Density Functional Theory Study on Equation of State and Vibration Properties of TATB Crystal

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Abstract: The equation of state and vibration properties of TATB crystal were investigated by using the density functional theory (DFT) and combining with van der Walls force correction(vdW-DF2). The partial vibration modes of TATB crystal were reassigned. Vibration mode coupling and the intermolecular interaction process were studied under pressure process up 8.5 GPa. Results show that the NO₂ and NH₂ vibrations were strongly coupled in TATB crystal. The vibration in the wave number range of 1100 cm⁻¹ to 1500 cm⁻¹ is particularly complex because of the coupling of NH₂ with NO₂ and benzene ring vibrations. With increasing pressure, TATB molecules from neighboring layers bend and close to each other, causing a coupling of NH₂ plane twist vibration or wag with NO₂ shear vibration, indicating a strengthening of intermolecular hydrogen bonding.

Key words: TATB; equation of state; vibration properties; density functional theory (DFT)

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含能材料基因科学学术研讨会在京召开

由中国工程物理研究院化工材料研究所发起主办,化工材料研究所含能材料基因科学研究中心承办的"含能材料基因科学学术研讨会"于 2016 年 5 月 20 日在北京召开。来自中国科协、中科院、军委装备发展部、北京理工大学、南京理工大学、中北大学、西南科技大学、中国钢研科技集团、航天科工集团、兵器科学研究院、中物院计算机应用研究所、中物院流体物理研究所和中物院化工材料研究所的三十余位院士、专家和学者齐聚一堂,就材料基因组计划的国内外发展动向、含能材料基因科学发展构想、含能材料基因表达与编码、数据库建设等前沿热点问题进行了热烈讨论。与会专家表示,将以此次研讨会为契机,在规划论证、联合攻关、数据库建设、人才培养和学术交流方面携手合作,共同推动含能材料基因科学在中国的发展壮大。

(化工材料研究所含能材料基因科学研究中心 供稿: