

Quantum-Chemical Description of Influence of the R-Groups on Formation of Peptide Bond

Jumber Kereselidze^{1*}, Marine Kvaraia², George Mikuchadze¹

¹I. Javakishvili Tbilisi State University, Tbilisi, Georgia

²Sukhumi State University, Tbilisi, Georgia

Email: *jumber.kereselidze@tsu.ge, mariqvaraia@yahoo.com, gmikuchadze@gmail.com

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Abstract

The description of influence of the R-groups on formation of the peptide bond by quantum-chemical method of density functional theory (DFT) is carried out. The criterion of probability of formation of the peptide bond has been constructed. In particular it is shown that the propensity to formation of the peptide bond is increased as a result of 1) decrease of the CO and NH bond orders (P_{CO} and P_{NH}) and of activation energy (ΔE^\ddagger), 2) increase of OH and CN bond orders (P_{OH} and P_{CN}), 3) exothermic property of this reaction ($\Delta E < 0$).

Keywords

Amino Acids Sequence, Peptide Bond, DFT Calculations

1. Introduction

One of greatest scientific achievements of the 20th century is the creation of recognition of a sequence of amino acids in proteins [1]-[6]. Sanger in the researches used quantum-chemical methods, however on an early stage of development these methods for the description of an electronic structure of biomolecules did not give authentic results and consequently the researches have been at a morphological level. At the end of century W. Kohn and J. Pople have created a modern method of quantum chemistry—Density Functional Theory (DFT) [7]. It is known, that the sequence of amino acids in the polypeptide chain of protein is coded by the sequence of nucleotide residues of DNA. On the other hand R-groups by means of their inductive and spatial effects promote or impede formation of the peptide bonds between of the amino acids pair. This effect, unlike genetic coding, can be used

*Corresponding author.

for the quantitative description of probability of formation of the peptide bonds. One may suppose, that these two phenomena do not contradict, but supplement each other. However the search of some quantitative correlation of given stage seems hopeless.

The detailed mechanism of the peptide bond synthesis that occurs on the large ribosomal subunit remains unknown. The most favorable mechanism is found not to involve any general acid-base catalysis by ribosomal groups but an intrareactant proton shuttling via the P-site adenine O2' oxygen, which follows the attack of the A-site—amino group of the P-site ester. The calculated rate enhancement for this mechanism is $\approx 10^5$, the catalytic effect is found to be entirely of entropic origin, in accordance with recent experimental data, and is associated with the reduction of solvent reorganization energy rather than with substrate alignment or proximity. This mechanism also explains the inability of 2'-deoxyadenine P-site substrates to promote peptidyl transfer. The observed H-bond network suggest an important structural of several universally conserved rRNA residues [8].

2. Methods

Density functional theory (DFT) [7] is a quantum-chemical modeling method used in physics and chemistry to investigate the electronic structure of many-body systems, in atoms, molecules, and the condensed phases. Within this theory, the properties of a many-electron system can be determined by using functionals to be which in this case is the spatially dependent electron density. Hence the name of density functional theory comes from the use of functionals of the electron density. DFT is among the most popular and versatile methods available in condensed-matter physics, computational physics, and computational chemistry. Using the B3LYP level of bases set [9]-[11] in the gas phase the complete geometry optimization. The calculation were performed by the “Priroda” software developed by Laikov and Ustinyuk [12] in the reaction coordinate.

3. Results and Discussed

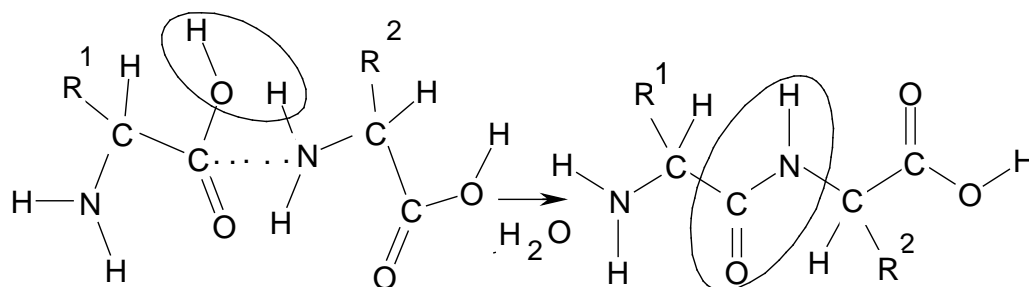
With use of the DFT method computational model, which represent a criteria of probability of formation of the peptide bonds has been created. As known the synthesis of protein catalyzed by ribosome is complex biochemical process. The ribosome mainly provokes excitement of the C-O bond of the oxycarbonyl group with purpose of pairing with amino group of another amino acid. On the other hand it is necessary to consider influence of a structure of amino acids on formation of the peptide bonds, which as against of a catalise role of ribosome varies depending on inductive and spatial properties of R-groups.

On the basis of successes of the computer description of micro-processes, we count that with application of an offered method is possible to receive more important results for an every possible combination of a sequence of amino acids. Without exaggeration one can noted, what by these researches the new era of quantum biochemistry will begin.

The purpose of the planned research is describe an electronic structure (orders of the broken off and formed bonds) and energetic condition (energy of activation and energy of reaction of formation peptide bonds) of amino acids, participating during formation of a natural and man-made sequence.

By means of comparison of the received results we can quantitatively estimate the probability of the formation of polypeptides and influences of the R-groups on this process (Scheme 1).

