11 \times 10^{-31} \text{ kg}$
$=\sqrt{\frac{K_{g}e^{2}c^{4}}{GK_{B}^{2}}} \div \sqrt{\frac{\hbar c^{5}}{GK_{B}^{2}}} = \sqrt{\frac{K_{g}e^{2}}{\hbar c}}$	Now, $F_1 = F_2$		= gm atomic mass of $H_2 \div 1837.N_A$
$= 1.211952 \times 10^{31} \div 1.41678 \times 10^{32}$	$\Rightarrow K_{\rm e}.e^2/r_{\rm e}^2 = m_{\rm e}v_{\rm e}^2/r_{\rm e}^2$	/r _e	= $[1.00784/(1837 \times 6.023 \times 10^{23})]$ gm
= 1/\137.036	\Rightarrow K _e .e ² = m _e v _e ² .r _e		Radius of the orbital:
1, 1 10, 1000	$\Rightarrow K_{e}.e^{2} = (m_{e}.v_{e}.r_{e}).$	Ve	Now, $2\pi r_e = h/mc\alpha = 2\pi\hbar/mc\alpha$
(Stoney charge ÷ Planck charge)	\Rightarrow K _e .e ² = L.v _e		$\therefore r_{\rm e} = \hbar/mc\alpha = 5.3 \times 10^{-11} \text{ m}$
b . w . ²	\Rightarrow K _e .e ² = ħ.v _e		The Learenge Fauntion:
$= e \div \sqrt{\frac{\hbar c}{K_e}} = \sqrt{\frac{K_e e}{\hbar c}} = \sqrt{\alpha}$	\Rightarrow v _e = K _e .e ² /ħ		$\mathbf{L} = \mathbf{K}\mathbf{E} - \mathbf{P}\mathbf{E} = (\frac{1}{2})\mathbf{m}_{1}\mathbf{v}_{2}^{2} - \mathbf{K}_{2}\mathbf{e}^{2}/\mathbf{r}_{2}$
$= 1.60217 \times 10^{-19} \div 1.8755 \times 10^{-18}$	$\Rightarrow v_e/c = K_e.e^2/\hbar c$		$\therefore \mathbf{L} = (1/2) \text{m}_{e} (c\alpha)^{2} - \text{K} e^{2} / (\hbar/mc\alpha)$
= 1/\sqrt{137.036}	$\div \mathbf{v}_{e}/c = \alpha$		(1) 2 2 2 2 2 2 2 2 2
			$= (\frac{1}{2})m_{e}c^{2}\alpha^{2} - m_{e}c^{2}\alpha^{2} = (\frac{1}{2})m_{e}c^{2}\alpha^{2}$

$\alpha = \frac{1}{2}$	$\frac{X_e e^2}{\hbar c} = \frac{\mu_o c e^2}{4\pi \hbar} = \frac{e^2}{2\varepsilon_o h c} = \frac{\mu_o c e^2}{2h} = \frac{\mu_o c}{2R_K} = \frac{e^2 Z_o}{2h} = \frac{e^2 Z_o}{4\pi \hbar} = \frac{v_e}{c} = \left(\frac{e}{Q_p}\right)^2 = \frac{2}{2\pi c}$	$\frac{\lambda_e}{\pi r_e} = \frac{\lambda_e}{\lambda} = \frac{2^{256}}{N_{Edd}} = \sqrt{\frac{r_0}{r_e}}$
		12
ε ₀	is the electric constant or permittivity in vacuum or free space	8.854 187 8128×10 ⁻¹² F/m
μ_0	is the magnetic constant or permeability in vacuum or free space	1.25663706212×10 ⁻⁶ N/A ²
Z ₀	is the vacuum impedance or impedance in free space $E/H = \mu_o c = 1/c\epsilon_0$	376.730313668 Ω.
R _K	is the Von Klitzing constant = h/e^2	25812.80745 Ω
λ	is the De Broglie's wavelength = h/mv_e	$3.33 \times 10^{-10} \text{ m}$
λ_{e}	is the Compton's wavelength = h/mc	$2.43 \times 10^{-12} \text{ m}$
Q _p	is the Planck charge (Ref: first page)	1.87555×10 ⁻¹⁸ C
N _{Edd}	Is the Eddington's Number, Total number of protons in the universe	1.57×10^{79}
r ₀	Classical Electron Radius (Not same as Bohr's radius as discussed later)	$2.818 \times 10^{-15} \text{ m}$
	2 2 2 2 2 2 2	

Again, P.E. $=\frac{K_e \cdot e^2}{r_0^2} \times r_0 = mc^2 \Rightarrow r_0 = \frac{K_e \cdot e^2}{mc^2} = \frac{\hbar \cdot K_e \cdot e^2}{\hbar \cdot mc^2} = \frac{K_e \cdot e^2}{\hbar c} \times \frac{\hbar}{mc} = \frac{\hbar \alpha}{mc}$. Now, $\frac{r_0}{r_e} = \frac{\hbar \alpha}{mc} \div \frac{\hbar}{mc\alpha} = \alpha^2$.

One of the prominent issue with the expression of Bohr's radius of orbital $r_e = \hbar/mc\alpha$ is, its discrepancy with the expression of the classical radius of orbital r_0 . Although the size of the electron is beyond the scope of ordinary quantum mechanics, one can think of its size as something the electron would need to have if its rest energy were only due to its electrostatic potential energy P.E. = $(F_1 \times r_0)$ instead of $(F_1 \times r_e)$. Also, $F_1 = K_e \cdot e^2/r_0^2$ instead of $K_e \cdot e^2/r_e^2$. But, P.E. = mc^2 . So, P.E. = $\frac{K_e \cdot e^2}{r_0^2} \times r_0 = mc^2 \Rightarrow r_0 = \frac{K_e \cdot e^2}{mc^2} = \frac{\hbar K_e \cdot e^2}{\hbar mc^2} \times \frac{\hbar}{mc} = \frac{\hbar\alpha}{mc}$. Now, $\frac{r_0}{r_e} = \frac{\hbar\alpha}{mc} \div \frac{\hbar}{mc\alpha} = \alpha^2$. Thus, the ratio of the classical radius of electron to the Bohr's radius is α^2 .

