Spectroscopy at the High-Energy Electron Beam Ion Trap (SuperEBIT)

Klaus Widmann
Peter Beiersdorfer
Jose R. Crespo López-Urrutia
Steve Elliott

Prepared for Submittal to
Proceedings of the 2nd Euroconference on Atomic Physics
with Stored Highly Charged Ions
June 2-7, 1996, Stockholm, Sweden

July 10, 1996

This is a preprint of a paper intended for publication in a journal or proceedings. Since changes may be made before publication, this preprint is made available with the understanding that it will not be cited or reproduced without the permission of the author.

DISTRIBUTION OF THIS DOCUMENT IS UNLIMITED

CONF-9606261-1
UCRL-JC-124408

RECEIVED
AUG 16 1996
OSTI

LAWRENCE
LIVERMORE
NATIONAL
LABORATORY

MASTER
DISCLAIMER

This document was prepared as an account of work sponsored by an agency of the United States Government. Neither the United States Government nor the University of California nor any of their employees, makes any warranty, express or implied, or assumes any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial products, process, or service by trade name, trademark, manufacturer, or otherwise, does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States Government or the University of California. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States Government or the University of California, and shall not be used for advertising or product endorsement purposes.
DISCLAIMER

Portions of this document may be illegible in electronic image products. Images are produced from the best available original document.
Spectroscopy at the High-Energy Electron Beam Ion Trap (SuperEBIT)

KLAUS WIDMANN¹,², PETER BEIERSDORFER¹, JOSÉ R. CRESPO LÓPEZ URRUTIA¹, AND STEVE R. ELLIOTT³

¹Lawrence Livermore National Laboratory, P.O.Box 808, Livermore CA 94551, USA
²Institut für Experimentalphysik, Technische Universität Graz, A-8010 Graz, AUSTRIA
³Department of Physics, University of Washington, P.O.Box 351560, Seattle, WA 98195-1560, USA

The following progress report presents some of the x-ray measurements performed during the last year on the Livermore SuperEBIT facility. The measurements include: direct observation of the spontaneous emission of the hyperfine transition in ground state hydrogenlike holmium, $^{165}$Ho$^{66+}$; measurements of the $n=2 \rightarrow 2$ transition energies in neonlike thorium, Th$^{80+}$, through lithiumlike thorium, Th$^{87+}$, testing the predictions of quantum electrodynamical contributions in high-Z ions up to the 0.4% level; measurements of the isotope shift of the $n=2 \rightarrow 2$ transition energies between lithiumlike through carbonlike uranium, $^{233}$U$^{89+...86+}$ and $^{238}$U$^{89+...86+}$, inferring the variation of the mean-square nuclear charge radius; and high-resolution measurements of the Kα radiation of heliumlike xenon, Xe$^{52+}$, using a transmission-type crystal spectrometer, resolving for the first time the $1s2p^3P_1 \rightarrow 1s^2S_0$ and $1s2s^3S_1 \rightarrow 1s^2S_0$ transitions individually.
1 Introduction

The electron beam ion trap (EBIT) facility at the Lawrence Livermore National Laboratory was especially designed for the study of atomic processes of highly charged ions[1]. The high-energy version, dubbed SuperEBIT, operates with an electron beam energy of up to 215 keV. Thus, SuperEBIT can produce and trap highly charged ions up to bare uranium, $^{92+}\text{U}$. The ions in the trap are relatively cold, just a few eV per charge[3], and are essentially at rest, which makes SuperEBIT an ideal source for spectroscopic diagnostic of few electron high-Z ions. A detailed description of SuperEBIT may be found elsewhere[4].

The SuperEBIT facility provides different options for injecting ions or neutrals into the trap region. One of the techniques to inject ions into SuperEBIT is based on a metal vapor vacuum arc (MeVVA)[5]. Depending on the operation mode, the MeVVA plasma contains atoms and ions from either the cathode or the trigger material. The ions are extracted from this plasma and focused into the trap region. A method to introduce neutrals into SuperEBIT is a gas injector. The pressure in the gas injector can be adjusted between $10^{-9}$ to $10^{-6}$ Torr, which is at least two to five orders of magnitude higher than the pressure in the trap itself. Precise adjustment of the gas injector ensures that the atoms or molecules intersect the electron beam, where they get ionized and trapped. A continuous injection of low-Z elements, such as nitrogen or neon, is very important for the evaporative cooling of the collisionally heated highly charged ions[6]. Recently, a technique has been developed which allows to inject materials which are available only in trace quantities[7]. This technique works by plating a small amount of the desired material on the pointed end of a wire probe. The wire probe is positioned near the compressed electron beam and material is removed from the sample by sputtering caused by trapped ions surrounding the electron beam. It enables to run experiments on SuperEBIT with only nanograms of feed material.

The ionization balance in SuperEBIT is optimized by choosing an appropriate elec-
tron beam energy, trap depth, cooling gas pressure, and the time during which the ions are kept in the trap before they are dumped and the trap is filled with "new" ions. The emission of the trapped ions is observed perpendicular to the electron beam. The confinement of the ions, along a 2–cm–long region within the 70–μm–diameter electron beam, makes SuperEBIT a perfect line source, which can be imaged by a spectrometer without applying an additional entrance slit.

During the last year the main research goal of the EBIT spectroscopy group was to extend the spectral range of our high–resolution measurements. With our flat–crystal low–energy and von Hámos–type medium–energy crystal spectrometer we have performed high–resolution measurements between 650 eV and 13000 eV[8, 9, 10, 11]. Adding a prism spectrograph and a transmission–type crystal spectrometer to the current EBIT instrumentation made the visible and near UV range, namely 1.5 eV to 6 eV, and the hard x–ray regime, beyond 13 keV, accessible to high–resolution spectroscopy.

In the following, we review our recent measurements in ascending order regarding the energy of the observed photons. In Section 2 we describe the direct observation of the hyperfine transition in hydrogenlike holmium, $^{165}$Ho$^{66+}$, which is situated in the visible range, i.e. around 2 eV. Section 3 contains investigations of 4–keV x rays and is divided into two parts: The first part presents the precision measurements of the n= 2 → 2 transition energies in neonlike thorium, Th$^{89+}$, through lithiumlike thorium, Th$^{87+}$, and a detailed comparison between the experimental values and theoretical predictions is given for berylliumlike thorium, Th$^{86+}$. The second part gives a short description of isotope shift measurements of the n= 2 → 2 transition energies comparing lithiumlike through carbonlike $^{233}$U and $^{238}$U. High–resolution measurements of hard x–rays, around 30 keV, are the topic of Section 4. In particular, it describes the implementation of a transmission–type crystal spectrometer at SuperEBIT and shows measurements of some n = 2 → 1 transitions of heliumlike xenon, Xe$^{52+}$. 
Visible regime: Direct observation of the hyperfine transition of hydrogenlike holmium, $^{165}\text{Ho}^{66+}$

Using a prism spectrograph we measured the $F=4$ to $F=3$ hyperfine transition of the $1s$ ground level of hydrogenlike $^{165}\text{Ho}^{66+}$ [12]. The experimental setup is shown in Fig. 1. It consists of a spectrograph with an aperture of f/9, and a cryogenically cooled CCD camera (1024×1024 pixels of 24μm side length). The spectral range covered in a single image was from 4175 Å to 6900 Å. The beam is stigmatically imaged by two achromats and, thus, spatially resolved in the vertical direction (0.1 mm resolution) allowing to distinguish between line radiation emitted by cooling gas atoms and by trapped ions. Radiation from the cooling gas is only emitted from the region where the atomic or molecular beam from the gas injector crosses the electron beam, whereas emission from trapped ions can be observed along the whole height of the trap. The spatial resolution is, therefore, important to determine the nature of the light emitting species. The ability to distinguish between atomic and ionic emission lines was also useful for the wavelength calibration of the optical spectra. In particular, we used lines emitted from neutral neon to calibrate the wavelength scale of our spectra.

