## LEAST SQUARES FITTING OF PERTURBED VIBRATIONAL POLYADS NEAR THE ISOMERIZATION BARRIER IN THE S $_1$ STATE OF $\rm C_2H_2$

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The S<sub>1</sub> electronic state of acetylene has recently been shown to have two potential minima, corresponding to cis- and trans-bent structures. The trans-bent isomer is the more stable, with the cis-bent isomer lying about 2670 cm<sup>-1</sup> higher; the barrier to isomerization lies roughly 5000 cm<sup>-1</sup> above the trans zero-point level. The "isomerization coordinate" (along which the molecule moves to get from the trans minimum to the barrier) is a combination of the  $\nu_3$  (trans bending) and  $\nu_6$  (cis bending) vibrational normal coordinates, but the spectrum is very confused because the  $\nu_6$  vibration interacts strongly with the  $\nu_4$  (torsion) vibration through Coriolis and Darling-Dennison resonances. Since the  $\nu_4$  and  $\nu_6$  fundamental frequencies are almost equal,<sup>*a*</sup> the bending vibrational structure consists of polyads. At low vibrational energies the polyads where these three vibrations are excited can be fitted by least squares almost to experimental accuracy with a simple model of Coriolis and Darling-Dennison interactions, but at higher energies the huge  $x_{36}$  cross-anharmonicity, which is a symptom that the levels are approaching the isomerization barrier, progressively destroys the polyad structure; in addition the levels show an increasing even-odd staggering of their *K*-rotational structures, as predicted by group theory. It is not possible to fit the levels near the barrier with a simple model, though some success has been achieved with extended models. Progress with the fitting of the polyads near the barrier will be reviewed.

<sup>a</sup>A. L. Utz, J. D. Tobiason, E. Carrasquillo M., L. J. Sanders and F. F. Crim, J. Chem. Phys. <u>98</u>, 2742, 1993.