Lifetimes of ultralong-range Strontium Rydberg molecules in a dense BEC

F. Camargo*, J. D. Whalen*, R. Ding*, T. C. Killian*, F. B. Dunning†, J. Pérez-Ríos‡, S. Yoshida‡, and J. Burgdörfer‡

* Department of Physics & Astronomy, Rice University, Houston, Texas, USA
† Department of Physics & Astronomy, Purdue University, West Lafayette, Indiana, USA
‡ Institute for Theoretical Physics, Vienna University of Technology, Vienna, Austria, EU

Synopsis The lifetimes and decay channels of ultralong-range strontium Rydberg molecules that contain tens to hundreds of ground-state atoms within the electron orbit are examined by monitoring the time evolution of the Rydberg population using field ionization.

Ultralong-range Rydberg molecules excited in a BEC can be used to explore collective phenomena in quantum degenerate gases such as the creation of polarons[1]. Their use as a probe, however, requires that their lifetimes be sufficient to allow interactions to produce measurable effects. Here we examine the lifetimes of strontium Rydberg molecules in a BEC and the reactions responsible for their destruction. 84Sr atoms cooled in a magneto-optical trap are loaded into an optical dipole trap where they are subject to evaporative cooling to create a BEC with a peak density of ~4x10^{14} cm–3. Two-photon excitation is used to create Rydberg molecules with values of principal quantum number n in the range 49 ≤ n ≤ 72. A typical excitation spectrum is included in Fig. 1, expressed as a function of detuning from the atomic line. The sharp peak at zero detuning results from excitation of thermal non-condensed atoms present in the trap, the broad feature extending to the red results from molecular excitation, the larger the detuning the larger the number of atoms present in the molecule. The evolution of the Rydberg population is monitored by field ionization. Analysis of the data indicates that the destruction of Rydberg molecules results from the same processes as identified in earlier studies with rubidium[2], namely associative ionization, leading to formation of Sr_2^+ molecules, and L-changing reactions.

Figure 1 shows the time evolution of the total Rydberg population. The initial rapid decrease in the Rydberg population results from associative ionization. The slower rate of loss seen at later times results from the decay of L-changed Rydberg atoms. The measured molecular lifetimes, ~3-10 µs, limit the timescales over which studies involving Rydberg molecules in cold, dense gases can be undertaken, and reduce the coherence time in such measurements.

Research supported by the AFOSR, the NSF, the Robert A. Welch Foundation, and the FWF (Austria)

References
[1] F. Camargo et al. to be published
[2] M. Schlagmüller et al. 2016 Phys. Rev. X 6 031020

E-mail: fbd@rice.edu