The Bohr radius uses the center of the proton as center, while the classical radius includes the fact that both the electron and the proton have mass, putting the center little away from the geometric center of the proton. If the electron clouds observed around the nuclei of atoms are purely statistical phenomena, then there should be no need for a radius. On the other hand, if the electron is moving in an orbit, like a moon around a planet, then the radius should be used on its own. The electron orbit is in other words neither a purely statistical phenomenon nor a conventional orbit. This is exactly what we should expect if the electron is bouncing on the atomic nucleus. Hence, electron would neither orbit, nor be entirely random it would be something in between.

The α^2 is also the ratio between the Harte energy (27.2 eV = 2 × Rydberg energy = 2 × its ionization energy) and the electron rest energy (511 keV). α is also the ratio of other two energies: (i) the energy needed to overcome the electrostatic repulsion between two electrons at a distance d, & (ii) the energy of a single photon of a wavelength $2\pi d$. if $\lambda = 2\pi d$, then $\left[\frac{e^2}{4\pi\epsilon_0 d} \div \frac{hc}{\lambda} = \frac{e^2}{4\pi\epsilon_0 d} \times \frac{2\pi d}{hc} = \frac{e^2}{4\pi\epsilon_0 hc} = \frac{e^2}{4\pi\epsilon_0 hc} = \frac{k_e e^2}{hc} = \alpha\right]$. Thus, the fine structure constant is not only the square root of the ratio of the classical radius of electron to the Bohr's radius, but also the ratio of the velocity of the electron in the first circular orbit of the Bohr model of the atom, to the speed of light in vacuum (v_e/c). This was the Summerfield's original physical interpretation. Therefore, α can similarly be expressed, as the ratio between the Compton's wavelength (h/mc) to the De Broglie's wavelength (h/mv_e) at ground state [4]. Enos Øye made the discovery that the Fine-structure constant is equal to the wavelength of the electron of a hydrogen atom, divided by half the wavelength of the photon required to kick it out of orbit, thus ionizing the hydrogen atom. The fine structure constant relates the energy of an electron in orbit around a proton with the energy of the photon required to free it from its orbit. Hence, α actually represents the probability that an electron will emit or absorb a photon.

We have already seen that $\sqrt{\alpha}$ is the conversion factor of Stoney units to Planck units. In this context, it must be pointed out that more than 25 years before Planck introduced his quantities, the Irish physicist Johnston Stoney in 1881 introduced the quantities of mass, length and time [5]. Thus, Stoney units came out in classical era while on the other hand Planck units introduced the quantum era.

II. LITERATURE REVIEW

The enigma of α remains as scientist were unsure about the fact that determines the value of α . As one of the students of Sommerfeld, Wolfgang Pauli wrote in 1948: "The theoretical interpretation of 137 is one of the most important unsolved problems of atomic physics". Hence, scientists began to mystify the number 137.

137 is the 33^{rd} prime number after 131 and before 139. It is also a Pythagorean prime: a prime number of the form 4n + 1, where n = 34 ($137 = 4 \times 34 + 1$) or the sum of two squares $11^2 + 4^2 = (121 + 16)$. Also, 137 is the only known primeval number whose sum of digits equals the number of primes "contained", it is the largest prime factor of 123456787654321 and also divides 11111111. It is the smallest prime with 3 distinct digits that remains prime if any one of its digits is removed. But we need to keep in mind the inverse of fine structure constant is almost 137.036, not 137 the full number. Again, $\alpha = 1/\sqrt{\pi^2 + 137^2} = 1/\sqrt{6\zeta(2) + 137^2}$. Which means, a triangle with base 137 and height π has the hypotenuse of a length that's very close to the measured inverse of the fine structure constant. Its close connection with π is uncanny as the sum of the squares of the first seven digits of π is also 137. As, $(3^2 + 1^2 + 4^2 + 1^2 + 5^2 + 9^2 + 2^2 = 137)$ [6]. Sum of the first 16 natural number is also one less than 137. Interestingly enough, $(\pi^{\pi} + e^e + \phi^{\emptyset})^{1.23456789...} = 137$. Also, the reciprocal of 137 is exactly 0.00 729 927 00 729 927 00 729 927 00 729 927 00... a palindromic number.

$\alpha = \frac{1}{4\pi^3 + \pi^2 + \pi} = \left(\frac{9}{16\pi^4}\right) \left(\frac{\pi^5}{5!}\right)^{\frac{1}{4}} = \left[\frac{3^2}{(2\pi)^4}\right] \left(\sqrt[4]{\frac{\pi^5}{5!}}\right)^{\frac{1}{4}}$	$\approx \frac{1}{\sqrt{2}\pi^4} \approx \frac{7\pi}{\pi^7} \approx \frac{5e\phi}{\pi^7} = \frac{1}{20\phi^4} = \frac{(5e)^4}{20(7\pi)^4} = \frac{5e^{-4}}{5e^{-4}} = \frac{1}{2}$	$\frac{36}{500\pi^2} = \frac{6}{500.\zeta(2)} \approx \frac{\phi^2}{360}$
$ie, \alpha = \frac{(2 \times 8 \times 18 \times 32)(\pi - 1) + 8}{8^2 [(2 \times 8 \times 18 \times 32)(\pi - 1)^2 + 8(\pi - 1) - 8]} = \frac{1}{8}$	$=\frac{1}{\frac{126-\frac{3}{200}}{1-\frac{5}{2\pi^{3}}}+\left[\frac{1}{10}\left\{\frac{1}{2}-\left(\frac{1}{5}\right)^{2}\right\}\right]^{2}+\left[\frac{1}{2}\left(\frac{1}{5}\right)^{4}\right]^{2}}$	$=\frac{-137+\sqrt{137^2+16}}{8}$

So, α is the positive root of the quadratic equation: $4x^2 + 137x - 1 = 0$ or, $x^2 + \left(2\sqrt{3}\pi^2 + \frac{1}{16}\right)x - \frac{1}{4} = 0$ [7]. Again, $\sqrt{e} \approx \Phi \approx 2^{\ln 2}$, hence, $\sqrt{e}/\Phi \approx 1 + \alpha \cdot \Phi^2$. Here, Φ is the golden ratio and e is the Euler's number [8, 9, 10].

