

Calculation of hydration enthalpies of aqueous transition metal cations using two coordination shells and central ion substitution

Uudsemaa, Merle; Tamm, Toomas Chemical physics letters 2004 / 1/3, lk. 54-58 : ill
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Calculations of hydrated titanium ion complexes : structure and influence of the first two coordination spheres

Uudsemaa, Merle; Tamm, Toomas Chemical physics letters 2001 / p. 667-672

Influence of protonation upon the conformations of biperidine, bismorpholine, and their derivatives

Uudsemaa, Merle; Laars, Marju; Kriis, Kadri; Tamm, Toomas; Lopp, Margus; Kanger, Tõnis Chemical physics letters 2009 / 1/3, p. 92-96

pKa calculation for monoprotonated biperidine, bismorpholine and their derivatives in H₂O and MeCN

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