Facial Isotope Differences of Carbon and Global Photosynthesis in the Frames of Global Carbon Cycle

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Abstract

Facial isotopic differences are usually regarded as the features of the carbon isotope composition of sedimentary organic matter, reflecting the conditions of formation and transformation in the location under study. In the present work it is shown that facial differences reflect the conditions of photosynthesis that took place in this location during the relevant period of geological time. These conditions are determined by the ratio of CO₂ and O₂ in the atmosphere, as well as by a set of other environmental parameters that affect on the ratio of these gases in photosynthesizing cells (illumination, water salinity, aeration, etc). The subsequent processes of organic matter transformation in the sediment do not significantly affect on the carbon isotope composition. The mechanism of formation of carbon isotope composition of photosynthetic part of "living matter" is suggested. The relationship of facial isotopic differences with isotope composition of photosynthetic biomass is traced.

Keywords: Facies; carbon isotope composition; Precambrian atmosphere; Photosynthesis; Assimilation; Photospiration; Oscillatory mechanism; Facial isotopic differences

Introduction

One of the geochronological characteristics of facies is a carbon isotope composition of sedimentary organic matter (δ¹³C). Considering that carbon isotope composition of coeval organic matter and carbonates are related and their difference is an analog of δ¹³C discrimination in photosynthesis in modern plants [1], both isotopic characteristics can be used to define facies. Oehlerl et al. [2] were the first, who having studied the most ancient sedimentary organic matter (Svaziland, 3, 3 Ga), found the jump between carbon isotope composition of organic matter in the overlaying younger strata and in the above strata of amounted to 12% on average (from -14.7/-19.5% to 35/40%). Following the actualism principle, the authors ascribed this jump to photosynthesis origin. This finding correlated well with the results of the studies on photosynthetic CO₂ assimilation where it was shown that CO₂ fixation, occurring at the stage of enzymatic carboxylation of ribulosebisphosphate (RuBP), causes the isotopic effect of the expected value [3]. The isotopic jump also witnessed the appearance in the course of evolution the key enzyme of photosynthesis mentioned above. The above conclusions were also supported by the subsequent examination of the mechanism of carbon isotope fractionation in photosynthesis [4,5]. Nevertheless, the obtained results were insufficient to define isotopic facial features, since isotopic composition of sedimentary organic carbon depends not only on isotope fractionation at the stage of "living matter", mainly related to photosynthesis, but also on the stage of organic matter oxidation after burial. First of all, it should be noted that carbon isotope composition of "living matter", which includes biomass of all living organisms on the Earth, is completely determined by carbon isotope composition of photosynthesizing biomass. It is explained by the fact that carbon isotope fractionation in heterotrophic assimilation is negligible relative to photosynthetic assimilation [6], combined with the notion that photosynthesizing organisms give rise to all trophic chains. One more notion should be done. Carbon isotope fractionation at the stage of "living matter" in non-oxidative conditions of environment is different as compared with isotope fractionation in oxidative atmosphere.

Facial carbon isotope differences in non-oxidative Precambrian atmosphere

It was established that Proterozoic carbonates in the period 2.3/1.9 Ga were abnormally enriched in δ¹³C (from 10% up to 20%) [7]. In the period under review the atmosphere was anoxic, because molecular oxygen, produced in photosynthesis hasn’t been collected, but completely was spent for oxidation of igneous rocks and other reduced forms. The atmosphere predominantly consisted of carbon dioxide and methane. The methanogens were the prevalent life form. It allows concluding that the cause of the abnormally δ¹³C-enriched carbonates could be bound to the microbial conversion of carbon dioxide into methane. This process is known to be followed by carbon isotope fractionation [7,8]. Protobionts, living in Precambrian, needed oxygen to supply their energy metabolism. However because of anoxic atmosphere in Precambrian they were enforced to take it from oxygen-containing compounds. Carbon dioxide was the most appropriate form in the primitive "atmosphere-hydrosphere" system to provide oxygen. It contributed to the occurrences of methanogens that produced methane [7]. The Raleigh effect, which commonly accompanies the isotope fractionation in natural conditions, strengthens isotopic discrepancies by multiplying one-off isotope effects of the reaction. Anaerobic photosynthesis continued until the ocean has not ceased to absorb oxygen and it appeared in the atmosphere in a molecular form. A toxic release of oxygen has happened in 2.5 to 2.3Ga interval [9]. Two billion years ago due to photosynthesis of cyanobacteria the atmosphere has become oxidative. It was the crucial point of evolution.

Facial carbon isotope differences in oxidative Precambrian atmosphere

In Late Proterozoic (0.8-0.6 Ga) the abnormally "heavy" carbonates were also found. In oxidative atmosphere of that time the reason for the appearance of unusually δ¹³C enriched sedimentary carbonates was...
quite different as compared with previous case. The $^{13}\text{C}$ enrichment was due to CO$_2$ consumption in photosynthesis followed by carbon isotope effect. It resulted in a depletion of CO$_2$ in "atmosphere-hydrosphere" system during orogenic cycle and was accompanied with Raleigh effect. The extent of depletion was abnormally high, since initial filling of the system with CO$_2$ in orogenic period of the cycle wasn't great because the amount of sedimentary organic matter, the source of CO$_2$, accumulated in the first orogenic cycles, was much less as compared with the subsequent periods of Phanerozoic [10]. The emergence of free oxygen in the atmosphere resulted in dramatic events in biosphere. Cyanobacteria got photorespiration. It means they adapted to oxygen and have learned to use it. The enzyme RuBP carboxylase has acquired a new oxygenase function and together with it the oscillatory mechanism of photosynthesis [5]. Notably, that assimilatory part of the photosynthetic mechanism was associated with carbon isotope fractionation resulting in 12C enrichment of the assimilated carbon. The photorespiratory part of the mechanism was also associated with carbon isotope fractionation but with the effect of opposite sign relative. The effect was bound to glycine decarboxylation reaction in photorespiratory chain of cell metabolism. Both CO$_2$ assimilation and photorespiration, contributed to carbon isotope composition of total biomass. The summary isotope composition of biomass depended on CO$_2$/O$_2$ concentration ratio in the environment. The increase of oxygen concentration intensified photorespiration and strengthened $^{13}\text{C}$ the enrichment of biomass. The growth of carbon dioxide stimulated assimilation and $^{13}\text{C}$ accumulation in biomass. Organisms with such an organization of photosynthesis are usually attributed to the C-3 type.