Although it was Arnold Sommerfeld who formally introduced the fine structure constant in 1916, its history can be traced back to Max Planck, as discussed previously in this article. Planck had noticed that the combination of $K_{e.}e^{2}/c$ has the same dimensions as the Planck constant h. He wondered if h was identical to $K_{e.}e^{2}/c$ and if this could somehow explain the value of the elementary charge. In 1909, while reviewing the status of the theory of blackbody radiation Albert Einstein tried to predict the value of "hc" from the value of K_{e} and e^{2} , but few decimals were missing. Lorentz reacted to Einstein's notes saying that, three missing decimals were too much and concluded that h had nothing to do with e. However, this agreement of prediction with the observed fine-structure splitting was bit accidental and led to considerable confusion in the early days of quantum theory. Although relativistic mass and momentum were used, the computed energy using classical mechanics led to a correction much larger than that actually due only to relativistic effects. Since, the fine structure is associated with a completely nonclassical property of the electron called spin. As α is a dimensionless number formed of universal constants, all observers will measure the same value for it. Therefore, several numerological experiments continued for some time, and these attempts are probably a measure of how desperate physicists were in their pursuit of a fundamental reason for the value of α [11].

Even before Bohr formally announced his model of hydrogen atom 1913, an Austrian physicist Arthur Erich Haas in 1910, observed that the different spectral red lines was actually a doublet, which was termed the 'fine structure' of lines. It means, the size of a hydrogen atom is a factor $\alpha^{-2} \approx 20000$ times the size of an electron. Arnold Sommerfeld thought he could improve upon the Bohr model by assuming that the orbits can be elliptical. In addition, he considered the effect of variation of mass with speed. He presented his calculations at the Bavarian Academy of Sciences in December 1915 & January 1916. The spectroscopist Friedrich Paschen soon set to work on comparing the prediction with observations. By May 1916 he reported to Sommerfeld that "my measurements are now finished, and they agree everywhere most beautifully with your fine structures". One month later, Paschen determined the value of α^{-1} as 137.9. This was when α got its name 'fine structure constant'. Sommerfeld's model was praised as a great progress. Einstein wrote to him a year later that, "Your investigation of the spectra belongs among my most beautiful experiences in physics. Only through it do Bohr's ideas become completely convincing." Planck went to the extent of comparing this work with that of the prediction of Neptune's orbit in astronomy [12].

However, all this work was superseded by the advent of wave mechanics of Schrödinger when the classical picture of fixed orbits of electrons was abandoned in favor of a probabilistic wave function. The uncertainly principle pointed out that the classical way of calculating the electron orbit was wrong because the

position and velocity could not be determined at any given time. These models could explain the fine structure and much more, without referring to elliptical orbits. For the fine structure of spectral lines, a new quantum number was invoked, that of the electron 'spin', which took the place of Sommerfeld's 'k' quantum number. But the role of the fine structure constant in the scheme of the subatomic world was already secured, and it keeps appearing in all expressions of energy levels in atoms. It is now viewed as one of the 'coupling constants' of Nature. The force of gravity couples all particles with the Newton's gravitational constant G. Similarly, one can think of the fine structure constant being a parameter that couples all charged particles [13].

Since the value of α is important for the electronic energy levels in atoms, scientists have wondered what would have happened if its value had been different. In the 1950s, astronomers Fred Hoyle and others worked out the detailed process with which stars produce heavy elements such as carbon, oxygen etc. They found that the abundance of carbon in the Universe could be explained only if the fine structure constant had this value. Hence, Richard Feynman famously quoted about α saying, "It's one of the greatest damn mysteries of physics: a magic number that comes to us with no understanding by man. You might say the hand of God wrote that number, and we don't know how He pushed his pencil. We know what kind of a dance to do experimentally to measure this number very accurately, but we don't know what kind of dance to do on the calculation to make this number come out" [14].

Arthur Stanley Eddington (1882–1944) argued that the value of the fine-structure constant, α , could be obtained by pure deduction. He related α to the Eddington number, which was his estimate of the number of protons in the observable universe. This led him in 1929 to conjecture that α was exactly 1/137. Other physicists did not adopt this conjecture and did not accept his argument [15]. In the late 1930s, the best experimental value of the fine-structure constant, α , was approximately 1/136. Eddington then argued, from aesthetic and numerological considerations, that α should be exactly 1/136. He devised a "proof" that N_{Edd} = 136 × 2²⁵⁶ or about 1.5747×10⁷⁹. Current estimates of N_{Edd} point to a value of about 10⁸⁰ [16]. These estimates assume that all matter can be considered to be hydrogen and require assumed values for the number and size of galaxies and stars in the universe. During a course of lectures that he delivered in 1938 as Lecturer at Trinity College, Cambridge, Eddington averred that: I believe there are N_{Edd} protons in the observable universe where, N_{Edd} = 15747724136275002577605653961181555468044717914527116709366231425076185631031296, as well as the same number of electrons [17]. This large number was soon named the "Eddington number". Shortly thereafter, improved measurements of α yielded values closer to 1/137, whereupon Eddington changed his proof to show that α had to be exactly 1/137 [18].

