

## *Chemical Science International Journal*

*30(10): 46-53, 2021; Article no.CSIJ.78363 ISSN: 2456-706X (Past name: American Chemical Science Journal, Past ISSN: 2249-0205)*

# **Synthesis and Preservation of Organic Molecules with Homochiral Excess by Adsorption on Carbon in Carbonaceous Chondrites**

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## *Author's contribution*

*The sole author designed, analyzed, interpreted and prepared the manuscript.*

### *Article Information*

DOI: 10.9734/CSJI/2021/v30i1030259 *Editor(s):* (1) Prof. Pradip K. Bhowmik, University of Nevada Las Vegas, USA. *Reviewers:* (1) Narges Omrani, Islamic Azad University, Iran. (2) Ninoska m. Paz palacios, Universidad Bolivariana De Venezuela, Venezuela. Complete Peer review History, details of the editor(s), Reviewers and additional Reviewers are available here: https://www.sdiarticle5.com/review-history/78363

> *Received 01 October 2021 Accepted 04 December 2021 Published 10 December 2021*

*Original Research Article*

## **ABSTRACT**

The nature of carbon, initial components, molecules of homochiral abiogenic synthesis and their preservation from decay and racemization for more than 4.5 billion years in carbonaceous chondrites has not been established. In the oxygen-free atmospheres of the nebula and early Earth, hydrogen and hydrogen-containing gases were oxidized with carbon monoxide and carbon dioxide to form carbon and water, as well as the intermediates of these reactions, formaldehyde and methane acid. Together with ammonia, they were the initial components of organic synthesis. According to the Rebinder rule, carbon adsorbs hydrogen well, including in organic molecules. In this connection, experiments with the assumed conditions of the early Earth were carried out by adsorption on carbon to obtain R-(rectus, Latin) ribose from formaldehyde, and S-(sinister) serine from formaldehyde, methane acid and ammonia. For other S-amino acids, a stereo chemical justification of their formation based on S-serine is given. For carbonaceous chondrites, the results of the above experiments were confirmed by the correlation of an increase in homochiral excess with an increase in the amount of hydrogen in aldonic acids and lactic acid with a coefficient of 0.94 and 0.85 in amino acids. The justification of the homochiral process will reduce the costs of searching for life on planets, for scientific research, for the production of medicines, perfumes, food, and so on. Doubts about the extraterrestrial

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origin of homochiral enantiomers in carbonaceous chondrites arise most often due to a lack of understanding of the reasons for their appearance. This work will significantly reduce such skepticism.

*Keywords: Nebula; chondrites; carbon; adsorption; enantiomers; homochirality; preservation.*

## **1. INTRODUCTION**

The problem of preservation of organic substances during the fall of meteorites was resolved after the detection of a thin glassy heatprotective crust, and the reasons for the appearance of carbon, initial components, homochiral molecules and their preservation from racemization and decay for more than 4.5 Ga have not been established. In the absence of oxygen, H2 and gases with H can be oxidized by CO2 and CO to form H2O and carbon (reactions 1-4, 7, Table 1) or intermediates H2CO (methanal) and HCOOH (methanoic acid) (reactions 5, 6, 8, 9, Table 1), which with NH3 were the initial components of abiogenic organic synthesis [1]. In experiments, formaldehyde was a universal starting component in the synthesis of amino acids and sugars. The formation of amino acids racemates has been established with a content from  $10^{-4}$  to  $5x10^{-3}$  mol in a solution with 2.5% H2CO (formaldehyde or methanal) and  $1.0 - 1.5%$  NH4NO3 or NH4C1 at a temperature of  $313 - 318$  K [2]. For the first time, sugars from H2CO were obtained in 1861 by A.M. Butlerov. Ribose from H2CO was synthesized with high yield in the presence of apatite as a catalyst in an Ar atmosphere at pH-7.3 and a temperature of 313 K and [3]. Previously, the possibility of the formation of fatty acids and nitrogenous bases was shown with the disproportionation of H2CO [1]. H2CO is installed in the atmosphere of Mars [4] in the interstellar medium and comets [5] in carbonaceous chondrites [6] and in volcanic gases.

