THE SEPARATION OF STABLE ISOTOPES OF CARBON BY THE COUNTER FLOW CHEMICAL EXCHANGE METHOD IN THE GASEOUS PHASE. I.

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THE SEPARATION OF STABLE ISOTOPES OF CARBON BY THE COUNTER FLOW CHEMICAL EXCHANGE METHOD IN THE GASEOUS PHASE. I.

This is a translation of an article written by G. M. Panchenkov, I. A. Semichkin, A. G. Maurina, and N. P. Yershova in Zhurnal Fizicheskoy Khimii (Journal of Physical Chemistry), Vol XXX, No 9, pages 2070-2076.

A very effective method of separating isotopes is that of chemical exchange carried out in a thermonuclear column. In this method, thermal diffusion is utilized only for the production of a counter flow between the molecules entering into the exchange reaction.

To separate isotopes by this method, it is necessary to select a convenient exchange reaction taking place in the gaseous phase with a constant of equilibrium different from one. For example, in the reaction:

\[ ^{12}C_2O_2 + ^{13}C_2O_2 \rightarrow ^{13}C_2O_2 + ^{12}C_2O, \]

the constant of equilibrium at 250°C is equal to 1.086. This reaction was studied in detail by Bernstein and Taylor. The exchange of \(^{13}\)C between CO\(_2\) and CO is carried out in a column of the type of Clusius and Dicket in which the cooler currents, directed downward by convection, became enriched with the heavier CO\(_2\) component. If the isotope exchange favors the transfer of a heavy isotope into a heavy molecule, then this isotope will be concentrated in the cold current and carried by convection to the bottom of the column. Bernstein and Taylor found that this enrichment of the carbon dioxide with the \(^{13}\)C isotope increases with the elevation of the temperature and pressure (the experiment was conducted in the presence of steam at P\(_{H_2O} = 20\) mm of Hg). Under identical conditions, greater enrichment was attained on a platinum filament than either on tungsten or on nichrome. The authors observed an increase in the effectiveness of the separation process in the presence of hydrogen and steam.

Urey and Brandner, and Norris and Ruben who studied the exchange between CO and CO\(_2\) concluded that nitrogen and oxygen also act as catalysts for the exchange.
The positive role of the addition was explained by their accelerating action on the surface of the catalyst \[3,4\], or by a change in the mechanism of the exchange in the mechanism of the exchange in comparison with the exchange without the addition \[1,3\]. These authors \[3,4\] carried out experiments under static conditions, different from those of the conduction process of separating isotopes where the concentration of additions exceeds that of the exchanging gases. Experiments by Bernstein and Taylor \[1,5\] in the study of the influence of additions of hydrogen and steam, though conducted directly in connection with the process of separating isotopes of carbon dioxide, were unsystematic because they were performed with casually selected concentrations of hydrogen and of steam. That presented an interesting opportunity for investigating the influence of the addition of various gases on the exchange. The present work is directed to this problem.

**Experimental Part.**

The study of the process of the separation of isotopes of carbon dioxide was conducted in a thermal diffusion column (1), with an inner tube 1.0 cm in diameter and a length of 165 cm. The tube was cooled externally with water at constant temperature. Inside the reaction tube was a tightly drawn platinum wire (2) 0.5 mm in diameter, to which were welded glass "crosses" to keep it in position. To keep the wire under tension, a small glass container (3) two-thirds filled with mercury, and having a diameter of 0.8 cm and a length of 20 cm, was soldered to it with two molybdenum wires. The contact was established by means of mercury poured in a trap (4). The filament was heated by an alternating current to 800-900°C. The temperature of the filament was determined by an OFFIR-45 optical pyrometer, and by the change in the resistance of the filament. As shown in the table (page 2072) both methods gave close by agreeing values.

As its upper part, the column was connected with a reservoir whose 15-l capacity guaranteed a supply of gas with a natural ratio of isotopes for the column.

To supply carbon dioxide from the reservoir to the column, a tubular furnace (5) was heated by an electric current to 800°C. At its lower end, the column was connected to a converter (6) in which reduction of CO₂ to CO was taking place to generate the carbon monoxide required for the exchange reaction. The converter consisted of a glass tube of 5 cm in diameter and 30 cm in length, equipped with a soldered pocket for a thermometer. Little balls prepared from zinc
dust and asbestos wadding were used as the reducing agent, according to the method of Bernstein and Taylor. 
Reduction of the carbon dioxide occurred at the temperature most favorable for the conversion (390-400°C). To supply carbon dioxide from the converter to the column, a tubular electric furnace was built in and heated to 450°C.

