About the Correspondence of Dark Nuclear Genetic Code and Ordinary Genetic Code

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Abstract

The basic problem in the understanding of the prebiotic evolution is how DNA, RNA, amino-acids and tRNA and perhaps even cell membrane and microtubules. The individual nucleotides and amino-acids emerge without the help of enzymes or ribozymes but the mystery is how their polymers emerged. If the dark variants of these molecules served as templates for their generation one avoids this hen-and-egg problem. The problem how just the biomolecules were picked up from a huge variety of candidates allowed by chemistry could be solved by the resonance condition making possible metabolic energy transfer between biomolecules and dark nuclei.

Simple scaling argument shows that the assumption that ordinary genetic code corresponds to \( h_{eff}/h = n = 2^{18} \) and therefore to the p-adic length scale \( L(141) \simeq .3 \text{ nm} \) corresponding to the distance between DNA and RNA bases predicts that the scale of dark nuclear excitation energies is .5 eV, the nominal value of metabolic energy quantum. This extends and modifies the vision about how prebiotic evolution led via RNA era to the recent biology. Unidentified infrared bands (UIBs) from interstellar space identified in terms of transition energies of dark nuclear physics support this vision and one can compare it to PAH world hypothesis.

p-Adic length scale hypothesis and thermodynamical considerations lead to ask whether cell membrane and microtubules could correspond to 2-D analogs of RNA strands associated with dark RNA codons forming lattice like structures. Thermal constraints allow cell membrane of thickness about 5 nm as a realization of \( k = 149 \) level with \( n = 2^{22} \) in terms of lipids as analogs of RNA codons. Metabolic energy quantum is predicted to be .04 eV, which corresponds to membrane potential. The thickness of neuronal membrane in the range 8-10 nm and could correspond to \( k = 151 \) and \( n = 2^{23} \) in accordance with the idea that it corresponds to higher level in the cellular evolution reflecting that of dark nuclear physics. The energy quantum of ordinary Josephson radiation is below the thermal energy for photons but the notion of generalized Josephson junction saves the situation. For massive particles associated with flux tubes the thermal energy \( T/2 \) is below the potential energy defined by action potential and that of metabolic energy quantum.

Also microtubules could correspond to \( k = 151 \) realization for which metabolic energy quantum is .02 eV slightly below thermal energy at room temperature: this could relate to the inherent instability of microtubules. Also a proposal for how microtubules could realize genetic code with the 2 conformations of tubulin dimers and 32 charges associated with ATP and ADP accompanying the dimer thus realizing the analogs of 64 analogs of RNA codons is made.

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1. Introduction

The idea about the realization of genetic code in terms of dark proton sequences giving rise to dark nuclei is one of the key ideas of TGD inspired quantum biology [L4]. This vision was inspired by the totally unexpected observation that the states of three dark protons (or quarks) can be classified to 4 classes in which the number of states are same as those of DNA, RNA, tRNA, and amino-acids. Even more, it is possible to identify genetic code as a natural correspondence between the dark counterparts of DNA/RNA codons and dark amino-acids and the numbers of DNAs/RNAs coding given amino-acid are same as in the vertebrate code [L4]. What is new is that the dark codons do not reduce to ordered products of letters.

1.1 Insights

The above observations inspire several speculative insights.

1. The emergence of dark nuclei identified as dark proton sequences would relate to Pollack’s effect in which irradiation of water generates in presence of gel phase bounding the water what Pollack calls exclusion zones (EZs). EZs are negatively charged and water has effective stoichiometry H\textsubscript{1.5}O. EZs deserve their name: somehow they manage to get rid of various impurities: this might be very important if EZs serve as regions carrying biologically important information. The protons of water molecules must go somewhere and the proposal is that they go to the magnetic body of some system consisting of flux tubes. The flux tubes contain the dark protons as sequences identifiable as dark nuclei.
2. Since nuclear physics precedes chemistry, one can argue that prebiotic life is based on these dark biomolecules serving as a template for ordinary biomolecules. To some degree biochemistry would be shadow dynamics and dark dynamics would be extremely simple as compared to the biochemistry induced by it. In particular, DNA replication, transcription, and translation would be induced by their dark variants. One can even extend this vision: perhaps also ordinary nuclear physics and its scaled up counterpart explaining “cold fusion” are parts of evolutionary hierarchy of nuclear physics in various scales.

3. Nature could have a kind of R&D lab allowing to test various new candidates for genes by using transcription and translation at the level of dark counterparts of the ordinary basic biomolecules.

### 1.2 Conditions on the model

The model must satisfy stringent conditions.

1. Both the basis A, T, C, G and A, U, C, G as basic chemical building bricks of RNA and DNA must have emerged without the help of enzymes and ribozymes. It is known that the biochemical pathway known as pentose-phosphate pathway (see [http://tinyurl.com/y9akkwok](http://tinyurl.com/y9akkwok)) generates both ribose and ribose-5-phosphate defining the basic building brick of RNA. In DNA ribose is replaced with de-oxiribose obtained by removing one oxygen. Pyrimidines U, T, and C with single aromatic ring are are reported by NASA to be generated under outer space conditions (see [http://tinyurl.com/y7sh9zk4](http://tinyurl.com/y7sh9zk4)). Carell et al [13] (see [http://tinyurl.com/z65kpyo](http://tinyurl.com/z65kpyo)) have identified a mechanism leading to the generation of purines A and G, which besides pyrimidines A,T (U) are the basic building bricks of DNA and RNA. The crucial step is to make the solution involved slightly acidic by adding protons. TGD inspired model for the mechanism involves dark protons [L6] [K4]. Basic amino-acids are generated in the Miller-Urey type experiments (see [http://tinyurl.com/4q2arv](http://tinyurl.com/4q2arv)). Also nucleobases have been generated in Miller-Urey type experiments [I5]. Therefore the basic building bricks can emerge without help of enzymes and ribozymes so that the presence of dark nuclei could lead to the emergence of the basic biopolymers and tRNA.

2. Genetic code as a correspondence between RNA and corresponding dark proton sequences must emerge. Same true for DNA and also amino-acids and their dark counterparts. The basic idea is that metabolic energy transfer between biomolecules and their dark variants must be possible. This requires transitions with same transition energies so that resonance becomes possible. This is also essential for the pairing of DNA and dark DNA and also for the pairing of say dark DNA and dark RNA. The resonance condition could explain why just the known basic biomolecules are selected from a huge variety of candidates possible in ordinary biochemistry and there would be no need to assume that life as we know it emerges as a random accident.

3. Metabolic energy transfer between molecules and their dark variants must be possible by resonance condition. The dark nuclear energy scale associated with biomolecule could correspond to the metabolic energy scale of .5 eV. This condition fixes the model to a high extent but also other dark nuclear scales with their own metabolic energy quanta are possible. In fact, the dark nuclear binding energy for $k = 151$ scaled up from the typical value of the ordinary nuclear binding energy about 1 MeV is .5 eV.

### 1.3 Vision

The basic problem in the understanding of the prebiotic evolution is how DNA, RNA, amino-acids and tRNA and perhaps even cell membrane and microtubules . The individual nucleotides and amino-acids emerge without the help of enzymes or ribozymes but the mystery is how their polymers emerged. If the dark variants of these molecules served as templates for their generation one avoids this hen-and-egg problem. The problem how just the biomolecules were picked up
from a huge variety of candidates allowed by chemistry could be solved by the resonance condition making possible metabolic energy transfer between biomolecules and dark nuclei.

The basic question is to what p-adic length scales $L(k)$ DNA, RNA and amino-acids correspond. The original hypothesis was that the p-adic length scale assignable to dark DNA is consistent with the radius of ordinary DNA. It however turned out that this implies that the binding energy scale of corresponding dark nuclear physics is too high for the recent biology. Also the assumption that the dark variant of DNA double strand is horizontally scaled up variant of ordinary DNA strand excludes this identification since it requires that the horizontal size scale of dark DNA strand is larger than that of ordinary DNA strand.

DNA coil has radius $L(151) = 10$ nm and this suggests that dark DNA radius does not correspond to the radius of ordinary DNA (as assumed in the original version of this text) but to the p-adic length scale $L(151)$, where $k = 151$ corresponds to first Gaussian Mersenne prime belonging to the group $k = 151, 157, 163, 167$. The primes $k > 151$ would correspond to higher level coilings of DNA. From this hypothesis one ends up to the proposal that RNA, tRNA, and amino-acids correspond to $k = 149$. This picture follows essentially from the contraints posed by various biological anomalies.

Also the smaller primes $k = 127, 131, 137, 139$ can be present in pre-biotic evolutions. This hierarchy of dark nuclear physics leads to a vision about how prebiotic evolution led via DNA era to the recent biology. Unidentified infrared bands (UIBs) from interstellar space identified in terms of transition energies of dark nuclear physics support this vision and one can compare it to PAH world hypothesis.

The vision about dark matter as a controller of biomatter leads to ask whether cell membrane and microtubules could correspond to 2-D analogs of RNA strands associated with dark RNA codons forming lattice like structures related to by radial scaling to their counterparts at the level of ordinary biomatter. This is supported by p-adic length scale hypothesis and thermodynamical considerations. These 2-D structures could represent 2-D variants of 1-D structures represented by DNA, RNA, and amino-acids with each node of lattice representing code letter.

Thermal constraints allow cell membrane of thickness about 5 nm as an additional realization of $k = 149$ level with $n = 2^{22}$ in terms of lipids as analogs of RNA codons. For $k = 149$ metabolic energy quantum is predicted to be $0.5$ eV. The thickness of neuronal membrane in the range 8-10 nm and could correspond to $k = 151$ and $n = 2^{23}$ in accordance with the idea that it corresponds to higher level in the cellular evolution reflecting that of dark nuclear physics. The energy quantum of ordinary Josephson radiation is just at the verge of thermal threshold. This could be understood in terms of minimization of metabolic resources. For bosonic singly charged ions the Josephson energy would be below the thermal threshold. The notion of generalized Josephson junction saves the situation. For massive particles associated with flux tubes the thermal energy $T/2$ is below the potential energy defined by action potential and that of metabolic energy quantum.

Also microtubules could correspond to $k = 151$ realization for which metabolic energy quantum is $E_{ex}(151) = 0.25eV$. Of course, the replacement of $E_{ex} = 1$ MeV for ordinary nuclei with $E_{ex} = 2$ MeV would give $E_{ex}(151) = 0.5$ eV so that one must take these estimates as order of magnitude estimates only. Also a proposal for how microtubules could realize genetic code with the 2 conformations of tubulin dimers and 32 charges associated with ATP and ADP accompanying the dimer thus realizing the analogs of 64 analogs of RNA codons is made.

The great vision would be that hierarchy of dark variants of DNA, RNA, amino-acids and their replication, transcription, and translation would be behind biological replication in various scales. Ordinary bio-chemistry would be shadow dynamics doing its best to mimic what happens at the level of dark matter. The reduction of bio-physics to that of dark matter level would mean a huge simplification of the vision about living matter.

2 About dark variants of DNA, RNA, and amino-acids

To make progress one must construct a concrete model for the dark nuclei. The recent picture relies strongly on various anomalies to which TGD provides a solution. The TGD inspired model for “cold fusion” leads to the notion of dark nuclear physics - actually hierarchy of them labelle by the values of $h_{eff}/h = n$ and corresponding p-adic length scales. Second basic idea [L2] is that cylindrical variants of EZs discovered by Pollack give rise to the dark counterparts of DNA,
2.1 Dark variant of DNA

RNA, and amino-acids as dark proton sequences. tRNAs would be analogs of tritium and $^3\text{He}$. Pollack effect serves as a strong constraint for the model. Also the effects of ELF em fields on vertebrate brain [?] combined with the rather recent finding about clustering of RNA II polymerase molecules [L14] exhibiting Comorosan effect [L8] provide valuable constraints on the model [L13]. The outcome of the arguments is that single strand of DNA, mRNA, tRNA and amino-acids most naturally correspond to $k = 149$ and double stranded DNA to $k = 151$.

**Remark:** The following argumentation is kind of Sherlock-Holmes-ing using all possible hints as constraints to select between imagined options rather than glorious march from axioms to theorems and thus not science in the usual sense.

