# High-resolution optical spectroscopy with a buffer-gas-cooled beam of BaH molecules

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Barium monohydride (BaH) is an attractive candidate for extending laser cooling and trapping techniques to diatomic hydrides. The apparatus and high-resolution optical spectroscopy presented here demonstrate progress toward this goal. A cryogenic buffer-gas-cooled molecular beam of BaH was constructed and characterized. Pulsed laser ablation into cryogenic helium buffer gas delivers  $\sim 1 \times 10^{10}$  molecules/sr/pulse in the  $X^2\Sigma^+$  (v''=0,N''=1) state of primary interest. Approximately  $4 \times 10^6$  of these molecules per pulse enter the downstream science region with forward velocities below 100 m/s and transverse temperature of 0.1 K. This molecular beam enabled high-resolution optical spectra of BaH in quantum states relevant to laser slowing and cooling. The reported measurements include hyperfine structure and magnetic g-factors in the  $X^2\Sigma^+$ ,  $B^2\Sigma^+$ , and  $A^2\Pi_{1/2}$  states.

## I. INTRODUCTION

Experiments with cold and ultracold molecules are on the forefront of table-top fundamental physics [1] and studies of quantum gases [2]. Diatomic molecules offer many features that are absent in cold atomic gases, such as dipole moments and rich dynamics of vibration and rotation. These properties make them appealing for a wide range of experiments including measurements of fundamental symmetries [3], investigations of anisotropic quantum gases [4], and demonstrating the quantum mechanical nature of basic chemical processes [5].

To enable control of diatomic molecules at the quantum state level, they must be trapped and cooled to ultracold temperatures, generally well below a millikelvin. In atomic physics experiments, the workhorse of ultracold science is the magneto-optical trap (MOT) which confines and simultaneously cools the particles with magnetic field gradients and optical cycling using laser light. While MOTs work best with two-level systems, there exists a set of diatomic (and even larger) molecules where one can isolate a quasi-two-level system despite the rovibrational level complexity, and carry out laser cooling and trapping [6, 7]. These ideas have enjoyed considerable success in the past several years, resulting in molecular MOTs of strontium monofluoride (SrF) and calcium monofluoride (CaF) [8, 9] and a two-dimensional MOT of vttrium monoxide (YO) [10].

A number of diatomic hydrides also offer promising laser cooling schemes [6, 11] but remain under-explored despite their amenability to theoretical treatment. A cold and slow beam of CaH has been reported [12] and cold beams of LiH have been investigated [13], both based on laser ablation of solid targets, but suffered relatively low molecule yields compared to some of the diatomic fluorides and oxides [14]. In this work, we demonstrate a bright and slow beam of BaH molecules with good stability and sufficient flux for optical slowing and cooling. BaH molecules present the challenges of insufficient spectroscopic information and the usual difficulty of hydride ablation, as well as a large mass, relatively long wavelengths of the optical cycling transitions, and naturally low photon scattering rates, all three of which suppress

laser cooling efficiency. On the other hand, BaH opens the door to ultracold diatomic hydrides, presents a low Doppler cooling limit < 40  $\mu$ K, and offers an opportunity to test new molecular MOT schemes due to its unusual magnetic moments. Further, it has a very large mass ratio of the constituent atoms which, from kinetic considerations, could yield ultracold dilute hydrogen samples upon dissociation [11].