In 2000, Kosinov suggested the more complex but more accurate formula $\alpha^{20} = (\pi.\Phi^{14})^{1/13}.10^{-43}$. He followed the footsteps of two American electrochemists, Lewis and Adams, who proposed back in 1914 that "all of the universal constants involve only integral numbers and π ". After applying cube root to the solution of Stefan-Boltzmann law (as it involves a 3D volume), Lewis derived [19]:

$$\alpha^{-1} = 32\pi \left(\frac{\pi^5}{5!}\right)^{\frac{2}{3}} = 137.35.$$

The Lewis–Adam's conjecture was discussed among physicists. In 1935, Heisenberg wrote to Dirac: "I do not believe at all any more in your conjecture that the Sommerfeld fine-structure constant may have something to do with the concept of temperature; that is, neither do I any more believe in the Lewis value". Indeed, Lewis' value is wrong, but his idea led to another dimensionless constant, involving the continued spectra of blackbody radiation [20]. Heisenberg wrote to Bohr with a joke formula suggested by Lunn in 1922, $\alpha^{-1} = 2^4.3^3/\pi$. Bohr replied, $\alpha^{-1} = 360/\Phi^2$. So back in 1935, after Heisenberg's letter to Dirac came into Pauli's notice, he then suggested that the five-dimensional Kaluza–Klein theory might help to understand the problem [21]. Following Pauli, Wyler came up with another formula in 1969 exposing a similar pattern with the Lewis formula, but in 4th root and in the reciprocal way [22].

$$\alpha^{-1} = \sqrt[4]{\frac{1^0 4^0 5^1 8^0 9^0}{2^{10} 3^7 6^0 7^0 10^0}} \pi^{11}$$

In 1989, Bailey and Ferguson used a supercomputer to check Wyler's formula, and automatically produced several "other relations of comparable complexity with even better accuracy". One example is $\alpha^{-5} = 150\pi (6^{5}5^2\pi^3)^8$, ie, $\alpha^{-1} = 137.036048362143$ [23]. This clearly showed that a Wyler-type formula could not be the unique answer for the fine structure constant. Wyler's formula is later discussed in the E8 lie groups. In 2006, Castro reviewed the coupling constant with the Complex Domains [24]. However, Wyler's work made people devise simpler ways to obtain the magic number, with no more care given to physical dimensional analysis. In this article a similar approach has been followed. Aether Theory in 1972 [25], Stoyan 2004 [26], Heyrovska 2005 [27], Naschia 2006 [28] Gilson 2007 [29], Lestone 2008 [30], Markovich 2009 [31], Rhodes 2010 [32], Kirakosyan 2011 [33], Code 2012 [34], Schonfeld 2013 [35–36], suggests that the pursuit never ended.

Nevertheless, among all the approximations, Michael J. Bucknum and Eduardo A. Castro came up with the most elegant solution in last year 2020 [37] with a convergent series, within a few terms, to better than 99999 parts in 100,000 of the true value of α . They suggested:

$$\sqrt{\alpha} = \sum_{n=0}^{\infty} \left(\frac{2n+1}{2n+3}\right)^{2n} \frac{(e\pi)^{n+1}}{10^{4n+2}} = \left(\frac{1}{3}\right)^0 \cdot \frac{(e\pi)^1}{10^2} + \left(\frac{3}{5}\right)^2 \cdot \frac{(e\pi)^2}{10^6} + \left(\frac{5}{7}\right)^4 \cdot \frac{(e\pi)^3}{10^{10}} + \left(\frac{7}{9}\right)^6 \cdot \frac{(e\pi)^4}{10^{14}} + \cdots \infty$$

Even after these countless efforts, Pauli's simplest question still remains unanswered: "Why 137?" [38–42]. In his Nobel Lecture delivered in Stockholm on 13 December 1946, Pauli expressed his goal was to establish a theory, "which will determine the value of the fine-structure constant and will thus explain the atomistic structure of electricity, which is such an essential quality of all atomic sources of electric fields actually occurring in nature" [43]. As the initialization, "from a physical point of view, that the existence of atomicity, in itself so simple and basic, should also be interpreted in a simple and elementary manner by theory and should not, so to speak, appear as a trick in analysis" [43]. His lifelong search for 137, a millennium puzzle, ended in hospital room 137 [44]. The difficulty of finding the correct α formula is partly due to the uncertainty of the experimental values - approximately 137.036. Some experimental data of the inverse of the fine structure constant is listed in the Table below [45–53].

Year	1/α	Source	Year	1/α	Source
1916	137.360563948	A. Sommerfeld	2000	137.03599976(50)	CODATA 1998
1929	137.29 ± 0.11	R. Birge	2002	137.03599911(46)	CODATA 2002
1930	136.94 ± 0.15	W. Bond	2007	137.035999070(98)	G. Gabrielse
1932	137.305 ± 0.005	R. Birge	2008	137.035999679(94)	CODATA 2006
1935	137.04 ± 0.02	F. Spedding et al.	2008	137.035999084(51)	G Gabrielse D Hanneke
1941	137.030 ± 0.016	R. Birge	2010	137.035999037(91)	R. Bouchendira
1943	137.033 ± 0.092	U. Stille	2010	137.03599913296(33)	T. Kinoshita
1949	137.027 ± 0.007	J. DuMond, E. Cohen	2011	137.035999074(44)	CODATA 2010
1949	137.041 ± 0.005	H Bethe, C Longmire	2015	137.035999139(31)	CODATA 2014
1957	137.0371 ± 0.0005	J.Bearden, J.Thomsen	2017	137.035999150(33)	Aoyama et al.
1969	137.03602(21)	CODATA 1969	2018	137.035999046(27)	Parker et al.
1973	137.03612(15)	CODATA 1973	2019	137.035999084(21)	CODATA 2018
1987	137.0359895(61)	CODATA 1986	2020	137.035999206(11)	Morel et al. 2020
1998	137.03599883(51)	T. Kinoshita			

III. RESULTS AND DISCUSSION

A simple VISUAL BASIC computer program was written to generate the factors for various particle pairs. The table below shows the largest common factors and multiples for electron-proton bonds that produce fine structure constants within a ± 3 standard error window of the measured fine structure constant value. All of the components are considered to be integer. The factor analysis is periodic, with several α candidates appearing within the search window. The factor analysis also shows that the α candidates with the highest common factors, all exhibited the same multiple, 472. This means that at every 472nd electron wave period, the electron and proton total energy waves overlap. The table below exhibits largest common factors for electron-proton bond & Fine Structure Constant (Brian Dale Nelson) retrieve from http://www.quantumpulse.com [55].