A MeVVA with a holmium cathode was used to inject holmium ions into SuperEBIT. Measurements with and without injecting holmium were done consecutively throughout the whole experiment in order to eliminate the background, e.g. thermal radiation from the heated electron–gun filament, as well as emission from cooling–gas atoms. To determine the ionic species and charge balance in SuperEBIT, we monitored the radiative recombination (RR) lines, produced when electrons from the beam are captured into open shells of the ions, with a high–purity germanium detector (see Fig. 1). A RR–spectrum with a holmium–filled trap, at an electron–beam energy of $\approx 133$ keV and an electron–beam current of 285 mA, is shown in Fig. 2 (a). In this case the holmium charge balance was: 0.5% bare $\text{Ho}^{67+}$, 6% hydrogenlike $\text{Ho}^{66+}$, 40% heliumlike $\text{Ho}^{65+}$, 25% lithiumlike $\text{Ho}^{64+}$, and 28.5% lower charge states.
The result of the measurements are shown in Fig. 2 (b), which displays two background-corrected spectra. The first was obtained when the trap was filled with hydrogenlike holmium, $\text{Ho}^{66+}$, and clearly shows a single feature at $5726.4 \pm 1.5 \text{ Å}$. The second spectrum was obtained by repeating the experiment with a much lower electron-beam energy, namely 63 keV, to avoid production of $\text{Ho}^{66+}$ but still keeping a similar abundance regarding the heliumlike and lower charge states. The fact that this spectrum does not show any indication of any line proves that hydrogenlike holmium accounts for the observed line. Our experimental result for the $F=4$ to $F=3$ hyperfine transition of the 1s ground level of hydrogenlike $^{165}\text{Ho}^{66+}$ is $5726.4 \pm 1.5 \text{ Å (air)}$, which corresponds to a vacuum wavelength of $5727.9 \pm 1.5 \text{ Å}$, and a transition energy of $2.1646(6) \text{ eV}$, respectively[12].

Figure 3 shows a comparison between our result and the values predicted by theory. Using Shabaev's formula[13] for the energy difference between two neighboring levels in a hydrogenlike ion $2.199 \text{ eV}$ can be derived. This value has an uncertainty of at least $15 \text{ meV}$ due to the uncertainty in the value for nuclear magnetic moment of $^{165}\text{Ho}$, $\mu_I = 4.173(27)\mu_N$, tabulated by Lederer and Shirley[14], and omits QED corrections. Our measured value significantly disagrees with this prediction. Based on recently published calculations regarding the two dominant QED corrections[15, 16], we estimate for Z=67 a $9.4\text{-meV}$ contribution due to the vacuum polarization[15] and a $-19.3\text{-meV}$ shift for the self-energy correction[16]. But even the QED-corrected value, i.e., $2.189 \text{ eV}$ disagrees with our measurement by $24 \text{ meV}$. The weak point of the calculation appears to be the nuclear magnetic moment. The value given by Lederer and Shirley was measured using the atomic beam resonance method[17], later recalculated[18], and is cited in the literature as $4.160(27)\mu_N$ or $4.173(27)\mu_N$, depending on the diamagnetic correction employed. A newer measurement was performed by Nachtsheim[19] with five times higher accuracy, and the result, $4.132(5)\mu_N$, is compiled in the reviews of Peker[20]. Using this value the theoretical value, including the QED corrections, for the
hyperfine transition energy is 2.1675(28) eV. This differs from our measured value of 2.1646(6) eV by only 0.13% and shows agreement within the 1–σ confidence level. Our measurement, thus, easily distinguishes between the two values of the nuclear magnetic moment, and clearly favors the one measured by Nachtsheim[19].

