Water nanostructures confined inside the quasi-1D channels of LTL zeolite

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Understanding the formation and evolution of confined water molecules is critical in understanding many chemical and biological processes as well as the water transport inside the Earth. It is often difficult, however, to probe such processes since the host-guest interactions are dynamic in nature. Using a well-defined zeolitic channel as an ideal host and hydrostatic pressure as a driving force, we show how water molecules are introduced and evolve into various confined nano-structures up to 3.37 GPa. In the initial stage of pressure-induced hydration (PIH) occurring inside the undulating 12-ring channels of a synthetic potassium gallosilicate with zeolite LTL topology, water molecules preferentially assemble into hydrogen-bonded clusters, which alternate with water layers. With increasing PIH (by \( \sim 50\% \)) at higher pressures, the interaction between the confined water molecules increases, and the water clusters and layers are interconnected to form hydrogen bonded water nanotubes inside the zeolitic channels. The confined water nanotube closes its maximum access diameter at further increasing pressures and gradually transform into isolated species interacting with the zeolitic host framework. The evolution of the confined water nano-structures is well coordinated by the concerted changes in the framework distortion and the reentrant cation migration, which appear to be driven by the gradual 'flattening' of the host 12-ring channels.

주요어 : 나노구조, 체융합, 다이아몬드엔멜, 방사광가속기, 결정구조, 고압화학

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