By object of the research was chosen a sequence of amino acids of insulin of cat: alanine-valine-histidine-alanine



Scheme 1. The formation of a peptide from two amino acid.

[13]. In this row only fourth amino acid are changed in species of others natural amino acids (see Table 1). The decrease of the C-O and N-H (P_{CO} , P_{NH}) bond orders and also values of energy of activation (ΔE^\ddagger), and the increase of the O-H and N-C bond orders (P_{OH} , P_{NC}) and exothermic property ($\Delta E < 0$) of formation of the peptide bonds there is an opportunity of the quantitative description of probability of a sequence of amino acids in proteins.

From the analysis of the tabular data, it is possible to make the conclusion, that the offered criterion, for transition state, mainly satisfies the avha (alanine, valine, histidine, alanine) sequence, what completely consent with the literary data [2]. Besides the high propensity to the peptide bond formation and for a sequence avhser (4) and avhoxylysin (12) is observed. On the other hand on the bases not suitable and high values of the bond orders and energetic characteristics the sequence avhglu (9), avhthr (10) and avhtrp (15) should not be realized.

4. Conclusion

The quantum-chemical model of the quantitative description of influence of the R-groups of amino acids on formation of the peptide bonds is constructed. It is shown, that decrease of the CO and NH bond orders (P_{CO} and P_{NH}) and increase of the OH and CN bond orders (P_{OH} and P_{CN}), and also decrease of energy of activation (ΔE^\ddagger) of the peptide bond formation and the exothermic property ($\Delta E < 0$) of processes the propensity to formation of peptide bonds are promoted.

Table 1. The values of the energy of the activation (ΔE^\ddagger) and reaction (ΔE), and the orders CO, OH, NH, and NC bonds (P_{CO} , P_{OH} , P_{NH} , and P_{NC}).

N	Peptide	ΔE^\ddagger kJ/mol	ΔE kJ/mol	P^1 CO	P^1 OH	P^1 NH	P^1 CN	P^* CO	P^* OH	P^* NH	P^* CN	P^2 CO	P^2 OH	P^2 NH	P^2 CN
1	avhala	10.5	-16.3	0.86	0.21	0.68	0.57	0.68	0.43	0.44	0.77	0.48	0.70	0.19	0.99
2	avhgly	24.5	-72.2	0.85	0.15	0.75	0.52	0.72	0.30	0.57	0.71	0.48	0.70	0.19	1.03
3	avhasn	11.8	-16.8	0.83	0.14	0.76	0.48	0.81	0.18	0.71	0.51	0.44	0.58	0.28	0.89
4	avhser	34.9	-0.02	0.91	0.18	0.72	0.51	0.65	0.50	0.38	0.84	0.50	0.70	0.20	1.00
5	avhcys	79.0	-36.7	0.79	0.28	0.58	0.70	0.67	0.40	0.46	0.83	0.46	0.69	0.19	1.01
6	avhmet	3.2	-60.9	0.76	0.18	0.71	0.53	0.58	0.36	0.51	0.77	0.39	0.70	0.20	1.00
7	avhasp	24.6	-131.2	0.88	0.16	0.72	0.53	0.84	0.19	0.68	0.58	0.45	0.70	0.19	0.93
8	avhgln	91.6	-139.1	0.84	0.13	0.76	0.43	0.65	0.31	0.54	0.60	0.33	0.72	0.20	1.01
9	avhglu	192.9	144.1	0.83	0.15	0.75	0.50	0.68	0.29	0.58	0.67	0.58	0.41	0.47	0.79
10	avhthr	86.9	67.2	0.84	0.14	0.74	0.57	0.56	0.46	0.41	0.86	0.43	0.69	0.23	0.89
11	avhlys	56.5	-78.1	0.83	0.13	0.75	0.50	0.58	0.35	0.51	0.71	0.47	0.69	0.19	0.97
12	avhoxylys	50.5	-7.2	0.91	0.15	0.74	0.54	0.55	0.50	0.35	0.90	0.47	0.68	0.19	1.00
13	avhphe	25.5	-48.1	0.89	0.15	0.74	0.56	0.64	0.38	0.48	0.79	0.46	0.69	0.19	1.00
14	avhtyr	10.9	-7.0	0.79	0.28	0.58	0.60	0.67	0.42	0.44	0.74	0.45	0.69	0.18	0.97
15	avhtrp	279.6	262.2	0.73	0.21	0.66	0.63	0.39	0.54	0.34	0.78	0.34	0.62	0.28	0.84
16	avhhis	1.1	-28.9	0.80	0.27	0.59	0.68	0.74	0.34	0.46	0.75	0.46	0.69	0.20	1.03
17	avhval	2.1	-22.9	0.79	0.28	0.61	0.62	0.68	0.42	0.48	0.76	0.48	0.71	0.20	0.98
18	avhleu	28.4	-60.7	0.86	0.11	0.79	0.42	0.77	0.22	0.64	0.61	0.46	0.69	0.19	0.97
19	avhileu	13.2	-157.6	0.83	0.14	0.75	0.54	0.77	0.17	0.70	0.60	0.50	0.69	0.18	0.97
20	avharg	34.1	-191.6	0.71	0.23	0.64	0.54	0.61	0.30	0.56	0.64	0.31	0.59	0.27	0.90

P^1 —basic state; P^* —transition state, P^2 —final state. The objects of research chose quartets of amino acids: avhx, where a = alanine, v = valine, h = histidine and x—others natural amino acids.

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