3 Low-energy x rays: Fine structure of L–shell transitions

For few-electron high-Z ions the n=2 intrashell transitions are in the low-energy x-ray regime. The effects of QED are enhanced in the spectra of the heaviest elements, because QED contributions scale with higher power of Z than the Dirac energies. For example, the Lamb shift for the 2s level in uranium is on the order of −70 eV of which about −40 eV are from QED contributions. Thus, the high-Z elements provide an ideal platform for sensitive tests of QED theory in intense nuclear fields. Additionally, the simplified atomic structure of few-electron high-Z systems allows to interpret the result of isotope shift measurements in terms of the variation in the mean nuclear radius.

3.1 Structure and Lamb shift of L–shell transitions in few-electron high-Z ions

High-precision measurements of the 2p_{3/2} → 2s_{1/2} transitions have been performed for lithiumlike through neonlike thorium[21] at the SuperEBIT facility. These transitions are around 4 keV for the thorium ions. The spectra were taken with a von Hámos-type crystal spectrometer, sketched in Fig. 4. The crystal, a LiF(200) with 2d = 4.027 Å, had a radius of curvature of 30 cm. The diffracted x rays were recorded with a position–sensitive proportional counter with a 10.0×3.0×0.4 cm³ active volume, filled with 70% xenon and 30% methane, 1.2 atm overpressure. The resolving power of this setup was E/ΔE ≈ 2200 and was mainly limited by 270-μm spatial resolution of the proportional counter. As in the case of the ^{165}Ho^{66+} measurement, the instrumentation
included a high-purity germanium detector, which was used to provide a survey of the x-ray emission and, thus, of the charge balance in SuperEBIT. The information about the charge balance was used to support the determination of the ionic species emitting the lines observed with the crystal spectrometer. High-resolution spectra of the \( 2P_{3/2} \rightarrow 2S_{1/2} \) transitions in the eight charge states from Th\(^{80+}\) to Th\(^{87+}\) are shown in Fig. 5. These spectra were taken under different ionization–balance conditions. Figure 5(a) represents a higher abundance of higher charge states than Fig. 5(b). The spectra were calibrated using transitions in hydrogenlike argon, Ar\(^{17+}\), namely Ly-\( \beta \), Ly-\( \gamma \), and Ly-\( \delta \). The uncertainty for the determination of the thorium transition energies is \( \pm 0.13 \) eV, i.e., about 30 ppm, for the strong lines and 0.20 to 0.43 eV, i.e., 45 to 100 ppm, for the weak lines. The accuracy of our measurements allows to test the QED contributions, which are predicted to be around \(-35\) eV for the highly charged thorium ions, within 0.4%.

Stimulated by earlier measurements of the same transitions in lithiumlike through neonlike uranium[22], also performed at the SuperEBIT facility, a lot of effort has been put in calculating the level scheme for these few-electron high–Z ions. Figure 6 shows a comparison between our measured values of the transition energies and several theoretical values for the berylliumlike thorium, Th\(^{86+}\), and uranium, U\(^{88+}\). In our comparison we include: multiconfiguration Dirac–Fock (MCDF) calculations, many-body perturbation theory (MBPT) calculations, and the relativistic configuration–interaction (CI) method. For the MCDF approach the code of Grant and co–workers was used and the QED corrections were taken from calculations in the extended–average level scheme[23, 24]. The result, 4071.7 eV for Th\(^{86+}\) and 4505.3 eV for U\(^{88+}\), significantly disagrees with our measured values, i.e., 4068.47(13) eV, and 4501.72(27), respectively[22, 21]. The MBPT calculations seem to produce better predictions than the MCDF method. In particular, we refer to two recently published papers[25, 26]. The first one[25] combines MBPT calculations of the atomic structure[27] with \( ab \) ini-
tio QED corrections[28, 29, 30]. To give a measure of the convergence of the MBPT method two different model potentials were used in this work to start the MBPT calculations, which were carried out to second order and include a quadrupole–shaped nuclear charge distribution[31]. The formalism was applied to highly charged uranium ions and, thus, two values for the berylliumlike U^{88+} are given, namely 4502.88 eV and 4504.28 eV. These calculated values, still, disagree with our measurements. The second MBPT result[26] we have included in Fig. 6 also has been carried out to second order and includes a QED estimate based on the one–electron Lamb shift data calculated in a (1s^2) potential[29]. The theoretical values for the transition energies obtained by applying spherical nuclear potentials are 4069.02 eV for Th^{86+} and 4501.60 eV for U^{88+}. In the case of berylliumlike uranium, the calculated and measured values agree but there remains significant disagreement for Th^{86+}. Taking the same theoretical method but including a non–spherical charge distribution for the nucleus leads to 4068.36 eV for the Th^{86+} transition energy, which, now, agrees with our experimentally obtained value. The third theoretical approach, presented in Fig. 6, to obtain the 2p_{3/2} \rightarrow 2s_{1/2} transitions energies combines CI energies with ab initio QED and mass polarization corrections[32]. The CI results for U^{88+} calculated with the N–electron Dirac–Slater(DS) and two–electron Dirac–Hartree(DH) potentials are 4501.53 eV (CI–DS) and 4501.61 eV (CI–DH), respectively. These provide excellent agreement with our measurement of 4501.72(27) eV.

