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In-source laser spectroscopy of dysprosium isotopes at the ISOLDE-RILIS

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ABSTRACT

A number of radiogenically produced dysprosium isotopes have been studied by in-source laser spectroscopy at ISOLDE using the Resonance Ionization Laser Ion Source (RILIS). Isotope shifts were measured relative to 152Dy in the 4f 154s 2I 1/2 (gs) → 4f 1566s6p (8, 1) 3/2 (418.8 nm, vac) resonance transition. The electronic factor, F, and mass shift factor, M, were extracted and used for determining the changes in mean-squared charge radii for 146Dy and 148Dy for the first time.

1. Introduction

The Resonance Ionization Laser Ion Source (RILIS) is the most selective of all ion sources available at the ISOLDE radioactive beam facility [1]. The selectivity is an intrinsic property of the ionization mechanism, based on stepwise resonance excitation and ionization via element-specific atomic levels. The isotope production takes place inside a thick target, on which protons, provided by CERN’s Proton Synchrotron Booster (PSB), impinge with an energy of 1.4 GeV. The reaction products are released from the target material and effuse via a transfer line into a resistively heated tubular cavity, where the atom-laser interaction takes place. The resulting ions are then extracted, accelerated up to 60 keV and mass separated by a dipole magnet according to their mass-to-charge ratio.

During so-called ‘in-source laser spectroscopy’, the RILIS lasers are used to probe a specific spectroscopic transition of the ionization scheme of different isotopes of one element. By determining the isotope shift (IS) of a chosen transition, changes in the nuclear mean-squared charge radii can be deduced. For states with nonzero nuclear spin I which exhibit a sufficiently large hyperfine structure (HFS), the nuclear moments (spin, magnetic dipole and electric quadrupole moments) can be extracted. Additionally, if the HFS of different isomers can be resolved (due to different spins and magnetic moments), isomer-selective ionization is possible. The spectral resolution of in-source measurements is limited by Doppler-broadening of the spectral lines inside the ion source (which is typically heated to ≈ 2100 °C). There have been several experimental campaigns, in which this in-source spectroscopy has been successfully applied (e.g. [2]) or where isomer separation was provided for higher selectivity during nuclear spectroscopy experiments (e.g. [3]).

Here, we report on the first in-source spectroscopy study of dysprosium radioisotopes, demonstrating the suitability of this method for a future extended study of IS in the dysprosium isotopic chain.

2. Experimental setup

The experiment was performed using beam provided by target #655 (target with tantalum rolls from mixed 25 and 6 μm foils at 1950 °C with a tungsten surface ion source at 1985 °C). No stable supply of dysprosium was available initially, so that the optimization was performed on radiogenically produced 152Dy. During the experiment, a proton current of 0.2 μA was used on target, providing a continuous supply of dysprosium.

The transition chosen for the spectroscopy leads from the [Xe]4f 154s 2I 1/2 ground state to the 4f 1566s6p (8, 1) 3/2 excited state at 23877.74 cm⁻¹ (≈ 418.8 nm) [4] (note: wavelengths given for vacuum).
A second laser, a non-tunable Nd:YVO4 laser (2nd harmonic, 532 nm), results in efficient ionization of Dy from the 4f106s6p, j = 8 excited level, despite the photon energy at 532 nm being below that required to reach the ionization continuum. The ionization efficiency saturates with an estimated 7 W of laser power in the ionization region (≈ 3 mm laser beam diameter). From this we conclude that the 532 nm light is coincidentally resonant with a second step transition to a high-lying level, from which a second 532 nm photon induces ionization via an auto-ionizing state. In the transition metals, the atomic level density, and the richness of the auto-ionizing spectrum, greatly increase the likelihood of such a coincidence in required transition wavelengths.

A newly developed narrow-linewidth intra-cavity frequency-doubled mode for the Ti:sapphire grating laser was applied for the first time, scanning across the 418.8 nm transition. It will be described in more detail in [5]. The wavelength was recorded with two High-Finesse/Angstrom WS7 wavelength meters installed in the RILIS laboratory. The wavemeters were calibrated before the measurements with a CW diode laser locked to the rubidium hyperfine structure. As the transition probability lies at A = 1.26 × 10^8 s^1 [6], the power of the first-step laser beam had to be reduced significantly, to < 1 mW in order to avoid saturation.

For the cases of 148,149,151,153,155 Dy, the ISOLDE Faraday cups were used for ion beam detection, as the resonant ion rates were sufficiently high (> 1 pA). For 145,147 Dy the ISOLDE tape station gamma detector was utilized (for more details see [7]). An overview over the yields measured with the tape station β-counting is given in Table 1.

### 3. Results

In the case of 147Dy, a long-lived isomeric state (T1/2 = 11/2+ with T1/2 = 55 s) exists in addition to the ground state (Ig = 1/2+ with T1/2 = 67 s). When monitoring the number of photoions in dependence of the first-step laser frequency by the intensity of the internal nuclear transition (678 keV, [12]) one obtains the optical spectrum of the pure high-spin isomer. Several γ-lines following the β-decay of 145Dy and 149Dy were also present in the collected γ-spectra. Corresponding optical spectra are the mixture of that for metastable and ground state. The yield of 147Dy is estimated to be ~ 5 times larger than the yield of 149Dy, following analysis of the γ-line intensities. As a result, it is impossible to estimate the IS of the ground state with reasonable accuracy.

