A $p$-Adic Model
of DNA Sequence and Genetic Code

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Abstract
Using basic properties of $p$-adic numbers, we consider a simple new approach to describe main aspects of DNA sequence and genetic code. Central role in our investigation plays an ultrametric $p$-adic information space which basic elements are nucleotides, codons and genes. We show that a 5-adic model is appropriate for DNA sequence. This 5-adic model, combined with 2-adic distance, is also suitable for genetic code and for a more advanced employment in genomics. We find that genetic code degeneracy is related to the $p$-adic distance between codons.

1 Introduction
It is well known that practically all genetic information in living systems is contained in the desoxyribonucleic acid (DNA) sequence. The DNA macromolecules are made of two polynucleotide chains with a double-helical structure. There are four nucleotides called: adenine (A), guanine (G), cytosine (C) and thymine (T). A and G belong to purine, while C and T to pyrimidine.

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The DNA is packaged into chromosome which is localized in the nucleus of the eukaryotic cells. One of the basic processes within DNA is its replication. The passage of its gene information to protein, called gene expression, performs by the messenger ribonucleic acid (mRNA), which is usually a single polynucleotide chain. In the first part of this process, known as transcription, the nucleotides A, G, C, T from DNA are respectively transcribed into the nucleotides U, C, G, A of mRNA, i.e. T is replaced by U, where U is the uracil. The next step is translation, when mRNA codon information is translated into synthesis of proteins. Codons are ordered sequences of three nucleotides of the A, G, C, U. Protein synthesis in all eukaryotic cells performs in the cytoplasm. The genes by their codons control amino-acid sequences in proteins. It is obvious that there are $4 \times 4 \times 4 = 64$ possible codons. However 61 of them specify the 20 different amino-acids and 3 correspond to stop-codons, which serve as termination signals. As a result most amino-acids are encoded by more than one codon. This degenerate correspondence between codons and amino-acids is known as genetic code, which is mostly universal for all living organisms. In almost all cells genetic information flows from DNA to RNA to protein. For a detail and comprehensive information on molecular biology aspects of DNA, RNA and genetic code one can see Ref. [1].

Processes within macromolecules can be regarded as quantum as classical depending on the scale we are interested in. Modeling of DNA, RNA and genetic code is a challenge as well as a chance for modern mathematical physics. An interesting model based on the quantum algebra $U_q(sl(2) \oplus sl(2))$ in the $q \to 0$ limit was proposed as a symmetry algebra for the genetic code (see [2], [3] and references therein). In a sense this approach mimics quark model of baryons. To describe correspondence between codons and amino-acids, it was constructed an operator which acts on the space of codons and its eigenvalues are related to amino-acids. Besides some successes of this approach, there is a problem with rather many parameters in the operator.

There are some very complex systems (e.g. spin glasses and some macromolecules) whose space of states has an ultrametric structure. The space of conformational states of proteins is such one. Processes on ultrametric spaces usually need new methods for their description. $p$-Adic models with pseudodifferential operators have been successfully applied to interbasin kinetics of proteins [4], [5], [6] (for a brief review see [7]). Ultrametricity is a suitable mathematical concept and a tool for description of systems with hierarchical structure. The first field of science where ultrametricity observed was taxonomy. The first review of ultrametricity in physics and biology was presented
twenty years ago [8]. A very significant and promising part of ultrametrics is p-adics.

$p$-Adic numbers are discovered at the end of the 19th century by German mathematician Kurt Hensel. They have been successfully employed in many parts of mathematics. Since 1987 they have been also used in construction of various physical models, especially in string theory, quantum mechanics, quantum cosmology and dynamical systems (for a review, see [9] and [10]). Some $p$-adic aspects of cognitive, psychological and social phenomena have been also considered [11]. The present status of application of $p$-adic numbers in physics and related branches of sciences is reflected in the proceedings of the 2nd International Conference on $p$-Adic Mathematical Physics [12].

A $p$-adic approach to genetics has not been tempted so far. The main aim of this paper is to make the first step towards $p$-adic genomics. Starting with a formulation of $p$-adic genetic information space, we propose a 5-adic model for DNA (and RNA) sequences and genetic code. A central mathematical tool to analyze classification of codons and structure of genetic code is $p$-adic distance between codons.