Proton Mass Component (Np)	Electron Mass Component (Ne)	Proton Electron Mass Ratio $(Np/Ne = 6\pi^5)$	Kinetic Energy Component Nv $= 1.45 \times 10^{-10}$	$\frac{1}{\alpha} = \sqrt{\frac{Ne}{2Nv}}$	Largest Common Factor	Multiple
553900587503	301663688	1836. 15267444121	8032	137. 035999065357	639135	472
434872647725	236839046	1836. 15267444119	6306	137. 035999080312	501791	472
810231325561	441265771	1836. 15267444118	11749	137. 035999110981	934910	472
375358677836	204426725	1836. 15267444117	5443	137. 035999124446	433119	472
691203385783	376441129	1836. 15267444116	10023	137. 035999140230	797566	472
315844707947	172014404	1836. 15267444115	4580	137. 035999158988	364447	472
572175446005	311616487	1836. 15267444113	8297	137. 035999181648	660222	472
828506184063	451218570	1836. 15267444113	12014	137. 035999190287	955997	472
256330738058	139602083	1836. 15267444111	3717	137. 035999209569	295775	472
709478244285	386393928	1836. 15267444110	10288	137. 035999232087	818653	472
453147506227	246791845	1836. 15267444109	6571	137. 035999244824	522878	472
649964274396	353981607	1836. 15267444108	9425	137. 035999258728	749981	472
846781042565	461171369	1836. 15267444107	12279	137. 035999266169	977084	472

While executing a fractal dimensional analysis, it was observed that, $8^3.4215 \approx 1230$. This finding is significant as, the number 1230 includes all the digits from 0 to 3 once, similarly the fractal dimension 3.4215 also includes all the digits from 1 to 5 once. Interestingly, $3.4215^{4} \approx 1/\alpha$. Thus, the relationship got devised. $[3.4215]^4 = 137.046$

 $[\log_8 1230]^4 = 137.043$

 $[\log_8 89e\pi\Phi]^4 = 137.028$

 $\left[\log_8 \frac{e\pi\Phi}{0.01123581321345589144233377\dots}\right]^4 = 137.029$

The value of $(1/\alpha - \alpha) = (137.036 - 1/137.036) = (137.036 - 0.0073) = 137.0287$

$$\left[\log_{2^3} \frac{e\pi \Phi}{0.0\ 1\ 1\ 2\ 3\ 5\ 8\ 13\ 21\ 34\ 55\ 89\ 144\ 233\ 377\ 610\ 987\ldots}\right]^{2^2} = \ \alpha^{-1} - \alpha$$

However, inspired by - $[e^{\phi^{\pi}} + \pi^{e^{\phi}} + \pi^{\phi^{e}}] = 22^{2}$, and $(\pi^{\pi} + e^{e} + \phi^{\phi})^{1.23456789...} = 137$.

$$[e^{\pi^{\phi}} + e^{\phi^{\pi}} + \pi^{e^{\phi}} + \pi^{\phi^{e}}]^{\frac{1}{\sqrt{2.01}}} = 137.0207 \approx \alpha^{-1} - \alpha$$

Hence, there is only 0.00022% error in this approximation. Again, the mean value of first four worth is $[137.046 + 137.043 + 137.028 + 137.029] \div 4 = 137.0365$. To check the acceptability of such error another table was created. And the hypothesis test and reliability analysis were done using the values of first two tables along with the next table that includes the values of α and $1/\alpha$ of similar kind of relations devised by other authors throughout the century.

^{*}Corresponding Author: Nafish Sarwar Islam

FORMULA	α	α^{-1}
1	0.0072973363440646	137.03630377587843
$\alpha = \frac{1}{4\pi^3 + \pi^2 + \pi}$	0.0052052401200210	127.02.002.4401.640.4
$\alpha = \left(\frac{9}{16\pi^4}\right) \left(\frac{\pi^3}{5!}\right)^{\frac{2}{4}}$	0.0072973481300318	137.03608244816494
$\alpha = [\frac{3^2}{(2\pi)^4}](\sqrt[4]{\frac{\pi^5}{5!}})$	0.0072973481300317835	137.03608244816525
$\alpha = \frac{1}{\sqrt{2}\pi^4}$	0.0072591456	137.7572576387
$\alpha = \frac{7\pi}{\pi^7}$	0.007281130313071	137.34131336789999
$\alpha = \frac{5\varepsilon\varphi}{\pi^7}$	0.0072812009592285801	137.33998080804884
$\alpha = \frac{1}{20\varphi^4}$	0.0072949016875157736	137.08203932499367
$\alpha = \frac{(5\sigma)^4}{20(7\pi)^4}$	0.007295184810741	137.07671922548952
$\alpha = \frac{36}{500\pi^2}$	0.0072951252222483	137.07783890401923
$\alpha^{-1} = \frac{360}{\varphi^2}$	0.0072723166354163741	137.50776405003786
$\alpha = \frac{(2 \times 8 \times 18 \times 32)(\pi - 1) + 8}{8^2 [(2 \times 8 \times 18 \times 32)(\pi - 1)^2 + 8(\pi - 1) - 8]}$	0.0072973525680816596	137.03599910657485
$\alpha = \frac{1}{\frac{126 - \frac{3}{200}}{1 - \frac{5}{2\pi^3}} + \left[\frac{1}{10}\left\{\frac{1}{2} - \left(\frac{1}{5}\right)^2\right\}\right]^2 + \left[\frac{1}{2}\left(\frac{1}{5}\right)^4\right]^2}$	0.0072973525692030473	137.03599908551646
$\alpha = \frac{-137 + \sqrt{137^2 + 16}}{8}$	0.007297715134418	137.02919086053788
$\alpha^{-1} = \sqrt{e^{\pi + \pi \varphi + \varphi}}$	0.0072888086608280953	137.19663206063473
$\alpha^{-1} = \frac{\varphi}{2} + e\pi^3\varphi$	0.0072895293666450360	137.18306761692137
$\alpha^{-1} = \frac{\pi^{12}}{5^3 \cdot e^4} + \varphi$	0.0072967839198649047	137.04667850689408
$\alpha^{-1} = (e + \pi + \varphi)^{\sqrt{6}} - 1$	0.00729188536340565	137.13874398224945
$\alpha^{-1} = \left(\frac{\pi\varphi}{\log\left(e\right)}\right)^2$	0.0072995081317228645	136.99553202141241
$\sqrt{\alpha} = (\frac{1}{3})^{0} \cdot \frac{(e\pi)^{1}}{10^{2}} + (\frac{3}{5})^{2} \cdot \frac{(e\pi)^{2}}{10^{6}} + (\frac{5}{7})^{4} \cdot \frac{(e\pi)^{3}}{10^{10}} + (\frac{7}{9})^{6}$	0.0072971935198860104	137.03898591627607