According to the Rebinder rule, carbon adsorbs hydrogen well, and in organic molecules its adsorption properties [7] are close to hydrogen. This made it possible to obtain R-ribose and Sserine (reaction 10, Table 1) in experiments conducted for the first time by adsorption on carbon with stereochemical justification [1, 8]. Other S-amino acids were formed on the basis of S-serine during the disproportionation of H2C(OH)2 (metanediol, H2CO in water) molecules, as shown by the example of the formation of alanine (reaction 11, Table 1) and

phenylalanine (reaction 12, Table 1). Homochirality in other optical isomers (for example, in glucose) could arise during evolution. Adsorption on carbon provided for organic synthesis: the concentration of the initial components, the formation of only R-ribose and S-amino acids, hydrophobic-hydrophilic<br>properties, protection from hydration. properties, protection from hydration, decomposition and racemization [1, 8]. The main purpose of this work is to identify the origin of abiogenic homochiral synthesis in carbonaceous chondrites and its similarities with the appearance on the early Earth.

#### **2. MATERIALS AND METHODS**

### **2.1 The Extraterrestrial Nature of Enantiomers with Homochiral Excess in Carbonaceous Chondrites**

Organic substances are known in the interstellar medium and comets [5], therefore their extraterrestrial nature in carbonaceous chondrites, and hence homochiral molecules, is beyond doubt. Doubts about the nature of enantiomers with homochiral excess in carbonaceous chondrites arise most often due to the lack of clarity of the processes of their appearance, which is justified below. In addition, the extraterrestrial nature of these enantiomers is confirmed by the isotopic ratios of C, N2 and H2  $[5, 9 - 11]$ . The same nature is indicated by the complete absence of high-molecular organic compounds in carbonaceous chondrites<br>(peptides, proteins, organophosphorus (peptides, proteins, organophosphorus compounds, RNA, etc.). The ice of Antarctica in meteorite impact sites and beyond contains organic matter in trillionths, which is most likely due to meteorite dust [12], and the content of various amino acids in carbonaceous chondrites is millions of times higher than in ice. Carbonaceous chondrites contain about 15 sugar acids and about 75 amino acids not found in the biosphere [10], as well as bio-inverse enantiomers. One meteorite may contain racemates, molecules with bio- and reverse excess, which is not typical of the biosphere.

## **2.2 Initial Components and Products of Homochiral Synthesis in Carbonaceous Chondrites**

The conditions in the nebula make it possible to study planetary regularities and the composition of carbonaceous chondrites. Differentiation of gases in the nebula caused the maxima of their relative abundances (mass of atmospheric gas / mass of the planet) with distance from the Sun for Venus (v) - CO2, Earth (e) - H2O (taking into account exospheres) and Jupiter (j) - H2 [13]. Mathematically, it is expressed in close ratios of the molecular masses of gases (M) to the distances to the Sun (R): (MCO2/MH2O)/(Re/Rv)  $=$   $(44/18)/(149.5 \times 10^6)/(108 \times 10^6) = 1.77$ ,  $(MH2O/MH2)/(Rj/Re) = (18/2)/(778 \times 10^6)/(149.5$  $\hat{x}$  10<sup>6</sup>) = 1.73 or (MH2O/MH2)/(MCO2/MH2O) = 3.69 and  $(Ri/Re)/(Re/Re) = 3.77$ .

Carbonaceous chondrites are divided into several types depending on the composition and the formation temperature from  $\sim$  1150 – 1200 K to the temperature of the existence of liquid water [14, 15]. The classifications of carbonaceous chondrites take into account the simultaneous increase in the content of carbon and organic substances [14,15] during the recovery of CO2, which reflects the relationship of their formation (reactions 1-7, Table 1). These chondrites contain carbonates [14,15] which, together with the atmospheres of the terrestrial planets, indicate the presence of CO2 in the nebula. The average contents of carbon and water in the main types of carbon chondrites [14] has an almost direct linear relationship. Such dependence could be realized in the nebula in the reactions of H2 and gases with H with CO and CO2 with their activation by solar radiation [16] and optimal temperatures. Carbon, H2O, or intermediates of H2CO and HCOOH were the products of these reactions (reactions 1-6, Table 1), the yield of which increased at temperatures below 970-940 K. Like reactions could occur in water (reactions  $7 - 9$ , Table 1).

H2CO and HCOOH together with NH3 were the initial components for organic synthesis during adsorption on carbon [1] and were detected in carbonaceous chondrites. Н2СО was found in 12 carbonaceous chondrites studied from 0.1 to 18 mg/g [6,11], and НСООН in chondrite of Lake Tagish from 209 to 378 mg/g [11] and in the composition of monocarboxylic acids in other chondrites [5]. Ammonia was identified in 6 carbonaceous chondrites from 0.5 to 10 mg/g [9] and in the composition of amino acids.