\section{$p$-Adic numbers}

Recall that numerical results of measurements in experiments and observations are rational numbers. The set of all rational numbers $\mathbb{Q}$, having usual properties of summation and multiplication, is algebraically a field. In addition to arithmetic operations it is often important to know also a distance between numbers. Distance can be defined by a norm. On $\mathbb{Q}$ there are two kinds of nontrivial norm: usual absolute value $|\cdot|_\infty$ and $p$-adic absolute value $|\cdot|_p$, where $p$ is any prime number. The usual absolute value is well known from elementary courses of mathematics and the corresponding distance between two real numbers $x$ and $y$ is $d_\infty(x, y) = |x - y|_\infty$. This distance also enables that all infinite decimal expansions of real numbers

$$x = \pm 10^n \sum_{k=0}^{-\infty} a_k 10^k, \quad a_k \in \{0, 1, \cdots, 9\}, \quad a_0 \neq 0, \quad n \in \mathbb{Z} \quad (1)$$

are convergent.

By definition, $p$-adic norm of a rational number $0 \neq x = p^\nu \frac{r}{s}$, where $\nu \in \mathbb{Z}$, and integers $r$ and $s$ are not divisible by given prime number $p$, is
This norm is a mapping from \( \mathbb{Q} \) into non-negative real numbers and has the following properties:

(i) \( |x|_p \geq 0 \), \( |x|_p = 0 \) if and only if \( x = 0 \),

(ii) \( |xy|_p = |x|_p |y|_p \),

(iii) \( |x + y|_p \leq \max \{|x|_p, |y|_p\} \leq |x|_p + |y|_p \) for all \( x, y \in \mathbb{Q} \).

Because of the strong triangle inequality \( |x + y|_p \leq \max \{|x|_p, |y|_p\} \) \( p \)-adic absolute value belongs to non-Archimedean (or ultrametric) norm.

\( p \)-Adic distance between two rational numbers \( x \) and \( y \) is

\[
d_p(x, y) = |x - y|_p.
\]  

Since \( p \)-adic absolute value is ultrametric, the \( p \)-adic distance \( d_p(x, y) \) is also ultrametric, i.e. it satisfies

\[
d_p(x, y) \leq \max \{d_p(x, z), d_p(z, y)\} \leq d_p(x, z) + d_p(z, y),
\]

where \( x, y \) and \( z \) are any three points of a \( p \)-adic space.

In direct analogy with the field \( \mathbb{R} \) of real numbers, the field \( \mathbb{Q}_p \) of \( p \)-adic numbers can be introduced by completion of \( \mathbb{Q} \) with respect to the distance \( d_p \). Note tat for each prime \( p \) there is one \( \mathbb{Q}_p \). Any \( x \in \mathbb{Q}_p \) has a unique expansion

\[
x = p^m \sum_{k=0}^{+\infty} a_k p^k, \quad a_k \in \{0, 1, \ldots, p-1\}, \quad a_0 \neq 0,
\]

where \( m \) is an ordinary integer.

In this paper we use only \( p \)-adic integers for which \( m = 0, 1, 2, \ldots \).

For a simple introduction into \( p \)-adic numbers one can see book [13].

\section{3 \( p \)-Adic Genetic Information Space}

We want to present now a mathematical formalism suitable for modeling genetic code and DNA sequence. Let us first introduce an information space \( \mathcal{I} \) as a subset of the set \( \mathbb{Z} \) of usual integer numbers, where to each \( m \in \mathcal{I} \) is attached an information. Different numbers \( a, b \in \mathcal{I} \) contain different information. Let be valid standard arithmetic operations (summation, subtraction and multiplication) on elements of \( \mathcal{I} \).

Since an information can be more or less similar (or dissimilar) to another, there is a sense to introduce a mathematical tool to measure similarity (or
dissimilarity). Such a tool is a distance between the corresponding integers. But now arises a question: What kind of distance we should take between integers to describe closeness on the information space? Recall that there are two kinds of distances for integers: usual real (Archimedean) and $p$-adic (non-Archimedean, ultrametric) distance. We propose, for a class of $\mathcal{I}$, to employ $p$-adic distance (defined in the preceding section), i.e. $d_p(a, b) = |a - b|_p$, $a, b \in \mathbb{Z}$. As a consequence one has a quite natural property: two information are closer, i.e. with smaller distance, if they have more equal first digits in their $p$-adic expansion. One has also that digits which come later in the expansion have smaller importance (for a similar treatment of information see [14]). In the sequel an information space with $p$-adic distance will be called $p$-adic information space. Some experimental properties of genetic code lead us to introduce $p$-adic genetic space $\mathcal{G}_p$ as a special case of $p$-adic $\mathcal{I}$. An element $m \in \mathcal{G}_p$ can be presented in the form