3.2 Isotope shift measurements

The 2p_{3/2} \rightarrow 2s_{1/2} transition energies in lithiumlike through carbonlike uranium, U^{89+} to U^{86+}, have been measured for two isotopes, ^{233}U and ^{238}U[33]. It was the first measurement to isolate the variation of nuclear effects in the x–ray transitions of few–electron heavy ions. The experimental setup, shown in Fig. 4, was similar to the one described in Section 3.1. For this measurement the radius of curvature of the LiF(200)
crystal was 75 cm. The resolving power of the von Hámos-type crystal spectrometer was approximately 5000. The radioactive $^{233}$U ions were introduced into the trap using a thin platinum wire. The tip of the wire was plated with a total mass of 100 ng of $^{233}$U, and less than 10 ng were consumed during the course of the experiment. A MeVVA source containing a $^{238}$U cathode provided the $^{238}$U ions. Data collection alternated between $^{233}$U and $^{238}$U spectra. Figure 7 shows the acquired spectra. Each spectrum represents 150 h of data accumulation at an electron-beam energy of 135 keV and electron-beam current of 180-220 mA. The dispersion of the spectrometer was determined using previous $^{238}$U transition energy measurements[22]. In order to infer the Coulomb shift ($\delta E_{\text{Coul}}$) and thus the variation in the mean-square nuclear charge radius ($\delta\langle r^2 \rangle$), the isotopic variation of the specific mass shift, of the QED terms, and of the nuclear polarization has to be estimated[34, 27, 28, 35, 36]. The uncertainty in this estimation is, however, small, which makes the use of highly charged ions ideal for measurements of the isotopic variation of the nuclear charge radii.

The measurements for the energy shift, $\Delta E$, are listed in Table 1 together with the inferred values for $\delta E_{\text{Coul}}$ and $\delta\langle r^2 \rangle$. The average value which can be derived for $\delta\langle r^2 \rangle_{233,238}$ is $-0.457(43)$ fm$^2$. This result can be compared with that of previous studies: neutral-atom optical isotope shift measurements obtaining a value of $-0.383(44)$ fm$^2$[37, 38], and muonic-atom x-ray observations achieving $-0.520(81)$ fm$^2$ for the variation of the mean-square nuclear charge radius[31]. The SuperEBIT measurement, thus, favors neither of the earlier measurements. It was the first time that precise isotope shift measurements have been performed in electronic transitions that are strongly affected by QED. The precision of our method rivals that of the "standard" methods allowing a reexamination of much of the nuclear radii data.
High-resolution x-ray measurements on EBIT have so far been performed only with reflection-type crystal spectrometers. These have worked efficiently for x-ray energies up to 13 keV, e.g., the K-shell radiation of heliumlike krypton[11]. In order to extend crystal spectrometer measurements to higher-energy x rays from higher-Z elements, we have designed a transmission-type crystal spectrometer[39]. It enabled us to observe individual transitions in the K-shell spectrum of high-Z ions well above krypton. In particular, we observed the $1s^2p^3p^1 \rightarrow 1s^21S_0$ and $1s^22S_1 \rightarrow 1s^21S_0$ transitions in heliumlike xenon, Xe$^{52+}$, which have an energy of about 30 keV and which had never before been spectroscopically resolved. Precision measurements of heliumlike and hydrogenlike xenon Kα transitions have been performed by Briand et al.[40] using a germanium detector with a resolution of 270 eV. The current setup improves the spectral resolution by more than an order of magnitude.

