Ordinary polyatomic molecules like H₂O are not supposed to exist on the surface of the sun; at the high temperature of ~5800 K, a water molecule should dissociate into a free-radical OH and an H atom or further into an O atom and two H atoms. I therefore found the title of the 1995 report by Wallace et al. (1) "Water on the Sun" quite incredible, although completely believable after seeing the data. This reaction was in contrast to the possible discovery of water on the moon, which was sensationalized reported in newspapers by way of a press release from the Pentagon. In this case, the scientists themselves were cautious (2), even though from a chemical point of view the presence of water on the moon is less surprising than that on the sun. The decisive difference between the two detective stories is the presence of an unmistakable fingerprint for the case of the sun: the laboratory infrared spectrum of hot water taken by Bernath and his co-workers matched exactly with the observed solar spectrum (1). On page 346 of this issue, Polyansky et al. (3) answer some of the theoretical questions about the spectra. Their results not only confirm the already solid evidence for H₂O on the sun but also demonstrate the power of a recent method of spectral analysis based on variational treatment of nuclear motions in a molecule.

Part of the solution to the puzzle is that H₂O is not found all over the surface of the sun but only in sunspot umbrae (central dark spots), where the temperature is lower, around 3200 K. The existence of molecules on "cold" stars—such as highly luminous red giants or obscure dwarfs, whose temperature in the photosphere is below 4000 K—has been well established observationally. Over the years, Tsuji has been studying chemical models to explain molecular abundance and opacity in such stars (4). His model predicts that water, methane, and ammonia are the most abundant species after hydrogen molecules in objects with temperature T < 1000 K, such as brown dwarfs. Nevertheless, the identification of H₂O in sunspots with the substantial column density of 2 × 10¹⁹ cm⁻² is still very exciting.

Wallace et al. (1) gave a comparison (see figure) of the observed infrared absorption spectra of a sunspot umbra and a penumbra (half dark spot), along with a laboratory emission spectrum. The three strong, broad features in the two lower curves are due to terrestrial water absorption and are incidental. The key feature is the extremely rich spectral structure in the umbra and laboratory spectra. Such intricate structure in the near-infrared region of 2.5 to 1.1 μm has been known for many years and was compiled in 1970 (5). Specialists of the water spectrum such as Benedict (cited in (5)), Flaud, and Camy-Peyret (6) proposed H₂O as the source of the spectrum and took laboratory flame spectra (at 2900 K), but somehow the identification of the spectrum had not been clearly claimed. The sunspot spectrum in the figure is not new either; it is a small portion of the spectrum recorded from 470 to 1233 cm⁻¹ by Noyes et al. in 1982 (7). It is really the new laboratory spectrum of hot H₂O that has led to the identification.

The two spectra may look different at a glance, but closer examination shows exactly coincidence in the frequencies of the spectral lines. Matching frequencies between astronomical and laboratory spectra has been the cornerstone in discoveries of molecules in space. The discriminative power of high-resolution spectroscopy is so high that many important radioastronomical discoveries have been claimed after the observation of one line (with few mistakes). The discovery of interstellar water by Townes and his colleagues was based on the only line of H₂O that appears in the centimeter-wavelength region (8). The matching of the frequencies of tens of spectral lines, as in the figure, is foolproof evidence of the H₂O detection. The two spectra look different because relative intensities of the lines are different and many spectral lines in the umbra spectrum are missing in the laboratory spectrum. This difference is because the temperature of Bernath's sample cell, 1820 K, was much lower than the umbra temperature of 3200 K, an example of the limit laboratory astrophysicists always experience in trying to mimic nature. This restriction sets a limit on empirical fingerprint matching by experiment; beyond this, we have to rely on theory.

The equilibrium structure of H₂O is an isosceles triangle with the apex angle of 104°, but in a real molecule, the nuclei do not stay still, deviating from this structure at any moment. The molecule vibrates and rotates, forming tens of thousands of quantized vibration-rotation levels. The spectrum (figure) is the result of transitions of H₂O between rotational levels and are more than 10 times stronger than the near-infrared transitions between vibrational states. As the temperature increases, higher quantum levels get populated and more spectral lines are observed. At room temperature (~300 K), the spectrum in the figure would be composed of the three strong atmospheric lines, whereas for the temperatures of 1820 and 3200 K, the number of lines increases by a factor of ~10 and ~30, respectively. Thus, in order to analyze a spectrum at high temperature, one needs to analyze higher energy quantum states. The traditional approach, in which vibrational and rotational motions are separated and interactions between them are treated by the perturbation method, does not work well.

Polyansky et al. (3) use a high-quality ab initio Born-Oppenheimer potential-energy surface given by Partridge and Schwenke (9) and solve the Schrödinger equation for the nuclear motion directly using a variational method. This type of brute force calculation made possible by the advent of modern computers is becoming the norm in high-resolution theoretical spectroscopy for simple and light polyatomic molecules. Such calculations by
Miller, Tennyson, and Sutcliffe and many others have played a great role in understanding the H$_3^+$ spectrum in the laboratory and in space (10). Polyansky et al. (3) have demonstrated the power of this new method in approximately doubling the energy range for which assignments of the quantum states of H$_2$O have been made. This has led to an understanding of the spectral lines in the figure and a great many more in other frequency regions. It also revealed new spectroscopic features of hot water, such as strong rotational difference transitions induced by Fermi resonance and large splittings of nearly degenerate levels.

Over a longer term, the following two points come to mind. (i) The discovery of water on the sun, together with the recent observation of H$_2$O on many objects based on data from the Infrared Space Observatory satellite (11), is a part of the ongoing development of molecular astrophysics. Molecules are found everywhere, and it is now accepted that half of visible interstellar matter in this galaxy is in the form of molecules (12). This interplay of astronomy and chemistry will no doubt increase. (ii) The major remaining approximation in calculations of the sort that Polyansky et al. have done is the Born-Oppenheimer separation of electronic and nuclear motion. We can hope that full quantum mechanical treatment of the whole system without this approxima-

References and Notes