The converter transformed almost all the CO₂ into CO, the latter moving as a stream (past the filament) upwards and exchanging with the down-moving stream of CO₂ isotopes. With the availability of a large-capacity reservoir at the top of the column, and with a complete return, the number of moles of the heavy component CO₂ entering into the column is equal to the number of moles of the light component CO leaving the upper part of the column. Under these conditions no drop in the concentration of both components occurs along the length of the column. At the same time there is a vertical isotope composition gradient. Upon attainment of equilibrium, this isotope composition gradient remains constant.

Chemically pure carbon dioxide, which occupied the column and the large reservoir prior to the start of the experiments, was prepared in an ordinary conventional type apparatus consisting of a drop funnel, flasks, two drying columns packed with CaCl₂ and P₂O₅, and a pressure gauge. The gas was made by the interaction of sulfuric acid and chemically pure Na₂CO₃. The system was evacuated to approximately 0.01 mm of Hg. To conduct experiments with steam in the system, a small trap was added in which a weighed amount of distilled water was introduced. Prior to evacuating the system (during the generation of the carbon dioxide), the water trap was cooled by means of liquid nitrogen.

To study the effects of the addition of various gases on the separation of the carbon isotopes, a special gas pipette was constructed (Fig 2) which allowed the entry of the additional gases into the system, in proportion to their partial pressures which was from 1 to 20 mm of Hg. Under the pressure of the carbon dioxide in the system (at 500 mm of Hg), the charging of the system with the additional gases was carried out by a branch tube of the apparatus for CO₂ generation, into which a capillary was welded for smooth entry of the gases.

Before every experiment, the entire system was evacuated to 0.01 mm, and maintained in this condition for 24 hours. Experiments were conducted only if this vacuum was maintained. The system was washed two to three times with carbon dioxide and evacuated each time to the indicated pressure, after which it was filled to the working pressure.
Fig 1. Apparatus for the separation of isotopes

The charging was conducted with heated furnaces and a converter (Fig 1, No. 6). The time was measured from the moment the current used to heat the filament to the necessary temperature was switched on. Each experiment was continued for four to five days. If the experiment was conducted with the addition of steam, then during the time of charging with carbon dioxide the trap was gradually heated to room temperature. The stream of dry carbon dioxide passing through the trap entrained water from it, in the form of steam, into the system. The charging of the system with the added quantities of gases to the specific pressure was done by means of a pipette (Fig 2), after filling the system with carbon dioxide to 500 mm Hg. (The reason for selecting this particular pressure will be explained below). After 2, 6, 12, 24, 48, 72, 96, and in some experiments 120 hours, samples of carbon dioxide were collected for analysis of $^{13}C$ content, and collected in
ampoules at the base of the column. The analysis was made with a mass spectrometer of the MC-3 type. The accuracy of the analysis was kept within one to two percent of the content of $^{13}\text{O}$ in the carbon dioxide.

Temperature of the Filament, $^\circ\text{C}$

<table>
<thead>
<tr>
<th>Temperature (F)</th>
<th>Value Determined from the Change in the Resistance of the Filament</th>
<th>Pyrometer OPPIR-45</th>
<th>Average Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>710</td>
<td>740</td>
<td></td>
<td>725</td>
</tr>
<tr>
<td>800</td>
<td>800</td>
<td></td>
<td>800</td>
</tr>
<tr>
<td>870</td>
<td>930</td>
<td></td>
<td>900</td>
</tr>
</tbody>
</table>

Prior to investigating the role played by the additions in the process of enrichment of the carbon dioxide with the $^{13}\text{O}$ isotope, we conducted experiments to clarify the influence of the exchange temperature, of the material in the filaments, and of the pressure in the column. Experiments were conducted at pressures of 350, 500 and 700 mm of Hg. We determined that the enrichment on the platinum filament was greater than on the nichrome. The enrichment increases with temperature. It was found that by changing the pressure within the above limits, maximum enrichment was obtained at 500 mm. For example, in experiments conducted at $900^\circ\text{C}$ (Fig 3) with a platinum filament, the enrichment was greater at 500 than at 350 and 700 mm. Similar effects were observed at $700^\circ\text{C}$, on both the platinum and nichrome filaments. These findings are somewhat contradictory to those of Bernstein and Taylor; with reference to the decrease in enrichment with increasing pressure.