The optical transitions that can potentially support laser cooling are  $X^2\Sigma^+ \leftrightarrow A^2\Pi_{1/2}$  at 1061 nm and  $X^2\Sigma^+$  $\leftrightarrow B^2\Sigma^+$  at 905 nm. The former transition is analogous to the ones used in most other molecular laser cooling experiments [9, 10, 15], while the latter transition allows an alternative optical cycling scheme that has only been shown for slowing CaF molecules [9]. Typically, the  $X^2\Sigma^+ \leftrightarrow B^2\Sigma^+$  transition involves a significant hyperfine splitting in the excited state which makes it difficult to ensure appropriate cooling laser detunings from all hyperfine levels. The  $X^2\Sigma^+ \leftrightarrow A^2\Pi_{1/2}$  transition is thus more commonly used, but this excited state tends to have a very small magnetic moment, leading to weak magnetooptical trapping forces [16]. In BaH, both of these challenges are mitigated. As we show here, the magnetic moment of the  $A^2\Pi_{1/2}$  state is of a similar magnitude to that of the ground state, permitting strong trapping forces. In addition, we have measured a relatively large hyperfine splitting of the  $B^2\Sigma^+$  state, and this feature, combined with a narrow 1.6-MHz natural linewidth of this transition [17], might allow efficient magneto-optical trapping on the  $X^2\Sigma^+ \leftrightarrow B^2\Sigma^+$  transition as well.

In Sec. II of this work we discuss the details of the cryogenic molecular beam source. In Sec. III we describe the properties of the resulting molecular beam. Section IV presents hyperfine-structure-resolved optical spectroscopy of the cryogenic beam, while Sec. V shows spectroscopy with applied magnetic fields and measurements of relevant magnetic g-factors. In Sec. VI we summarize the results and present conclusions.

#### II. EXPERIMENTAL APPARATUS

The cryogenic molecular beam source of BaH is shown in Fig. 1(a). BaH molecules are produced via laser ablation of a solid BaH<sub>2</sub> target inside a copper cell held at  $\sim$  6 K. The cell is at the center of a cryostat cooled by a pulse tube refrigerator (PTR). Helium buffer gas flows into the cell and past the target at approximately 10 sccm via an inlet pipe at the back of the cell. The ablated molecules partially thermalize with the He before being swept out of a 7-mm diameter beam aperture. In order to maintain vacuum in the cryogenic source region ( $\sim 10^{-6}$  torr while He is flowing), the interior of the 4 K shield is coated with activated coconut charcoal (Calgon Carbon OLC  $12 \times 30$ ) which acts as a fast cryopump for He. The charcoal is attached by liberally coating the copper shields with thermal epoxy (Loctite Stycast 2850FT). The charcoal-coated panels are regenerated by warming up the source region to  $\sim 80$  K every 10 hours of active operation. In addition, the panels are weekly cycled to room temperature and occasionally baked to > 100 °C. The source region takes < 14 hours to cool down to the minimum cell temperature or to fully warm up to room temperature.

The buffer-gas cell, shown in Fig. 1(b), consists of a cylindrical cavity (1.25" long, 1" diameter) cut out of a solid copper block. Smaller cutouts with windows provide optical access for ablation as well as for absorption spectroscopy to monitor the molecule production process. A woven copper cloth  $(100 \times 100 \text{ mesh size})$  glued to the helium inlet separates the inlet from the interior of the cell. To prevent dust buildup on the ablation window, it is extruded by an extended narrow section. The target holder can screw in and out of the cell wall to facilitate quick and minimally invasive ablation target changes. Pulsed ablation of a typical target [18] yields approximately 1000 shots, after which the signal begins to deteriorate but can be recovered by manually moving to a different spot on the same target. The solid targets are produced by cleaving commercial BaH<sub>2</sub> rocks 1-3 mm across (Materion DP-532-12-1) to expose a smooth surface, and gluing 4-6 of them to the holder with enough superglue (Loctite 414) to wick up the sides of the rocks. The rocks are chosen for their flatness, and no significant difference in ablation yield was observed after annealing the rocks. The Nd:YAG ablation laser (BigSky Ultra) emits 8-ns 50-mJ pulses at 1064 nm with a 2 Hz rate and is focused to a 0.5 mm diameter spot on the target.