A sample of 50 numbers were taken from these abovementioned three tables. While $[50 \times 137] = 6850$ the total value of those 50 $(1/\alpha)$ values were 6854.12, which produces an average score of 137.0824 and a standard deviation of 0.13320284. Keeping this output in mind a more elegant yet less accurate relationship with 0.00077% error can be formed by introducing a new constant $K = (\pi + \pi \Phi + \Phi)$. This introduction of K helps to avoid the Fibonacci sequence as well as any other number except than natural constants.



Now with n = 50, μ = 137.082, σ = 0.13320284 & \bar{x} = 137.035 a value of Z = - 0.2495 is obtain. From which the following P values were found for the 0.00077% errored result: P (x<Z) = 0.0063, P (x>Z) = 0.9937, P (Z<x<0) = 0.4937, P (-Z<x<Z) = 0.9874, P (x<-Z or x>Z) = 0.0126. It is necessary to figure out the P values because the theory does not predict its value. Therefore, α must be determined experimentally. In fact, α is one of the empirical parameters in the Standard Model of particle physics, whose value is not determined within the Standard Model. Also, it is evident from the diagram, $\theta = \tan^{-1}(\alpha) = 0.418^{\circ} = 0.007297$ rad = (α) rad. 2002 paper by Eliahou Thousson entitled, "The discreate four spatial dimensional space representation of electro-discrete charges" also claims that, the various interaction between electro-discrete charges cannot be described alone by the interaction factors as the Fine-Structure constant could have been resulted in by an appropriate correlation angle of 0.418°. [54]. However, the simplest of mathematical relationship that can be established among all these constants by using the Stirling's approximation n! = [{(2π n)^(1/2)} × {(n/e)^(n)}] is (π . Φ)! - (e/ Φ)!! = (π . Φ)! - (1.68)!! = 137.04 = $\alpha^{-1} + (\alpha/2)$.

Nevertheless,

$$\frac{\log_{1/\alpha} \left[e^{\pi^{\phi}} + e^{\phi^{\pi}} + \pi^{e^{\phi}} + \pi^{\phi^{e}} \right] = \sqrt{2.01} \approx \sqrt{2} \qquad (\pi + e + \phi)^{\sqrt{2 \times 3}} - \frac{\pi}{3^{2}} = 137.045$$
As well as,

$$\sqrt{\left[(\pi + e + \phi) + \frac{1 + \sqrt{2}}{2(\pi + e + \phi)} \right]!} = 137.022$$
W CONCLUSION

To realize the significance of the value of α we need to look into the 137th element of the periodic table. The element is Feynmanium, an undiscovered hypothetical element with the symbol Fy & atomic number 137. It is named in honor of Richard Feynman. The outer most electron of this element of the periodic table is supposed to move nearly at the speed of light. The idea is quite simple, as 1/FSC is the odds that an electron will absorb a single photon. Protons and electrons are bound by interactions with photons. So when we get 137 protons, we get 137 photons, and we get (137/137.036)×100% chance of absorption and electron in the ground state is supposed to orbit at the speed of light. This is the electromagnetic equivalent of a black hole. But for the element number 138 the g orbital get fully occupied for the very first time. For this reason it is the most unstable and a temporarily observable hypothetical element. There is (138/137.036)×100% probability that an electron will absorb or emit a photon. As per the Aufbau principal when the g orbital gets fully occupied for the first time then it is supposed to get an atomic number of 138. The maximum occupancy level of s, p, d, f, & g orbitals are 2, 6, 10, 14, & 18 respectively, [2(2n + 1)]; where n = 0, 1, 2, 3, & 4.

According to the Aufbau principal till the element number 120 we do not observe the presence of g orbital. Unbinilium, is the hypothetical chemical element in the periodic table with atomic number of 120. After this hypothetical 120th element for the first time the g orbital comes into existence. Which means, even the g orbital itself is a hypothetical one. When it gets fully occupied with the allotted 18 electrons, then the total number of electrons in the element becomes (120 + 18) = 138. Hence, more than 100% probability that an electron will absorb or emit photon. So, α is directly related to the coupling constant determining the strength of the interaction between electrons–photons. Therefore, it is the "coupling constant" or measure of the strength of the strong nuclear force is 137.036 times stronger than the electromagnetic force, which implies it is impossible to pack more than 137.036 proton inside a nucleus as their combined electromagnetic repulsion will be stronger enough to overcome the strong nuclear force binding them inside it. Though, the recent extended periodic table hypothesize the existence up to 172 elements, while the Aufbau principle hypothesize existence of 168 elements (means it is possible to discover the entire 8th row with 50 elements, as till now only 118 elements are found).

The idea is quite simple. Inert gases follows the following of sequence for their atomic numbers. He: $(2.1^2) = 2$, Ne: $(2 + 2.2^2) = 10$, Ar: $(10 + 2.2^2) = 18$, Kr: $(18 + 2.3^2) = 36$, Xe: $(36 + 2.3^2) = 54$, Rn: $(54 + 2.4^2) = 86$, & Og: $(86 + 2.4^2) = 118$, this is what we have so far. Hence if the entire eighth row gets filled up the last inert particle is supposed to be $(118 + 2.5^2) = 168^{\text{th}}$. While with the nuclei magic number sequence for stability (2, 8, 20, 28, 50, 82), the valley of stability is found among these elements: He: $2 = 2.(^1C_1)$, O: $8 = 2.(^2C_1 + ^2P_2)$, Ca: $20 = 2.(^3C_1 + ^3P_2 + ^3C_3)$, Ni: $28 = 2.(^4C_1 + ^4C_2 + ^4C_3)$, Sn: $50 = 2.(^5C_1 + ^5C_2 + ^5C_3)$ and finally, Pb: $82 = 2.(^6C_1 + ^6C_2 + ^6C_3)$. Hence, the next stable element is supposed to be $2.(^7C_1 + ^7C_2 + ^7C_3) = 126^{\text{th}}$ element. Coincidentally the average of these three numbers (120, 126, 168) is also [(120 + 126 + 168) ÷ 3] = 138.