2.1 Dark variant of DNA

Concerning the identification of the size scale of dark DNA one can consider several options. The first guess was that the scale is same as for ordinary DNA: $L(141)=.34$ nm obtained by scaling from the distance of protons in the $k = 127$ dark nucleus implicated by the findings of Holmlid et al [?, ?] [L3]. It however turns out that the p-adic length scale assignable to dark DNA is most naturally $k = 151$ corresponding to the thickness 10 nm of DNA coil. The hypothesis that the integer $k$ labelling p-adic length scale is prime is attractive working hypothesis leaving very few options under consideration. The options $k = 137$ and $k = 149$ are excluded since the pairing of dark DNA and ordinary DNA would not be possible without the coiling of ordinary RNA around dark DNA. This leaves only options for which $k \geq 149$ for prime values of $k$.

**Remark:** The p-adic length scale associated with a system is defined to be $L(k)$ if the size of the system is in the half open interval $[L(k), L(k+1))$. One can also consider the possibility that p-adic length scale corresponds to the upper end of $[L(k-1), L(k))$.

2.1.1 General considerations

Consider first some background.

1. The TGD based model leads to the proposal for a formation of this kind of dark nuclear strings such that the distance between protons is rather precisely electron Compton length $L_e \approx 4 \times 10^{-12}$ meters explains “cold fusion” in terms of dark nucleosynthesis which should have preceded ordinary nucleosynthesis by heating the material to the temperature required by it [L8] [K18].

Dark nucleosynthesis would have produced part of heavier nuclei outside stars. The binding energy scale for dark nuclear physics would be scaled down like 1/length and 2.6 MeV binding energy per nucleon for $^3\text{He}$ of the ordinary nuclei would be scaled down by a factor $2^{-11}$ to 1.3 keV. Note however that it is excitation energies of order 1 MeV what matters and would scale down to .5 keV. This level does not yet correspond to biology as we know it but could be one step in the evolutionary hierarchy leading from nuclear physics also based on nuclear strings to biology involving increase of Planck constant $h_{eff}/h = n$ identifiably as the dimension of algebraic extension of rationals characterizing the complexity of the dynamics.

2. These dark nuclei have $h_{eff}/h = n = 2^{11}$ (or near to it) and cannot be those responsible for the dark variants of biomolecules since the distances of dark protons given by electron Compton length are much shorter than the distance between DNA nucleotides about .34 nm, which is roughly 142 times the electron Compton length $2.4 \times 10^{-3}$ nm.

3. The distance between the dark protons appearing as counterparts of DNA nucleotides should be larger than that between ordinary DNA nucleotides. The simplest assumption that dark DNA coil is a horizontally scaled variant of DNA coil with same twisting angle so that DNA nucleotides are projected horizontally to their dark counterparts at the surface of a cylinder. Once the p-adic length scale of this cylinder is given, the distance between dark protons is fixed by p-adic scaling from the distance between dark protons for $k = 127$ case - that is electron Compton length. In the case of uncoiled RNA/AA one could have also a coil rotating around the ordinary RNA/AA.
The distance between dark nucleotides must be longer than the distance $3 \times .34 \sim 1$ nm taken by single ordinary DNA codon. If $k$ is prime this leaves only $k = 149$ or $k = 151$ into consideration.

4. The negative charge of DNA and RNA assignable to one oxygen of phosphate combining with ribose and DNA/RNA base could come from the tubular EZ formed in the formation of DNA. The negative charge of phosphates and the positive charge of dark protons could guarantee the stability of pairs of dark proton sequences and ordinary RNA and DNA. DNA strand has radius of $R = 1$ nm. The Debye length $R_D$ of DNA gives rough idea about the scale above which the negative charge of DNA nucleotides associated with the phosphates screened. $R_D$ should be longer than $R$; otherwise it is not possible to speak about charge of DNA only atomic length scales. One should have $R_D > R$: otherwise it does not make sense to assign negative DNA charge except in atomic length scales. The simplest option is that dark DNA has size scale $L(151)$.

Remark: The rough estimates depend on how one identifies p-adic length scale. For the identification as $L(k) = \sqrt[5]{L_e(k)}$ motivated by the mass formula for electron, one would have $L(149) = \sqrt[5]{L_e(149)} = 0.67$ nm. With this interpretation the estimate for the screening radius would be still shorter than $R$.

Remark: Scaled up hadron physics would be associated with flux tubes of the magnetic body of the codon at which one would have nucleons as 3-quark color singlets. I have already earlier proposed that scaled variants of hadron physics [K5] appear in TGD inspired biology. One motivation comes from honeybee dance [A1].

The pairing dark AAs with positive charge with ordinary AAs might lead to problems since 16 AAs are neutral. The only charged AA residues are Lys (+), Arg (+), Asp (-) and Glu (-).

1. The formation mechanism for dark proton sequences gives for dark AAs a large positive charge. AAs are however not accompanied by negatively charged phosphate ions. Does charge neutrality require that the dark bonds between dark proton has negative charge so that one has effectively neutron?

2. Proteins in water environment have a charge due to protons or electrons attaching to them. This charge depends on pH and becomes negative above certain critical pH. One might think that the limit of very large pH (no protons) corresponds to the situation in which the electrons of EZ attach to AAs.

Dark codons do not have decomposition to letters whereas ordinary codons have. In a well-defined sense one could say that dark code is “holistic” whereas the ordinary code is “reductionistic”.

1. This brings in mind western written language in which words decompose to letters. In some eastern languages the symbols of written language correspond to entire words. Do these differences correspond at deeper level to ordinary and dark genes. Could the analytic and holistic aspects of cognition relate to the differences between ordinary and dark code.

2. One cannot exclude the entanglement between codons and evolution as emergence of entanglement even suggests this. Could this kind of entanglement give rise to basic units of DNA, in particular genes and introns. Could the decomposition of gene into coding regions and introns correspond to a decomposition to unentangled products of internally entangled pieces. This would increase exponentially the degrees of freedom involved and explain why organisms with practically the same code can be at so different evolutionary levels. In the splicing process when intronic portions are cut out from DNA sequence. Do the remaining
pieces of RNA get entangled or does the decomposition of dark RNA to unentangled pieces have some meaning? Note that also ordinary RNA would be entangled or entangled. Could introns provide the means for decomposing the coding RNA to unentangled pieces.

3. The most natural possibility is that entanglement contains superposition of codon sequences in which each sequence codes for the same AA. The chemical codons appearing in the superposition have different masses and chemical properties but in zero energy ontology (ZEO) this is possible. Situation would be like for a superconductor in which coherent state means superposition of states with different numbers of Cooper pairs and thus different fermion number in standard ontology but in ZEO this problem disappears.

2.1.2 Why one must have $k = 151$ for dark DNA

It was already found that for prime values of $k$ the options $k < 149$ are not possible for dark DNA since ordinary DNA should coil around dark DNA. There is also second objection against prime $k < 149$ from energetics inspiring the hypothesis DNA corresponds to $k = 151$.

1. The scaling of the dark nuclear binding energy $E_b \sim 7$ MeV per nucleon as $L(107)/L(k)$ predicts very high binding energies for primes $k < 149$. For instance, $k = 139$ would correspond to the scaled binding energy $E_b(139) = E_b L(107)/L(139)$, $E_b \sim 7$ MeV, which is typical nuclear binding energy. This gives $E_b(139) = E_b/2^{(139−107)/2} = .14$ keV. For $k = 139$ the typical nuclear excitation energy $E_{ex} = 1$ MeV scales down to 20 eV, which is still very high but could correspond to energies of atomic transitions. For $k = 151$ it $E_b$ scales down to 3.5 eV. The typical dark excitation energy for $k = 151$ is $E_{ex}(151) = .5$ eV and the identification as a nominal value of metabolic energy quantum is attractive. Dark nuclear physics might therefore control biochemistry using dark nuclear transitions as a tool to provide desire energy currency.

2. The TGD based explanation of Pollack effect provides a consistency test for the idea [12]. In Pollack effect IR light (besides either kinds of energy feeds) induces the formation of negative charged exclusion zones (EZs) in water bounded by gel phase. In TGD based model this would correspond to the formation of dark proton sequences at magnetic flux tubes. The scale of dark nuclear binding energy would be most naturally in eV scale. The binding energy scale of hydrogen atoms in water molecules is about 5 eV which suggests that the binding energy scale for dark protons sequences is smaller since otherwise energy would be liberated. This would suggest $k = 149$ as will be found.

3. One can imagine that an external perturbation induces

(a) a transition in which the proton bound to water molecule transforms to its dark variant in higher energy state or
(b) that the proton goes over a potential wall, whose height is measured in eV:s.

If the dark nuclear binding energy is higher than the binding energy of proton in water molecule, the process should liberate energy and could occur spontaneously unless high potential wall prevents it. Hence the first option seems the only realistic one. Note that one could consider the cancellation of dark nuclear binding energy and repulsive Coulomb energy which scale in the same manner as function of $p$-adic length scale so that still the net energy would scale increase in shorter $p$-adic length scales.

Pollack effect suggests that if $k$ is prime, one must have $k = 149$ for dark proton sequences formed in Pollack effect.

1. For $k = 149$ one has $E_b(151) \sim E_b/2^{(149−107)/2} = 3.5$ eV for $E_b = 7$ MeV, which is in UV range slightly above the visible range. The binding energy of hydrogen atom in water is about 5 eV which would require the incoming radiation to have energy 1.5 eV which is indeed in IR range. This option looks therefore realistic.
2.1 Dark variant of DNA

For \( k = 151 \) one would have \( E_b(151) \sim 7MeV/2^{(151-107)/2} \approx 1.75 \text{ eV} \), which just above the IR energy range. Now the energy needed to transform ordinary protons to dark protons in Pollack effect would be in UV range so that this options seems to be excluded.

This argument suggests that dark proton sequences generated in Pollack effect are analogs of single DNA strand, which would naturally correspond to \( L(149) = L(151)/2 \). Also RNA would naturally correspond to this scale.

1. \( L(151) \approx 10 \text{ nm} \) is the thickness of coiled DNA double strand. The size scale of dark nucleons would be \( L(151) \) and the dark DNA strand should be horizontally scaled variant of ordinary DNA strand by a scaling factor \( \lambda \sim L(151)/33 \text{ nm} = 30 \). DNA double strand would be obtained by a transversal scaling from the ordinary DNA double strand.

2. The higher coilings of DNA could correspond to higher horizontally scaled variants of DNA corresponding to \( k = 157, 163, 167 \). \( k = 167 \) would correspond to nuclear membrane length scale of 2.5 \( \mu \text{m} \). The emergence of nuclear membrane in \( k = 151 \) length scale would have been accompanied by the emergence of dark DNA in this scale. Cell membrane could correspond to \( k = 173 \) and \( p \)-adic length scale 17.6 \( \mu \text{m} \). Neurons have size varying from 4-100 micrometers (the definition of size depends on whether one includes axons) and might correspond to \( k = 179, 181 \) and length scales of .16 mm and perhaps even .32 mm.

The only justification for this speculative picture is that it is consistent with the other basic ideas about TGD inspired quantum biology.

1. Cisse et al [I4] found that RNA II polymerase molecules cluster during transcription and their dynamics involves multiples of the time scale \( \tau = 5 \text{ seconds} \). Comorosan reported long time ago that just these time scales are universal bio-catalysis [I8]. The TGD inspired model [L13] for the findings of Cisse et al allows to sharpen the TGD based view about quantum biology considerably.

2. The basic parameter of the model is the value of gravitational Planck constant \( h_{gr} = GM_Dv_0/\text{v}_0 \) assigned to magnetic flux tubes mediating gravitational interactions. Already earlier work gives estimates for the value \( M_D \) of dark mass and velocity parameter \( v_0 \) and the model leads to the same estimates. The identification of the values of \( \tau \) as Josephson periods assuming the potential difference \( V \) along flux tubes connecting reacting molecules is universal and same as over neuronal membrane fixed the value of \( h_{gr} \). The value of \( V \) along flux tube serving as Josephson junction would be universal and equal to membrane potential. Josephson radiation would have energies coming as multiples of \( ZeV \) just above the thermal energy at physiological temperatures fixed by the membrane potential.

3. The model forces the conclusion that the endogenous magnetic field \( B_{end} \) has at its upper bound \( B_{end} = .2 \text{ Gauss} \) deduced from the findings of Blackman about effects of ELF em fields on vertebrate brain [?]. The earlier ad hoc hypothesis was that \( B_{end} = .2 \text{ Gauss} \) is minimum value of \( B_{end} \). Furthermore, for the required value of \( h_{gr} \), \( B_{end} = .2 \text{ Gauss} \) corresponds to dark cyclotron energy of .12 keV, which is surprisingly large energy at the upper end of UV band: the earlier intuitive guess was that energy scale is in visible range. Also harmonics of cyclotron frequencies were found to have effects so that really large energy scales are involved with the interaction of ELF radiation and one can ask whether this picture really makes sense. This raises a question about the mechanism of the interaction of ELF em radiation with living matter. One also wonder why the ELF radiation has effects on both behavior and physiology.