We have observed that the rotational temperature of BaH in the buffer-gas cell thermalizes to  $\sim 9$  K with a time constant of 0.3 ms for typical buffer gas flow rates. The extraction of BaH from the cell occurs between 0.5 and 2 ms after ablation, ensuring good rotational thermalization of the molecules. This equilibrated rotational temperature is optimal for maximizing population in the  $X^2\Sigma^+$  (v''=0,N''=1) state of interest for laser slowing and cooling. Furthermore, the relatively quick extraction times can be beneficial for laser slowing [19].

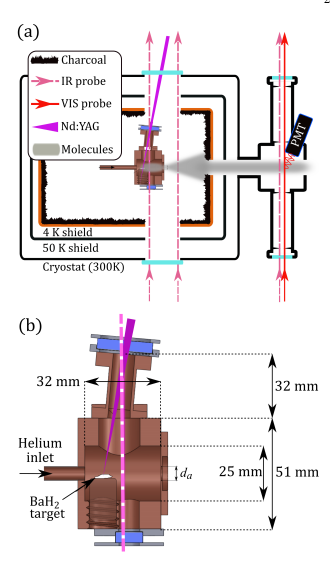


FIG. 1. a) Diagram of the buffer-gas-cooled beam source of BaH. The diatomic molecules are created through pulsed laser ablation of a BaH<sub>2</sub> rock inside a cryogenic copper cell filled with flowing He gas. As the molecules are swept out of the cell, they can be optically probed at various locations along the beam: in the cell and just after the cell via absorption, and in the downstream science region which is furnished with a PMT for fluorescence detection using visible light. Coconut charcoal coating on the inner shield as well as on additional copper fins (not shown) acts as a fast cryopump for excess He. The inner copper shield is nominally at 4 K and is surrounded by a 50 K aluminum shield. The entire assembly is enclosed in a vacuum-tight aluminum chamber with large optical windows on both sides. b) Details and dimensions of the cryogenic buffer-gas cell. The beam aperture is  $d_a=7$  mm.

Directly outside the buffer-gas cell, molecular beam extraction can be quantified via absorption spectroscopy. Further downstream, geometric apertures imposed on the molecular beam consist of a charcoal-coated plate with a 1-cm diameter hole located 11 cm from the cell exit, and again by a 3.8-cm diameter vacuum flange at the

output port of the cryostat 53 cm from the cell. Here the molecules enter a fluorescence detection region made by a six-way 2.75" ConFlat vacuum cube with a black coating (Acktar Metal Velvet) and blackened copper baffles to reduce light scattering [20]. This region is pumped by a 700 l/s turbomolecular pump (550 l/s for He), and the vacuum here is currently limited by the vacuum level in the cryopumped beam source to  $\sim 10^{-6}$  torr.

The optical cycling transitions for BaH lie in the near-infrared, which is an inconvenient wavelength range for fluorescence detection with photomultiplier tubes (PMTs). As an alternative, molecules are detected via a higher-energy transition  $X^2\Sigma^+ \leftarrow E^2\Pi_{1/2}$  with a 684 nm wavelength. The 8.6 GHz spin-rotation splitting of the N'' = 1 ground-state rotational level [18] cannot be easily addressed by a single laser, and without repumping and remixing of dark magnetic sublevels, the expected detection rate is about one photon per molecule even for quasi-cycling transitions. However, this is sufficient for precisely characterizing molecular beam parameters and spectroscopic properties. For measurements of hyperfine energy splittings and magnetic g-factors, probe lasers on the  $X^2\Sigma^+ \to A^2\Pi_{1/2}$  or  $X^2\Sigma^+ \to B^2\Sigma^+$  transitions interact with the molecular beam a few millimeters upstream from the 684-nm detection laser, allowing for resonant depletion or enhancement of the populations in the detected ground states.

### III. MOLECULAR BEAM

In the context of creating samples of trapped ultracold molecules, the most important molecular beam properties are the flux, forward velocity distribution, and transverse temperature.