Organophosphorus compounds with H3PO4 have not yet been detected. These initial components were in the nebula at pressures from  $\lg P = -3$  to  $-5$  bar [14] in gaseous state or in solution in carbonaceous chondrites with condensed water.

Carbonaceous chondrites contain pentoses, sugar alcohols, acids, and more than 100 types of amino acids, as well as other organic compounds. Among them, a homochiral excess is often found. For example, amino acids with Sexcess are found in 24 out of 24 studied carbonaceous chondrites, and in one of them Ralanine predominates by 12% [5]. 33 amino acids were found in Lake Tagish chondrite, but homochiral S-excess was studied and revealed in 10 types, and its excess is in alanine (%) 22, in isovaline - 13 and in valine - 97.7 [11]. According to the review [10], out of 101 identified amino acids in carbonaceous chondrites S-excess was found in 23 from 1 to 99% of 28 studied and 5 are represented by racemates, 13 were achiral, homochirality not studied in 60. In the composition of 101 amino acids, 75 types are characteristic only for meteorites. Homochiral Rexcess found in 2 aldonic acids with 3 CH (or CH2) radicals from 33 to 55%, in 4 with 4 radicals from 47 to 100%, and in 6 with 5 radicals of 100% each. S-excess of 3 to 12% is noted in lactic acid. S-excess of 7% and an R-excess of 3% identified in the alanine of one meteorite. Rexcess is present in allo-isotreonine up to 8% and from 12 to 60% in allo-isoleucine. The listed compounds with a biologically inverse homochiral excess contain an asymmetric CH3 radical relative to hydrogen at the central carbon atom  $(H^1,$  Fig 1. a) and two radicals in alloisoleucine.

## **2.3 Initial Components and Conditions of the Origin of Life on Earth**

Planetary regularities show that the Earth's atmosphere in the early Archean did not qualitatively differ from the atmospheres of Venus and Mars and consisted of CO2, N2 and other gases in descending order, as on Venus:  $SO_2$ , CO, Cl<sub>2</sub>, HCl, H<sub>2</sub>S, CH<sub>4</sub>, H<sub>2</sub>, NH<sub>3</sub> [17]. The Martian atmosphere contains H2, CH4, NH3, and H2CO [4]. The volcanic gases of the Earth have a similar composition. Of 108 samples for 6 volcanoes [18], the average content of gases without H2O is (%): CO2 - 58.7, SO2 - 20.1, H2 - 7.7, HCl - 6.7, CO - 5.1, HF - 0.58, H2S - 0.41, NH3 - 0.16, CH4 - 0.016, sometimes OCS, СS2 Н2СО and others are noted. Carbon,

N <sub>2</sub>	<b>Reactions</b>	IgKT at T, K
		298 323 348
	$COg+H2Og=CO2g+H2g$	5.1 4.6 4.1
	$CO2g+2H2g = 2H2Oliq$	14.0 11.6 8.9
3	CO2g+CH4g=2H2Oliq+2Cgr	5.1 3.7 2.5
4	3/2CO2g+2NH3g=3H2Oliq+3/2Cgr+N2g	15.1 12.7 10.7
5	3/2CO2g+2NH3g=H2COg+2H <sub>2</sub> Oliq+1/2Cgr+N2g	7.17.06.9
6	2CO2g+2NH3g=HCOOHfl+2H2Oliq+Cgr+N2g	$2.71.1 - 0.3$
	CO2s+2H2Ss=2H2Oliq+Cgr+2Scr	5.74.63.7
8	HCO3 <sup>-</sup> +2HS <sup>-</sup> +H <sup>+</sup> = H2COs+2H2Oliq+2Scr	7.17.27.3
9	2HCO3 <sup>-</sup> +HS <sup>-</sup> +2H <sup>+</sup> =HCOO <sup>-</sup> +CO2s+2H2Oliq+Scr	8.78.68.5
10	CH2(OH)CH(OH)2a+HCOONH4= <sup>+</sup> NH3CHCH2OHCOOa <sup>□</sup> +2H2Oliq	
11	$-CH2OH+ H2C(OH)2+HCO3 = -CH3+HCOO+H2Oliq+H2CO3$	pH 6.3-7.5
12 <sup>°</sup>	$-CH2OH+12 H2C(OH)2+12HCO3 =$	pH 6.3-7.5
	-CH2C6H5+6HCOO <sup>-</sup> +13H2Olig+12H2CO3	