Assume

(a) that dark photons with energies coming as multiples of .12 keV are in question,
(b) that these dark photons excite dark cyclotron states in the cellular length scale deduced from flux quantization and
(c) that the dark cyclotron photons radiated as the excited cyclotron states return to the ground states perform some control action on ordinary DNA coil - this is in accordance with the basic vision about the role of magnetic body.
X rays have energy range varying from 100 eV to 100 keV and wavelengths varying from 10 nm to .01 nm. The wavelength of an ordinary photon resulting from dark photon with energy of .12 keV would be of order 10 nm, the radius of DNA coil for \( k = 151 \).

Could this energy induce an analog of standing EM wave in transversal degrees of freedom of DNA perhaps transformable to many phonon state with very large number of photos and thus classical acoustic wave? This would allow to understand how cyclotron harmonics can have non-trivial effects. The effects of ELF radiation on behavior and physiology could be understood as gene expression induced by the irradiation.

Both dark cyclotron radiation and radiation generated in dark nuclear transitions could have biological effects.

1. Can one relate energy scale of .12 keV associated with dark cyclotron radiation to atomic physics? The ionization energies behave as \( Z_{\text{eff}}^2/n^2 \), where \( Z_{\text{eff}} \) is nuclear charge minus the charge of the closed shells. \( Z_{\text{eff}} \) is also reduced by electronic screening by other valence electrons. The binding energies of valence electrons decrease with the principal quantum number \( n \) so that only \( n = 2 \) row of the periodic table might allow so high ionization energies for valence electrons.

Oxygen is certainly the first candidate to consider. The ionization energy for oxygen is .12 eV from an estimate assuming that the effective nuclear charge is 6 (with the contribution of 2 valence electrons subtracted). The actual value is 68.9 eV: the reduction is due to electron screening. This value is smaller than the estimate estimate for \( E_b = .12 \text{ keV} \) and since harmonics of this energy are involved, the interpretation in terms of ionization does not make sense.

2. Not only oxygen but also heavier elements are ionized in living matter and at least to me this has remained more or less a mystery. Could dark photons emitted by dark nuclei of MB perform control by inducing the transitions and even ionization of oxygen and other biologically important atoms. The process could proceed also in opposite direction. The energy scale would correspond to that of nuclear excitations scaled down by the above ratio of p-adic length scales. If the energy scale of ordinary nuclear excitations is taken to be about 1 MeV, the dark energy scale for \( k = 127 \) assignable to the dark nuclei created in “cold fusion” is keV. For \( k = 131 \) the scale would be 250 eV and above the ionization energy scales for valence electrons. For \( k = 137 \) the scale would be 17 keV. These dark nuclear transitions could generate dark photons inducing transitions of atoms and even ionizations.

2.2 What about dark variants of RNA, tRNA, and AAs?

Also RNA and AAs should have dark variants and one should understand their role. Suppose that the integer \( k \) characterizing the p-adic length scale is prime. The vision about RNA era preceding DNA era suggests that RNA accompanying dark RNA is at lower level in the evolution, and hence the value of \( h_{\text{eff}} \) is smaller for dark RNA than for dark DNA. Also the p-adic length scale for RNA would be shorter.

1. The most natural option is that RNA corresponds to \( k = 149 \) as also single DNA strand. This would conform with the above suggestion that the Pollack effect generates \( k = 149 \) dark proton sequence (dark RNA?). DNA double strand would correspond to \( k = 151 \).

The emergence of \( k = 151 \) level would mean the emergence of structures with scale characterized by \( L(151) \). This includes DNA double strand forming a coil with thickness \( L(151) \) and nuclear and cell membranes. During RNA era these structures would have been absent. Both DNA double strand and cell membrane have binary structures. Therefore single DNA strand and lipid layer could correspond to \( k = 149 \). In transcription DNA opens and double strand becomes pair of strands having naturally \( k = 149 \). Therefore mRNA should have also \( k = 149 \).

2. If AAs correspond to \( k = 149 \) then also tRNA should correspond to \( k = 149 \). On the other hand, tRNA does not form strands and should be more elementary structure than RNA.
Could tRNA corresponds to $k = 139$ or $k = 137$? This would require that also the attached AA would correspond to $k = 139$ or $k = 137$, which does not look plausible.

**Remark:** TGD vision assumes tRNA was present already at RNA era and the role of AA in tRNA was to catalyze RNA replication. In fact, RNA could have been just tRNA at very early stages.

What about AAs? The following arguments suggest that one has $k = 149$ for both AAs and RNA.

1. For dark AAs one can imagine p-adic evolutionary hierarchy analogous to that for DNA. In TGD inspired vision AA sequences emerged together with DNA. Proteins can appear also as coils. Since mRNA pairs with single DNA strand and AAs with mRNA, it seems that AAs should correspond to $k \geq 149$?

2. One could however argue that AAs are building bricks rather than information molecules and $k$ could be rather small for dark AAs. Dark AAs should pair with proteins. Pairing without coiling is possible only if the length per letter is same as the length per AA and thus same as for DNA letter, which is longer than the length taken by $k = 139$ dark proton. Also this suggests $k = 149$ for dark AAs and their coiling around the ordinary AAs.

### 2.3 Clustering of RNA polymerase molecules and Comorosan effect

Once again I had good luck: I received a link (see [http://tinyurl.com/y7bego83](http://tinyurl.com/y7bego83)) to a highly interesting popular article telling about the work by Ibrahim Cisse at MIT and his colleagues [I4] (see [http://tinyurl.com/y9wzt5yl](http://tinyurl.com/y9wzt5yl)) about the clustering of RNA polymerase proteins in the transcription of RNA. Similar clustering has been observed already earlier and interpreted as a phase separation giving rise to protein droplets [L17]. Now this interpretation is not proposed by experiments but they say that it is quite possible but they cannot prove it.

I have already earlier discussed the coalescence of proteins into droplets as this kind of process in TGD framework [K19] [L17]. The basic TGD based ideas is that proteins - and biomolecules in general - are connected by flux tubes characterized by the value of Planck constant $h_{\text{eff}} = n \times h_0$ for the dark particles at the flux tube. The higher the value of $n$ is the larger the energy of given state. For instance, the binding energies of atoms decrease like $1/n^2$. Therefore the formation of the molecular cluster liberates energy usable as metabolic energy.

**Remark:** $h_0$ is the minimal value of $h_{\text{eff}}$. The best guess is that ordinary Planck constant equals to $h = 6h_0$ [L3] [L14] (see [http://tinyurl.com/goruuzm](http://tinyurl.com/goruuzm) and [http://tinyurl.com/y9jxyjns](http://tinyurl.com/y9jxyjns)).

### 2.3.1 TGD view about the findings

Gene control switches - such as RNA II polymerases in DNA transcription to RNA - are found to form clusters called super-enhancers. Also so called Mediator proteins form clusters. In both cases the number of members is in the range 200-400. The clusters are stable but individual molecules spend very brief time in them. Clusters have average lifetime of $5.1 \pm 4$ seconds.

Why the clustering should take place? Why large number of these proteins are present although single one would be enough in the standard picture. In TGD framework one can imagine several explanations. One can imagine at least following reasons.

1. If the initiation of transcription is quantum process involving state function reduction, clustering could allow to make this process process deterministic at the level of single gene in spite of the non-determinism of state function reduction. Suppose that the initiation of transcription is one particular outcome of state function reduction. If there is only single RNA II polymerase able to make only single trial, the changes to initiate the transcription are low. This could be the case if the protein provides metabolic energy to initiate the process and becomes too “tired” to try again immediately. In nerve pulse transmission there is analogous situation: after the passing of the nerve pulse generation the neuron has dead time period. As a matter of fact, it turns out that the analogy could be much deeper.
2.3 Clustering of RNA polymerase molecules and Comorosan effect

How to achieve the initiation with certainty in this kind of situation? Suppose that the other outcomes do not affect the situation appreciably. If one particular RNA polymerase fails to initiate it, the others can try. If the number of RNA transcriptase molecule is large enough, the transcription is bound to begin eventually! This is much like in fairy tales about princess and suitors trying to kill the dragon to get the hand of princess. Eventually comes the penniless swineherd.

2. If the initiation of transcription requires large amount of metabolic energy then only some minimal number of $N$ of RNA II polymerase molecules might be able to provide it collectively. The collective formed by $N$ molecules could correspond to a formation of magnetic body (MB) with a large value of $h_{\text{eff}} = n \times h_0$ and controlling the molecules and inducing its coherent behavior. The molecules would be connected by magnetic flux tubes.

3. If the rate for occurrence is determined by an amplitude which is superposition of amplitudes assignable to individual proteins the rate is proportional to $N^2$, $N$ the number of RNA II polymerase molecules. The process for the cluster is reported to to be surprisingly fast as compared to the expectations - something like 20 seconds. The earlier studies have suggests that single RNA polymerase stays at the DNA for minutes to hours.

Clustering could allow to speed up bio-catalysis besides the mechanism allowing to find molecules to find by a reduction of $h_{\text{eff}}/h = n$ for the bonds connecting the reactants and the associated liberation of metabolic energy allowing to kick the reactants over the potential wall hindering the reaction.

Concerning the process of clustering there are two alternative options both relying on the model of liquid phase explaining Maxwell’s rule assuming the presence of flux tube bonds in liquid and of water explaining its numerous anomalies in terms of flux tubes which can be also dark (see http://tinyurl.com/ydhknc2c).

1. **Option I**: Molecules could form in the initial situation a phase analogous to vapour phase and there would be very few flux tube bonds between them. The phase transition would create liquid phase as flux tube loops assignable to molecules would reconnect form flux tube pairs connecting the molecules to a tensor network giving rise to quantum liquid phase. The larger then value of $n$, the longer the bonds between molecules would be. This kind of model [?] (see http://tinyurl.com/yassnhzb) is used to explain the strange findings that a system consisting of plastic balls seems to show primitive features of life such as metabolism.

2. **Option II**: The molecules are in the initial state connected by flux tubes and form a kind of liquid phase and the clustering reduces the value of $h_{\text{eff}}/h = n$ and therefore the lengths of flux tubes. This would liberate dark energy as metabolic energy going to the initiation of the transcription. One could indeed argue that connectedness in the initial state with large enough value of $n$ is necessary since the protein cluster must have high enough “IQ” to perform intelligent intentional actions.

Protein blobs are said to be drawn together by the “floppiness” bits (pieces) of intrinsically disordered proteins. What could this mean in the proposed picture? Disorder would mean absence of correlations between building bricks of floppy parts of the proteins in translational degrees of freedom.

1. Could floppiness correspond to low string tension assignable to long flux loops with large $n$ assignable to the building bricks of “floppy” pieces of protein? Could reconnection for these loops give rise to pairs of flux tubes connecting the proteins in the transition to liquid phase (Option I)? Floppiness would also make possible to scan the environment by flux loops to get in touch with the flux loops of other molecules and in the case of hit (cyclotron resonance) induce reconnection.

2. In spite of floppiness in this sense, one could have quantum correlations between the internal quantum numbers of the building bricks of the floppy pieces. This would also increase the value of $n$ serving as molecular IQ and provide molecule with higher metabolic energy liberated in the catalysis.
2.3.2 About Comorosan effect and clustering of RNA II polymerase proteins

What about the interpretation of the time scales $\tau$ equal 5, 10, and 20 seconds appearing in the clustering of RNA II polymerase proteins and Mediator proteins? What is intriguing that so called Comorosan effect \cite{18,12} involves time scale of 5 seconds and its multiples claimed by Comorosan long time ago to be universal time scales in biology. The origin of these time scales has remained more or less a mystery although I have considered several TGD inspired explanations for this time scale is based on the notion of gravitational Planck constant \cite{15} (see http://tinyurl.com/yb8fw3kq).

One can consider several starting point ideas, which need not be mutually exclusive.

1. The time scales $\tau$ associated with RNA II polymerase and perhaps more general bio-catalytic systems as Comorosan’s claims suggest could correspond to the durations of processes ending with “big” state function reduction. In zero energy ontology (ZEO) there are two kinds of state function reductions \cite{11}. “Small” state function reductions - analogs of weak measurements - leave the passive boundary of causal diamond (CD) unaffected and thus give rise to self as generalized Zeno effect. The states at the active boundary change by a sequence of unitary time evolutions followed by measurements inducing also time localization of the active boundary of CD but not affecting passive boundary. The size of CD increases and gives rise to flow of time defined as the temporal distance between the tips of CD. Large reductions change the roles of the passive and active boundaries and mean death of self. The process with duration of $\tau$ could correspond to a life-time of self assignable to CD.