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Now,
$$E = \frac{1}{2} mv^2 + P.E.(U) = \frac{m^2 v^2}{2m} + U = \frac{(P)^2}{2m} + U$$

Now, $E = mc^2 = h f = \frac{h}{t} = \frac{h}{2\pi} \cdot \frac{2\pi}{t} = \hbar.\omega$
Again, $mc^2 = h f = \frac{h}{t} \Rightarrow (mc) \cdot c = \frac{h}{t} \Rightarrow (P) \cdot \frac{\lambda}{t} = \frac{h}{t}$
i.e., $p = \frac{h}{\lambda} = \frac{h}{2\pi} \cdot \frac{2\pi}{\lambda} = \hbar.k$
Now, $E = \frac{p^2}{2m} + U \Rightarrow E\psi = \frac{p^2}{2m}\psi + U\psi$
 $\Rightarrow \hbar\omega\psi = \frac{\hbar^2 k^2\psi}{2m} + U\psi$
 $\Rightarrow i\hbar.(-i\omega\psi) = -\frac{\hbar^2(-k^2\psi)}{2m} + U\psi$
 $\Rightarrow i\hbar.\frac{d\psi}{dt} = -\frac{\hbar^2}{2m} \cdot \left[\frac{d^2\psi}{dx^2}\right] + U\psi$

For Three Dimensions Schrodinger Equation

$$i\hbar\frac{d\psi}{dt} = -\frac{\hbar^2}{2m} \cdot \left[\frac{d^2\psi}{dx^2} + \frac{d^2\psi}{dy^2} + \frac{d^2\psi}{dz^2}\right] + U\psi \Rightarrow i\hbar\frac{d\psi}{dt} = -\frac{\hbar^2}{2m} \cdot \nabla^2\psi + U\psi$$

Problem: 1st derivative of time 2nd derivative of space. Solution: E² = P²c² + m²c⁴ Therefore, E² ψ = c² (P² ψ) + m²c⁴ ψ \Rightarrow - E² ψ = -c² (P² ψ) - m²c⁴ ψ \Rightarrow - h² $\omega^{2} \psi$ = -c² h² k² ψ - m² c⁴ ψ \Rightarrow h² $\cdot \frac{d^{2} \psi}{dt^{2}} = h^{2}c^{2} \cdot \frac{d^{2} \psi}{dx^{2}} - m^{2}c^{4}\psi$ $\Rightarrow \frac{1}{c^{2}} \cdot \frac{d^{2} \psi}{dt^{2}} = \frac{d^{2} \psi}{dx^{2}} - \left(\frac{mc}{\hbar}\right)^{2} \psi$ For 3D: $\frac{1}{c^{2}} \cdot \frac{d^{2} \psi}{dt^{2}} = \frac{d^{2} \psi}{dx^{2}} + \frac{d^{2} \psi}{dy^{2}} + \frac{d^{2} \psi}{dz^{2}} - \left(\frac{mc}{\hbar}\right)^{2} \psi$

For a photon m = 0, and the above equation converts into a classical wave equation.

Now,
$$E = \sqrt{p^{2}c^{2} + m^{2}c^{4}} = s\frac{B}{c} = \sqrt{p^{2} + m^{2}c^{2}}$$

Say $\sqrt{p^{2}c^{2} + m^{2}c^{4}} = \alpha_{x}P_{x} + \alpha_{y}P_{y} + \alpha_{z}P_{z} + \beta mc$
 $=> \left(\frac{B}{c}\right)^{2} = p^{2} + (mc)^{2} = (\alpha_{x}P_{x} + \alpha_{y}P_{y} + \alpha_{z}P_{z} + \beta mc)^{2}$
 $=> \left(\frac{B}{c}\right)^{2} = (\alpha_{x}P_{x} + \alpha_{y}P_{y} + \alpha_{z}P_{z} + \beta mc) \times (\alpha_{x}P_{x} + \alpha_{y}P_{y} + \alpha_{z}P_{z} + \beta mc)$
 $=> \left(\frac{B}{c}\right)^{2} = \alpha_{x}^{2}P_{x}^{2} + \alpha_{y}^{2}P_{y}^{2} + \alpha_{z}^{2}P_{z}^{2} + \beta^{2}(mc)^{2}$
 $+ (\alpha_{x}\alpha_{y} + \alpha_{y}\alpha_{x})P_{x}P_{y} + (\alpha_{y}\alpha_{z} + \alpha_{z}\alpha_{y})P_{y}P_{z} + (\alpha_{z}\alpha_{x} + \alpha_{x}\alpha_{z})P_{z}P_{x}$
 $+ (\alpha_{x}\beta + \beta\alpha_{x})P_{x}mc + (\alpha_{y}\beta + \beta\alpha_{y})P_{y}mc + (\alpha_{z}\beta + \beta\alpha_{z})P_{z}mc$
But, as the vector $P^{2} = P_{x}^{2} + P_{y}^{2} + P_{z}^{2}$, Therefore, $\alpha_{x}^{2} = \alpha_{y}^{2} = \alpha_{z}^{2} = \beta^{2} = 1$ & $\alpha_{x}\alpha_{y} + \alpha_{y}\alpha_{x} = \alpha_{y}\alpha_{z} + \alpha_{z}\alpha_{y} = \alpha_{z}\alpha_{x} + \alpha_{x}\alpha_{z} = \alpha_{x}\beta + \beta\alpha_{x} = \alpha_{y}\beta + \beta\alpha_{z} = 0$
 $a_{x} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & -1 & 0 \\ 0 & 0 & 0 & -1 \end{bmatrix} a_{xy} = \begin{bmatrix} 0 & 0 & 0 & 1 \\ 0 & 1 & 0 & 0 \\ 1 & 1 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{bmatrix} \alpha_{z} = \begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & -1 \\ 1 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 \end{bmatrix} \beta_{B} = \begin{bmatrix} 0 & 0 & 0 & -i \\ 0 & 0 & i & 0 \\ 0 & -i & 0 & 0 \end{bmatrix}$
Now, $\frac{B}{c} = \alpha_{x}P_{x} + \alpha_{y}P_{y} + \alpha_{z}P_{z} + \beta mc$
 $=> \frac{h\omega}{c} = \alpha_{x}hK_{x} + \alpha_{y}hK_{y} + \alpha_{z}hK_{z} + \beta mc$
 $=> \omega = [\alpha_{x}K_{x} + \alpha_{y}K_{y} + \alpha_{z}hK_{z} + \beta mc$
 $=> \omega = [\alpha_{x}(iK_{x}) + \alpha_{y}(iK_{y}) + \alpha_{z}(iK_{z})]c - \frac{i\beta mc^{2}}{h}$
 $=> -i\omega\Psi = - [\alpha_{x}(iK_{x}) + \alpha_{y}(iK_{y}) + \alpha_{z}(iK_{z})]c - \frac{i\beta mc^{2}}{h}$
 $=> -i\omega\Psi = - [\alpha_{x}(iK_{x}) + \alpha_{y}\frac{\partial\Psi}{\partial_{y}} + \alpha_{z}\frac{\partial\Psi}{\partial_{z}}].c - \frac{i}{h}\beta mc^{2}\Psi = 0$