The molecular flux downstream from the source is detected via fluorescence with a PMT as shown in Fig. 1(a). For characterizing the molecular beam we use a singlefrequency laser resonantly driving the N'' = 1, J'' = 1/2spin-rotation level of the electronic ground state to the N'=0, J'=1/2 level of the  $\mathrm{E}^2\Pi_{1/2}$  excited state. Assuming that the molecules are equally distributed in each hyperfine magnetic sublevel, we detect one third of the total molecules present in the N''=1 rotational state. In addition, the probe laser intersects slightly less than a tenth of the molecular beam cross-sectional area. The detection efficiency of the system is the product of the 2% PMT quantum efficiency at 684 nm and the 2.5% geometric collection efficiency of the detection optics. Each of these factors combined with the signal size of  $\sim 500$  PMT counts in the detection region per ablation pulse yields approximately  $4 \times 10^7$  molecules in the  $X^2\Sigma^+$ (v''=0,N''=1) state per pulse. Some of our ablation targets yield up to three times as many molecules.

To determine the forward velocity distribution of the molecules, we make time-resolved molecular density measurements 2 cm away from the cell exit via absorption spectroscopy and, simultaneously, downstream in

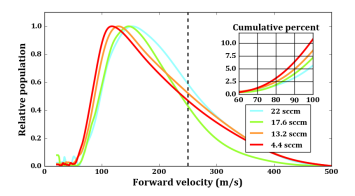


FIG. 2. Measured forward velocity distributions of BaH molecules at a range of buffer gas flow rates. Each distribution is normalized to its peak value. The total molecule number does not significantly vary in this range of flow rates. Velocity distributions above  $\sim 250$  m/s (dashed line) are slightly less reliable due to a moderate sensitivity on specific data cuts made to reject fluorescence noise from ablation light. The small-scale structure at low velocities is an artifact of deconvolution. The distributions show little variation with He flow rate for high velocities, but there is a clear enhancement of the slow molecule number for lower flow rates. Some variation is seen for different ablation samples or spots, as in the 17.6 sccm trace, indicating incomplete translational thermalization. The inset shows the percentages of molecules below a given velocity under 100 m/s.

the fluorescence detection region, using the  $X^2\Sigma^+$  (v''=0, N'' = 1) ground state. These two measurements of beam density as a function of time are deconvolved to yield the forward velocity distribution in a process analogous to spatial time-of-flight analysis in quantum gas experiments. The results of this analysis are given in Fig. 2 for various He flow rates from 4.4 to 22 sccm. While the overall number of detected molecules per ablations pulse does not significantly vary in this range of flow rates, Fig. 2 and its inset show that the number of slow molecules with forward velocities below 100 m/s doubles as the He flow is reduced from 22 to 4.4 sccm, reaching a fraction of  $\sim 11\%$ . At even lower flow rates the molecular flux begins to degrade. It is evident from the data in Fig. 2 that while the low-velocity behavior is stable and systematic in its dependence on the flow rate, the overall shape of the velocity distribution can be affected by the particular choice of target or ablation spot, as is the case for the 17 sccm trace. The velocity distributions above  $\sim 250$  m/s are slightly less reliable than those for lower velocities since this signal comes from faster molecules arriving in the detection region very shortly after ablation, while there is a minimum waiting time after ablation to begin detecting in order to avoid the stray fluorescence. The relatively broad and asymmetric nature of the velocity distributions is the result of incomplete thermalization of ablated BaH with cryogenic He. The present configuration of the beam source was chosen to produce a maximized number of molecules with velocities below

100 m/s, but could be further optimized by making additional changes to the buffer-gas cell geometry.

In the fluorescence detection region we can limit the transverse temperature of the molecules to 0.1 K by comparing the expected natural linewidth of the  $B^2\Sigma^+$  state [17] to the measured spectra. This cold transverse temperature is consistent with geometric constraints on the beam and allows us to characterize the relevant properties of BaH at a higher optical resolution than was previously possible, enabling direct measurements of hyperfine structure and molecular g-factors.

