DETERMINATION OF THE STRUCTURE OF CYCLOPENTENEOXIDE AND THE ARGON CYCLOPENTENEOXIDE VAN DER WAALS COMPLEX

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Rotational spectra of cyclopenteneoxide and the argon cyclopenteneoxide van der Waals complex were studied using pulsed-jet Fabry-Perot Fourier transform microwave spectroscopy. Spectra of the normal isotopomer along with those of the $^{13}$C singly substituted isotopomers for both the monomer and the complex were measured in the frequency region of 6-20 GHz. The boat structure for cyclopenteneoxide was confirmed with naturally abundant $^{13}$C isotopic substitution. For the argon cyclopenteneoxide complex, both $a$ and $b$-type transitions were observed and the rotational constants for the all $^{12}$C isotopomer were determined to be $A = 3268.2537(5)$, $B = 993.3458(2)$, and $C = 950.4300(2)$ MHz. The coordinates of the argon in the principal axis system of cyclopenteneoxide are $a = 0.27$, $b = 0.42$, and $c = 3.91\,\text{Å}$, such that the argon atom is exo to the bridged oxygen of the ring. An investigation into whether or not the argon atom undergoes large amplitude cross-ring motion will be presented.