Remark: It is not quite clear whether CD can disappear and generated from vacuum. In principle this is possible and the generation of mental images as sub selves and sub CDs could correspond to this kind of process.

2. In \cite{15} I proposed that Josephson junctions are formed between reacting molecules in bio-catalysis. These could correspond to the shortened flux tubes. The difference $E_J = ZeV$ of Coulomb energy of Cooper pair over flux tube defining Josephson junction between molecules would correspond to Josephson frequency $f_J = 2eV/h_{\text{eff}}$. If this frequency corresponds to $\tau_J = 5$ seconds, $h_{\text{eff}}$ should be rather large since $E_J$ is expected to be above thermal energy at physiological temperature.

Could Josephson radiation serve as a kind of of synchronizing clock for the state function reductions so that its role would be analogous to that of EEG in case of brain? A more plausible option is that Josephson radiation is a reaction to the presence of cyclotron radiation generated at MB and performing control actions at the biological body (BB) defined in very general sense. In the case of brain dark cyclotron radiation would generate EEG rhythms responsible for control via genome and dark generalized Josephson radiation modulated by nerve pulse patterns would mediate sensory input to the MB at EEG frequencies.

A good guess motivated by the proposed universality of the Comorosan periods is that the energy in question does not depend on the catalytic system and corresponds to Josephson energy for protein through cell membrane acting as Josephson junction and giving to ionic channel or pump. The flux tubes themselves have universal properties.

3. The hypothesis $h_{\text{eff}} = h_{gr} = GMm/\beta_0 c$ of Nottale \cite{[13]} for the value of gravitational Planck constant \cite{13,8,20,19} gives large $h$. Here $v_0 = \beta_0 c$ has dimensions of velocity. For dark cyclotron photons this gives large energy $E_c \propto h_{gr}$ and for dark Josephson photons small frequency $f_J \propto 1/h_{gr}$. Josephson time scale $\tau_J$ would be proportional to the mass $m$ of the charged particle and therefore to mass number $A$ of ion involved: $f_J \propto A$ possibly explaining the appearance of multiples of 5 second time scale. Cyclotron time scale does not depend on the mass of the charged particle at all and now sub-harmonics of $\tau_J$ are natural.

The time scales assignable to CD or the lifetime-time of self in question could correspond to either cyclotron of Josephson time scale $\tau$.

1. If one requires that the multiplies of the time scale 5 seconds are possible, Josephson radiation is favoured since the Josephson time scale proportional to $h_{gr} \propto m \propto A$, $A$ mass number of ion.
The problem is that the values \( A = 2, 3, 4, 5 \) are not plausible for ordinary nuclei in living matter. Dark nuclei at magnetic flux tubes consisting of dark proton sequences could however have arbitrary number of dark protons and if dark nuclei appear at flux tubes defining Josephson junctions, one would have the desired hierarchy.

2. Although cyclotron frequencies do not have sub-harmonics naturally, MB could adapt to the situation by changing the thickness of its flux tubes and by flux conservation the magnetic field strength to which \( f_c \) is proportional to. This would allow MB to produce cyclotron radiation with the same frequency as Josephson radiation and MB and BB would be in resonant coupling.

Consider now the model quantitatively.

1. For \( h_{eff} = h_qr \) one has
   \[
   r = \frac{h_q}{\hbar} = \frac{G M_D m_p}{c b_0} = 4.5 \times 10^{14} \times \frac{m_p y}{m_p b_0}. 
   \]
   Here \( y = M_D / M_E \) gives the ratio of dark mass \( M_D \) to the Earth mass \( M_E \). One can consider 2 favoured values for \( m \) corresponding to proton mass \( m_p \) and electron mass \( m_e \).

2. \( E = h_{eff} f \) gives the concrete relationship \( f = (E / eV) \times 2.4 \times 10^{14} \times (h / h_{eff}) \) Hz between frequencies and energies. This gives
   \[
   x = \frac{E}{eV} = 0.4 \times r \times \frac{f}{10^{14} \text{Hz}}. 
   \]
3. If the cyclotron frequency \( f_c \) = 300 Hz of proton for \( B_{end} = .2 \) Gauss corresponds to biophoton energy of \( x \ eV \), one obtains the condition
   \[
   r = \frac{G M_D m_p}{h b_0} \approx .83 \times 10^{12} x. 
   \]
   Note that the cyclotron energy does not depend on the mass of the charged particle. One obtains for the relation between Josephson energy and Josephson frequency the condition
   \[
   x = \frac{E_j}{eV} = 0.4 \times .83 \times 10^{-2} \times \frac{m_p}{m_p} \times \frac{f_j}{H z} \times \frac{f_j}{H z} \times E_j = \text{ZeV}. 
   \]
   One should not confuse \( eV \) in \( \text{ZeV} \) with unit of energy. Note also that the value of Josephson energy does not depend on \( h_{eff} \) so that there is no actual mass dependence involved.

For proton one would give a hierarchy of time scales as \( A \)-multiples of \( \tau(p) \) and is therefore more natural so that it is natural to consider this case first.

1. For \( f_j = 2 \) Hz corresponding to the Comorosan time scale of \( \tau = 5 \) seconds this would give \( \text{ZeV} = .66x \) meV. This is above thermal energy \( E_{th} = T = 27.5 \) meV at \( T = 25 \) Celsius for \( x > 42 \). For ordinary photon \( (h_{eff} = h) \) proton cyclotron frequency \( f_c(p) \) would correspond for \( x > 42 \) to EUV energy \( E > 42 \) eV and to wavelength of \( \lambda < 31 \) nm.

The energy scale of Josephson junctions formed by proteins through cell membrane of thickness \( L(151) = 10 \) nm is slightly above thermal energy, which suggests \( x \approx 120 \) allowing to identify \( L(151) = 10 \) nm as the length scale of the flux tube portion connecting the reactants. This would give \( E \approx 120 \) eV - the upper bound of EUV range. For \( x = 120 \) one would have \( GM_D m_p / v_0 \approx 10^{14} \) requiring \( \beta_0 / y \approx 2.2 \). The earlier estimates \([K19]\) for the mass \( M_D \) give \( y \approx 2 \times 10^{-4} \) giving \( \beta_0 \approx 4.4 \times 10^{-4} \). This is rather near to \( \beta_0 \approx 2^{-11} \approx m_e / m_p \) obtained also in the model for the orbits of inner planets as Bohr orbits.

For ion with mass number \( A \) this would predict \( \tau_A = A \times \tau_p = A \times 5 \) seconds so that also multiples of the 5 second time scale would appear. These multiples were indeed found by Comoran and appear also in the case of RNA II polymerase.
2. For proton one would thus have 2 biological extremes - EUV energy scale associated with cyclotron radiation and thermal energy scale assignable to Josephson radiation. Both would be assignable to dark photons with \( h_{\text{eff}} = h_{gr} \) with very long wavelength. Dark and ordinary photons of both kind would be able to transform to each other meaning a coupling between very long lengths scales assignable to MB and short wavelengths/time scales assignable to BB.

The energy scale of dark Josephson photons would be that assignable with Josephson junctions of length 10 nm with long wavelengths and energies slightly above \( E_{\text{th}} \) at physiological temperature. The EUV energy scale would be 120 eV for dark cyclotron photons of highest energy would be fixed by flux tube length of 10 nm.

For lower cyclotron energies forced by the presence of bio-photons in the range containing visible \([K16, K17]\) and UV and obtained for \( B_{\text{end}} \) below .2 Gauss, the Josephson photons would have energies below \( E_{\text{th}} \). That the possible values of \( B_{\text{end}} \) are below the nominal value \( B_{\text{end}} = .2 \) Gauss deduced from the experiments of Blackman \([?]\) does not conform with the earlier ad hoc assumption that \( B_{\text{end}} \) represents lower bound. This does not change the earlier conclusions.

Could the 120 eV energy scale have some physical meaning in TGD framework? The corresponding wavelength for ordinary photons corresponds to the scale \( L(151) = 10 \text{ nm} \) which correspond to the thickness of DNA double strand. Dark DNA having dark proton triplets as codons could correspond to either \( k = 149 \) or \( k = 151 \). The energetics of Pollack effect suggests that \( k = 149 \) is realized in water even during prebiotic period \([L12]\) (see [htp://tinyurl.com/yalny39]). In the effect discovered by Blackman the ELF photons would transform dark cyclotron photons having \( h_{\text{eff}} = h_{gr} \) and energy about .12 keV. They would induce cyclotron transitions at flux tubes of \( B_{\text{end}} \) with thickness of order cell size scale. These states would decay back to previous states and the dark photons transformed to ordinary photons absorbed by ordinary DNA with coil structure with thickness of 10 nm. Kind of standing waves would be formed. These waves could transform to acoustic waves and induce the observed effects. Quite generally, dark cyclotron photons would control the dynamics of ordinary DNA by this mechanism.

It is natural to assume that \( B_{\text{end}} = .2 \) Gauss corresponds to the upper bound for \( B_{\text{end}} \) since magnetic fields are expected to weaken farther from the Earth’s surface: weakening could correspond to thickening of flux tubes reducing the field intensity by flux conservation. The model for hearing \([K11]\) requires cyclotron frequencies considerably above proton’s cyclotron frequency in \( B_{\text{end}} = .2 \) Gauss. This requires that audible frequencies are mapped to electron’s cyclotron frequency having upper bound \( f_c = (m_e/m_p)f_c(p) \approx 6 \times 10^5 \text{ Hz} \). This frequency is indeed above the range of audible frequencies even for bats.

For electron one has \( h_{gr}(e) = (m_e/m_p) \times h_{gr}(p) \approx 5.3 \times 10^{-4}h_{gr}(p), h_{gr}(p)/h = 4.5 \times 10^{14}/\beta_0 \). Since Josephson energy remains invariant, the Josephson time scales up from \( \tau(p) = 5 \text{ seconds} \) to \( \tau(e) = (m_e/m_p)\tau(p) \approx 2.5 \text{ milliseconds} \), which is the time scale assignable to nerve pulses \([K12, K2]\).

To sum up, the model suggests that the idealization of flux tubes as kind of universal Josephson junctions. The model is consistent with bio-photon hypothesis. The constraints on \( h_{gr} = GMiptm/v_0 \) are consistent with the earlier views and allows to assign Comorosan time scale 5 seconds to proton and nerve pulse time scale to electron as Josephson time scales. This inspires the question whether the dynamics of bio-catalysis and nerve pulse generation be seen as scaled variants of each other at quantum level? This would not be surprising if MB controls the dynamics. The earlier assumption that \( B_{\text{end}} = 0.2 \) Gauss is minimal value for \( B_{\text{end}} \) must be replaced with the assumption that it is maximal value of \( B_{\text{end}} \).

3 TGD view about the emergence of chemical life

Consider first the basic assumptions.

1. Dark DNA, RNA,... emerged before chemistry and serve as templates for ordinary DNA,
3.1 The quantum vision about the prebiotic evolution

RNA... The replication, transcription, and translation for ordinary DNA, RNA,... are induced by the corresponding processes for their dark counterparts.

2. Dark proton sequences are associated with tubular E\text{Z}s in water generated by Pollack effect.

3. The amount of entanglement measured by entanglement negentropy (having a well-defined meaning in adelic physics \([\text{L10}]\)) is expected to increase gradually during evolution. Hence one expects generation of more and more entangled sequences of dark nucleons. At the bottom - perhaps ordinary nuclear physics - one would have the product states of dark nucleons. Perhaps dark nuclear physics with \(n = 2^{11}\) came next. After that came \(n = 2^{18}\) dark nuclear physics. But which came first: dark variants amino-acids, tRNA, RNA, or DNA and their chemical counterparts? And could one see even genes as entangled codon sequences coding for the same protein?

3.1 The quantum vision about the prebiotic evolution

The following vision about quantal prebiotic evolution beginning from amino-acids suggests itself. The basic idea is that all processes took place at dark level and induced the processes for ordinary biomolecules in water environment. Even the enzyme and ribozyme actions essential in recent biology would be replaced with corresponding actions at dark level and biochemistry would reduce to shadow dynamics.

1. Amino-acids are easiest to produce (as Miller-Urey experiment demonstrated (see \(\text{http://tinyurl.com/4q2arv}\)) requiring no enzymatic action and there is just single chemical amino-acid per dark RNAs coding for it. Therefore the pairs of amino-acids and their dark variants could have emerged first. Note that proteins were not yet present.