# IV. HYPERFINE STRUCTURE MEASUREMENTS

To explore the possibility of optical radiation pressure experiments with BaH, it is necessary to understand the ground and excited state hyperfine structure. While BaH has been studied intermittently for over 100 years, it is less well explored than other molecular candidates for laser cooling. The cold molecular beam described here is well suited for optical spectroscopy that reveals hyperfine structure of the ground and excited states.

The transverse Doppler width of the molecular beam in the fluorescence detection region, in combination with our signal-to-noise ratio, allows us to resolve hyperfine energy splittings at the  $\sim 4$  MHz level. It is challenging to directly detect fluorescence from the  $A^2\Pi_{1/2}$  or  $B^2\Sigma^+$  excited states due to the PMT wavelength limitations. However, these states can be probed by optically pumping molecules between the two ground-state spin-rotation levels (J''=3/2 and J''=1/2) while detecting population in one of them via the  $X^2\Sigma^+\leftarrow E^2\Pi_{1/2}$  transition. This is possible due to the highly favorable vibrational branching ratios of the  $A^2\Pi_{1/2}$  and  $B^2\Sigma^+$  excited states into the ground state [11, 18].

TABLE I. Measured hyperfine intervals for the electronic states of BaH that are relevant to laser cooling. Negative values denote 'flipped' hyperfine structure.

State	Measured hyperfine spacing (MHz)	
$X^2 \Sigma^+ (J'' = 1/2)$	-0(4)	
$X^2 \Sigma^+ (J'' = 3/2)$	39(4)	
$B^2 \Sigma^+ (J' = 1/2)$	-52(5)	
$A^2\Pi_{1/2}(J'=1/2)$	0(4)	

Hyperfine structure measurements are outlined in Fig. 3 and the results are reported in Table I. Four types of optical spectra were collected to fix the values of the four unknown hyperfine intervals. These are labeled A, B, C, and D in Fig. 3. In each of these experiments, the  $X^2\Sigma^+(J''=1/2)$  population is monitored by sending a 683.7268 nm laser beam through the detection region and recording fluorescence from the spontaneous decay of the  $E^2\Pi_{1/2}$  state. Several beam waists upstream, a

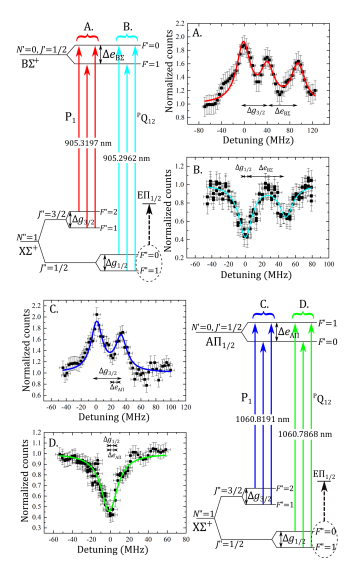


FIG. 3. Hyperfine-structure-resolved energy levels and measurements for BaH electronic states relevant to laser cooling. Four types of spectra are collected (labeled A, B, C, and D). In all cases, detection is made by monitoring the fluorescence of molecules excited from the  $X^2\Sigma^+(N''=1,J''=1/2)$  ground state to the  $E^2\Pi_{1/2}$  excited state, indicated by the dashed arrows. Transitions labeled A and C show population enhancement due to pumping from the  $X^2\Sigma^+(N''=1,J''=3/2)$  state via  $B^2\Sigma^+$  or  $A^2\Pi_{1/2}$ . Transitions labeled B and D show population depletion due to pumping out of the  $X^2\Sigma^+(N''=1,J''=1/2)$  state. The studied ground-state and excited-state hyperfine intervals are marked as  $\Delta g$  or  $\Delta e$  in the diagrams and in the spectra.