However, the fact that the conclusions of these authors were based on experiments at pressures of 0.5, 1 and 1.5 atmospheres, and at intervals at which the process was not studied, allows us to hope that the accuracy of our conclusions are sound. In this connection, all our further experiments were conducted only at a pressure of 500 mm of Hg.
(Fig 2. Gas pipette for measuring doses of gas)

The general coefficient of enrichment was calculated according to the formula:

\[ S_t = \frac{R_0}{R_t}, \]  

where \( R_0 \) is the ratio of the peaks obtained by mass spectrometric analysis of the existing carbon dioxide, and is equal to the ratio of mole fractions of \( \frac{[^{12}\text{O}_2]}{[^{12}\text{O}_2]} \) to \( \text{CO}_2 \) in carbon dioxide at \( t = 0 \); \( R_t \) is the corresponding ratio in a sample of carbon dioxide taken from the lower part of the column at some time \( t \) after the start of the experiment.

It is not difficult to show that under equilibrium conditions of a column with a large reservoir, the common coefficient of enrichment, calculated by the above method, is equal to the general coefficient of separation. Indeed,

\[ S_{t=\infty} = \frac{R_0}{R_{t=\infty}} = \frac{1}{\frac{1}{R_0} \cdot \frac{[^{12}\text{O}_2]_{t=\infty}}{[^{12}\text{O}_2]_{t=\infty}} \cdot \frac{[^{13}\text{O}_2]_{t=\infty}}{[^{12}\text{O}_2]_{t=\infty}}} \]

where \( h \) is the length of the column from which the samples are taken.

Taking into account the fact that at the upper end
of the column, when the system is in equilibrium, the concentration of $[\text{C}^{18}\text{O}]_{t=\infty; h=H}$ is equal to the concentration of $[\text{C}^{18}\text{O}]$ in the reservoir (that is, is equal approximately to $[\text{C}^{18}\text{O}]_{t=0}$, it is possible to express $S_{t=\infty}$ in the following manner:

$$S_{t=\infty} = \frac{[\text{C}^{18}\text{O}]_{t=\infty; h=0}}{[\text{C}^{18}\text{O}]_{t=\infty; h=H}} / \frac{[\text{C}^{18}\text{O}]_{t=\infty; h=0}}{[\text{C}^{18}\text{O}]_{t=\infty; h=H}}.$$

At equilibrium, there is no gradient of concentration of both components CO and CO$_2$ along the column at any given height:

$$[\text{CO}] = [\text{CO}_2]; \quad [\text{C}^{18}\text{O}] = [\text{C}^{18}\text{O}_2] = [\text{C}^{18}\text{O}] = [\text{C}^{18}\text{O}_2].$$

If we assume this to be correct also for the upper end of the column (where $h$ equals $H$), then the expression for $S_{t=\infty}$ could be given by the expression:

$$S_{t=\infty} = \frac{[\text{C}^{18}\text{O}_2]_{t=\infty; h=0}}{[\text{C}^{18}\text{O}_2]_{t=\infty; h=H}} / \frac{[\text{C}^{18}\text{O}_2]_{t=\infty; h=0}}{[\text{C}^{18}\text{O}_2]_{t=\infty; h=H}}.$$

That is, we arrive at the expression of the general coefficient for the separation of isotopes.

Therefore, in further discussion, the meaning of $S_{\text{static}} = S_{t=\infty}$, calculated according to equation (2) after reaching a steady state, will be considered as the equilibrium general coefficient of separation.

![Graph](image.png)

Fig 3. Dependency of the general coefficient of enrichment on the pressure in the system (Platinum filament at 900° C.) 1 = 350 mm, 2 = 700 mm, 3 = 500 mm of Hg.
Effect of steam. We conducted four experiments at steam concentrations in the system corresponding to partial pressures of 1, 3, 10, and 15 mm of Hg. The pressure of CO₂ was 500 mm. The calculations for the water necessary to develop these pressures were made according to the ideal gas laws. The duration of each experiment was four days; the temperature of the platinum kilament was 800°C. As can be seen in Figs 4 and 5, the equilibrium general coefficient of separation (of isotopes in our column) increases with an increase of steam, from S₀ = 1.13 (without additions) to S₀ = 1.20 (at a water pressure of 15 mm of Hg). The time needed to reach equilibrium decreases from four to five days (without additions) to one day (in the presence of steam), while remaining independent of the concentration of steam in the system.

The addition of steam increases the speed of exchange, which leads to an increase in the efficiency of the column, and to a change in the number of theoretical plates.