$$m = \pm p^N \sum_{i=0}^{n} m_i p^i, \quad m_i \in \{0, 1, \cdots, p - 1\},$$

(5)

where $N, n$ are nonnegative integers and $m_i$ are digits. For a given $p$ and $N$, information $m$ is characterized by the sequence of digits $m_0, m_1, \cdots, m_n$. In other words, information is coded by ordered sequence of digits $m_0, m_1, \cdots, m_n$. If integers $a, b \in \mathcal{G}_p$ have expansions

$$a = a_0 + a_1 p + a_2 p^2 + \cdots, \quad b = b_0 + b_1 p + b_2 p^2 + \cdots,$$

(6)

then $d_p(a, b) = p^{-k}$ if $a_0 = b_0, \cdots, a_{k-1} = b_{k-1}$ and $a_k \neq b_k$. Accordingly $d_p(a, b) = p^{-k}$ is smaller as $k$ is larger and $a, b$ are closer (i.e. more similar). This $p$-adic closeness will be later exploited in analysis of genetic code degeneration, but now let us turn to the $p$-adic modeling of DNA.

4 $p$-Adic model of the DNA sequence

To have an appropriate $p$-adic genetic space $\mathcal{G}_p$ that can describe DNA sequence and genetic code, one has to choose the corresponding prime number $p$ which will be used as a base for expansion. For the base in expansion of genetic information we choose $p = 5$, because 5 is the smallest prime number which contains four nucleotides (A, T, G, C) in DNA, or (A, U, G, C) in RNA, in the form of four different digits. At the first glance, because
there are four nucleotides, one could start to think that a 4-adic expansion, which has just four digits, might be more appropriate. However, note that 4 is a composite integer and that related expansion is not suitable since the corresponding $| \cdot |_4$ absolute value is not a norm but a pseudonorm and it makes a problem with uniqueness of the distance between two points. To illustrate this problem let us consider, for instance, a distance between numbers 4 and 0. Then we have $d_4(0, 4) = |4|_4 = \frac{1}{4}$, but on the other hand $d_4(0, 4) = |2|_4 |2|_4 = 1$.

Thus for four nucleotides, which appear in the strict complementarity between the two DNA strands, i.e. make two base pairs ($A, T$) and ($C, G$), we choose the corresponding 5-adic integer numbers to construct the corresponding DNA sequence model. Namely, we attach digits (1, 2, 3, 4) to nucleotides ($C, A, T, G$) in the following way:

$$C = 1, \ A = 2, \ T = 3, \ G = 4. \quad (7)$$

Recall that there are $p$ digits in representation of a $p$-adic number. According to this approach, the digit 0 does not play a role in the representation of single helicoidal chain and in the RNA coding. It is worth noting that we also considered some other choices of possible connection between nucleotides and four of the above five digits. However, we find that the choice (7) is the most suitable and attractive.

In this way any of the DNA chains can be presented as a 5-adic number in the form

$$x = 5^N(x_0 + x_1 5 + x_2 5^2 + \cdots + x_n 5^n), \quad x_i \neq 0, \quad N \in \mathbb{N} \cup \{0\}, \quad n \in \mathbb{N}, \quad (8)$$

where $x_i = 1, 2, 3, 4$ and $n$ is an enough large natural number. This chain can be also presented as

$$x = \sum_{j=1}^{\omega} 5^{N_j}(x_0 + x_1 5 + x_2 5^2 + \cdots + x_{n_j} 5^{n_j}), \quad N_1 < N_2 < \cdots N_\omega, \quad (9)$$

where $\omega$ is a number of subsequences, which encode and those which do not encode proteins, in a chain of the DNA. One can introduce 5-adic distance between genes and it will be characterized by $5^{-N_j}$.

For a simple illustrative example ($N = 0, n = 10$), to a chain of nucleotides

$$a = ATGCAAGTG\quad (10)$$
corresponds 5-adic number
\[ a = 2 + 3 \cdot 5 + 4 \cdot 5^2 + 1 \cdot 5^3 + 2 \cdot 5^4 + 2 \cdot 5^5 + 4 \cdot 5^6 + 3 \cdot 5^7 + 4 \cdot 5^8 + 2 \cdot 5^9 , \] (11)
which can be written also using only its digits
\[ a = 2 3 4 1 2 2 4 3 4 2 . \] (12)