probe laser drives transitions from one of the ground electronic spin-rotation levels through the  $\mathrm{B}^2\Sigma^+$  or  $\mathrm{A}^2\Pi_{1/2}$  excited state. The A and C transitions (905.3197 nm and 1060.8191 nm) enhance the detected fluorescence as the molecules get pumped from the J''=3/2 ground-state level to J''=1/2, while the B and D transitions (905.2962 nm and 1060.7868 nm) decrease the fluores-

cence as the molecules are pumped out of the J''=1/2 level. Each data point in the spectra of Fig. 3 is an average of 5 ablation shots. The peaks and their spacings in the spectra can be identified by assuming an ordering for the excited-state hyperfine structure and fitting the peak positions. The relative peak amplitudes are given by the hyperfine level degeneracies. A correctly chosen ordering yields peak spacings and heights that are consistent for all four data sets in Fig. 3. The shown fits to the fluorescence spectra are constrained only by the expected peak height ratios.

The hyperfine structure data can be analyzed to extract molecular hyperfine constants which have been previously measured for ground-state BaH in cryogenic solid argon [21]. Excluding negligible terms, the hyperfine structure of  $X^2\Sigma^+$  is described by the Hamiltonian

$$H_{\rm hf} = b_F \mathbf{S} \cdot \mathbf{I} + cI_z S_z \tag{1}$$

where  $b_F$  is the Fermi contact interaction constant, c is the dipolar coupling constant, and  $\mathbf{S}$  and  $\mathbf{I}$  are the electronic and nuclear spin angular momenta. Each term in Eq. (1) can be evaluated as in Sec. 9.5 of Ref. [22]. The resulting hyperfine interaction matrix elements for  ${}^2\Sigma^+$  states are

$$H_{\rm hf} = \begin{pmatrix} \frac{b_F}{4} + \frac{c}{20} & 0 & 0 & 0\\ 0 & -\frac{b_F 5}{12} - \frac{c}{12} & \frac{b_F \sqrt{2}}{3} + \frac{c\sqrt{2}}{6} & 0\\ 0 & \frac{b_F \sqrt{2}}{3} + \frac{c\sqrt{2}}{6} & -\frac{b_F}{12} + \frac{c}{12} & 0\\ 0 & 0 & 0 & \frac{b_F}{4} - \frac{c}{4} \end{pmatrix}.$$

Our hyperfine structure measurements yield  $b_F = 50(7)$  MHz and c = 39(8) MHz. The value for  $b_F$  is consistent with previous measurements of 47(2) MHz [21]. The value for c, while lacking previous reliable measurements, is consistent with those for other alkaline-earthmetal monohydrides [21].

The hyperfine structure results in Table I can guide experiments on radiation-pressure slowing and cooling of BaH. For the lower spin-rotation level of the ground state (J''=1/2) hyperfine structure is unresolved, while for the higher level (J'' = 3/2) the splitting is 39(4) MHz and can be easily covered by sidebands imprinted on the laser light with standard acousto-optical techniques. Hyperfine structure is also small, or on the order of the natural linewidth, in the  $A^2\Pi_{1/2}$  excited state as is the case for other diatomic molecules that have been investigated as laser cooling candidates. This feature allows all excited-state sublevels to participate in optical cycling, thus maximizing radiation pressure forces. The 52(5) MHz hyperfine interval in the  $B^2\Sigma^+$  excited state is  $\sim 30$  times larger than the natural linewidth and is of a similar magnitude to that of the ground-state J'' = 3/2level, such that the combination of the two can be managed with acousto-optical techniques.

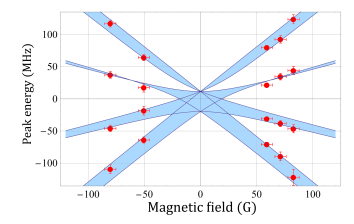


FIG. 4. Measured Zeeman shifts of the  $X^2\Sigma^+(J''=3/2)$  magnetic sublevels, overlayed with a prediction from the Zeeman Hamiltonian. Solid lines represent energies of the  $m_F$  sublevels. Shaded areas between pairs of sublevels emphasize structure that is spectroscopically unresolved even if selection rules allow both sublevels to couple to the excited state.