$$\therefore \alpha_{x}^{2} = \alpha_{y}^{2} = \alpha_{z}^{2} = \beta^{2} = \begin{bmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{bmatrix} = 1$$

$$\alpha_{x}\alpha_{y} + \alpha_{y}\alpha_{x} = \begin{bmatrix} 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \\ 0 & -1 & 0 & 0 \\ -1 & 0 & 0 & 0 \end{bmatrix} + \begin{bmatrix} 0 & 0 & 0 & -1 \\ 0 & 0 & -1 & 0 \\ 0 & 1 & 0 & 0 \\ 1 & 0 & 0 & 0 \end{bmatrix} = 0$$

$$\alpha_{y}\alpha_{z} + \alpha_{z}\alpha_{y} = \begin{bmatrix} 0 & -1 & 0 & 0 \\ 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & -1 \\ 0 & 0 & 1 & 0 \end{bmatrix} + \begin{bmatrix} 0 & 1 & 0 & 0 \\ -1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & -1 & 0 \end{bmatrix} = 0$$

$$\alpha_{z}\alpha_{x} + \alpha_{x}\alpha_{z} = \begin{bmatrix} 0 & 0 & -1 & 0 \\ 0 & 0 & 0 & -1 \\ 1 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 \end{bmatrix} + \begin{bmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & -1 \\ -1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{bmatrix} = 0$$

$$\therefore \alpha_{x}\alpha_{y} + \alpha_{y}\alpha_{x} = \alpha_{y}\alpha_{z} + \alpha_{z}\alpha_{y} = \alpha_{z}\alpha_{x} + \alpha_{x}\alpha_{z} = 0$$

$$\begin{bmatrix} 0 & 0 & 0 & -i \end{bmatrix} \begin{bmatrix} 0 & 0 & 0 & -i \end{bmatrix}$$

$$\begin{aligned} \alpha_{x}\beta + \beta\alpha_{x} &= \begin{bmatrix} 0 & -i & 0 & 0 \\ 0 & i & 0 & 0 \\ -i & 0 & 0 & 0 \end{bmatrix} + \begin{bmatrix} 0 & 0 & i & 0 \\ 0 & -i & 0 & 0 \\ i & 0 & 0 & 0 \end{bmatrix} = 0 \\ \alpha_{y}\beta + \beta\alpha_{y} &= \begin{bmatrix} i & 0 & 0 & 0 \\ 0 & -i & 0 & 0 \\ 0 & 0 & i & 0 \\ 0 & 0 & 0 & -i \end{bmatrix} + \begin{bmatrix} -i & 0 & 0 & 0 \\ 0 & i & 0 & 0 \\ 0 & 0 & -i & 0 \\ 0 & 0 & 0 & i \end{bmatrix} = 0 \\ \alpha_{z}\beta + \beta\alpha_{z} &= \begin{bmatrix} 0 & -i & 0 & 0 \\ -i & 0 & 0 & 0 \\ 0 & 0 & 0 & -i \\ 0 & 0 & -i & 0 \end{bmatrix} + \begin{bmatrix} 0 & i & 0 & 0 \\ i & 0 & 0 & 0 \\ 0 & 0 & 0 & i \\ 0 & 0 & 0 & i \end{bmatrix} = 0 \end{aligned}$$

 $\therefore \alpha_{x}\beta + \beta\alpha_{x} = \alpha_{y}\beta + \beta\alpha_{y} = \alpha_{z}\beta + \beta\alpha_{z} = 0$

$$e^{i\pi} = \cos(\pi) + i.\sin(\pi) = -1 + i(0) = -1 = i^2 = (i)^{\frac{2i}{i}} = (i)^{\frac{2ii}{-1}} = (i)^{-2i.i}$$

$$\therefore e^{\pi} = (i)^{-2i} = 23.140692632779269005729086367948547380...$$

Now,

 $e^{\pi} = 23.1406926328$ Again, $\pi^{e} = 22.4591577184$ Therefore, $e^{\pi} + \pi^{e} = 45.6$ Hence, $3(e^{\pi} + \pi^{e}) = 136.8$ And, $\phi e = 4.4$ So, $e^{\pi} + \pi^{e} + \phi e = 50$ Means, $2(e^{\pi} + \pi^{e} + \phi e) = 100$ Thus,



Implies,

$$3(e^{\pi} + \pi^{e}) + \frac{e^{\pi}}{2(e^{\pi} + \pi^{e} + \phi e)} + \frac{1}{2 \times 10^{2}} = \alpha^{-1}$$

And this relationship above has zero percent error, means it exhibits the exact value. However, if we consider the previous relation (137.031) then the error becomes only 0.00365%.



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