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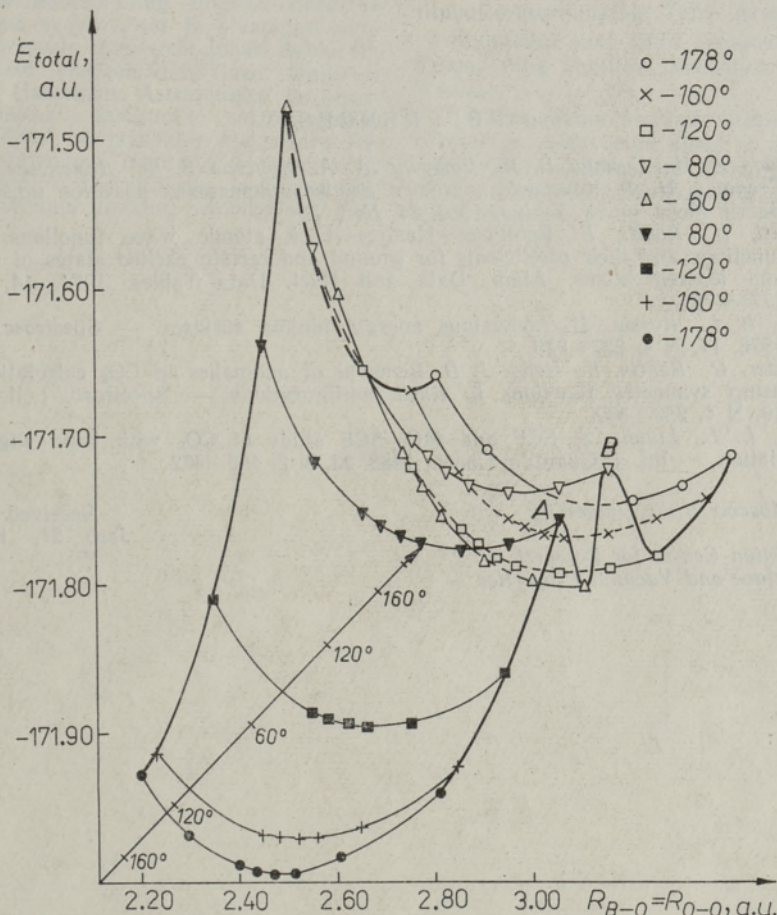
## THEORETICAL INVESTIGATION OF THE O—B—O → B—O—O REARRANGEMENT

P.-E. LOMP, V. SIMKIN, A. DEMENTIEV, N. STEPANOV. O—B—O → B—O—O ÜMBERASSETUSE  
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 ПЕРЕГРУППИРОВКИ O—B—O → B—O—O

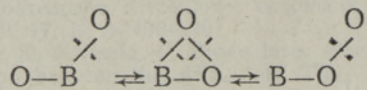
(Presented by M. Veiderma)

To investigate the O—B—O → B—O—O rearrangement, *ab initio* restricted SCF calculations of fragments of potential surfaces in  $C_s$  point



Potential surface of O—B—O → B—O—O rearrangement. SCF calculations in  $C_s$  point group in STO-3G basis set.

group for  $X_2\Pi$  ( $^2A'$ ) and  $^2A''$  states using the SPUSH program system [1] were carried out ( $R_{B-O}$  and  $R_{O-O}$  here varied between 2.00 and 3.20 a.u., bond angle between 180° and 60°). The calculations in the standard STO-3G basis set [2] yield OBO to be more stable. The rearrangement is proposed to go over a triangular structure:



The potential surface of this rearrangement for the ground state is presented in the Figure. The surface for the  $^2A''$  state is similar.

It is remarkable that the total energy minimum corresponds to approximately  $R_{B-O}=R_{O-O}$  in BOO. This condition is also used in the Figure.

The calculations show an existence of peaks approximately at 80° ( $A$  and  $B$ ). So the triangular structure seems to have a stability. Similar results have been published for  $X^1\Sigma^+$  ( $^1A_1$ ) state of  $\text{CO}_2$  [3, 4] where the second minimum was found at  $\angle \text{OCO}=80^\circ$  and maximum at 105° while MCSCF treatment [5] between 60 and 180° showed that a cyclic isomer is unstable.

For the energy of rearrangement a value of 320 kJ/mol (0.136 a.u.) has been found.

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