How does CO$_2$/O$_2$ concentration ratio in the environment determines the carbon isotope composition of "living matter"

The "living matter" consists of two parts – photosynthetic and heterotrophic. Since photosynthesizing organisms give rise to all trophic chains carbon isotope composition of them determines isotope composition of whole "living matter". It means to understand carbon isotope composition of "living matter", it is necessary to find out how carbon isotope composition of photosynthesizing biomass is formed. Basing on previous studies [4], let's see, how it is formed carbon isotope composition of biomass of a generalized cell of C-3 organisms. (Figure 1) illustrates the mechanism of formation of carbon isotope composition of biomass of C-3 organisms. Carbon dioxide enters the cell and, fixed by RuBP acceptor, is involved into Calvin cycle. The fixation is followed by carbon isotope effect resulting in $^{13}\text{C}$ enrichment of the assimilated carbon. Then a part of the pool of assimilated carbon is used to provide glycolate cycle with substrates, where photorespiration occurs. In glycine dehydrogenase reaction of the glycolate cycle a part of the carbon flow is subjected to oxidative decarboxylation and is evolved from the cell. Due to iso tope effect of decarboxylation the photosynthetic flow of the cycle is enriched in $^{13}\text{C}$. Both flows of assimilatory and photorespiratory carbon are separated in time due to oscillatory mechanism and to a strict temporal cell organization. Thus, carbon of assimilatory pool enriched in $^{13}\text{C}$ and carbon of photorespiratory pool enriched in $^{12}\text{C}$, make the total biomass. The contribution of each pool to biomass determines its carbon isotope composition and depends on CO$_2$/O$_2$ concentration ratio in the environment. In the accordance with the suggested oscillatory mechanism of photosynthesis [5], the alternation of assimilation and photorespiration is controlled by the switching of enzyme RuBP carboxylase/oxygenase from carboxylase to oxygenase function and back depending on the CO$_2$/O$_2$ ratio in the cell. The latter in turn depends on the ratio in the environment. Thus carbon isotope composition of total biomass is a function of CO$_2$/O$_2$ ratio. Basing on actualism principle, we believe that oscillatory mechanism of photosynthesis is typical to photosynthesis of C-3 type organisms at present and in the past. Passing from the photosynthesis of the individual organism to the concept of global photosynthesis, which characterizes the photosynthesis in the global carbon cycle, it should be underlined that the latter has all the characteristics of photosynthesis of an individual organism, excepting ontogenesis, i.e. it doesn't depend on time [1].
What are the facial isotopic differences?

To proceed to the definition of facial isotopic differences it is necessary to estimate isotopic shifts in the transformation of “living matter” into sedimentary organic matter. We assumed that no carbon isotope fractionation occurs in microbial oxidation of biogenic molecules since at present there are no data evidencing to the contrary. The isotopic shifts might appear as a result of partial oxidation of biochemical fractions because they slightly differ in carbon isotope composition due to intracellular isotope fractionation. But this fractionation isn't great. Maximal isotopic differences in the range of 5% characterize lipid fraction. It is known that lipid fraction at the same time is most resistant to oxidation. So the maximal isotopic shifts due to partial oxidation can’t exceed 5%. But in real situation it should be much less because full oxidation of other fractions is a low probable case. Indeed, many researchers, who studied carbon isotope composition of organic matter in sedimentation and diagenesis, observed slight 13C enrichment about 2%-3% with transformation depth of organic matter [11,12]. It is evidenced in favor of increase of lipid fraction in the composition of the remaining organic matter. If to compare the isotopic shifts for the partial oxidation of organic matter with the shifts due to photosynthesis in different environmental conditions, which exceeds 20%, it becomes clear that the effects of photosynthesis should play a dominant role. As said, oxidation doesn’t contribute to carbon isotope composition of sedimentary organic matter. It is left to find out how chemical destruction of organic matter can change its carbon isotope composition. Obviously, carbon isotope composition could change, if the parts of organic matter, mainly carbon products of its transformation with the other isotopic composition, leave organic matter. It might be petroleum, as a whole, and gas (methane). The 13C enrichment of petroleum is a result of their formation, which is bound to lipid fraction. As to methane, its massive removal occurs at high degree of catagenetic transformation.

Conclusion

From balance consideration it is clear, that even in the cases isotopic shifts should be small. Thus one can conclude that at all stages of organic matter transformation its carbon isotope composition reflects isotope fractionation emerged in photosynthesis. By the other words, facial isotopic differences are inherited from the “living matter” and reflect the environmental conditions of photosynthesis (illumination, water salinity, water availability, temperature, mixing, etc.). A set of the environmental parameters forms isotopic peculiarities of the locality, i.e. facial isotopic differences.

Available data confirm these assertions. It is well established fact that the organic matter from marine, fresh water, salt marsh and terrigeneous sediments distinctively different in carbon isotope ratio. The cause is in that all these localities are characterized by different CO2/O2 ratios determining the photosynthesis conditions. The more this ratio is the more organic matter is enriched in 13C.

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