   \textbf{Remark:} Vivo-vitro difference could mean that dark partner of biomolecule is present in vivo and missing in vitro.

2. DNA requires cell membrane. This requires RNA emerged after amino-acids. This implies that dark variants of dark tRNA, their pairing with tRNA and the pairing of dark RNA with RNA emerged next?

   This picture supports that the old TGD inspired idea about the role of tRNA during RNA era. Dark tRNA would have made possible the replication of dark RNA sequences (rather than the translation of RNA to amino-acid sequence) during this era. The dark amino-acid of dark tRNA would have served as a catalyst inducing the addition of dark RNA codon to the growing RNA sequence. No chemical transcription machinery nor DNA was needed at this stage. This would solve one hen-or-egg problem.

3. After that a revolution would have occurred. For some reason dark amino-acids began to attach to the growing sequence of amino-acids and dark RNA codon was left alone. What prevented dark RNA codon to attach to the growing dark RNA sequence? Was it the emerging entanglement between dark codons giving rise to genes as entangled pieces of DNA that made this impossible.

   This means entanglement also between the ordinary codons, which makes sense only in ZEO. If possible at all this entanglement should respect genetic code so that entangled superposition would involve only codons coding for the same amino-acid so that the translation to a single amino-acid sequence rather than their quantum superposition is possible. If more general superpositions are allowed the translation process would be like state function reduction to amino-acid sequence.

4. At this step the replication of both dark and ordinary RNA was lost and it seems that dark DNA-DNA pairs replicating dark DNA and transcribing it to dark RNA and inducing corresponding process at the level of chemistry must have emerged at the same time.

   The emergence of DNA requires also the emergence of cell membrane. Could the emergence of cell membrane relate to the emergence of dark nuclei in the \(p\)-adic length scale \(L(k), k = 149\) and could the double layered structure of cell membrane serve as an analog for that of DNA double strand? Could lipid layers correspond to 2-D analogs of DNA strand with lipids taking the role of codons?
5. Could the full genetic code emerged in step-wise manner as proposed earlier \[K3 \ K14]\? Genetic code can be seen in a good approximation as a fusion of 16-letter code and 4-letter code. This might be understood if the entanglement of dark codons emerges first as entanglement of only two first letters.

What gave rise to the correspondences between dark DNA, RNA, tRNA, amino-acids and their dark variants? How the amino-acids and nucleotide bases were selected?

1. The basic principle would be the condition that metabolic energy can be transferred between chemical and dark levels. This is possible if there identical transition energies in the spectra of biomolecules and their dark variants making possible resonance.

2. Metabolic energy quantum in the range \[.4 - .5\ eV\] should correspond to the excitation energy scale of dark dark nuclear physics if \[E_{ex} = 1\ MeV\] is taken as the estimate for a typical nuclear excitation energy. Hydrogen bonds also correspond to this energy scale but this might be just what is needed to give rise to coherent metabolic activity.

The original proposal was that dark DNA associated with ordinary DNA corresponds to \[k = 141\] assignable to the ordinary DNA but this proposal predicts \[E_{ex}(141) = 16\ eV\]. This proposal turned out to be unrealistic also in other respects. \[k = 149\] assignable to dark RNA predicts \[E_{ex}(149) = .5\ eV\] and is a more plausible option in many other aspects. Also lower values of \[k\] than \[k = 149, 151\] might be present - at least during the prebiotic stage. Pollack’s findings however support the view that the irradiation of water with IR light generates dark proton sequences with \[k = 149\]. Does this mean that the evolutionary level of water is raised to \[k = 151\] in presence of gel phase binding the water sample? Note that “cold fusion” \[L3 \ L8\] might be interpreted as creation of \[k = 127\] dark proton sequences.

To sum up: for DNA,RNA, and tRNA the emergence of entanglement would have created the chemical counterparts of quantum superpositions: ZEO is necessary since in positive energy ontology superpositions are highly implausible.

There are some questions to ponder.

1. Why the decomposition into triplets? Does resonance condition for the metabolic energy transfer select triplets as basic units and also the RNA-amino-acid correspondence? Do also intronic regions have triplets as basic units?

   One ends up to a prediction of vertebrate genetic code also from a model of music harmony \[L1\]. In fact, the model explains also its slight variation and the 2 additional amino-acids. Could this help to understand why the triplet code is so unique.

2. Could one imagine that also quarks and antiquarks were involved? Could dark nucleon pair with dark quark with same spin and isospin and color confinement forces dark proton triplets? Dark quarks indeed define a representation for A,T,C, G. In the model of topological computation \[K3 \ K14\], I have actually speculated with the possibility that dark quarks and antiquarks are paired with ordinary DNA codons.

3. Could dark conjugate protons or their triplets of parallel dark DNA strands form Cooper pairs or does pairing of dark protons triplets (their conjugates) with dark quarks (anti-quarks) give rise to bosonic states?

3.2 Unidentified Infrared Bands as a test for the proposal

Unidentified Infrared Bands (UIBs) are an ill-understood phenomenon associated with radiation coming from interstellar space. There are also other analogous phenomena having no explanation in terms of molecular transitions \[K1\] and one can ask whether they could be seen as signatures of dark nuclear physics.

1. UIBs are observed around bands around IR energies \[E \in \{.11, .20, .375\}\ eV\].

2. Poly-aromatic hydrocarbons (PAHs) (see \[http://tinyurl.com/atx4t9a\]) are known to generate UIBs \[K1\]. Therefore the UIBs from interstellar space could originate from PAHs.
### 3.2 Unidentified Infrared Bands as a test for the proposal

#### 3.2.1 TGD based models for UIBs

TGD suggests several explanations for UIBs involving new physics related to the p-adic length scale hypothesis and \( \hbar_{eff}/\hbar = n \) hierarchy.

1. For years ago I discussed a model for UIBs based on p-adic length scale hypothesis [K1]. The idea was that protons “drop” from atomic space-time sheet with \( k = 137 \) to a larger space-time sheet to \( k_1 > 137 \) space-time sheet and the difference of zero point kinetic energies is liberated as radiation [K1]. The proposal was that the zero point kinetic energies give rise to a hierarchy of metabolic energy quanta.

   Second possibility is phase transition in which the size of the \( k = 137 \) space-time sheet increases to \( k_1 > 137 \) and liberates the difference of zero point kinetic energy. For the third option energy preserving phase transition increasing \( \hbar_{eff}/\hbar = n \) by a factor \( (k_1 - k)/2 \) followed by a phase transition reducing the value of \( h_{eff} \) back to the initial one but without change of the size of the space-time sheet would liberate the difference of zero point kinetic energies.

2. Could dark nuclear transitions explain UIBs? For \( k = 149 \) as the p-adic length scale of DNA letters would give nuclear energy scale \( E = .5 \) eV equal to the metabolic energy quantum by scaling 1 MeV for the ordinary nuclei by factor \( 2^{149-107}/2 = 2^{21} \) (here the original version of text contained error: this claim was made for \( k = 141 \)). This energy has correct order of magnitude but is too high an energy for UIBs but there are of course also smaller energies possible for the nuclear excitations possibly explaining the UIBs.

3. What about hydrogen bonds? The strength of hydrogen bond - essentially the bond energy - is in the range \( .4-.5 \) eV -, which as such does not correspond to the average UIB energy, which come approximately as three lowest powers of two. The range of bond energies is \( .1 \) eV is smaller than the smallest UIB energy \( .11 \) eV.

   UIBs can be associated with hydrogen bonds if there are states of bond with higher bond energy. They could correspond to higher values of \( n = h_{eff}/h \) for the de-localized dark proton associated with the bond (analogous to de-localized valence electron). For instance, if the energy of the bond corresponds to the cyclotron energy of proton in a magnetic field associated with the bond, it is proportional to \( n \).

   The photon energies come approximately as powers of 2. If the favored values of \( n \) are in bands around \( n = 2^k \) favored by the p-adic length scale hypothesis, one has hopes of understanding the band structure in terms of transitions reducing the value of \( k \).

Membrane potential (see [http://tinyurl.com/chylve9](http://tinyurl.com/chylve9)) plays a key role in metabolism and one can wonder whether UIBs might relate to the potential energies defining energies \( E_J = ZeV \) of Josephson photons associated with membrane if it acts like Josephson junction like structures associated with the prebiotic lifeforms.

1. Membrane potential energy varies in the range \( (.04, .08) \) eV (cell interior is negatively charged). Excitable cells (able to generate action potentials) include neurons, muscle cells, endocrine cells, and some plant cells. The average value for them is around \( .06 \) eV and further depolarization makes these cell more excitable. This suggests that the instability is caused by thermal radiation with nearly the same energy. The threshold for the generation of the action potential \( E_{act} \) is in the range \( (.050, .055) \) eV. Interestingly, during ageing neurons become more hyperpolarized and therefore less excitable. In photoreceptors the resting potential energy can be as low as \( .03 \) eV making them very sensitive to light.

2. In TGD inspired quantum biology azonal membrane can be seen as a generalized Josephson junction [K9, K10, K12] decomposing nanoscopically to Josephson junctions defined by cell membrane proteins. The protein as junction would correspond to a magnetic flux tube along which various charged particles with \( h_{eff} = n \times h \) flow possibly as supra currents. As a special case cell membrane acts like an ordinary Josephson junction. In this case the increment of the electrostatic energy of the Cooper pair over membrane given by \( E_J = 2eV \) defines the energy of the smallest quantum of Josephson radiation.
3.2 Unidentified Infrared Bands as a test for the proposal

The intensity of thermal radiation at temperature \( T \) as function of photon energy \( E \) has a peak at \( E \approx 3T \), which for room temperature about \( T = .03 \) eV gives \( E_{\text{max}} = .09 \) eV. The energy ZeV of Cooper pair should be larger than \( E_{\text{max}} \). For critical action potential one has \( E_{\text{act}} = 0.1 \) eV, which is slightly above \( E_{\text{max}} = .09 \) eV so that the action potential has minimal value and thus minimizes metabolic energy costs and implies quantum criticality with temperature as a critical parameter.

Note however that for energies below \( E_{\text{max}} \) the intensity of thermal radiation decreases so that also these energies might serve as Josephson energies: this and the fact that incoming photons have intensity higher than thermal background at this energy could explain why some photoreceptors can have \( eV = .03 \) eV.

3. Could also Josephson radiation relate to UIBs? The Josephson energy of Cooper pair for the membrane potential is around \( E_f = 0.1 \) eV, which corresponds to the lowest UIB band, which could thus correspond to action potential .05 eV of excitable membrane. The higher bands would correspond roughly to two octaves suggesting that the action potentials in these case are roughly .1 eV and .2 eV. Quantum criticality would suggest that temperatures scale like the energies of the bands slightly higher than \( E_{\text{max}} \approx 3T \).

Metabolic energy transfer between magnetic body and biological body (defined in very general sense for any system) is possible if the spectra of transition energies share common transition energies. Therefore the spectrum of transition energies assignable to hydrogen bonds could have many transition energies common with that assignable to dark nuclear transitions and second and third explanation could be consistent with each other.

3.2.2 Model for hydrogen bond

The explanations of UIBs in terms of hydrogen bonds encourages to consider a concrete model for the hydrogen bond as flux tube. This suggests a connection with metabolism at cellular level involving transfer of protons through cell membrane against potential gradient assumed to take place as dark protons carrying the metabolic energy and providing it to ADP-ATP process after their return.

1. The simplest model for the proton inside flux tube is as particle in 1-D flux tube with magnetic field. Unless the magnetic field strength and/or \( n \) is very large, the kinetic energy in the direction of flux tube dominates and phase transition would change the scale of kinetic energy proportional to \( n^2 \) for fixed flux tube length. For \( n = 2^k \) this would give too strong dependence of photon energies on \( k \).

2. On the other hand, if the flux tubes are flux loops of the magnetic body of molecule their lengths naturally scale as \( n \) and the longitudinal kinetic energy is not affected in the transition. The cyclotron energy proportional to \( n \) would change and for \( n \approx 2^k \) one obtains qualitatively correct behavior.