**Table 1. The most likely reactions the formation organic molecules and carbon**

*Notes: g-gas, s-solution, gr-graphite, liq-liquid, cr-crystal a-adsorbed.\*Approximate values. ▬chemical bond. ▬CH2OH serine radical. (Thermodynamic potentials [19 ‒ 21]*

carbohydrates and amino acids were found in the products of volcanic eruptions with an a biogenic nature according to isotopic data [18]. Volcanoes during the degassing of terrestrial planets determined the composition of the atmospheres in the Archean. The composition of volcanic gases is similar to the composition of comets [5], gases in carbonaceous chondrites and their minerals [14, 15], therefore, the initial components of the abiogenic synthesis of the Earth and the nebula were identical.

Organic synthesis could have arisen on Earth with the appearance of sedimentary rocks and water basins  $3.7 - 3.8$  billion years ago. Typical sedimentary rocks of this period contain carbon and carbonates and are rare [22], apparently due to the small amount of water and pools on the planet. The CO2 pressure in the atmosphere was about  $50 - 60$  kPa, and the temperature was close to  $310 - 320$  K [23, 24]. The height of the Archean troposphere could be about 6.5 km, which is 1.7 times less than its current average height of 11 km (from the comparison of the modern homogeneous atmosphere with the atmosphere from CO2 at pressure of 55 kPa by the formula H=P0/ρ x g, where P0 and ρ are the pressure and density at sea level, g - is the acceleration of gravity). These factors provided sharp daily temperature changes up to  $25 - 35$  K (as in tropical deserts) with change in the solubility of gases and pH (hydrogen index) in small pools.

The totality of the above data allows us to identify the main factors the emergence of life on Earth. During the accretion of planets, the distance of the Earth from the Sun determined the maximum concentration of water on it and, together with the content of greenhouse gases in the atmosphere, the optimal temperature for its existence in liquid form. The appearance of H2CO and HCOOH was facilitated by the mass of the planet, which prevented dispersion of H2 into space, as well as the predominance of CO2 pressure in the atmosphere over the sum of pressures of H2 and gases with H (but with PH2 exceeding the minimum for reactions with CO2, at PCO2 =  $0.55$  bar, PH2 >  $10^{-6.8}$  from reaction 2 of Table 1).

#### **3. RESULTS AND DISCUSSION**

## **3.1 Synthesis and Preservation of Enantiomers with Homochiral Excess by Adsorption on Carbon in Carbonaceous Chondrites**

The age of chondrites varies from 4.5 to 4.575 billion years [14]. The time of racemization of asparagine at 252 K is estimated at 20 million years, and in the shell of mollusks with an age of one million years, the ratio of R to S amino acids was: for asparagine - 0.88, alanine - 0.95, valine - 0.85 [10]. Half of the racemization cycle varies from 500 to 180,000 years for 11 amino acids from the composition of proteins [25]. Long-term preservation of homochirality in enantiomers from decay and racemization could be achieved by fixing them on a solid material and at low temperatures. The most accessible method of fixation is adsorption, and the most effective adsorbent in carbonaceous chondrites is carbon.

An increase in the number of hydrogen atoms in 3 alkanes and 5 alcohols was accompanied by an increase in their adsorption from solutions (experiments with alkanes were performed 4 times on 4 brands of coal, and with alcohols 5 times on 5 brands of coal) [7]. Therefore, in the homochiral synthesis of molecules by adsorption on carbon, there must be a correlation between the number of hydrogen atoms and the homochiral excess.

Significant correlation coefficients of 0.94 (with a significance level of 0.001) were established between the number of hydrogen atoms in aldonic acids and lactic acid and the average values of homochiral excess and 0.85 (with 0.05) for amino acids (Table 2). The correlation coefficients show that the homochiral synthesis of enantiomers was carried out by adsorption on carbon. Correlation is also noted for homochiral excesses and molecular weights of aldonic acids and lactic acid due to their proportionality to the amount of hydrogen, but in its absence there is no correlation in amino acids. Enantiomers with homochiral excess could not appear and persist

in the nebula without carbon protection from radiation and collisions. The absence of complex organic molecules (sugars, amino acids, etc.) in the interstellar medium and comets [5] is a confirmation of this.

