

Simulation of oriented NMR spectra : combining molecular dynamics and chemical shift tensor calculations

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Theoretical prediction and assignment of vicinal 1H–1H coupling constants of diastereomeric 3-alkoxy-6,7-epoxy-2-oxabicyclo[3.3.0]octanes

Aav, Riina; Pehk, Tõnis; **Tamp, Sven**; **Tamm, Toomas**; **Kudrjašova, Marina**; **Parve, Omar**; **Lopp, Margus** Magnetic resonance in chemistry 2011 / p. 76-82 : ill