For proton in magnetic field of \( B_{\text{end}} = .2 \) Gauss the cyclotron frequency is 300 Hz and corresponds to \( E_c(B_{\text{end}}) = 1.2 \times 10^{-12} \) eV. The identification of \( E_c(B) = .5 \) eVs would give \( E_c(B) = n(B/B_{\text{end}}) \times E_c(B_{\text{end}}) = E_c(B) = .5 \) eV. An estimate for \( B \) for the flux tube of hydrogen bond comes from flux quantization: \( eBS = 1 \) holds true for unit quantum of flux and for flux tube radius of one Angstrom this would give \( B/B_{\text{end}} \approx 5 \times 10^8 \). This gives the estimate \( n \approx 10^8 \approx 2^{27} \). The rather large value conforms with the general vision for the values of \( n \) for dark protons whereas dark electrons of valence bonds would have much smaller values. The emergence of dark protons could be seen as the transition from chemistry already involving \( n \) as characterizer of valence bonds [L9] to bio-chemistry.

3. The identification of the metabolic energy quantum in terms of cyclotron energy could apply also in the case of cellular metabolism. The model for the generation of ATP from ADP assumes that protons are pumped by the energy coming from nutrient molecules against the membrane potential.

The membrane potential correspond to energy of .05 eV but metabolic energy quantum is 10 times larger. This looks like an inconsistency, which in thermodynamical approach is resolved.
by introducing of chemical potentials. In genuine quantum approach the introduction of thermodynamics quantities is not allowed.

The general vision about metabolic energy as a tool to increase $h_{eff}/h = n$ defining kind of molecular IQ suggests that the transformation to dark proton at magnetic flux tube along which proton can travel through the membrane is responsible for the most of the energy needed for pumping. After the dark proton has returned through cell membrane it transforms to ordinary proton and liberates the metabolic energy and makes possible ADP-APT transformation.

The above model assumes that the lengths of hydrogen bonds as flux loops scale like $n$. This makes possible the reconnection of flux loops coming from opposite sides of the membrane to pair of flux tubes along which dark protons can flow. Similar picture applies also to other biologically important ions.

The general view about superconductivity in TGD Universe suggests that reconnection can give rise to a Cooper pairs of protons with members at separate flux tubes. Also Cooper pairs of electrons and biologically important ions could form by the same mechanism.

3.3 PAH world hypothesis from TGD point of view

The so called PAH world hypothesis (see [http://tinyurl.com/ycxm9zes](http://tinyurl.com/ycxm9zes)) has been proposed as a prebiotic era preceding RNA world. As a matter of fact, PAH world hypothesis inspired more a detailed development of TGD based model for dark nuclei.

Let us first list some properties of poly-aromatic hydrocarbons (PAHs) (see [http://tinyurl.com/atx4t9a](http://tinyurl.com/atx4t9a)).

1. PAHS consist of aromatic rings glued together along sides. By definition aromatic rings have delocalized electrons. In benzene, which is the classical and simplest example of PAH, the electronic state is quantum superposition of states in which bonds and double bonds alternate along the ring but are shifted by 60 degrees with respect to each other. Naphtalene has two aromatic rings and anthracene and pnenanthtrene have 3 rings.

2. PAHs are very stable non-charged non-polar molecules and are very common in Earth. They are found in coal and tar deposits and produced in an incomplete combustion of organic matter. PAHs are poisonous. For instance, tobacco smoke contains PAHs with carcinogenic effects. The stability of PAHs motivates the belief that a large fraction of carbon in the interstellar space consists of PAHs.

3. Benzene in difficult to detect in in the interstellar space since the rotational symmetry does not allow to detect rotational transitions. Recently however nitrobenzene was detected so that benzene and more complex PAHs presumably exist in interstellar space (see [http://tinyurl.com/yap9ksrg](http://tinyurl.com/yap9ksrg)).

Benzene and more complex PAHs can give rise to more complex aromatic by hydrogenation, oxidation, carboxylation, and nitrogenation and led also to the basic building bricks of DNA and amino-acids and PAHs are proposed to have played important role in prebiotic life.

1. PAH world hypothesis states that the polymer like sequences of PAHs serve as scaffolding for the formation of RNA like polymers (see [http://tinyurl.com/ycxm9zes](http://tinyurl.com/ycxm9zes)). The key motivation is that the distances between PAHs are same as between RNA and DNA bases: 3.4 nm. The proposal is that during PAH era RNA nucleosides A, U, C, G were attached to PAH scaffolding.

2. Second hypothesis is that formaldehyde molecules [(H2C)=O] formed valence bonds with RNA bases and with each other giving rise to sequences analogous to the phosphate-ribose backbone of RNA. The sequence of disjoint CO=O was replaced with the sequence ..(C-R)-O-(C-R)-O-.. with R denoting the RNA nucleoside. After this hydrogen bonds were split and the predecessor of RNA was detached from the PAH scaffolding. Later the pre-RNA strands were folded to form double pre-RNA strands similar to ribozymes. The problem is to understand how the formaldehyde backbone was replaces with more stable phosphate-ribose backbone.
In TGD framework dark nuclei would serve as scaffolding, which however does not detach from the corresponding biomolecules. The distances between dark variants of biomolecules would explain why the two distances are the same. Very many molecules, including PAHs, can attach around dark RNA/DNA and the periodic structure would reflect the properties of dark nuclei. This could explain UIBs as emission bands of both dark nuclei and hydrogen bonds essential for the pairing and the transfer of metabolic energy between ordinary and dark biomolecules. Also in DNA double strand hydrogen bonds could serve similar function. If thermal radiation excites higher energy states of nuclei, the emission of UIBs depends on temperature. Perhaps this could be tested.

UIBs could therefore serve as a direct signature of dark nuclear physics. If dark nuclei are not associated with PAHs in vitro or in an environment not containing water, UIBs would be absent.

**3.4 Did RNA replicate in codon-wise manner during RNA era?**

There was an interesting popular article in Spacedaily with title “Scientists crack how primordial life on Earth might have replicated itself” (see [http://tinyurl.com/y92ng5vd](http://tinyurl.com/y92ng5vd)). The research paper [I7] is titled “Ribozyme-catalysed RNA synthesis using triplet building blocks” and published in eLife (see [http://tinyurl.com/ya5qyjfn](http://tinyurl.com/ya5qyjfn)).

It is possible to replicate unfolded RNA strands in Lab by using enzymes known as ribozymes, which are RNA counterparts of enzymes, which are amino-adic sequences. In the presence of folding the replication is however impossible. Since ribozymes are in general folded, they cannot thus catalyze their own replication in this manner. The researchers however discovered that the replication using RNA triplets - genetic codons - as basic unit can be carried out in laboratory even for the folded RNA strands and with rather low error rate. Also the ribozyme involved can thus replicate in codon-wise manner. For units longer than 3 nucleotides the replication becomes prone to errors.

These findings are highly interesting in TGD framework. In TGD the chemical realization of genetic code is not fundamental. Rather, dark matter level would provide the fundamental realizations of analogs of DNA, RNA, tRNA, and amino-acids as dark proton sequences giving rise to dark nuclei at magnetic flux tubes [L12] (see [http://tinyurl.com/yalny39x](http://tinyurl.com/yalny39x)). Also ordinary nuclei correspond in TGD Universe to sequences of protons and neutrons forming string like entities assignable to magnetic flux tubes.

The basic unit representing DNA, RNA and tRNA codon and amino-acid would consist of 3 entangled dark protons. The essential aspect is that by entanglement the dark codons do not decompose to products of letters. This is like words of some languages, which do not allow decomposition to letters. This representation is holistic. As we learn to read and write, we learn the more analytic western view about words as letter sequences. Could the same hold true in evolution so that RNA triplets would have come first as entities pairing with dark RNA codons from from dark proton triplets as a whole? Later DNA codons would have emerged and paired with dark DNA codons. Now the coupling would have been letter by letter in DNA replication and transcription to mRNA.

It is intriguing that tRNA consists of RNA triplets combined from amino-acids and analogs of mRNA triplets! The translation of mRNA to amino-acids having no 3-letter decomposition alone forces the holistic view but one can ask whether something deeper is involved. This might be the case. I have been wondering whether during RNA era RNA replicated using a prebiotic form of translational machinery, which replicated mRNA rather than translated RNA to protein formed from amino-acids (AAs) with AA serving as a catalyst.

1. During RNA era amino-acids associated with pre-tRNA molecules would served as catalysts for replication of RNA codons. The linguistic mode would have been “holistic” during RNA era in accordance with the findings of the above experiments. RNA codon would have been the basic unit.

2. This would have led to a smaller number of RNAs since RNA and RNA like molecules in tRNA are not in 1-1 correspondence. A more realistic option could have been replication of subset of RNA molecules appearing in tRNA in this manner.
3. Then a great evolutionary leap leading from RNA era to DNA era would have occurred. AA catalyzed replication of RNA would have transformed to a translation of RNA to proteins and the roles of RNA and AA in tRNA would have changed. [Perhaps the increase of \( h_{\text{eff}} \) in some relevant structure as quantum criticality was reached led to the revolution]

4. At this step also (subset of) DNA and its transcription to (a subset of) mRNA corresponding to tRNA had to emerge to produce mRNA in transcription. In the recent biology DNA replicates and is transcribed nucleotide by nucleotide rather than using codon as a unit so that helicases and DNA and RNA polymerases catalyzing replication and transcription should have emerged at this step. The ability of DNA to unwind with the help of helicase enzyme helping DNA to unwind is essential for the transcription and translation of DNA. Therefore helicase must have emerged together with the “analytic linguistic mode” as an analog of written language (DNA) decomposing codons to triplets of letters. This would been a crucial step in evolution comparable to the emergence of written language based on letters. Also the counterpart of RNA polymerase and separate RNA nucleotides for transcription should have emerged if not already present.

An alternative option would involve “tDNA” as the analog of tRNA and the emergence of helicase and polymerases later as the transition from holistic to analytic mode took place.

The minimal picture would be emergence of a subset of DNA codons corresponding to RNAs associated with pre-tRNA and the emergence of the analogs of helicase and DNA and RNA polymerases as the roles of amino-acid and RNA codon in tRNA were changed.

5. How DNA could have emerged from RNA? The chemical change would have been essentially the replacement of ribose with de-oxiribose to get DNA from RNA and U \( \rightarrow \) T. Single O-H in ribose was replaced with H. O forms hydrogen bonds with water and this had to change the hydrogen bonding characteristics of RNA.

If the change of \( h_{\text{eff}} = n \times h_0 \) was involved, could it have led to stabilization of DNA? Did cell membrane emerge and allow to achieve this? I have proposed \([L12]\) (see http://tinyurl.com/yalny39x) that the emergence of cell membrane meant the emergence of new representation of dark genetic code based on dark nuclei with larger value of \( h_{\text{eff}} \).

**Remark:** One has \( h = 6 \times h_0 \) in the most plausible scenario \([L5, L11]\) (see http://tinyurl.com/goruuzm and http://tinyurl.com/y9jxyjns).

The communication between dark and ordinary variants of biomolecules involves resonance mechanism and would also involve genetic code represented as 3-chords, music of light, and it is interesting to see whether this model provides additional insights.

1. The proposal is that 3-chords assignable to nucleotides as music of light with allowed 64 chords defining what I have called bio-harmony is essential for the resonance \([L13, L16, L14]\) (see http://tinyurl.com/ydhexen4g, http://tinyurl.com/yd5t82gd and http://tinyurl.com/y9jxyjns). The 3 frequencies must be identical in the resonance: this is like turning 3 knobs in radio. This 3-fold resonance would correspond to the analytic mode. The second mode could be holistic in the sense that it would involve only the sum only the sum of the 3 frequencies modulo octave equivalence assigning a melody to a sequence of 3-chords.

2. The proposal is that amino-acids having no triplet decomposition are holistic and couple to the sum of 3 frequencies assignable to tRNA and mRNA in this manner. Also the RNAs in tRNA could couple to mRNA in this manner. One could perhaps say that tRNA, mRNA and amino-acids codons sing whereas DNA provides the accompaniment proceeding as 3-chords. The couplings of DNA nucleotides to RNA nucleotides would rely on the frequencies assignable to nucleotides.

3. If the sum of any 3 frequencies associated with mRNA codons is not the same except when the codons code for the same amino-acids, the representation of 3-chords with the sum of the notes is faithful. The frequencies to DNA and RNA nucleotides cannot be however independent of codons since the codons differing only by a permutation of letters would correspond to the same frequency and therefore code for the same amino-acid. Hence the
information about the entire codon would be needed also in transcription and translation
and could be provided either by dark DNA strand associated with DNA strand or by the
interactions between the nucleotides of the DNA codon.

4. The DNA codon itself would know that it is associated with dark codon and the frequencies
 assignable to nucleotides could be determined by the dark DNA codon. It would be enough
that the frequency of the letter depends on its position in the codon so that there would be 3
frequencies for every letter: 12 frequencies altogether.

What puts bells ringing is that this the number of notes in 12-note scale for which the
com/ydhxen4g) based on the fusion of icosahedral (12 vertices and 20 triangular faces) and
tetrahedral geometries by gluing icosahedron and tetrahedron along one face, provides a
model as Hamiltonian cycle and produces genetic code as a by-product. Different Hamiltonian
cycles define different harmonies identified as correlates for molecular moods.

Does each DNA nucleotide respond to 3 different frequencies coding for its position in the
codon and do the 4 nucleotides give rise to the 12 notes of 12-note scale? There are many
choices for the triplets but a good guess is that the intervals between the notes of triplet
are same and that fourth note added to the triplet would be the first one to realize octave
equivalence. This gives uniquely \( CEG^\sharp, C^\sharp FA, DF^\sharp B^\flat, \) and \( DG^\sharp B \) as the triplets assignable
to the nucleotides. The emergence of 12-note scale in this manner would be a new element
in the model of bio-harmony.

There are \( 4! = 24 \) options for the correspondence between \( \{A, T, C, G\} \) as the first letter and
\( \{C, C^\sharp, D, D^\sharp\} \). One can reduce this number by a simple argument.

(a) Letters and their conjugates form pyrimidine-purine pairs \( T, A \) and \( C, G \). The square
of conjugation is identity transformation. The replacement of note with note defining
at distance of half-octave satisfies this condition (half-octave - tritonus - was a cursed
interval in ancient music and the sound of ambulance realizes it). Conjugation could
correspond to a transformation of 3-chords defined as

\[
CEG^\sharp \leftrightarrow DF^\sharp B^\flat, \quad C^\sharp FA \leftrightarrow D^\sharp GB.
\]

(b) One could have

\[
\{T, C\} \leftrightarrow \{CEG^\sharp, C^\sharp FA\}, \quad \{A, G\} \leftrightarrow \{DF^\sharp B^\flat, D^\sharp GB\},
\]

or

\[
\{T, C\} \leftrightarrow \{DF^\sharp B^\flat, D^\sharp GB\}, \quad \{A, G\} \leftrightarrow \{CEG^\sharp, C^\sharp FA\}.
\]

(c) One can permute \( T \) and \( C \) and \( A \) and \( G \) in these correspondences. This leaves 8
alternative options. Fixing the order of the image of \( (T, C) \) to say \( (C, C^\sharp) \) fixes the
order of the image of \( (A, G) \) to \( (D, D^\sharp) \) by the half-octave conjugation. This leaves 4
choices. Given the bio-harmony and having chosen one of these 4 options one could
therefore check what given DNA sequence sounds as a sequence of 3-chords [L1].

That the position the frequency associated with the nucleotide depends on its position in the
codon would also reflect the biochemistry of the codon and this kind of dependence would be
natural. In particular, different frequencies associated with the first and third codon would
reflect the parity breaking defining orientation for DNA.

4 Improved reckless speculation about higher level variants of dark genetic code

In an earlier article I represented what I called reckless speculations about higher level variants of
genetic code (see [L12] for the updated version of the original article). The speculations turned out
to be not only reckless but to contain besides an unrealistic working hypothesis for p-adic length
scale of dark DNA also a numerical error in the estimate of dark nuclear excitation energy scale leading to a wrong track.

The wrong working hypothesis was the assumption that ordinary DNA, RNA, etc correspond to same p-adic length scale as their dark variants. Simple argument shows that the dark scales must result via radial scaling of the typically linear structures such as DNA, RNA, etc and also 2-D structures such as membranes and microtubules giving rise to 2-D lattice like realizations of genetic code generalizing the ordinary 1-D realizations.

Also new improved picture conforms with the vision that dark realizations of genetic code at various p-adic length scales serve as controllers of the ordinary biochemistry, which is kind of shadow dynamics. Replication, certainly one of the most mysterious feats of living matter, would reduce to the replication at the level of dark DNA in various p-adic length scales involved. This would be a huge simplification.

A hierarchy of dark nuclear physics with hierarchy of \( n = \frac{h_{\text{eff}}}{h_0} = n \) coming as certain powers of two so that the corresponding length scales correspond to p-adic length scales is an attractive idea. I have speculated with this idea already earlier. A hierarchy of dark nuclear physics with hierarchy of \( n = \frac{h_{\text{eff}}}{h} = n \) coming as certain powers of two so that the corresponding length scales correspond to p-adic length scales is an attractive idea. I have speculated with this idea already earlier \([K7]\).

### 4.1 Ideas

Consider first the general ideas.

1. The assumption of prime values for \( k \) in \( L(k) \) would pose extremely tight constraints on the allowed p-adic length scales and values of \( h_{\text{eff}}/h_0 \). One would have \( k \in \{127, 131, 137, 139, 149\} \) and \( k \in \{151, 157, 163, 167\} \) and \( k \in \{173, \ldots\} \) at least at the level of dark matter. So predictive an idea deserves to be killed, if not anything else.

   A further motivation for these speculations is that the Gaussian Mersenne primes \( M_{G,k}= (1+i)^k - 1 \) for \( k \in \{151, 157, 163, 167\} \) define p-adic length scale \( L(k) \propto 2^{k/2} \) between 10 nm assignable to the neuronal membrane and 2.5 \( \mu \text{m} \) assignable to cell nucleus: so many Gaussian Mersenne in so short length scale range is a number theoretical miracle.

2. Cell membrane consisting of two lipid layers (see [http://tinyurl.com/h9a2hso](http://tinyurl.com/h9a2hso)) is a binary structure as also DNA double strand. DNAs replicate as would do also RNAs during RNA era. Also cells and therefore also cell membranes replicate so that the analogy might make sense. Since processes like translation and transcription do not occur, cell membrane might serve as 2-D as analog of RNA: the counterpart of RNA era might prevail at these levels. Neuronal membrane might correspond to 2-D analog of DNA.

   So: could various 2-D structures such as nuclear membrane, cell membrane, neuronal membrane, and microtubuli correspond to a new level in the hierarchy of dark codes for which genes and their dark variants would be 2-D rather than 1-D structures? One would have 2-D lattices of codons. Could there be entire hierarchy of them assignable to certain p-adic length scales? As 2-D realizations could be paired with their dark variants so that one could speak of dark variants of various membrane like structures. This applies also to microtubuli.

   The idea that dark variants of DNA, RNA, and AAs are their radially scaled up variants generalizes also. The processes like replication of cell could be induced by a much simpler replication of 2-D dark DNA. This kind of pairing hierarchy could be behind miraculous looking replication of entire organisms. p-Adic fractality and hierarchy of dark DNAs could lurk behind the curtains.

3. The structures of ordinary bio-matter and also their dark variants assumed to control them are characterized by p-adic length scales. How these p-adic length scales could relate? The natural idea inspired by scaling invariance is that the dark variants of 1-D linear structure and 2-D structures formed from ordinary bio-matter are obtained by radial scaling consistent with p-adic length scale hypothesis, and guaranteeing that the distances between building bricks are scaled to the size scales of dark variants of DNA and other basic molecules. This rule makes sense also for the 2-D structures. For instance, it would scale up the p-adic length

4.2 Could cell membrane and neuronal membrane realize genetic codons as 2-D structures?

In the sequel I discuss in more quantitative level the idea that cell membrane and neuronal membrane realize analogs of genes as 2-D structures.

4.2.1 The p-adic length scales associated with the dark variants of 2-D structures?

Consider next the p-adic length scales associated with the structures considered.

1. The thickness of ordinary cell membrane corresponds roughly to \( L_e(149) = 5 \text{ nm} \) whereas the coiling associated with the cell membrane corresponds to \( L_e(151) \). Also neurons correspond to \( L_e(151) \). Could \( k = 149 \) resp. \( k = 151 \) define levels of ordinary cell resp. neuron in the hierarchy of dark nuclear physics?

2. Cell membrane consists of lipid bilayer. The lipid layer has three parts (see http://tinyurl.com/h9a2hsq).
4.2 Could cell membrane and neuronal membrane realize genetic codons as 2-D structures?

- The totally hydrated layer nearest to water is hydrophilic head group, which in the case of phospholipids contains negatively charged phosphate. This phosphate layer has thickness \(0.7\) or \(1.0\) nm.
- Below it is a partially hydrated layer of thickness \(0.3\) nm, which corresponds to \(L(141)\): this of course puts bells ringing!
- Hydrophobic lipid tail layer below it is dehydrated. The thickness of single lipid layer is \(1.25-1.75\) nm and would correspond to the \(p\)-adic length scale \(L_e(145) = 1.2\) nm. \(k = 145\) is not prime.

3. The phosphate layer analogous to phosphate-ribose backbone and the thickness \(L(141)\) of partially hydrated layer suggests that it corresponds to EZ created in Pollack effect so that there would be parallel dark RNA sequence along axon (possibly helical as for microtubules). In the case of cell membrane would have lattice like system formed from dark protons, and maybe even dark neutrons (as an analog for the neutron halo in some nuclei).

4. If the recent biology is the analog of RNA era for \(k = 149\) codes, their manifestations could be seen as analogs of RNAs and the number of different lipids associated with the cell membrane could give some idea about their number. Cell membrane could perhaps be seen as a 2-D analog of RNA polymer. Cell division implying membrane replication would be induced by dark RNA replication. Even the analogs of tRNA and AAs but not proteins might be present if one takes the analogy very seriously. Could one identify pairs of lipids and some molecules analogous to proteins appearing in cell division?

What kind of general conditions can one pose on the dark variants of DNA, RNA, and AAs?

1. Dark variant of 2-D variants of DNA, RNA, or AAs realizing the hierarchy of dark codes should control their analogues or possibly some other molecules coded by them. The coupling would be by resonance. This suggest the hierarchy of codes uses as building bricks simpler structures by starting from 1-D structures and building from them more complex structures. Hence the natural hypothesis is that the 2-D variants of proteins consisting of a 2-D lattice like structure formed from proteins is in question.

2. The geometric aspect of membrane dynamics would be determined by basic dynamics of TGD determined by action, which is a generalization of charged point-like particle coupling to Maxwell field by replacing the particle orbit with 4-D surfaces. This allows as special case minimal surfaces such as deformations of cosmic strings giving magnetic flux tubes. Cell membranes should correspond to extremals for which coupling to Kähler force is non-trivial as it indeed is by membrane potential. This because static closed surfaces, in particular spherical layers, are not possible as minimal surfaces. Remarkably, these extremals are not analogs of external particles (geodesic lines) but correspond to interaction regions. This conforms with the fact that cell membrane is a self-organization pattern requiring a continual feed of metabolic energy.

The 2-D dark variants of DNA, RNA, and AAs would be involved mostly with the control the electro-chemistry of membrane like structures. Of course their geometrodynamics would induce also morphogenesis of ordinary bio-matter.

Also enzymes and ribozymes would have dark variants controlling their behavior. Folded protein represents an interesting example about possibly 3-dimensional graph like structure in which the protein forms an analog of Hamilton’s cycle going through all points of the graph defined as a lattice with nearest neighbors connected by edges without self-intersections. This hypothesis is rather powerful since for Hamiltonian cycle do not necessarily exist for an arbitrary graph.

3. In the case of cell membrane membrane proteins are the natural candidate for the building bricks. They indeed have an active role and serve as both channels and pumps and in the case of the neural membrane this role is especially important. Membrane proteins are identified in TGD framework as generalized Josephson junctions. In the case of cell membranes membrane proteins having length of about 5 nm (5 AAs) or 10 nm (10 AAs) going through the membrane
4.2 Could cell membrane and neuronal membrane realize genetic codons as 2-D structures?

are an excellent candidate for the basic building brick. One could see the basic structure either as 2-D structure built from membrane proteins or 3-D structure build from AAs. Membrane proteins would form kind of generalized protein as a 2-D lattice of proteins and accompanied by their dark variants or of 2-D dark variants of RNA or DNA coding for them and identifiable as radial scalings of these proteins to \( k = 149 \) or \( k = 151 \).

The model for topological quantum computation [K3] suggesting that DNA codons of are connected to lipids of cell membrane could be modified so that that dark DNA, RNA, or AAs associated with membrane proteins are connected to them by flux tubes which can get braided. This would allow the quantum control of the 2-D protein like structure and make it effectively single quantum coherent Josephson junction as suggested in the quantum model for nerve pulse [K12].

The original proposal was that that there might exist an analog of genetic code for lipids. The number of different lipids is however too high to allow any simple correspondence. Lipids have also rather passive role in the dynamics of the cell membrane: their serve as signal pathways, provide metabolic energy, and serve as signal pathways (see [http://tinyurl.com/z7d7osm](http://tinyurl.com/z7d7osm)). The proposal however deserves to be explained.

