

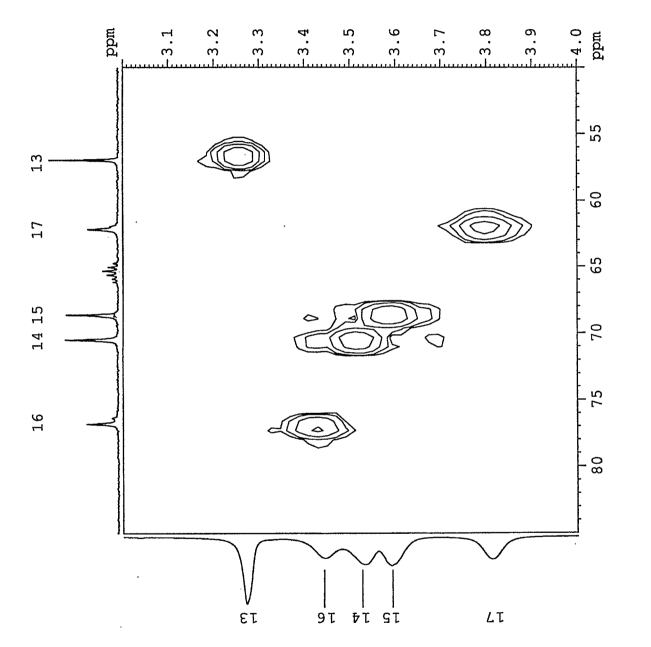
Macromolecules, 1998, 31(9), 2731-2743, DOI:10.1021/ma971368s

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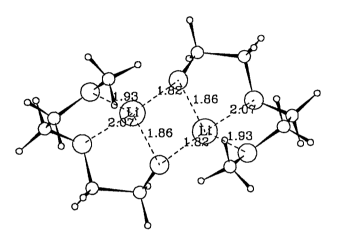


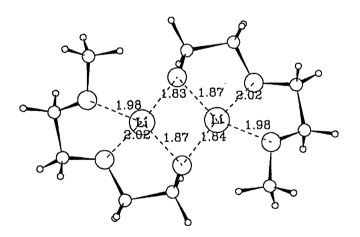
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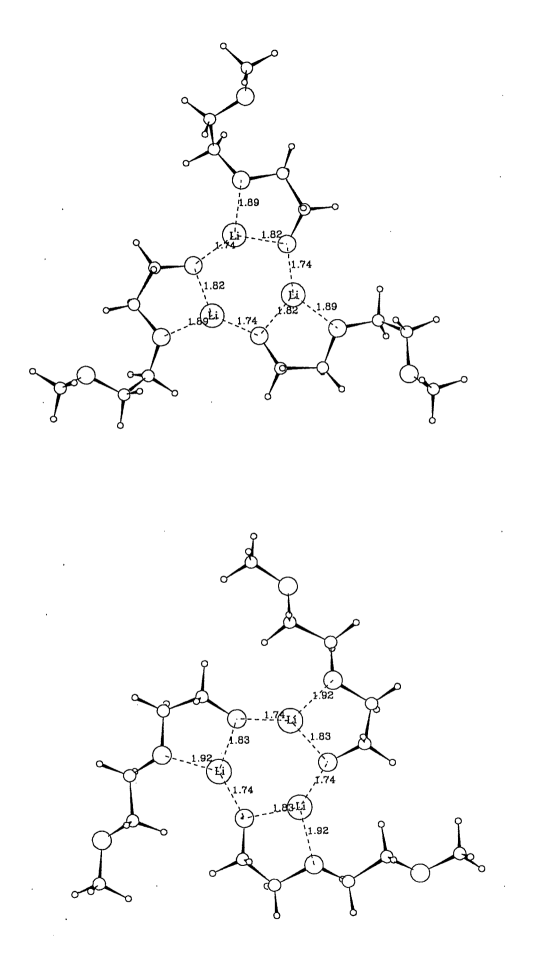
Page 1: :  ${}^{1}H{}^{-13}C$  COSY(HETCOR) NMR spectrum of the 1M solution of LiOEEM in THFd<sub>8</sub> at 253 K.

Page 1



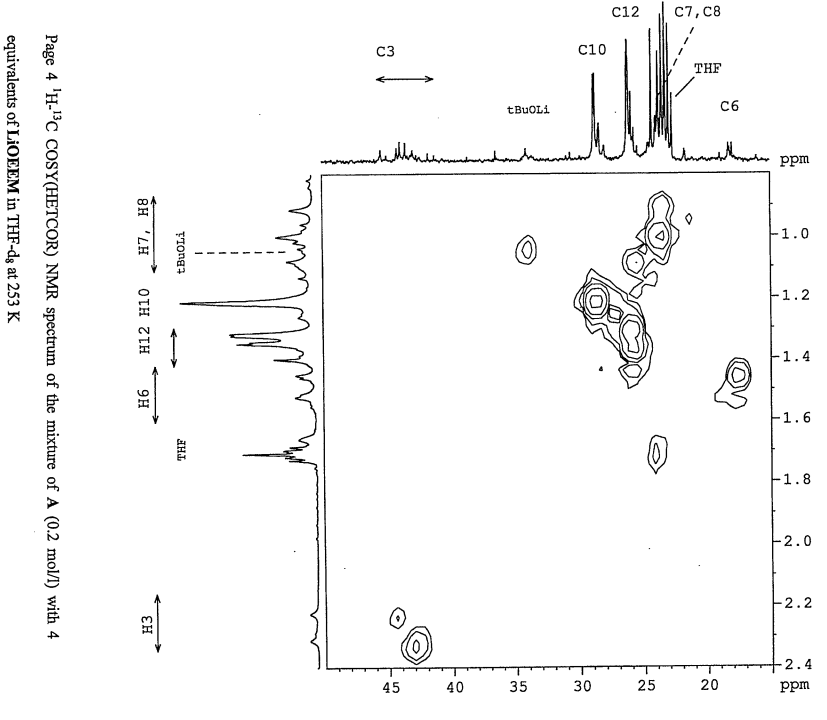


Page 2: Structures of two most stable dimeric aggregates of LiOEEM as predicted by *ab initio* SCF 3-21G calculations



Page 3: Structures of two most (and almost equally) stable trimeric aggregates of LiOEEM as

predicted by ab initio SCF 3-21G calculations



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