Our transmission-type spectrometer design is based on the DuMond geometry[41] employing a cylindrically bent crystal. The radius of curvature of the crystal is the diameter of the so-called Rowland circle (see Fig. 8). Placing the x-ray source, i.e., the trap region of SuperEBIT, on the Rowland circle strongly reduces the bandwidth of the diffracted x rays with the advantage that the throughput is tremendously increased at a given wavelength of interest. If the opening angle of the crystal is small enough, the diffracted x rays are quasimonochromatic. Therefore, no position sensitive detector is necessary, and solid state detectors, such as high-purity germanium detectors, can be used with almost 100% counting efficiency for 30 keV photons. To obtain a spectrum we rotate the crystal using a stepper motor mounted on a modified rotation stage. At each crystal position we count the number of photons reaching the detector and tag this number with the current position of the stepper motor. We normalized each step using the countrate of directly excited lines, emitted by the highly charged ions in the trap, which were measured with a second germanium detector.
A layout of the transmission-type crystal spectrometer for SuperEBIT is shown in Fig. 8. In order to observe the Kα radiation of heliumlike xenon, Xe⁵²⁺, which is situated near 30 keV, we use a Quartz crystal cut perpendicular to the (1340) planes, $2d = 2.3604\ \text{Å}$. Thus, the nominal Bragg angle is around $\theta = 9.9^\circ$. The change in the Bragg angle is $0.000741^\circ \pm 0.000001^\circ$ per step which is equivalent to an energy change of about 2.2 eV for the diffracted photons at these x-ray energies. The radius of curvature of the crystal is $R_c = (2713.8 \pm 3.2)\ \text{mm}$ and was measured using an optical setup. The illuminated area of the crystal is $(60 \times 40)\ \text{mm}^2$. Positioning the crystal so that the electron beam is part of the Rowland circle is non-trivial. At perfect alignment the energy spread of the diffracted x-ray photons reaching the detector is less than 0.5 eV across the detector area. If the distance between the crystal and SuperEBIT is off by ±10 mm, the bandwidth of the diffracted x-rays is increased to $\approx 3.5\ \text{eV}$. The influence of the finite width of the source, i.e., the width of the electron beam $\Delta x_{\text{beam}} = 60\ \mu m$, is about 4 eV for every diffracted x-ray photon. Therefore, the nominal resolution of our transmission-type spectrometer cannot be better than $\Delta E = 4.5\ \text{eV}$, and the resolving power not better than $E/\Delta E = 6800$ for this setup. Further limitation of the resolving power is due to the quality of the focus of the bent crystal.

Figure 9 presents the result of the high-resolution measurement of some heliumlike Kα transitions using the transmission-type crystal spectrometer. The spectrum shows the $1s2s^2S_1 \rightarrow 1s^22^1S_0$, and $1s2p^3P_1 \rightarrow 1s^22^1S_0$, transitions in heliumlike Xe⁵²⁺. It took 131 hours to collect the amount of counts shown in Fig. 9. One channel represents the sum of the counts obtained during three consecutive steps. Thus, the dispersion is 6.6 eV per channel. Using a simple Gaussian fit we obtain a full width at half maximum of $(20 \pm 5)\ \text{eV}$ for these transitions. Therefore, the measured resolving power of our transmission-type crystal spectrometer is about 1500. This shows that the resolving power is limited by the quality of the focus of the crystal.
shows the feasibility of resolving individual lines in the spectra of very highly charged ions and sets the stage for further such measurements in the future.

