New insights into boron's chemistry at room temperature
29 March 2013, by Anne M Stark

A ball-and-stick structural model of rhombohedral boron is shown in the foreground and a picture of Badwater Basin in California is shown in the background. The Badwater Basin salt flats contain high concentrations of evaporative minerals such as borax, an important boron-containing compound. Credit: Tadashi Ogitsu; Liam Krauss/Livermore Computing.

(Phys.org) —Livermore researchers have described in detail the properties of the room temperature form of the element boron.

In the periodic table, boron occupies a peculiar, transitional position. It sits on the first row, and has metallic elements to its left, and non-metals to its right. Furthermore, it is the only non-metal in the third column of the periodic table.

It is not surprising that the crystallographic structure and topology of boron's stable form at room temperature (β-boron) are not shared by any other element, and are extremely complex. The formidable intricacy of β-boron, characterized by interconnecting icosahedra (a regular polyhedron with 20 identical equilateral triangular faces) partially occupied sites, and an unusually large number of atoms per unit cell (more than 300), has been known for more than 40 years.

The bonding orbitals (red and blue surfaces) in B-boron demonstrate how vacancies and self-interstitials can stabilize the structure. Left: Part of the stable form of boron called the B28 unit (gold ball-and-stick) has a local instability that leads to the introduction of B13 vacancies with unoccupied orbitals (red surfaces). Right: The system is stabilized as two interstitials boron atoms (B17 and B18) are introduced as a pair, which transforms the unoccupied orbitals (red surfaces on the left) to nearly complete chemical bonds (blue surfaces on the right near the B17 and B18 interstitials).

Boron remains the only element purified in macroscopic quantities for which the ground state geometry has not been completely determined by experiments. Theoretical progress over the last decade has shed light on numerous properties of elemental boron, leading to a thorough characterization of its structure at ambient conditions, as well as of its electronic and thermodynamic properties.

In the March 8 online edition of Chemical Reviews, LLNL researchers Tadashi Ogitsu and Eric Schwegler along with Giulia Galli of University of California, Davis, discuss in detail starting from the history of boron research, and the properties of β-boron, as inferred from experiments and the ab-initio theories developed over the last decade.