### V. MAGNETIC g-FACTOR MEASUREMENTS

Magnetic g-factors are crucial for understanding magneto-optical trapping forces on molecules. In particular, the trapping forces depend strongly on the ratios of the g-factors in the ground and excited states [16]. Here we report predictions and measurements of the relevant magnetic g-factors in BaH.

The predictions can be made by diagonalizing the Zeeman interaction Hamiltonian for each molecular state. For  $^2\Sigma^+$  states, the electronic and nuclear Zeeman Hamiltonians are expressed in Eqs. (8.183) and (8.185) of Ref. [22], and involve 12 hyperfine-structure magnetic sublevels  $m_F$  for the  $X^2\Sigma^+$  state and 4 sublevels for the  $B^2\Sigma^+$  state of BaH.

The Zeeman shifts for <sup>2</sup>Π states are strongly influenced by the parity dependent contributions  $g'_l \approx \frac{p}{2B}$  and  $g_r^{e'} \approx \frac{q}{R}$ , where p and q are the  $\Lambda$ -doubling constants and B is the rotational constant. These constants have been measured in BaH for the  $A^2\Pi_{1/2}$  excited state [23] and for the detection state  $E^2\Pi_{1/2}$  [24]. The matrix elements of the applicable Zeeman Hamiltonian are expressed in Eq. (9.71) of Ref. [22], and the dominant parity dependent term can be described with a single effective g-factor as  $g_{\text{eff}} = (g_l' - g_r^{e'})/3$ . This contribution alone would result in a g-factor value of -0.27 for  $A^2\Pi_{1/2}$ , while a purely semiclassical prediction would yield a value of 0. The predicted q-factor, however, is -0.44 in this case, because additional contributions to  $g_{\text{eff}}$  result from interactions between electronic states. In BaH, the lowest 5d excited states form an interacting complex [25] where the strongest mixing is between  $B^2\Sigma^+$  and  $A^2\Pi_{1/2}$ . This results in an enhancement of the  $A^2\Pi_{1/2}$  state g-factor and a slight reduction of the  $B^2\Sigma^+$  state g-factor.

While at very low magnetic fields it is natural to

TABLE II. Measured and predicted effective g-factors in the  $m_J$  basis which is most pertinent to the field regimes used in magneto-optical trapping.

State	Measured $g_{\text{eff}}$	Prediction
$X^{2}\Sigma^{+}(N''=1,J''=1/2)$	-1.37(10)	-1.4
$X^{2}\Sigma^{+}(N''=1,J''=3/2)$	+0.56(19)	+0.50
$B^2 \Sigma^+ (N' = 0, J' = 1/2)$	+2.76(6)	+2.86
$A^{2}\Pi_{1/2}(N'=0, J'=1/2)$	-0.56(1)	-0.44
$E^2\Pi_{1/2}(N'=0, J'=1/2)$	-0.16(10)	-0.04