For the case of 145Dy (Ig = 11/2+ with T1/2 = 14 s and I1/2 = 1/2+ with T1/2 = 6/3), only the γ-line resulting from the 145Dy β-decay at 639 keV was observed. The missing observation of other lines is attributed to the high background from the β-decay of the surface ionized isotobars. Correspondingly, only results for the isomeric state were obtained.

The optical spectra are summarized in Fig. 1 and the IS are summarized in Table 2. Due to high background levels, stemming from the surface ionization of dysprosium and other isobaric (lanthanide) contaminants, the uncertainties of the IS are rather large (200-400 MHz). The signal-to-background ratio in case of e.g. 148Dy is only 0.014. The same is true for the γ-spectra, where close lying, more intense γ-lines from isobaric contaminants dominate.

For extracting variations of the mean-squared charge radii Δ(r²), results of the IS measurements with the 421.3 nm transitions from [13]...
were used for comparison. A ‘standard’ King-plot procedure (see e.g. [11]) is not possible, due to missing IS data for the light dysprosium isotopes (only $\delta(r^2)$ are cited in [13,11]). A modified approach was used. Starting with the well-known relation that

$$\delta_{\text{IS},A_0} = F \cdot \delta(r^2)_{A_0} + M \cdot \frac{A - A_0}{A_0},$$

it follows that the modified IS

$$\sigma_v = \delta_{\text{IS},A_0} \cdot \frac{A - A_0}{A_0}$$

is linearly dependent on the modified $\delta(r^2)$

$$\sigma_v = \delta(r^2)_{A_0} \cdot \frac{A - A_0}{A_0},$$

with the slope equal to the electronic factor $F$ and the intercept equal to the mass-shift constant $M$:

$$\sigma_v = F \cdot \sigma_e + M.$$  

The nuclear masses $A$ and $A_0$ used in the calculations by Eqs. (1)–(3) were taken from [14].

As shown in Fig. 2, all newly measured modified IS for the 418.8 nm transition, as well as the previously measured $\delta_{\text{IS},A_0}$ for $A = 164, 160$ [15] over modified $\delta(r^2)$ lie on a straight line, testifying to the consistency of the newly obtained data. From this plot, the electronic factor $F$ and mass-shift factor $M$ were determined to be $f^{418.8 \text{nm}} = -3580(110) \text{ MHz fm}^{-2}$ and $N_{418.8 \text{nm}} = -60(360) \text{ GHz amu}$ (note, that the uncertainty of the $F$ and $M$ factors for the previously studied 421-nm transition are not taken into account. These values were cited in [11] without uncertainties). With the derived $F$ and $M$ factors, changes in $\delta(r^2)$ for the high-spin isomers in $^{147,148}$Dy were derived for the first time (see Table 2).

Fig. 3 shows the newly obtained values for $\delta(r^2)_{418.8,148}$ and $\delta(r^2)_{418.8,147}$ together with the data taken from [13] for $N = 78, 83 - 86$ relative to $^{148}$Dy. The shell effect in the $\delta(r^2)$ (kink at $N = 82$) is evident for odd- and even-$N$ isotopes. It was found previously that there is a marked isomer shift between 1/2+ ground states and 11/2− isomers in $\alpha$Sm and $\alpha$Gd nuclei at $N < 82$ [16,17]. This isomer shift leads to the disappearance of the odd-even staggering (OES) in $\delta(r^2)$ of the 11/2− isomers. The results obtained in the present work for 11/2− dysprosium isotopes do not contradict this observation, although no definite conclusion can be inferred due to the large experimental uncertainties.

4. Outlook

In order to better investigate the disappearance of normal OES in the vicinity of $N = 82$ for the high-spin isomers in dysprosium, expected to be influenced by the $\nu_{1/2}$ state, further studies with dedicated beam time are necessary. A better resolution of the $\gamma$-spectra and detection efficiency, using the ISOLDE Decay Station (IDS), would help to separate the ground from isomeric state. The relative uncertainties could be additionally reduced by using the transition to the $\gamma^{148}$Gd, $J = 9$ state at 23736.61 cm$^{-1}$ (≈ 421.3 nm). This transition has been shown to have an isotope-shift sensitivity twice the size of the 418.8 nm transition [11]. As seen in Table 2 and Fig. 1, at the signal to background ratio larger than ~1 and sufficient statistics the uncertainty of the IS determination can be reduced to 100 MHz ($^{148}$Dy) and lower (taking into account reduction of the uncertainty also for CoG measurement for the reference isotope). This accuracy is expected to be sufficient to investigate the evolution of OES (see results for similar 11/2-state in Sm [16]). However, more accurate results may be achieved with better resolution which would enable reliable analysis of the odd Dy isotope HFS.

It is estimated that dysprosium isotopes down to around $A = 141$ are accessible for IS measurements by the in-source spectroscopy method, provided sufficient suppression of isobaric background is achieved (e.g. with the Laser Ion Source and Trap (LIST) [18]). It is worth to note that dysprosium isotopes with $A < 146$ have noticeable decay rate branching and proton current monitoring by delayed proton detection might give more favorable background conditions. Near this point, a strong onset of deformation is expected which would be reflected in the IS values.
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