5 Conclusion

The spectroscopy measurements at SuperEBIT have been developed to a point where high-precision measurements are possible over a wide range of photon energy, from the visible light to hard x rays. This includes very detailed tests of relativistic QED theory in few-electron high-Z systems, i.e., in intense nuclear fields, where theory is tested at the sub 1% level. SuperEBIT measurements not only test atomic theory but are also sensitive to nuclear parameters. We are able to measure nuclear magnetic momenta at an accuracy level of 0.2%. Moreover, we have been able to measure the isotope variations in the nuclear charge distribution. the measurements are competitive with results obtained with nuclear physics techniques. This is a new area for spectroscopy, and more detailed investigations will be done in the future.

Acknowledgements

We would like to thank Gregory Brown, Vincent Decaux, and Daniel Savin for their participation on all the experiments. We also thank Ed Magee and Dan Nelson for their technical support. This work was performed under the auspices of U.S.DOE by Lawrence Livermore National Laboratory under contract # W-7405-ENG-48 and was supported by the Office of Basic Energy Sciences, Division of Chemical Science.

References

157–163.


observation of the spontaneous emission of the hyperfine transition F=4 to F=3 in ground state hydrogen–like $^{165}$Ho$^{66+}$ in an electron beam ion trap, submitted to Phys. Rev. Lett.


[22] P. Beiersdorfer, D. Knapp, R.E. Marrs, S.R. Elliott, and M.H. Chen, Structure and lamb shift of $2p_{3/2}–2s_{1/2}$ levels in lithiumlike U$^{89+}$ through neonlike U$^{82+}$,


Table 1: Measured energy shifts between the $^{233}\text{U}$ and $^{238}\text{U}$ ions for the transitions shown in Fig. 7, and inferred values for the Coulomb shift ($\delta E_{\text{Coul}}$) and the variation in the mean-square nuclear charge radius ($\delta\langle r^2\rangle_{233,238}$)

<table>
<thead>
<tr>
<th>Key</th>
<th>$E_{233\text{U}} - E_{238\text{U}}$</th>
<th>$\delta E_{\text{Coul}}$</th>
<th>$\delta\langle r^2\rangle_{233,238}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>meV</td>
<td>meV</td>
<td>fm$^2$</td>
</tr>
<tr>
<td>Li</td>
<td>$256 \pm 118$</td>
<td>$280 \pm 118$</td>
<td>$-0.364 \pm 0.153$</td>
</tr>
<tr>
<td>Be</td>
<td>$300 \pm 61$</td>
<td>$324 \pm 61$</td>
<td>$-0.436 \pm 0.081$</td>
</tr>
<tr>
<td>B-1,2</td>
<td>$320 \pm 52$</td>
<td>$344 \pm 52$</td>
<td>$-0.455 \pm 0.068$</td>
</tr>
<tr>
<td>C</td>
<td>$362 \pm 62$</td>
<td>$386 \pm 62$</td>
<td>$-0.515 \pm 0.083$</td>
</tr>
</tbody>
</table>
Figure 1: Layout of the prism spectrograph in the horizontal plane of EBIT. The electron beam is perpendicular to the page.

Figure 2: (a): Radiative recombination spectrum from SuperEBIT with a holmium-filled trap showing the presence of the bare Ho$^{67+}$ and hydrogenic Ho$^{66+}$ ions in the trap. The electron-beam energy was set to 133 keV. (b): Spectra, recorded with the CCD camera, showing the F=4 to F=3 hyperfine transition of the 1s ground level of hydrogenlike $^{165}$Ho$^{66+}$ and the result when there is only heliumlike and lower charge states of holmium in the trap.