When the change in the coefficient of separation was determined by the velocity of isotope exchange and not by diffusion, Bernstein and Taylor [1] obtained the following ratio between the number of theoretical plates n and the constant velocity of the exchange k:

\[ n = \frac{k}{v} H \]

where v represents average convection velocity (in cm/perm sec), and H is the height of the column.

In drawing conclusions from this relationship, the authors assumed that the reactions of isotope exchange are subject to first order equation relative to the presence of the rare isotope [7].
Fig 4. Dependency of the general coefficient of enrichment upon time, at the various partial pressures of steam. 1 = without addition of steam; 2 = 5 mm. steam pressure; respectively, 1, 3, 10 and 15 mm. of Hg.

![Graph showing dependency of Enrichment upon Time](image)

Fig 4. Dependency of the equilibrium general coefficient of separation on the partial pressure of the steam.

The acceleration in the rate of exchange of isotopes of carbon dioxide between CO₂ and CO by means of steam depends upon the predominance of the reaction:

\[
\text{CO} + \text{H}_2\text{O} \rightarrow \text{CO}_2 + \text{H}_2
\]

which, at temperatures around 800° becomes important in the oxidation of CO (8).

It is possible, too, that the accelerating action of steam on the surface of the platinum filament is a
contributing factor.*

Effects of hydrogen. We conducted four series of experiments with additions of hydrogen, at partial pressures in the system of 1, 3, 5 and 10 mm of Hg. The conditions of the experiments were the same as those with steam. Repeated efforts to conduct the experiment at partial pressures of hydrogen of 20 mm were unsuccessful because the platinum filament burned out 20-30 minutes after the start of the experiment.

Apparently, due to the high adsorbeny of the hydrogen by the platinum wire, the latter disintegrated to such an extent that it was unable to withstand the tension, and hence fractured.

![Graph](image)

Fig 6. Dependency of the general coefficient of enrichment on time at various partial pressures of hydrogen. 1 = 1 mm; 2 = 3 mm; 3 = 5 mm; 4 = 10 mm Hg; 5 = without addition of H₂.

The experiment demonstrated (Fig 6) that the time for attaining an equilibrium at the pressure of hydrogen diminishes from 100-120 hours (in the absence of any additions to the mixture), to approximately 12 hours. The coefficient of separation increases from 1.3 (in the absence of additions) to 1.20 (with pressure of hydrogen equal to one mm of Hg). However, in case of an increase

* In experiments with the addition of steam and various gases, we were not able to determine their effect upon thermal diffusion and the consequent effect upon the column, even though such influences, in all probability occur.
in concentration of hydrogen in the system, the coefficient of separation gradually diminishes to 1.10 (at a pressure of hydrogen equal to 10 mm), i.e., it appears to be even smaller than in the absence of additions.

![Graph showing the dependency of the equilibrium general coefficient of separation on partial pressures of hydrogen.](image)

Fig 7. Dependency of the equilibrium general coefficient of separation on partial pressures of hydrogen.

The curve shown in Fig 7, indicating the dependency of the equilibrium general coefficient of separation on the partial pressure of hydrogen, convinces us that it is necessary to accept the following: in the process of exchange of isotopes between CO and CO2 an important role appears to be played by the reaction:

\[ \text{CO}_2 + \text{H}_2 \rightleftharpoons \text{CO} + \text{H}_2\text{O} \]

which takes place on the surface of the platinum filament. In support of this assumption, there is the presence of a maximum on the curve Fig 7, the exact location of which is in the interval of 0-3 mm of Hg. Indeed, considering the expression for the velocity of the reaction of two gases on a surface, derived on the basis of the Langmuir equation:

\[ v = \frac{k_{PAPB}}{(1 + k_{PAPB})^x} \]

it is not difficult to note that the expression with the greater adsorbency of the gases (in our case, hydrogen) inevitably leads to the existence of a maximum. Similar effects were observed earlier with the hydrogenation of carbon dioxide by hydrogen on platinum.
Conclusions

1. The reaction of isotope exchange (1) taking place in a diffusion column was studied.
2. The optimum pressure of carbon dioxide for the separation of the isotopes of carbon was found to be 500 mm of Hg.
3. An increased effect in separation was obtained by the action of steam.
4. It was determined that the introduction of hydrogen into the system (in small quantities on pressure of hydrogen of the order of one mm of Hg) increases the effectiveness of the isotope separation, but that it then decreases with increased concentration of hydrogen. A theory explaining similar effects was advanced.

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Submitted
20 November 1955

Bibliography

FOR REASONS OF SPEED AND ECONOMY
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