

CONFORMATIONAL ASSIGNMENTS OF ROTATIONALLY RESOLVED SPECTRA OF 1-HEXOXY AND 1-HEPTOXY

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In recent work we have used ab initio calculations and rotational analyses to make conformational assignments for a number of bands in the laser induced fluorescence (LIF) spectra of the 1-propoxy, 1-butoxy and 1-pentoxyl radicals. For larger alkoxy radicals (number of carbon atoms greater than 5), the number of conformers becomes so great and the rotational structure so complex, that this method is very time-consuming and unattractive. Rather than pursue detailed rotational analysis of the 1-hexaoxy and 1-heptoxy spectra we have made conformational assignments to individual spectra lines by using a guide to the corresponding conformational assignments of smaller alkoxy radicals and their rotational constants and parameters from ab initio calculations. Refined assignments were made by using the GUI program SpecView.