use the  $m_F$  basis for the Zeeman interaction Hamiltonian, at fields exceeding  $\sim 10$  G the Zeeman shifts are best described in the  $m_J$  basis. As a result, the reported factors  $g_{\text{eff}}$  describe the measured energy shifts for magnetic field strengths of tens of gauss. These shifts are  $\Delta E = g_{\text{eff}} \mu_B m_J B$ , where B is the applied field and  $\mu_B$  is the Bohr magneton. High-resolution Zeeman spectra were collected for both excited states, for both spin-rotation levels of the ground state, and for the  $\mathrm{E}^2\Pi_{1/2}(N'=0,J'=1/2)$  state used in the detection scheme. Figure 4 shows data for Zeeman shifts of the  $X^2\Sigma^+(J''=3/2)$  ground state sublevels, along with the prediction from the Zeeman Hamiltonian. The experimental results together with the predictions are listed in Table II. For all experiments, magnetic field was applied perpendicularly to the probe laser propagation direction and calibrated in situ with a commercial gaussmeter. Measurements of the ground-state Zeeman shifts were done via the  $X^2\Sigma^+ \leftarrow E^2\Pi_{1/2}$  transition, where we could separately identify the ground and excited state splittings, as well as the relative signs of their g-factors, by switching the probe laser polarization between  $\pi$  and  $\sigma^{\pm}$ transitions. The Zeeman shifts in the  $B^2\Sigma^+$  and  $A^2\Pi_{1/2}$ excited states were measured via the fluorescence depletion method as in the hyperfine structure studies, and polarization was again switched to drive  $\pi$  and  $\sigma^{\pm}$  transitions in order to determine the relative signs of all the g-factors. To fix the absolute signs, a calibration measurement was made by applying a magnetic field along the laser propagation axis and using circularly polarized light.

The results in Table II highlight an interesting difference in laser cooling prospects between BaH and other diatomic molecules currently in use such as SrF [8] and CaF [9]. Unlike the fluorides, BaH has a large magnetic moment in both excited states that could be used for optical cycling,  $B^2\Sigma^+$  and  $A^2\Pi_{1/2}$ . This should allow several approaches to magneto-optical trapping using simpler MOT schemes than what is needed for the fluorides [16]. The excited-state g-factors with magnitudes reported in Table II can lead to trapping forces  $\sim 5$  times stronger than near-zero magnetic moments [16, 26].

# VI. CONCLUSIONS

We have built and characterized a molecular beam of BaH that is cooled by cryogenic He buffer gas and delivers packets of molecules to a downstream interaction region at a rate of a few hertz. Approximately  $4 \times 10^6$ molecules in each packet are transversely cold (0.1 K) and have forward velocities below 100 m/s. Various improvements to the setup could potentially boost this number further, particularly by squeezing the high-velocity tail of the forward velocity distribution. Some of the projected improvements include achieving a higher level of vacuum in the beam region, and systematically varying the buffer-gas cell parameters such as its inner diameter as well as its aperture diameter and shape, while tracking the velocity distribution. Additionally, it is possible that producing sintered BaH<sub>2</sub> pellets or other types of targets could allow us to reduce the intensity of the ablation laser, which in turn might lead to more complete thermalization of BaH with the buffer gas.

The molecule numbers and velocities that were achieved here are a starting point for experiments in which radiation pressure forces are applied to BaH for laser slowing, cooling, and trapping. The critical spectroscopic parameters were determined in this work both theoretically and experimentally, since the cold molecular beam allows optical spectroscopy with a resolution of a few megahertz. Hyperfine structure was measured in the ground electronic state and in both excited states that could be used for optical cycling. Modest magnetic fields were applied to the molecules to measure all relevant magnetic moments of the ground and excited electronic states. The resulting effective g-factors point to several promising laser cooling and trapping schemes that could result in relatively large trapping forces.

As the experimental field of ultracold molecules gains momentum, it is important to attempt laser-cooling experiments with different types of molecules, and especially with under-explored species such as diatomic hydrides. BaH is particularly appealing because of its low Doppler cooling limit, good optical cycling properties [11, 18], unusual excited-state magnetic moments as was shown here, and a large mass ratio between its atomic constituents [11]. The results of this work are an advance toward these goals.

### ACKNOWLEDGMENTS

We thank M. G. Tarallo and L. Abbih for their contributions to this work, and S. Truppe and M. R. Tarbutt for helpful discussions. We acknowledge partial support by the ONR grant N00014-16-1-2224. R.L.M. and G.Z.I. acknowledge support by the NSF IGERT Grant DGE-1069240.

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