1. Both sides of the lipid bilayer of cell membrane could pair with 2-D lattice of dark RNA whose size scale would be obtained by radial scaling giving rise to what might be called dark cell membrane. In the case of neuronal membrane the dark lattice would consist of pairs of dark DNA codon and its conjugate. In the case of axon one could have the analog of dark DNA strand extended to a cylinder containing bundles of these strands at its surface. Lipid layers would be 2-D analogs of 1-D DNA strands in this case.

2. Lipids would be analogs of ordinary RNA codons and dark RNA codons would code for them: this would predict 64 different lipids in cell membrane. Single dark RNA would correspond to the size scale of single lipid given by \( L(143) = 2L(141) = .625 \text{ nm} \). The dark nuclear physics would correspond to \( k = 149 \). The number \( N \) of parallel dark RNA strands would be roughly the circumference of the axonal lipid layer divided by the size of single lipid about \( L(143) = .625 \text{ nm} \) given by \( N \approx 2\pi \times L_e(167)/L_e(143) = \pi \times 2^{24} \sim 5 \times 10^6 \).

4.2.2 Thermodynamical constraints

Could this totally irresponsible speculation about p-adic hierarchy of dark nuclear physics and genetic codes survive thermodynamical constraints?

1. The condition that metabolic energy quantum is not below thermal energy at physiological temperatures poses constrains on the model. I have considered several identifications of the metabolic energy quantum. These identifications need not be mutually exclusive.

   - One interpretation is as 1-D zero point kinetic energy of proton at tubular space-time sheet of atomic size with transversal length scale \( L(137) \). This energy is invariant under scalings induce by increase of \( h_{eff} \) since \( h_{eff}^2/L^2 \) is not changed.
   - Second identification of metabolic quanta would be as energies assignable to hydrogen bond and its dark variants.
   - Third identification of the metabolic energy quantum would be as scaled variant of \( E_b(k) = 2(k-107)/2E_b \) of typical dark nuclear binding energy \( E_b \approx 1 \text{ MeV} \). The value would be about .5 eV for \( k = 149 \) and .25 eV for \( k = 151 \).

2. Note that the action potential assignable to \( k = 151 \) neuronal membrane is around .05 eV (the membrane potential for some photoreceptors is .03 eV). In TGD Universe the cell membrane can be seen as Josephson junction decomposing in an improved resolution to membrane proteins acting as Josephson junctions [K9, K10]. Josephson energy of Cooper pair is twice this - that is \( E_J = 0.1 \text{ eV} \) slightly above the maximum \( E_{max} = 3T = .09 \text{ eV} \) of the thermal distribution at physiological temperature.
### 4.3 DNA packing problem and p-adic length scales

DNA manages to pack huge amount of DNA to single cell nucleus. For instance, human DNA as length of about 1 meter. This is achieved by a hierarchical coiling structure involving 3 levels with highest level identifiable as chromatides and the lowest level defined by nucleosomes (see [http://tinyurl.com/yat5cm4y](http://tinyurl.com/yat5cm4y)) wound around histon isomers linked together by straight portions of DNA. One can find a detailed representation of the 4-levelled packing of DNA (see [http://tinyurl.com/ybxv6w4v](http://tinyurl.com/ybxv6w4v)).

There are 4 levels involved. Could they relate to the Gaussian miracle primes $k = 151, 157, 163, 167$? The general proposal is that the products of powers of small primes define the scale hierarchy. There
is evidence that at least the powers of 2 and 3 define p-adic length scales, which would correspond also to dark scales. The simple guess is that the dark scales are identical to the ordinary p-adic scales.

- The diameter of the nucleosome is 11 nm = 1.1L(151), which suggests $k = 151$. Chromosome consists of histone $H_1$ plus nucleosome.
- Nucleosomes coil to form a fiber of diameter $d = 30$ nm. This scale is $3L(151)$.
- At the next level loops of average length $300$ nm = $30L(151) \sim 32L(151)$. This level is only intermediate level in packing.
- These loops compress and fold to $250$ nm = $25L(151) \simeq 3 \times L(157)$, $L(157) = 8L(151)$ wide fiber. Thus third harmonic of also the miracle length scale $L(157)$ would be involved.
- This fiber compresses a tight coil of radius $700$ nm = $70L(151) \simeq 64L(151) = L(163) = 640$ nm giving rise to the chromatid fiber of chromosome. $k = 163$ is the third miracle length scale.
- Chromosomes have width $1400$ nm which corresponds to the scale $L(165)$.

The 3 levels $k = 131, 157, 163$ seem to be realized although not in the simplest manner. Nuclear membrane would correspond to $L(k = 167) = 2.5$ µm. For $n = h_{eff}/h_0$ these levels would correspond to the values $n$ of form $n = 2^3 3^4$.

Consider next nucleosome.

1. DNA wraps of around histone octamers forming a cubical structure consisting of 8 smaller cubes (octamers). There are $2 \times 4$ histones forming two identical layers. The 4 histones $H_{2A}$, $H_{2B}$, $H_3$, $H_4$ of given layer are not identical. There is also histone $H_1$ attached to the entire structure. The incoming DNA double strand enters to the upper end of $H_1$ and leaves from its lower end. $H_1$ is related to the secondary coiling. The wrapping gives rise nucleosomes as helices with two turns and containing about 146 base pairs making 48 codons plus 2 base pairs.

2. According to the standard model of nucleosome double DNA strand wraps around the analog of a spool formed from an octamer consisting of two identical units above each other consisting of 4 different histones. The incoming DNA strand enters the upper 4-histone unit and winds once around it and then does the same for the lower unit before leaving the nucleosome.

One can construct a rough TGD inspired model for this structure (not completely realistic) to get a concrete idea about what is involved.

1. The size scale of the cube like structure is $L(151) = 10$ nm so that single histone corresponds to a cube with side roughly about $L(149) = 5$ nm. One can estimate the total length $L$ of the wire from the equation $z = xR\phi/\pi$, $R \sim L(149)$, $\phi \in [0, 4\pi]$, as $L = \sqrt{1 + \pi^2 4\pi R}$. For $R \sim L(149)$ and $h = L(151)$ this gives $L \sim 66$ nm. There are roughly 146 DNA base pairs and 48 whole codons ($144 = 3 \times 48$ base pairs) and each codon has length about 1 nm. This gives total length of 48 nm. The reduction of radius $R$ by factor $r = 48/66 = 3/4$ to $R = 3L(149)/4$ would give a correct value of $L$.

According to the representation for the hierarchy of packings (see [http://tinyurl.com/ybxv6w4v](http://tinyurl.com/ybxv6w4v)), the diameter of the structure is $d = 1.1L(151)$ rather than small and the height of the structure is smaller in the illustration. This width is however not consistent with the helix structure for any value of the height.

2. If the double DNA strand is accompanied by a dark double strand of radius $L(149)$, the situation is like having a band of width $L(151)$ going around the spool. The dark double strand covers an area, which is $4/3$ times the spool area. The horizontal thickness of the entire dark structure is about $d_D = (7/4) L(151)$. If the radius of DNA double strand is $r = L(151)$ the area covered by the double strand is roughly twice the area of the spool. This suggests that one should identify the p-adic length scale of DNA double strand as its diameter about $L(151)$ rather than its radius.
4.4 Microtubules as quantum critical systems

Also microtubules (see \url{http://tinyurl.com/y8km9vve}) are 2-D structures having a strong resemblance with the lipid layers of cell membrane. Could a higher level representation of genetic code similar to the one proposed for cell membranes make sense for them. Also now one can imagine that the microtubular surface is accompanied by its dark variant realizing 2-D dark genes, dark RNA, or dark proteins with scaled up size. The p-adic prime should correspond to \(k > 151\) so that higher level realization of genetic code would be in question. In the case of axons a possible identification for the dark scale would be as the radius of the axonal membrane.

1. Microtubules are hollow cylinders with outer resp. inner diameter equal to 24 resp. 12 nm (the scales differ by factor 2) so that their thickness is 12 nm is same as the inner radius and would correspond to \(L(151) = 10\) nm. They decompose to 13 parallel helical filaments consisting of 13 tubulin proteins having size scale of order \(L_e(151)\).

2. Tubulins are dimers of \(\alpha\) and \(\beta\) tubulin and the pairs are oriented along the helical filament. One can estimate the size of \(\alpha\) and \(\beta\) tubulin by diving the circumference of 24 nm of the microtubule with the number of filaments, which is 13. This gives for the size scale of tubulin the estimate \(R_{tub} \sim 12\) nm not far from \(L(151)\). This supports the view that p-adic length scale \(L(151)\).

The size scale of the transversal volume associated with lipid is roughly .62 nm that is \(L(143) = 2L(141)\) so that they could correspond to \(k \in \{141, 143\}\), presumably \(k = 141\). Therefore one could see microtubules as scaled up variants of cell membrane with scaling factor \(2^{(151-141)/2} = 2^5 = 32\). Similar scaling would take place for the value of \(n = h_{eff}/\hbar\) giving \(n = 2^{23}\) so that microtubules would represent a higher level of evolution identified as increase of \(n\). Microtubules have indeed emerged after cell membrane.

3. It has been proposed that the \(\alpha\) and \(\beta\) conformations of tubulin give rise to bit or even qubit. If this were the case, single helical filament rotating one full turn would have \(2^{13}\) states and carry 13 bits of information. 13 independent filaments would have \(2^{26} \approx 64 \times 10^6\) states and carry 26 bits of information. One could also think of codon as sequence of 13 filaments with the states of filaments representing \(2^{13}\) letters of the code.
4. Microtubular surface has rather high charge density and is polarized: the almost stationary end has negative local charge density roughly equal to that of DNA whereas the growing end has lower surface charge density. One manner to control the charge of the tubulin dimer is in terms of the charge states of GDP and GTP by ionization of the phosphates. Maximal negative charge for tubulin dimer would be 5 units.

Microtubules are highly dynamical objects with inherent instability and have varying length: one might say that microtubules are quantum critical objects. Quantum criticality and thus instability might relate to the fact that the metabolic energy quantum is very near to thermal energy at room temperature.

The dynamics for the length of microtubule could be induced from the dynamics of EZ involving the flow of protons between microtubule and its magnetic body defined by dark DNA. The gradient in charge density would make possible positive net charge density at the growing end of the microtubule.

In ZEO it looks reasonable to argue that the dynamical patters are coded by a generalization of genetic code just as computer programs code for deterministic dynamical patterns.

5. What could the dark code behind the dynamics be? The $\alpha$- and $\beta$ tubulins of tubulin dimer involve GTP (see \url{http://tinyurl.com/ybtjluaf}) resp. GDP (see \url{http://tinyurl.com/y8uok7kq}). In the case of DNA one has $XMP$, $X = A,T,C,G$. The analogs of dark RNA sequences would contain mere $G$ and the information coded by the tubulin would be determined by the conformation of the tubulin dimer giving 1-bit code. This looks somewhat disappointing.

If the charge states of the phosphates of GDP and GTP can vary and all charge combinations for phosphates are possible, one has $2^3$ charge states for GTP and $2^2$ charge states for GDP. Together with the bit associated with the tubulin conformation this would give $2^6$ states and realize 6 bits of the ordinary genetic code! One would have 2-D realization of the genetic code analogous to that proposed for the lipid layer with the state of tubulin analogous to RNA codon.

This coding together with thermal criticality would make microtubule a dynamical object since the deviation of the tubulin charge from -1 units would spoil charge local charge neutrality of tubulin-dark RNA pair.

I have proposed that flux tubes connecting tubulins to the lipids of the axonal lipid layer could give rise to topological quantum computation $\mathbb{K3}[\mathbb{K3}]$. The size scale of lipid is about $L_e(141)$ and that of tubulin about $L_e(151) = 32L_e(141)$, and the radius of axonal membrane is by two orders of magnitude larger than microtubular surface. Hence this proposal does not look realistic unless one assumes that sub-structures of cell membrane with size scale of order $L_e(167)/L_e(151) = 2^8$ larger than tubulin size represented as space-time sheets with cell nucleus size $L(167)$ have flux tube connections to tubulins.

This kind of map would give rise to a kind of abstraction about what happens at the level of axonal membrane integrating out un-necessary details. This abstraction is natural since microtubules would indeed correspond to a higher level of cognitive hierarchy. Roughly $N = 2^{16}$ lipids would contribute to the information received by single tubulin. Could nerve pulse patterns can induce braiding of the flux tubes in this scale?

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