According to this approach a DNA sequence can be presented as a sum of two 5-adic integers. Let us denote DNA sequences by Greek letters \( \alpha, \beta, \cdots \) and their chain components by Latin ones \( a, b, \cdots \). Then an \( \alpha = a + b \). In fact \( a \) and \( b \) are firmly correlated because of complementarity, i.e. \( b = \bar{a} \), where \( \bar{a} \) obtains from \( a \) replacing digits \((1, 2, 3, 4)\) by \((4, 3, 2, 1)\), respectively. The corresponding \( \alpha \) related to (10) is
\[ \alpha = a + \bar{a} = 2 3 4 1 2 2 4 3 4 2 + 3 2 1 4 3 3 1 2 1 3 = 0 1 1 1 1 1 1 1 1 1 , \] (13)
where we performed summation of digits from the left to the right, taking \( 1 + 4 = 0 + 1 \cdot 5 \) and \( 2 + 3 = 0 + 1 \cdot 5 \). In this way the sum (13), which corresponds to an example of DNA, is presented in the very simple form: it is quite definite sequence of the digit 1, which is of the same length as DNA and shifted at one place on the right.

One can easily check that integers \( a, \bar{a} \) and \( \alpha \) in (13) form vertices of an equilateral triangle whose all three sides have the same 5-adic length equal to 1.

It is worth mentioning that human genome, which presents all genetic information of the organism, is composed of more than three billion base pairs and contains more than 30,000 genes.

5 \( p \)-Adic genetic code

A living cell is a very complex system composed mainly of protein macromolecules playing various roles. All those proteins are made of only 20 amino-acids, which are the same for all living world on the Earth. Different sequences of amino-acids form different proteins. An intensive study of connection between ordering of nucleotides in the DNA (and RNA) and ordering of amino-acids in proteins led to the discovery of genetic code.

At the end of the 50th and beginning of the 60th of the last century many basic properties of genetic code were obtained. Genetic code is understood
as a dictionary for translation of information from the DNA (through RNA) to production of proteins by amino-acids. The information is contained in codons, which are ordered sequences of three nucleotides. There are three stop codons, and 61 codons are related to 20 amino-acids. There are various multiplicity (one, two, three, four and six) of codons which correspond to amino-acids in proteins, i.e. genetic code is degenerate. This is an well established experimental fact.

However, there is no simple theoretical understanding of genetic coding. In particular, it is not clear why genetic code is just in the known way and not in many other possible ways. What is a principle (or principles) used in fixing mitochondrial and eukaryotic codes? What are properties of codons responsible for their appearance in quadruplets, sextets, doublets, and even in a triplet and a singlet. These are only some of many questions which can be asked about genetic code. Recall that the ribosome performs synthesis of proteins and it knows somehow very firmly which amino-acid corresponds to a given codon. In fact, the ribosome is a molecular machine which performs multiple functions, and one of them should be a computing of codon properties.

Let us consider now possible answers to the above questions on genetic code starting from the 5-adic model. According to our approach, a codon in RNA is an integer number of the following form:

\[ c = c_0 + c_1 5 + c_2 5^2, \quad c_0, c_1, c_2 \in \{1, 2, 3, 4\}, \]

where, without loss of generality, we take \( N = 0 \). In the RNA the nucleotide T is replaced by U and we remain the same digit (T=3) and take U=3. In this way there is no digit 0 used in presentation of codons.

Having the above choice of digits (i.e. C=1, A=2, U=3, G=4) we can now look at the Tables 1 and 2, and observe the corresponding ultrametric (5-adic and 2-adic) reason for formation of quadruplets and doublets. Codons are simultaneously denoted by three digits and capital letters. The corresponding amino-acids are presented in the usual three letters form.
|   |   |   |   |   |
|---|---|---|---|---|
| 111 | CCC Pro | 211 | ACC Thr | 311 | UCC Ser | 411 | GCC Ala |
| 112 | CCA Pro | 212 | ACA Thr | 312 | UCA Ser | 412 | GCA Ala |
| 113 | CCU Pro | 213 | ACU Thr | 313 | UCU Ser | 413 | GCU Ala |
| 114 | CCG Pro | 214 | ACG Thr | 314 | UCG Ser | 414 | GCG Ala |
|   |   |   |   |   |
| 121 | CAC His | 221 | AAC Asn | 321 | UAC Tyr | 421 | GAC Asp |
| 122 | CAA Gln | 222 | UAA Lys | 322 | UAA Ter | 422 | GAA Glu |
| 123 | CAU His | 223 | AAU Asn | 323 | UAU Tyr | 423 | GAU Asp |
| 124 | CAG Gln | 224 | AAG Lys | 324 | UAG Ter | 424 | GAG Glu |
|   |   |   |   |   |
| 131 | CUC Leu | 231 | AUC Ile | 331 | UUC Phe | 431 | GUC Val |
| 132 | CUA Leu | 232 | AUA Met | 332 | UUA Leu | 432 | GUA Val |
| 133 | CUU Leu | 233 | AUU Ile | 333 | UUU Phe | 433 | GUU Val |
| 134 | CUG Leu | 234 | AUG Met | 334 | UUG Leu | 434 | GUG Val |
|   |   |   |   |   |
| 141 | CGC Arg | 241 | AGC Ser | 341 | UGC Cys | 441 | GGC Gly |
| 142 | CGA Arg | 242 | AGA Ter | 342 | UGA Trp | 442 | GGA Gly |
| 143 | CGU Arg | 243 | AGU Ser | 343 | UGU Cys | 443 | GGU Gly |
| 144 | CGG Arg | 244 | AGG Ter | 344 | UGG Trp | 444 | GGG Gly |