The increase in homochiral excesses and the number of hydrogen atoms in the molecules indicates the fixation of hydrogen on carbon, what ensured their preservation for more than 4.5 Ga in carbonaceous chondrites at low temperatures. The vast majority of meteorites (99.8%) originate from the asteroid belt, where the temperature varies depending on the distance of the orbit to the Sun from 165 to 190 K. It was even lower at the early luminosity of the Sun. At these temperatures, racemization of enantiomers could occur in molecules with weak adsorption adhesion after desorption, which depends on the number of adsorbed hydrogen atoms. Desorption could occur due to the penetration of high-energy cosmic rays and internal radioactive decay, or due to strong heating of the frontal part and impact during falls.





*Notes. \*Sample from 24 lactic and aldonic acids, and 53 amino acids. \*\*The accuracy of the definition is not given. \*\*\*Values were excluded if the accuracy of the determination was close to racemates*



**Fig. Structural diagram of the formation of R-alanine during adsorption on carbon. (a) Salanine at the time of formation of H<sup>4</sup> (dash-dotted lines - directions of interaction, dotted line direction of electron transfer at the time of formation hydroxyl O<sup>1</sup>H 1 ), (b) R-alanine after inversion of group C<sup>3</sup>**

## **3.2 Synthesis and Preservation of Enantiomers with Biologically Reverse Homochirality**

The asymmetric position of CH3 radicals in the structure of enantiomers relative to  $H^1$ , as in alanine (Fig.1a), could lead to bio-inversion and fixation of the reverse homochiral excess only on non-mobile molecules, that is, adsorbed on carbon. Otherwise, only equilibrium racemates would be preserved. The effect of asymmetry in the structure of amino acids on their inversion is confirmed by an increase in the homochiral Rexcess from 3-12% with one asymmetric CH3 radical in alanine to 60% with two such radicals in allo-isoleucine. In valine, the S-excess reaches 97.7% [11] with the symmetrical position of two CH3 radicals. The radical appears after all groups in amino acids upon adsorption on carbon  $[1, 8]$ . H<sup>4</sup> was formed instead of serine hydroxyl (Fig. 1b, reaction 11, Table1) with an asymmetric position relative to  $H^1$ . The asymmetric position of  $H^4$  in the carbon adsorption force field and the dipole effect of  $O<sup>1</sup>$ could lead to the rotation of groups  $C^1$  and  $C^2$ counterclockwise (from  $C^2$ ) with the convergence of  $O^1$  with H<sup>1</sup>. due to the delay of H<sup>1</sup> adsorption on carbon. This led to the homolytic break of the  $C^1$ -H<sup>1</sup> bond with the formation of hydroxyl O<sup>1</sup>-H<sup>1</sup> and the  $C^1$ - $C^3$  bond instead of  $C^1$ -H1. At the same time, an excess electron appeared, the transfer of which from  $O^1$  to  $O^2$  determined the occurrence of a  $C^3$ -O<sup>2</sup> bond and a C<sup>1</sup>-H bond instead of  $C^1$ - $C^3$  (Fig. b), probably as during racemization, but with the fixation and preservation of a new structure by adsorption on carbon.

#### **4. CONCLUSIONS**

The processes of abiogenic homochiral synthesis on the early Earth and in carbonaceous chondrites were carried out by adsorption on carbon and were identical, but differed in a small amount of water and the absence of organophosphorus compounds in carbonaceous chondrites. The adsorption nature of abiogenic homochiral synthesis on carbon in carbonaceous chondrites is confirmed by significant correlation coefficients between the amount of hydrogen and homochiral excess, their preservation from decay and racemization for more than 4.5 Ga, the formation of only R-ribose and S-amino acids with a rare bio-inversion confirming this nature.

#### **ACKNOWLEDGMENTS**

The author expresses gratitude for the discussion of the article and useful advice to Doctor of Pharmacological Sciences Budko E.V. and Doctor of Geological and Mineralogical Sciences Yu.T. Usikov. The work was carried out on the initiative and at the expense of the author, without a conflict of interests. Living organisms were not involved for experiments.

#### **COMPETING INTERESTS**

Author has declared that no competing interests exist.

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*Zhmakin; CSIJ, 30(10): 46-53, 2021; Article no.CSIJ.78363*

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> *Peer-review history: The peer review history for this paper can be accessed here: https://www.sdiarticle5.com/review-history/78363*