Figure 3: Comparison of our measured value for the F=4 to F=3 hyperfine transition and theoretical predictions. The hatched area marks the 1–σ confidence limit of our measurement. Open symbols (Δ without, and ○ with QED contributions) refer to calculated values using the nuclear magnetic moment ($\mu_I$) tabulated by Lederer and Shirley[14]. The ●–value is based on the same calculations as ○ but using a more recent and much more accurate measurement for $\mu_I$ of $^{165}$Ho[19]. The error bars reflect the uncertainty of the $\mu_I$ measurements.

Figure 4: Schematic layout of the van Hámós-type crystal spectrometer on SuperEBIT. A high-purity germanium detector concurrently monitors the x-ray emission. The wire probe was used to introduce radioactive $^{233}$U ions into the trap (see Section 3.2). The electron beam is perpendicular to the page.

Figure 5: Spectra obtained with the van Hámós-type crystal spectrometer of the $2p_{3/2} \rightarrow 2s_{1/2}$ transitions in lithiumlike Th$^{87+}$ through neonlike Th$^{80+}$. Spectrum (a) was taken at a higher ionization balance conditions than spectrum (b), which favors the lower charge states, i.e. carbonlike through neonlike. The labels show the isoelectronic sequence. If more lines are affiliated to the same ionic species, they are numbered in respect to their transition energies.
Figure 6: Comparison between our measurement and calculated values for the
\((2s_1/2p_3/2)_J=1 \rightarrow (2s^2)_J=0\) transition in berylliumlike Th\(^{66+}\) and U\(^{88+}\). The figure consists of two graphs, one for \(E_{\text{theo}} > E_{\text{expt}}\), which is on top of the graph representing the case \(E_{\text{theo}} < E_{\text{expt}}\). The hatched area refers to the 68% confidence limit of our measured values. The dashed lines show the energies obtained by theory when neglecting QED contributions. MCDF'93 (+)…multiconfiguration Dirac–Fock calculations[22, 21]. MBPT'95 (ο)…many–body perturbation theory calculations[25]. The two different results are obtained by starting the calculation with different model potentials. MBPT'96 (Ω)…Ref.[26]. The open symbols mark calculations using a spherical nuclear charge distribution, the values marked by filled symbols were obtained after feeding a non–spherical nuclear potential into the formalism. CI'96 (ο, χ)...relativistic configuration interaction calculations with the N–electron Dirac–Slater (DS) and two–electron Dirac–Hartree (DH) potentials, respectively[32].

Figure 7: Crystal–spectrometer spectra of the \(2p_{3/2} \rightarrow 2s_{1/2}\) transitions in lithiumlike U\(^{89+}\) through oxygenlike U\(^{35+}\) for the two isotopes \(^{233}\)U and \(^{238}\)U. The labels show the isoelectronic sequence. The \(^{233}\)U spectrum is offset by 500 counts/channel. The dashed lines indicate the center of gravity of the \(^{238}\)U lines.

Figure 8: Layout of the transmission–type crystal spectrometer in the horizontal plane of SuperEBIT. The electron beam is perpendicular to the page. GeSE and GeX are high–purity germanium detectors; \(\theta\)…Bragg angle;

Figure 9: First resolved spectrum of the \(1s2p^3P_1 \rightarrow 1s^21S_0\) and \(1s2s^3S_1 \rightarrow 1s^21S_0\) transitions in heliumlike Xe\(^{52+}\) observed with the transmission–type crystal spectrometer.
Figure 2
Figure 3

$E_{\text{theo}} - E_{\text{expt}}$ (meV)
Figure 4

Gas injector

LN2 shield

LHe shield

Wire probe

Position sensitive detector

X rays

He atmosphere

LiF Crystal

Ge detector

Super EBIT

X rays
Figure 5
Figure 6
He-like $\text{Xe}^{52+}$

$1s2s^3S_1 \rightarrow 1s^2 \, ^1S_0$

$1s2p^3P_1 \rightarrow 1s^2 \, ^1S_0$

X-ray Energy (eV)

Counts

$20 \text{ eV}$

$\pm 5 \text{ eV}$

Figure 9