Table 1: The vertebral mitochondrial code
Our observations are as follows.

(i) Codons with the same first two digits have the same 5-adic distance equal to $\frac{1}{25}$. This property leads to clustering of 64 codons into their 16 quadruplets. Namely, any two codons $a$ and $b$ whose the first two digits are mutually equal and the third one is different, have 5-adic distance

$$d_5(a, b) = |a_0 + a_1 5 + a_2 5^2 - (a_0 + a_1 5 + b_2 5^2)|_5 = |(a_2 - b_2) 5^2|_5 = 5^{-2}, \quad (15)$$

where $a_0, a_1, a_2, b_2 \in \{1, 2, 3, 4\}$ and $a_2 \neq b_2$. Since $a_0$ and $a_1$ may have four values, there are 16 quadruplets.

(ii) With respect to 2-adic distance, the above clusters may be regarded as composed of two doublets: $a = a_0 a_1 1$ and $b = a_0 a_1 3$ make the first doublet, and $c = a_0 a_1 2$ and $d = a_0 a_1 4$ form the second one. 2-Adic distance between codons within each of these doublets is $\frac{1}{2}$, i.e.

$$d_2(a, b) = |(3 - 1) 5^2|_2 = \frac{1}{2}, \quad d_2(c, d) = |(4 - 2) 5^2|_2 = \frac{1}{2}. \quad (16)$$
(iii) Quadruplets which have at the second position digit 1 do not decay into two doublets. Each of these four quadruplets corresponds to the one of four different amino-acids.

(iv) Quadruplets which have at the second position digit 2 decay into two doublets mentioned in (ii). Each of these eight doublets corresponds to the one of the new eighth different amino-acids.

(v) The doublet structure of quadruplets which have at the second position digit 3 or 4 becomes more complex and depend also on digit at the first place. Quadruplets with digits $13i$, $43i$, $14i$ and $44i$, where $i \in \{1, 2, 3, 4\}$, are stable and have not substructure. However, for other four combinations of the first two digits the situation depends on the kind (mitochondrial or eukaryotic) of coding. The situation is simple for the vertebral mitochondrial code: quadruplets with digits $23i$, $33i$, $24i$ and $44i$, where $i \in \{1, 2, 3, 4\}$, are not stable and decay into doublets. In the case of the eukaryotic (universal) code one has: quadruplet with digits $23i$ decays into one Ile-triplet $(231, 232, 233)$ and one Met-singlet $234$, while the quadruplet $34i$ separates into one doublet and two different singlets.

We would like to emphasize that codons ending on digits 1 and 3, and having 2-adic distance $\frac{1}{2}$, appear always together and determine the same amino-acid.

6 Concluding remarks

In this paper we proposed a new and simple model to investigate information aspects of DNA, RNA and genetic code. To this end, we introduced the corresponding $p$-adic information space and connected it with DNA when $p = 5$.

An essential property of any $p$-adic space is ultrametric behavior of distances between its elements, which radically differs from usual distances on a space of real numbers. It is significant that we attached just 5-adic integer numbers to the sequence of codons and not real integers in base 5.

Classification of any set of objects is an ordering them into groups according to some their relations. Using 5-adic and 2-adic distances between codons we obtained their classification into quadruplets and doublets, respectively. As a result of the above analysis one obtains the following principle of genetic coding: $p$-adically close codons correspond to the same amino-acid.

We plan to continue research on this model and to develop its formalism
as well as to apply it to more concrete cases.

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