

扩展功能

## 重水中固氮酶催化还原乙炔产物的<sup>1</sup>H NMR研究

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摘要 用<sup>1</sup>HNMR研究了固氮酶在重水中催化还原乙炔的反应产物氘代乙烯。

种用群对称性对<sup>1</sup>HNMR谱图进行了归属,计算了几种可能的C~2H~2D~2

结构以及C~2H~3D结构的NMR谱线频和强度,得出了理论谱,通过理论谱与实验谱的比较,

表明固氮酶在重水中催化还原乙炔的产物主要以顺式结构C~2H~2D~2为主,并含有较多的单氘代烯。

单氘代乙烯相对乙烯的化学位移往高场移动约4.0Hz,而双氘代乙烯向高场的位移大约是单氘代乙烯的2倍左右。

关键词 [乙炔](#) [氘代乙烯](#) [催化反应](#) [固氮酶](#) [重水](#) [化学位移](#) [质子磁共振谱法](#)

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## **<sup>1</sup>H NMR study on the products of the catalytic reduction of ethyne by nitrogenase in D~2O**

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**Abstract** The products of the catalytic reduction ethyne by nitrogenase in D~2O were studied by <sup>1</sup>H NMR. In order to assign the <sup>1</sup>H NMR spectrum, the theoretical spectra of C~2H~2D~2 were calculated using group symmetry. The final eigenfunctions,eigenvalues,allowed transitions and intensities of C~2H~2D~2 and C~2H~2D were obtained. It can be concluded that the products comprised mainly of cis-C~2H~2D~2. There was also some C~2H~3D. <sup>1</sup>H chemical shift in C~2H~3D moved by ca. 4. 0Hz to the upfield compared with that in C~2H~4. The upfield shift in C~2H~2D~2 was about twice as large as that in C~2H~3D. The theorical results are in good agreement with experimental ones.

**Key words** [ACETYLENE](#) [CATALYTIC REACTION](#) [NITROGENASE](#) [HEAVY WATER](#) [CHEMICAL SHIFT](#)  
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